

Full length Research paper

Integrated structure of rearing *Mugil* species in a crude oil treated saltwater using an oceanic *Pseudomonas aeruginosa* damage elnaggar

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This examination intended to make an incorporated model framework to bio-remediate got away raw petroleum in marine condition and concentrate the effect of the bioremediation forms on the oceanic living life forms utilizing *Mugil cephalus* fries as a living contextual analysis. Three unrefined petroleum fixations (100, 300 and 500 ppm) were tried utilizing glass aquaria vaccinated with *M. cephalus* fries. The substance treatment was done utilizing a business oil disperser and a microbial treatment was done by a marine bacterial seclude (*Pseudomonas aeruginosa* strain elnaggar1), the trial was stretched out for 45 days. The effects of these medicines on the development execution and the survival percent of these *M. cephalus* fries were recorded by the adjustments in their clinical signs, lengths and weights along the raising time frame. The outcomes demonstrated that the microbial treatment utilizing *P. aeruginosa* is more compelling for the remediation of the raw petroleum defiled seawater and furthermore to keep the development execution of the tried fish as comparative as the untreated fries. Then again, the utilized substance treatment prompted increment the vulnerability of these tried fries towards the decay balance illness.

Key words: Bioremediation, *Mugil cephalus*, crude oil, *Pseudomonas aeruginosa*, marine bacteria.

INTRODUCTION

Petroleum production and operation produce serious ecological problems. Pollution of environment due to accidental oil spillage, seepage and rupture pipelines had been well reported (Okerentugba and Ezeronye, 2003). Moreover, oily wastewater, especially from oil field, has posed a great hazard for terrestrial and marine ecosystems. The traditional treatments of oily wastewater, such as containment and collection using floating booms, adsorption by natural or synthetic materials, etc., cannot degrade the crude oil thoroughly. So far, the biodegradation processes suggest an effective method where the crude oil considered as a carbon source for microbial growth, which results in the breakdown of the oil to lower molecular weight compounds (Guo-liang et al., 2005).

However, on creation a system include three major components oil, fish and bacteria many challenges appear,

firstly, Giles et al. (2006) determine that the exposure of the mullet (*Mugil cephalus*) to crude oils had relationship with the fin rot disease. In addition, rearing of *M. cephalus* in presence of some bacterial strains such as *Vibrio anguillarum* and *Aeromonas hydrophila* led to a fish disease known as 'red spot' (Burke and Rodgers, 2006; Rodgers and Burke, 2006). Moreover, the bacterial utilization of the crude oil components requires complex cell surface adaptation to allow adherence to oil and precede the degradation process. Meanwhile, it was mentioned that the use of probiotics for disease control in aquaculture is an area of increasing interest. Probiotics have been defined by the World Health Organization's-Food and Agriculture Organization (FAO, 2001) as "live microorganisms which when administered in adequate amounts confer a health benefit on the host." In the last decade, several gram-negative and gram-positive bacteria have been evaluated *in vitro* or *in vivo* for their potential to inhibit fish-pathogenic organisms and overcome infections in fish and larvae in aquaculture (Irianto and Austin, 2002). On the other hand, many authors used *Pseudomonas* sp. for the decontamination

of wastes and such biodegradation processes proved to result in little or no impact on environment (Cernigilia, 1992; Obayori et al., 2009).

Thus, the goal of this study is to create an integrated system using a marine local bacterial isolate *Pseudomonas aeruginosa* strain *elnaggar1*, which isolated from Sidi Kerir, West Alexandria, to enable the rearing of *M. cephalus* fries in presence of crude oil contaminated seawater. In addition, to evaluate the use of this marine bacterial isolate as a probiotic agent for controlling the rot fin disease which attack the *M. cephalus* when exposed to the crude oils.

MATERIALS AND METHODS

Experiments were carried out at National Institute of Oceanography and Fisheries, Alexandria, Egypt. *M. cephalus* post larvae were captured using fishing nets (1 mm mesh) from the eastern harbor of Alexandria. They were transported and acclimated in two 100 L-glass tanks filled with seawater from the collecting site (33 salinity and 20°C temperature). The tanks were equipped with constant aeration and kept under natural photoperiod (13 L: 11 D). The post larvae remained in the tanks for two weeks, according to (Sampaio et al., 1998).

