## BACTERIOLOGIGAL AND GENETIC STUDY OF Pseudomonas aeruginosa ISOLATES S. M. Tawfiq

Dept. Soil &Water Sci.,Coll. Agric.,Univ. of Sallahaddin- Erbil- Kurdistan Region e.mail.dr.shamerantawfiq@gmail.com

#### ABSTRACT

Fifty P. aeruginosa isolates were isolated from patients suffered of wound, ear and chronic pulmonary infections to assess the presence of nan1 and nan2 genes .P. aeruginosa isolates were 10(20%) from chronic pulmonary infection ,20(40%) from each of ear and wound infections. isolates exhibted highly sensitivity to (Cefepime ,Meropenem ,Azitheromycin) and Gentamicin, version resistance to Cefotaxime, Ceftazidim .Clorithromycin, Tobramicin, Ciprofluxacin In percentages (70%), (60%), (50%), (40%), (36%) and (34%) respectively. Seventeen isolates were choosen to searching nan1 and nan2 genes by PCR technique depending on the resist to four groups of antibiotics (Aminoglycoside, Third generation of beta-lactam ,quinolones ,Macrolides , (29.4%) possess both genes.nan1 were distributed in(41.2%) in percentages (50%) for each of wound and ear isolates and (20%)for chronic pulmonary isolates, while nan2 was dissemination (70.6%) in percentages100%, 66.6% and 50% for chronic pulmonary, ear and wound isolates respectively In conclusion, these data suggest that the prevalence of *nan2* in all infections ,that suggest apossible role of this gene in disease evolution especially in pulmonary patients while nan1 was low prevalence in isolates may be related to infection site.

Key words: P. aeruginosa, nan1, nan2 genes, neuraminidase, infections.

مجلة العلوم الزراعية العراقية – 27- 35 :(1)49 /49 يتوفيق دراسة باكتيرولوجية وجينية لعزلات P.aeruginosa شاميران محمود توفيق مدرس قسم التربة والمياه-كلية الزراعة-جامعة صلاح الدين-اربيل- اقليم كردستان المستخلص

من اصابات الجروح والاذن والتهابات الرئة المزمنة وذلك P.aeruginosa عزلت خمسون عزلة من بكتريا ،توزعت العزلات الى 10(20%) من التهابات الرئة المزمنة،20(40%) nan1,nan2 للتحري عن الجينين، Cefepim,Meropenem للتحري عن الجينين، Ceftazidim،Clorithromycin لكل من اصابات الاذن الوسطى والجروح،اظهرت العزلات حساسية عالية تجاه ( 43%) Ceftazidim، Clorithromycin ( ( 36%) لكل من اصابات الاذن الوسطى والجروح،اظهرت العزلات حساسية عالية تجاه ( 64%) Azitheromycin ( 36%) و (36%) من الصابات الاذن الوسطى والجروح،اظهرت العزلات حساسية عالية تجاه ( 64%) Azitheromycin ( 36%) ( 36%) ( 36%) و (36%) ( 36%)، (36%)، (36%)، (36%)، (36%)، (36%)، (36%)، الترتيب ( 34%)، من حمواميع معلم المترابعة مجاميع من حمواميه المروكة الله المروكة التحري عن الجن الجينين المضادات الحياتية ( 34%)، (36%)، (36%)، الكينولية للتحري عن الجن الجينين المضادات الحياتية ( 34%)، (36%)، (36%)، المعروبية للتحري عن الجن الجينين المضادات الحياتية ( 34%)، (36%)، (36%)، المعروبية للتحري عن الجن الجينين المضادات الحياتية ( 34%)، (36%)، (36%)، الكينولونات، الماكروليتات)، اظهرت ا nan1 الحينين المضادات الحياتية ( 34%)، الكينولونات، الماكروليتات)، اظهرت ا nan1 الحينين المضادات الحياتية ( 34%)، (36%) من عزلات التضاعفي المتسلسل ان ( 24.4%) من العزلات كانت موجبة للجينين. توزعت الى (12%)، (36%)، في كل من عزلات البنا الحروح والاذن و (20%) في عزلات الرئة المزمن، بينما كانت (36.6%)، (30%)، (36%)، و (35%) لكل من عزلات الرئة المزمن والاذن المسبة الدين المنه، التشار 14 في معرود في تطور الاصابة وخاصة في 200%)، لكل من عزلات الرئة المزمن والاذن المسبة التشار 14 في موقع الاصابة وخاصة في 20%) الكل من عزلات الرئة المزمن والاذن المسبة التشار منخفض ربيا ما دور في تطور الاصابة وخاصة في 20%) لكل من عزلات الرئة المزمن والاذن المسبة الرئة المزمن، بينما كانت (م8.6%)، وموقع الاصابة وخاصة في 20%) لكل من عزلات الرئة المزمن والاذن المسبة النشار منخفض ربيا البينا البين الجين المور الاصابة وخاصة في 20%) الحرف بينما الحين المور الحين المور الاصابة الرئة المزمن بينما المزمن بينما الحمود في تطور الاصابة وحاصة مي 20% مالمي مي من ينما الحين المود الحيام الحيام مولحاصة في لامام الحمود مي المود اللموم اللهم مولحاصة مي 20%

كلمات مفتاحية: التهاب الرئة، مقاومة، حساسية، عزلات.

\*Received:28/11/2017, Accepted:15/2/2018

## **INTRODUCTION**

Pseudomonas aeruginosa is a ubiquitous can micro-organism infect that immunocompromised individuals and is responsible for nosocomial infections. Pseudomonas aeruginosa has many virulence factors such as formation of biofilm(colonies of *P. aeruginosa* composed of alginates) protects the bacterium from the host's immune response and from antibiotics (13,21). Other virulence factors are Exoenzyme S is an ADPribosyltransferase that is secreted by a type-III secretion system directly into the cytosol of epithelial cells (23). Exotoxin A which inhibits protein biosynthesis. LasB elastase has an elastolytic activity, The adhesion of apathogen to epithelial surface is the first step in an infection. *P*. aeruginosa exoproducts, particularly proteinases and neuraminidase (sialidases), are noted in many studies to enhance bacterial adherence and to act as virulence factors and may increase the avaialability of such receptors by cleaving terminal sialic acid residues from cell surface gan gliosides(24) Bacterial adherence to mucosal surfaces is a significant aetiological factor in infections(2), The importance of proteinases is facilitating attachment, Extracellular neuraminidase was secreted in growth medium during the early stationary phase. The enzyme hydrolyzed the alpha 2 leads to 3 glycosidic linkages from Nacetylneuraminlactose and fetuin. Р aeruginosa strains, with detectable proteinases during adherence assay, have the greatest adherence(2).the contribution of bacterial neuraminidase to the pathogenesis of infection is not as clearly defined. Neuraminidaseproducing species such as Hemophilus S. pneumoniae (9), and **Pseudomonas** aeruginosa share a common ecological niche, colonizing the heavily asialylated secretions and surfaces of the upper respiratory tract. Although each can bind to asialylated glycolipids exposed by neuraminidase activity, they differ substantially in their ability to either metabolize or incorporate sialic acid into surface structures, Thus it is likely that bacterial neuraminidases interact with both prokaryotic and eukaryotic glycoconjugates, adherence to tissue is result of bacterial interactions between surface molecules on the

the increasing adherence of P aeruginosa to epithelial cells after exposure to P aeruginosa neuraminidase(3). The identification of putative neuraminidase genes in P. aeruginosa The sequence of the P. aeruginosa PAO1 genome was published recently (accession no. AE004091;. The strategy used to detect putative neuraminidase genes was based on the observation that all described bacterial sialidases have four or five copies of an aspartate box ('Asp-box') motif, Ser-X-Asp-X-Gly-X-Thr-Trp, where X is any amino acid By using this peculiarity and information from the Pseudomonas Genome Project two genes containing four Asp-boxes were identified in the P. aeruginosa PAO1 genome;

bacteria and plasma membrane receptors on

host cell(22,18) for examplles the ability of *P*.