Before the post larvae were distributed in the experimental aquaria, 50 individuals were randomly selected for the detection of the initial weight and length. Then the post larvae were transferred to ten 30 L glass aquaria (30 × 30 × 50 cm) filled with 27 L seawater at densities 10 mullet/aquarium.

investigational design

The impact of three crude oil concentrations 100, 300 and 500 ppm was determined in these aquaria O&F100, O&F300 and O&F500 respectively. The impact of 10% inoculums of the marine bacterial strain *P. aeruginosa* was estimated in an aquarium (B&F). Moreover, the bacterial treatments of 100, 300 and 500 ppm crude oil concentrations were carried out in these aquaria B&F&O100, B&F&O300 and B&F&O500, respectively, using 10% inoculums of the marine bacterial strain *P. aeruginosa* strain *elnaggar1*.

In addition, the chemical treatments of 300 and 500 ppm crude oil were carried out in the aquaria chem.300 and chem.500, respectively, using a commercial oil disperser. A control was made using untreated seawater without any additives (C) as shown in Figure 1. The experimental period was extended for 45 days; it was divided into 15 days for adaption and acclimatization, 15 days for crude oil treatments and 15 days for washing process.

The feeding regime was applied at 5% body weight per day throughout the experiment; the frequency of feeding was maintained as twice a day for six days a week. The

artificial diet was analyzed for moisture, crude protein, ether extract and ash according to standard AOAC methods (AOAC, 1980) (Table 1).

Bacterial damage

P. aeruginosa was isolated from Sidi Kerir bottom water sample through enrichment process on crude oil. The isolate has been described as a gram-negative, rod-shaped and aerobic bacterium. The partial identification process was carried out according to Bergey's manual, section-4 "gram negative aerobic rods and cocci" (Sneath et al., 1986).

Molecular detection process

This process was carried out at Mubarak City for Scientific Re-search and Technology Applications, Arid Land Institute, Molecular Plant Pathology Department, New Borg El Arab City, 21934, Alexandria, Egypt.

DNA extraction

DNA was extracted from overnight pure culture of this marine isolate using Qiagen DNeasy kit (QIAGEN-Inc., Germany) and Genomic DNA purification kit (Promega). The procedure was identical to that recommended by the manual instructions. The preparations were analyzed on a 0.7% agarose gel and then determined spectrophotometrically (Sambrook et al., 1989).

PCR-amplification and sequencing of 16S rDNA gene.

The amplification of the genomic DNA was carried out by a PCR using *Pseudomonas* 16S rDNA gene primer PA-SS-F' GGGGGATCTTCGGACCTCA, PA-SS-R' TCCTTAGAGTGCCAC CCG, according to Spilker et al. (2004).

Nucleotide sequence accession number

The GenBank accession number for the 16S rDNA sequences generated in this study is GQ505252, the isolate was identified as *P. aeruginosa* strain *elnaggar1*.

Culture circumstances

The culture medium used for isolation and activation of *P. aeruginosa* was modified from that of Guo-liang et al. (2005), where the seawater was used instead of distilled water. The composition of this basal mineral salt medium was as follow (g/l): NaNO₃ 4.0, NaCl 1.0, KCl 1.0, CaCl₂·2H₂O 0.1, KH₂PO₄ 3.0, Na₂HPO₄·12H₂O 3.0, MgSO₄ 0.2, FeSO₄·7H₂O 0.001; 2 ml trace element stock solution composed of (g/L): FeCl₃·6H₂O 0.08, ZnSO₄·7H₂O 0.75, CoCl₂·6H₂O 0.08, CuSO₄·5H₂O 0.075, MnSO₄·H₂O 0.75, H₃BO₃ 0.15, Na₂MoO₄·2H₂O

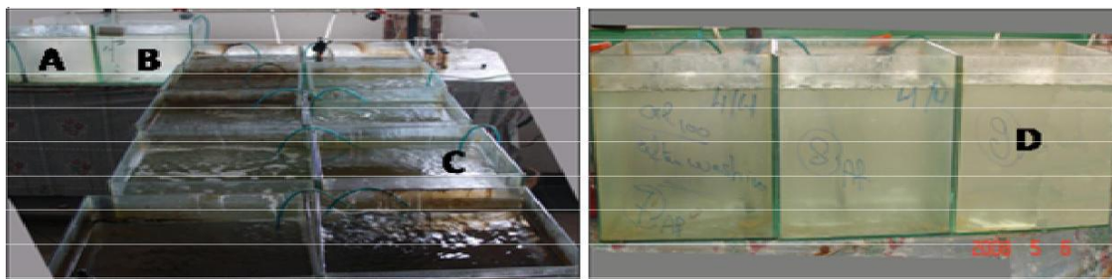


Figure 1. Photographs shows experimental frame work and the used glass aquaria filled with untreated seawater (control) (A); the *P. aeruginosa* in contact with *M. cephalus* fries - blank (B); the oily contaminated seawater under treatments (C) and; the glass aquaria of the washing process (D).The feeding regime.