aeruginosa to persist in the respiratory tract is

correlated with the organism's ability to

adhere to buccal epithelial cells (BEC), *nan1* gene initiate colonization and facilitate long-

term infection, P. aeruginosa cells adhere to

BECs via pili, which recognize a GalNAcβ1-4

glycolipids, such as asialo-GM1 (25). Several

arguments suggest that the *nan1* gene encodes

neuraminidase, an enzyme able to release

terminal sialic acid residues from sialylated

gangliosides, thus increasing the amount of

potential bacterial receptors and consequently

enhancing adhesion (11). The removal of sialic acid residues from cell surface proteins also

interactions through reduction of the net

negative surface charge of one of the distinct

neuraminidaseis 1000 fold more active than

realeasig sialic acid this effect correlated with

*Clostridium perfringens* enzyme

higher

sialic

in

in

cell-to-cell

by enhancing protein-protein

acid .P aeruginosa

exposed

disaccharide

results

the

presumably

features of

# we named them *nan*1 and *nan*2 (11). **MATERIAL AND METHODS**

# (I) Bacterial isolation and identification

Pseudomonas aeruginosa isolated from different human infections (Ear. Wond. Sputum) samples activated by culturing in brain heart infusion broth medium and incubation at 37 (18-24) h, After activation inoculated on the MacConKey agar (positive isolates does not ferment lactose), a colonies were selected. for more purification,

asialvlated

contact.

in

inoculated on the selective medium cetrimide agar 0.03% cetrimide, suspected isolates identified by growth at 4°C and biochemical tests were [oxidase(+), indol(-), ureas(-,+), citrate(+),catalase(+)] and tested by microscope which showed Gram negative rod shape, stored at -20°C in tubes contained 4ml brain heart infusion broth+1ml Glecerol, till other bacteriological and molecular tests were done(5,6).

# Antimicrobial susceptibility test

This test was done according to standard methods by disk diffusion 2009 and 2010 CLSI guidelines. *P. aeruginosa* isolates were tested for their susceptibility: Cefepime FEP 30µg, Meropenem MEM 10µg, Azitheromycin ATM 30µg, Cefotaxime CTX 10µg, Gentamicin CN 30µg, Ceftazidim CAZ 30µg, Clorithromycin CLR 15µg, Tobramicin ToB 30µg,Ciprofluxacin Cip 5µg).

# (II) Molecular Biology Studies

Seventeen isolates of *P. aeruginosa* five from chronic pulmonary infection and six from each of ear and wound infections which had appeared resist aginst four groups of antibiotics were chosen to study the presence of two virulence genes *nan1,nan2* (both genes encoding for neuraminidase).

# **DNA extraction:**

The selected isolates were grown in brain heart infusion broth at 37°C for 24 hr, and DNA was extracted by using extraction and purification kit provided from Promega Company (USA). 3 ml was transferred to ml test tube. The growing cells were deposited by centrifugation at 4000 rpm for 10 min then the pellet was resuspended with 100 µl TB buffer vortex to completely suspended the pellet, 10 µl was added lysozyme, incubated at 37 °C for 10 min., added 100 µl RNaseA was added then incubated in room tempreture for 5 min. centrifuged at 10000rpm/min for 2 min. then supernatedwas transferred to 1.5 ml Eppendrof tube, 220 µl BDL buffer was added, incubated in65°C for 10 min. 220 µl ethanol (96-100%)was added and vortexed for 20 sec.at maximum speed, inserted HiBind DNA mini column into a 2 ml collection tube, transferred. The entire sample was added to HiBind including any precipitate, the centrifuged at 10000 rpm/min for 1 min.,

discarded the filtrated and the collection tube, inserted the HiBind into new 2ml collection tube, 500 µl HB buffer was added, centrifuged at 10000 rpm for 1 min., discarded the filtrate and reused collection tube. 700 µl DNA wash buffer was added then centrifuged at 10000 rpm/min for 1 min., discarded the filtrate and reused collection tube, centrifuged the empty HiBind at 10000 rpm for 2 min. inserted HiBind DNA mini column into a 2 ml collection tube, added 50-100 µl preheated elution buffer(65°C) to the HiBind into anew nuclease-free 1.5 ml Eppendrof tube, incubated HiBind DNAcolumn for 3-5 min. at room tempreture, centrifuged at 10000 rpm/min for 1 to eluted the DNA, incubated overnight at 4  $^{\circ}$ C then stored at – 20  $^{\circ}$ C till use.

# Detection of *nan*1 and *nan*2 genes:

Nan1 and nan2 genes determind by PCR (Applied Biosystem, Singapore) using specific primers (Alpha DNA) The frequency of the gene encoding neuraminidase virulence nan1and nan2 was determined by PCR (11). Amplification was performed with specific primers F(5'nan1 ATGAATACTTATTTTGATAT-3') and nan1R (5'-CTAAATCCATGC TC TGA CCC -3') yielding a 1316 bp product and primers nan2 -F. (5'- ACAACAACGGGGACGGTAT-(5'-GTTTTGCTGAT-3') and nan2R vielding a 1161 bp GCTGGTTCA-3'), product, PCR was carried out with 2 µl template DNA, 0.25 µM of each primer, 0.2 mM deoxyribonucleoside triphosphates, 1x reaction buffer, 2 mM MgCl<sub>2</sub> and 1.5 U Prime Taq DNA polymerase (Promega Company ) in a total volume of 25µl shown in Table1. The DNA was amplified using the following protocol: using the following protocol: 94°C for 3 min, 30 cycles of 94°C for 30 s, 55°C for 1 min, and 72°C for 1.5 min, and finally 72°C for5 min. PCR products were separated in 1.5% agarose gel for 50- 110 min at 130 V, stained with ethidium bromide  $(0.5\mu gl^{-1})$  and The electrophoresis products were visualized by UV transilluminator and photographed by Gel Documentation.

Table.1	PCR mixture total volume 25	
Volume of matter		Sequence
added	Component	
13.8µl	<b>Distilled water</b>	1
5µl	PCR Buffer1X	2
1.5µl	DNTPs	3
1.2µl	Primer-F	4
1.2µl	Primer-R	5
0.3µl	(1.5U)TaqDNA Polymerase	6
2µl	DNA	7
25µl		<b>Total volum</b>

~-

Table.2 PCR mix	ture total volume 25
-----------------	----------------------

Volume of matter		Sequence
added	Component	
13.8µl	Distilled water	1
5µl	PCR Buffer1X	2
1.5µl	DNTPs	3
1.2µl	Pr	rimer-F 4
1.2µl	Primer-R	5
0.3µl	(1.5U)TaqDNA Polymerase	6
2µl	DNA	7
25µl	Total volum	

#### **RESULTS AND DISCUSSION**

Seventeen isolates were selected to evaluate the prevalence of two virulence genes nan1 and nan2 genes among 50 isolates of P. aeruginosa which had appeared resist to four groups of antibiotics (Aminoglycoside, Macrolides, quinolone, Third generation of betalactam) (Table 3 shows the antimicrobial patterns) , In current study resistance seventeen isolates we choosed according to (12) whom result that the proportion of multi drug-resistance nosocominal P.aeruginosa isolates containing alarg number of virolence genes (38.1%) was higher than the proportion of non multi drug -resistant isolates (17.6%).and according to (10) with result that most virulence factors were produced at higher rate in ESBLs producing than in no ESBLs Pseudomonas producing aeruginosa are isolates Table 4, figures1,2,3,4,5,6 illustrate the profiling of the amplification products of nan1 and nan2 virulence genes in each of the 17 tested isolates),(29.4%) of isolates possess both genes, As shown figures.1,2,3,nan1 was detected in 7(41.2%) PCR products of the(17) tested isolates of Pseudomonas aeruginosa possess this gene, the result showed that this gene in pulmonary infection was (20%) and (50%) in each of wound and ear infections,

since gel electrophoresis results showed DNA bands with molecular size 1316 bp in comparison with DNA marker. While nan2 gene was detected in the (70.6%) in all studied isolates, with differences to site of infection, the result showed higher spread this gene among pulmonary isolates(100%) and(66.6%) (50%) in each of ear and wound isolates respectively in these isolates, since gel electrophoresis result showed DNA bands with molecular size 1161bp in comparision with DNA marker for all tested isolates as shown in figure4,5,6. Our results were not in agreement with the results reported by (8) reported that the nan1 gene was detected in 57.9% of P. aeruginosa isolates causing bloodstream infections, and with (23) whom detected nan1 gene in 52 (83.9%) from 62 of hospital P. aeruginosa strains and suggested that there was no significant difference in the frequency of this gene among P. aeruginosa strains from sources, different isolated they significantly observed higher adhesion of these bacteria to human buccal epithelial cells when the *nan*1 gene was present than when the same gene was absent, they found in their study that the *nan1* gene may play a role in the binding of clinical P. aeruginosa strains to buccal cells, and with (15) whom result reported low