Table 1. The composition of the artificial diet used for *M. cephalus* feeding regime.

Ingredients	Percentage
Fish meal	25.00
Soybean meal	12.00
Broken corn	17.00
Wheat milling by-product	45.00
Mineral mixture	0.70
Vitamin premix	0.30
Chemical analysis	
Dry matter	87.12
Crude protein	32.1
Either extract	2.63
Crude fiber	2.46
Ash	11.61

0.05. The initial pH was adjusted to 7.7 and the incubation was done for 24 – 48 h at 25°C.

Sampling processes and microbial examination processes in the oily contaminated seawater

Water samples for microbiological analyses were taken regularly from the rearing fish aquaria after zero, 2, 3, 4, 7 and 14 days of the treatment and in addition, after one week of the washing process. 1 ml of these seawater samples was used aseptically to inoculate 9 ml of sterilized culture medium and then incubated for 24 – 48 h at 25°C. Then, the dry weight of the bacterial growth was estimated in mg/100 ml according to Dalgaard et al. (1994).

Examination of skin

The skin of three treated fish per aquarium were swapped and resuspended in 5 ml sterile phosphate-buffered saline (PBS), it composed of; 8.0 g of NaCl, 0.3 g of KCl, 0.73 g of NaH₂PO₄ and 0.2 g of K₂HPO₄ to 1 liter of de-ionized water, pH 7.4. All samples were 10-fold diluted, squeezed by hand for few minutes and then

spread onto nutrient agar plates which composed of (g/l): 3; beef extract, 5; peptone and 20; agar using a glass spreader (5 cm). The total bacterial count, colony forming unit (cfu/ml) was estimated in each sample according to Buller (2004). Sterile gloves, bags, swabs, and glass beakers were used for sampling.

Examination of muscle

The head, internal organs and the tail of three examined fish per aquarium were removed under sterile conditions. Then 1 g of the muscle part was transferred to sterile tubes contain 1 ml PBS. All samples were 10-fold diluted, squeezed by hand for a few minutes and spread onto nutrient agar plates as mentioned above. The total bacterial count, colony forming unit (cfu/ml) was estimated in each sample according to Buller (2004).

Examination of internal organ

The internal organs of each examined fish were removed under sterile condition using sterile forceps and transferred to sterile tubes contain 1 ml PBS. All samples were 10-fold diluted, squeezed by hand for a few minutes

Table 2. The effect of the microbial and chemical crude oil treatments on the growth performance, survival percentage and feed utilization parameters of *M.*

Estimated parameters	Treatments						
	Control	B&F	O&F100	O&F300	O&F500	O&B&F100	O&B&F300
At zero time (Stocking data)							
Av. Initial weight (g) ± SD	0.04 ± 0.21	0.04 ± 0.21	0.04 ± 0.21	0.04 ± 0.21	0.04 ± 0.21	0.04 ± 0.21	0.04 ± 0.21
Av. Initial length (cm) ± SD	1.1 ± 0.32	1.1 ± 0.32	1.1 ± 0.32	1.1 ± 0.32	1.1 ± 0.32	1.1 ± 0.32	1.1 ± 0.32
After 30 days of rearing							
Av. final weight (g)	0.32	0.29	0.1	0.1	0.12	0.3	0.29
Av. final length (cm)	2.6	2.5	1.9	1.8	1.8	2.5	2.3
Gain in weight (g/fish)	0.28	0.25	0.08	0.06	0.06	0.26	0.25
Daily weight gain (mg/fish/day)	18.67	16.67	5.33	4	4	17.33	16.67
Daily length gain (mm/fish/day)	1.5	1.4	0.8	0.8	0.7	1.4	1.2
After 45 days of rearing							
Av. final weight (g) ± SD	2.1 ± 0.04	1.86 ± 0.01	0.39 ± 0.02	0.28 ± 0.07	0.24 ± 0.01	1.75 ± 0.28	2.00 ± 0.66
Av. final length (cm) ± SD	5.35 ± 0.67	4.25 ± 0.33	2.85 ± 0.22	2.5 ± 0.38	2.2 ± 0.11	5.1 ± 0.87	5.1 ± 0.33
Gain in weight (g/fish)	2.06	1.77	0.35	0.24	0.2	1.71	1.96
Daily weight gain (mg/fish/day)	45.78	39.33	7.78	5.33	4.44	38	43.56
Daily length gain (mm/fish/day)	0.94	0.7	0.39	0.31	0.24	0.89	0.89
Instant daily growth (IDG)	3.71	3.59	2.85	2.78	2.76	3.56	3.67
Survival rate (%)	90	80	50	40	30	80	70
Feed utilization data							
Feed conversion ratio (FCR)	2.16	2.23	2.98	3.1	3.76	3.41	3.53
Protein efficiency ratio (PER)	2.96	1.99	1.65	1.41	1.23	1.33	1.06

Table 3. The statistical analysis of the growth parameters of *M. cephalus* along 45 days of the rearing process under microbial and chemical treatments.

Estimated Parameters	Treatment									
	Control	B&F	O&F100	O&F300	O&F500	O&B&F100	O&B&F300	O&B&F500	Chem300	LSD
Daily weight gain (mg/ fish/day)	32.2 ^{a†}	28.05 ^c	6.56 ^t	4.67 ^g	4.21 ^g	27.67 ^c	30.08 ^b	23.24 ^d	8.45 ^e	0.263
Daily length gain (mg/fish/day)	1.27 ^a	1.05 ^b	0.59 ^{de}	0.55 ^{ef}	0.47 ^f	1.2 ^a	1.05 ^b	0.9 ^c	0.69 ^d	0.118

*Mean values in the same row which have the same letters are insignificantly.

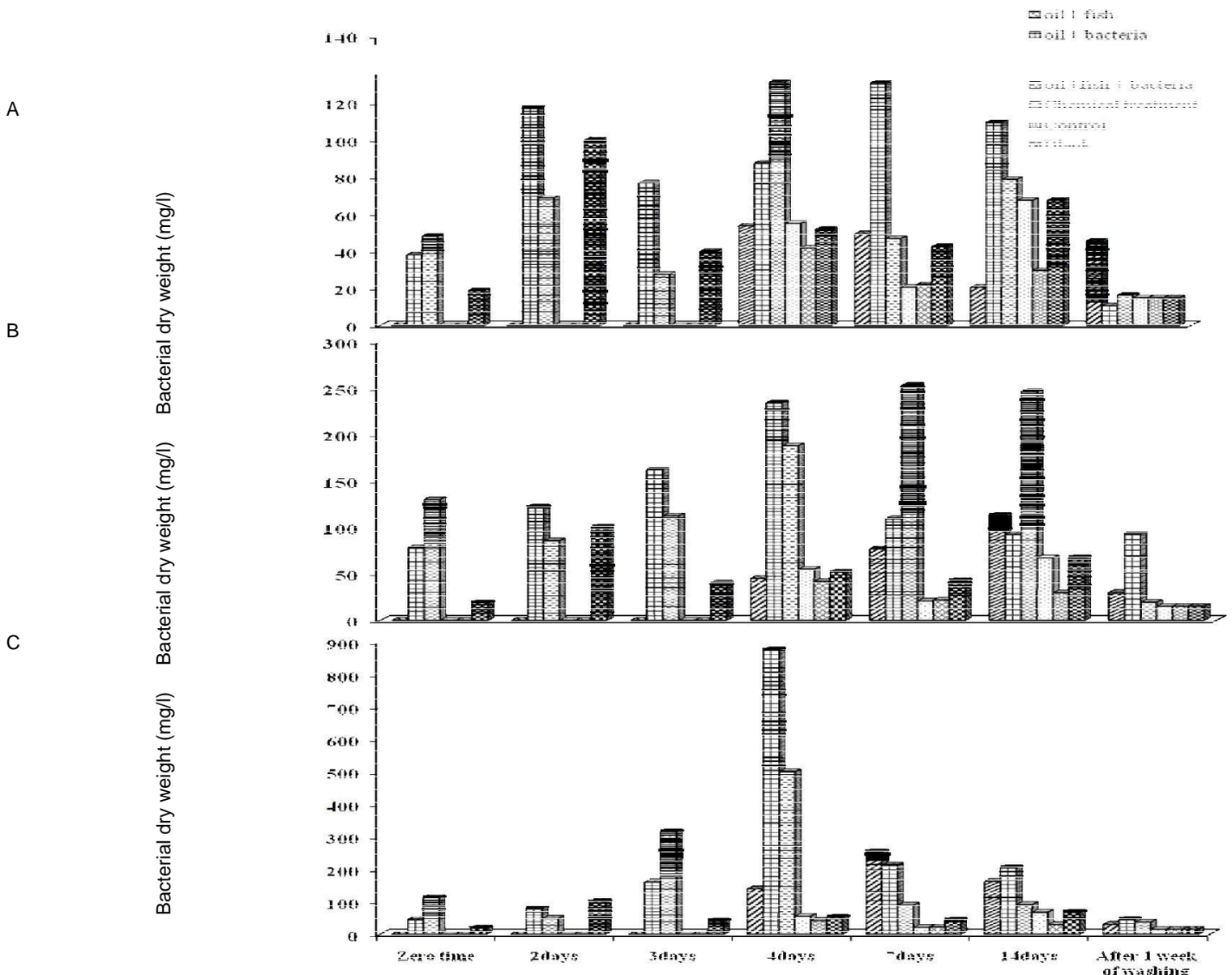


Figure 2. The bacterial dry weight in the surrounding media of the contaminated aquaria (O&F, B&F, O&F&B, Chem., F&B and the control) during 15 days of 100ppm (A), 300 ppm (B) and 500 ppm (C) crude oil treatment and after one week of washing process.

and spread onto nutrient agar plates as mentioned above. The total bacterial count, colony forming unit (cfu/ml) was estimated in each sample according to Buller (2004).

Chemical treatment

The chemical treatment of the used crude oil was carried out using 100 ppm of a commercial OIL DISPERSER. The physical properties of this disperser were estimated as follows: Clear pale yellow liquid, Sp; Gravity at 25°C =

Table 4. The remediation rates of the soluble crude oil fractions by microbial and chemical treatments.

Time (day)	Remediation rate (ppm/day)					Mean*
	Microbial treatment			Chemical treatment		
	O&F&B ₁₀₀	O&F&B ₃₀₀	O&F&B ₅₀₀	Chem ₃₀₀	Chem ₅₀₀	
1	1.78	12.5	7.9	3.2	9.56	6.99 ^d
4	5.3	27	36.3	6.3	20.8	19.14 ^a
7	3.4	3.6	19.7	5.6	4.9	7.44 ^b
14	2.6	0.36	6.2	0.057	0.36	1.92 ^c
Mean*	3.27 ^c	10.86 ^b	17.53 ^a	3.79 ^c	8.91 ^b	