frequency of this gene(15%) among their study isolates. (12) reported that the frequency of this gene is low(16.9%) in non-cystic fibrosis isolates of *Pseudomonas aeruginosa* which isnot in agreement with our results and the role of this factor is not known in the pathogenicity of this bacterium in non- cystic fibrosis patients. Astudy performed by (10) on 44 MDRPA isolates, 12 isolates representing different resistance profiles and sources of samples were selected for further molecular studies showed that only 4(33.3%) of them possess *nan1* gene which was relatively low,While (18).detected nan1 gene in39(38.2%) among 102 cystic fibrosis P.aeruginosa isolates, and were relatively agreed with our results. (11) used PCR to assess the prevalence of eight virulence genes (algD, lasB, toxA, plcH, plcN and nan2), PCR detected algD, lasB, toxA, plcH, plcN and nan2 in all of the 162 isolates used to nan1 detected in isolates (44.4 %). was demonstrated that the *nan1* gene was present in 61.7% of CF isolates and 44.4% of non-CF isolates. The frequency of this gene in CF isolates tended to increase as the clinical status worsened. Indeed, nan1 was detected in 57% of isolates from patients with an excellent or good clinical status, in 63% of isolates from patients with a moderate status, and in 71% of isolates from patients with a poor or weak clinical status. and suggested that the prevalence of this gene in CF isolates tended to increase as the clinical status worsened. Indeed, nan1 was detected in 57 % (23/40 isolates) of isolates from patients with an excellent or good clinical status, in 63 % (17/27) of isolates from patients with a moderate status and in 71 % (10/14 isolates) of isolates from patients with a poor or weak clinical status and the distribution of nan1 was significantly related to strain origin and the role of this factor is not known in the pathogenicity of this bacterium in non-CF patients, Lanotte results were agreed with present results except for *nan*2, present study showed (100%)pulmonary isolates possess this gene followed (66.6%) and (50%) for ear and woud isolates may be that related to the virulence of P. aeruginosa is multifactorial and caused by several extracellular enzymes and other substances. The importance of these

human P. aeruginosa infections is dependent on the type of infection (7). The differences in the distributions of virulence factor genes in the populations strengthen the probability that some P. aeruginosa strains are better adapted to the pulmonary conditions found in CF patients (1). (16) reported that the PA2794 neuraminidase locus (Delta 2794) mutant of P. aeruginosa PAO1 was unable to colonize the respiratory tract. They suggest that inhibition of bacterial neuraminidase could provide a novel mechanism to prevent colonization of the respiratory tract by this important pathogen(nan1,nan2) genes is one of many virulence factors that can play an important role in the pathogenesis of P. aeruginosa infection (1). Nan1, nan2The first argument is that the deduced bacterial protein encoded by nan1, nan2 possesses four Asp-boxes, a characteristic of bacterial sialidases (20). Secondly, the deduced protein contains conserved amino acids at key sites that are probably part of the catalytic site. Although there is a low degree of similarity between nan1, nan2 and previously identified bacterial sialidases, this is usual for bacterial sialidase genes (20). Nevertheless, part of the deduced protein encoded by nan1 has 26 % identity to part of the Clostridium tertium sialidase protein, The G+C content of nan1 (48.2 mol%) differs notably from the G+C content of the whole PAO1 genome (66.7 mol%) (17). Thus, the gene was probably acquired by horizontal transfer, as observed previously for some other bacterial sialidase genes. Several arguments suggest that nan1, nan2 encodes a sialidase, an enzyme theoretically able to release sialic acid from sialylated gangliosides, thus increasing the amount of asialoGM1, a major receptor for adherence to the respiratory tract(22), increased virulence in environmental strains through acquisition of new virulence genes (such as exoU) via horizontal transmission is a feasible explanation (11). P. aeruginosa is not an obligate parasite of humans, many factors harmful to humans expressed by this organism will most probably perform a more innocuous function within the organism's natural habitat. Virulence factors in P. aeruginosa are divided into specific groups dependent upon their mode of action or

virulence factors for the pathogenesis of

## Tawfiq

method of delivery to the host cell. Thus, these virulence factors may be described as belonging to adhesins and secreted toxins and enzymes dependent or independent of the type I secretion system (T1SS), type II secretion system (T2SS) and type III secretion system (T3SS)(2). The GalNAc $\beta$ 1-4Gal sequence present in asialylated gangliosides can act as areceptor for several pathogens of the respiratory or gastrointestinal tract including *P. aeuroginosa*, Neuraminidase has been implicated as a virulence factor and may serve as a marker for determining virulence of *P*.

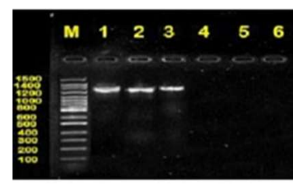
*aeuroginosa* strains. This enzyme has a key role in the initial stages of pulmonary, urinary and gastrointestinal tract infections by targeting bacterial glyco-conjugates and contributing to the formation biofilm(19). present study concluded high spread nan2 gene among clinical isolates compared with nan1 gene, (100% vs.20%, 66.6% vs.50%, 50% vs.50%) in pulmonary, ear, wound infections respectively, the distribution of virulence genes encoding adhesins (neuraminidase) varied in respect to the infection localization in hospitalized patients.

Table 3.sensitivity tests percentages of P.aeruginosa isolates for selected antibiotics

Seq	Antibiotics	symb ol	R- Resista nt	R%	S- Sensitive	S%
1.	Tobramicin	То	18	36	32	64
2.	Gentamicin	GM	30	60	20	40
3.	Cefepime	CEF	Zero	zero	50	100%
4.	Clorithromycin	CLR	25	50	25	50
5.	Cefotaxime	CFm	35	70	15	30
6.	Ceftazidime	CFX	20	40	30	60
7.	Meropeneme	MPM	Zero	zero	50	100%
8.	Azithromycin	ATM	Zero	zero	50	100%
9.	Ciprofloxacin	CIP	17	34	33	66

#### Table.4 nan1 and nan2 genes diversity in isolates from different source

Nan2	Nan1	Isolate site	Seq.
5(100%)	1(20%)	Sputum(5)	1-
	3(50%)	Wound(6)	2-
3(50%)	3(50%)	<b>Ear(6)</b>	3-
	7(41.2%)	17	Total
4(2066.6%)			
12(70.6%)			



#### Figure1.result amplification of nan1 of *Pseudomonas aeruginosa* isolated from wound

Agarose gel electrophoresis(1.5% concentration of agarose,60 v voltage,2 h) of specific PCR product (1316bp) by specific primers to *nan1* gene. Lane M represents 100 bp DNA ladder markers. Lanes 1,2,3 represent positive amplification process of the *nan1* 

gene in DNA of *P. aeruginosa* isolate, Lanes 4,5,6 represent negative amplification process of the *nan1* gene in DNA of *P. aeruginosa* isolate. Lanes from 1 to 6 represents isolates respectively

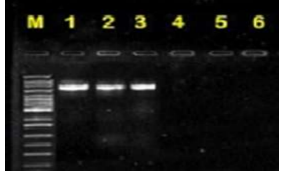


Figure 2. result from amplification of nan1 of *Pseudomonas aeruginosa* isolated from ear Agarose gel electrophoresis (1.5% concentration of agarose,60 v voltage,2 h) of specific PCR product (1316 bp) by specific primers to *nan1* gene. Lane M represents 100 bp DNA ladder markers. Lanes 1,2,3 represent positive amplification process of the *nan1* gene in DNA of *P. aeruginosa* isolate, Lanes 4,5,6 represent negative amplification process of the *nan1* gene in DNA of *P. aeruginosa* isolate. Lanes from 1 to 6 represents isolates respectively

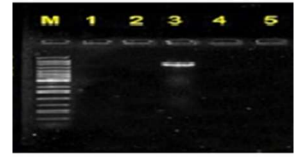


Figure 3. result from amplification of nan1 of *Pseudomonas aeruginosa* isolated from sputum

electrophoresis Agarose gel (1.5%)concentration of agarose,60 v voltage,2 h) of specific PCR product (1316 bp) by specific primers to nan1 gene. Lane M represents 100 bp DNA ladder markers. Lane 3 represent positive amplification process of the nan1 gene in DNA of P. aeruginosa isolate, Lanes 1.2.4.5 represent negative amplification process of the nanl gene in DNA of P. aeruginosa isolate. Lanes from 1 to 5 represents isolates respectively

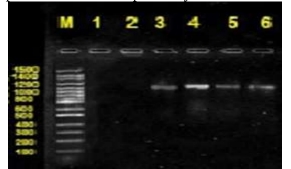


Figure 4. result from amplification of nan2 of *Pseudomonas aeruginosa* isolated from ear

Agarose gel electrophoresis (1.5% concentration of agarose, 60 v voltage, 2 h) of specific PCR product (1161 bp) by specific primers to *nan2* gene. Lane M represents 100 bp DNA ladder markers. Lanes 3, 4, 5, 6 represent positive amplification process of the *nan2* gene in DNA of *P. aeruginosa* isolate, Lanes 1,2 represent negative amplification process of the *nan2* gene in DNA of *P. aeruginosa* isolate, Lanes 1,2 represent negative amplification process of the *nan2* gene in DNA of *P.* 

*aeruginosa* isolate Lanes from 1 to 6 represents isolates respectively

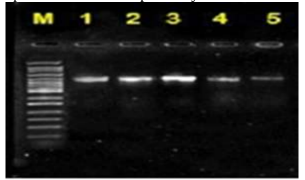
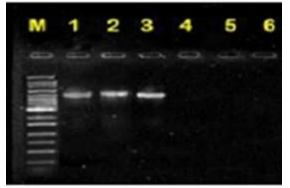


Figure5. Result from amplification of nan2 of *Pseudomonas aeruginosa* isolated from sputum

Agarose gel electrophoresis(1.5% concentration of agarose, 60 v voltage,2 h) of specific PCR product (1160 bp) by specific primers to *nan2* gene. Lane M represents 100 bp DNA ladder markers. Lanes 1, 2, 3, 4, 5 represent positive amplification process of the *nan2* gene in DNA of *P. aeruginosa* isolate, Lanes from 1 to 5 represents isolates respectively



## Figure 6. resulted from amplification of nan2 of *Pseudomonas aeruginosa* isolated from wound

Agarose gel electrophoresis(1.5% concentration of agarose, 60 v voltage, 2 h) of specific PCR product (1160 bp) by specific primers to *nan2* gene. Lane M represents 100 bp DNA ladder markers. Lanes 1, 2, 3 represent positive amplification process of the *nan2* gene in DNA of *P. aeruginosa* isolate, Lanes 4,5,6 represent negative amplification process of the *nan2*gene in DNA of *P. aeruginosa* isolate. Lanes from 1 to 6 represents isolates respectively

#### REFERENCES

1. Antonov V. A.; V.V,Altukhova;S.S. Savchenko; G.A. Tkachenko;V. Zamaraev V.; S.I. Zhukova; O.G. Kramar; N.L. Matveeva;

O.V. Ostrovskiĭ and Z.G. Dudchenko. 2010. Molecular genetic analysis of *Pseudomonas aeruginosa* strains isolated from environment and patients in health care facilities. Mikrobiol Epidemiol Immunobiol (2):8-13

2. Bradbury,L.; A. Roddam; D.Merritt,; M. Reid; and A. Sanre.2010.ChampionVirulence gene distribution in clinical, nosocomial and environmental isolates of *Pseudomonas aeruginosa*.J. Medical Microbiology., 84 (5-6): 499-510

3. Cacalano, G., M. Kays, L. Saiman; and A.Prince. 1992. Production of the *Pseudomonas aeruginosa* neuraminidase is increased under hyperosmolar conditions and is regulated by genes involved in alginate expression. J Clin Invest 89, 1866–1874

4. Clinical and labratory standards institute CLSI. *Performance* Standards for Antimicrobial Susceptibility Testing:. Wayne. 2009. 181 p. M100-S-18 Eighteenth Informational Supplement

5. Collee, J.G.;A. Fraser; B.P.Marmion and A.Simmons. 1996. Mackie and McCartney Practical Medical Microbiology". 14th ed., Churchill Livingstone, New York, pp. 413-423 6. Cruickshank, R.; J. P. Duguid and R. H. A. Swain. 1975. Medical Microbiology, a Guide to the Laboratory Diagnosis and Control of Infection. 11th ed., Churchill Livingstone, Edinburgh and London and New York

7. Doring G. 1987. Significance of Pseudomonas aeruginosa virulence factors for acute and chronic Pseudomonas aeruginosa infections. Infection.15 Suppl 2: S38-41

8. Endimani, A.; B.Pini,;A.Boj and A.Toniolo. 2006.Blood stream infection due top aclinical outcome associated with pathogemesis-related gene.ESCMID.European society of clinical Microbology and Infections disease., 3(7):1164-1180.

9. Ghazaei, C,; M. Ahma and N.Hosseini. 2010. Optimization and comparative characterization of neuraminidase activities from *Pseudomonas aeruginosa* with *Klebsiella pneumoniae*, Hep-2 cell, sheep kidney and rat liver lysosome,Iranian.J.Microbiology.2(1):30-17.

10. Khalil,M.; F.Ibrahim ;M.Sonbol; F.Badr and S.Ali. 2015. Comparative study of virulence factors among ESβLs producers and non producers *Pseudomonas aeruginosa*  clinical isolates. Turkish Journal of Medical Sciences, 45: 568-577

11. Lanotte, P.; S.Watt.;L. Mereghetti.;N. Dartiguelongue.; A.Rastegar-Lari; A.Goudeau and R.Quentin. 2004. Genetic features of *Pseudomonas aeruginosa* isolates from cystic fibrosis patients compared with those of isolates from other origins. J. Med. Microbiol. 53 (1): 73-81

12. Mitov, I., T.Strateva and B. Markova. 2010. Prevalence of virulence genes among bulgarian nosocomial and cystic fibrosis isolates of *Pseudomonas aeruginosa* .Braz. J. Microbiol .41 ; 3

13. Moradian, F.; E. Perdosi and Z. Molana. 2012.Molecular detiction of integron genes and pattern of antibiotic in *P. aeruginosa* strains isolates from intensive unit.Iran .J. MAM.,1(4):424.

14. National Committee for Clinical Laboratory Standards 2010. "Peformance Standards for Antimicrobial Susceptibility Testing ; 18 th Informational Supplement. J. Clin. Microbiol., 28(10): 2331-2334

15. Raoof,W. 2011.Distribution of *algD,lasB,pilB* and *nan1* genes among MDR clinical isolates of *P.aeruginosa* in respect to site of infection.

16. Soong G.; A.Mui; M.Gomez; J.Waks; B.Reddy; P.Planet;P.K. Singh.;Y. Kanetko; M.Wolfgang; Y.Hsiao; L.Tong and A.Prince. 2006. Bacterial neuraminidase facilitates mucosal infection by participating in biofilm production. J. Clin. Invest. 116: 2297–2305. pathogen. Nature

17. Stover, K.; Q.Pham and A.Erwin 2000. Complete genome sequence of *Pseudomonas aeruginosa* PAO1, an opportunistic pathogen. Nature 406, 959–964

18. Strateva T.; G.Petrova ;P. Perenovska and I.Mitov. 2009. Bulgarian cystic fibrosis *Pseudomonas aeruginosa* isolates: antimicrobial susceptibility and neuraminidase-encoding gene distribution .J Med Microbiol . 58 , 5 : 690-692

19. Tang, HB.;E. Dimango; R.Bryan.; M.Gambello;BH. Iglewski and JB.Goldberg 1996. Contribution of specific *Pseudomonas aeruginosa* virulence factors to pathogenesis of pneumonia in a neonatal mouse model of infection *Infect Immun* . 64: 37-43 20. Taylor, G. 1996. Sialidases: structures, biological significance and therapeutic potential. Curr Opin Struct Biol 6, 830–837

21. Wilson, A.;T.Treasure; M.Sturridge and R.Gruneberg. 1986. A scoring method(ASEPSIS) for postoperative wound infections for use in clinical trials of antibiotic prophylaxis. *Lancet* : 311-313

22. Wolska K.;P Rudaś and A.Jakubc zak. 2005. Reduction in the adherence of *Pseudomonas aeruginosa* to human buccal epithelial cells with neuraminidase inhibition. Pol. J. Microbiol. 54: 73–76 pathogen. Nature 23.Wolska,K,;B.Koti;C.Mioduszewska;L.Bork owskaandK.Rymuza. 2012.Occurrence of the *nan1* gene and adhesion of *Pseudomonas aeruginosa* isolates to human buccal epithelial cells Biological let. 49(1): 59–64

24. Wolska, K. and P.Szweda. 2009. Genetic features of clinical *Ps. aeruginosa* strains. J. Bacteriol., 58, 3, 255-260

25. Woods, D. W.Stras,; J. Johanson; M.Berry and J.Bass. 1980.Rol of pili in adherence of *P.aeruginosa* to mammalian buccal epethlial cell. infect, immune, 29, 1146-15.