*Mean values in the same column or in the same row which have the same letter are insignificantly different at $P < 0.01$ while the mean values with different letters are significantly different at $P < 0.01$.

Table 5. The microbiological examinations of *M. cephalus* after 45 days of rearing.

Treatments	Bacterial count CFU $\times 10^2$ /100 ml			Mean*
	Muscle	Skin	internal organs	
Control	2	0.8	2	1.6 ^d
B&F	1.3	3.8	3.8	2.97 ^c
O&F&B ₁₀₀	2.2	3.6	2.3	2.7 ^c
O&F&B ₃₀₀	2.5	6	4	4.16 ^b
O&F&B ₅₀₀	1.5	4.7	5	3.73 ^b
Chem ₃₀₀	4.8	10	8.8	7.87 ^a
Mean*	2.38 ^c	4.81 ^a	4.31 ^b	

Mean values in the same column or in the same row which have the same letter are insignificantly different at $P < 0.01$ while the mean values with different letters are significantly different at $P < 0.01$.

1.02 - 1.025 g/cm³; Solubility, soluble in water; Flash point, no; Flammable limits in air, no and; the pH for 1% aqueous solution = 8.25.

Determination of crude oil using gas chromatography extraction

1000 ml of each collected water samples were extracted three times with 100 ml of dichloromethane in a separating funnel. Sample extracts were combined and concentrated to 5 ml using a rotary evaporator under reduced pressure. Finally, samples were concentrated under a gentle stream of pure nitrogen to a volume of 1 ml (UNEP/ IOC/ IAEA, 1992).

Clean-up and fractionation process

Clean-up and fractionation was performed prior to gas chromatograph/flame ionization detector (GC/FID). The extracted volume was passed through the silica column prepared by slurry packing with 10 g of silica, followed by 10 g of alumina and finally 1 g of anhydrous sodium sulphate. The aliphatic fraction (F1) was sequentially eluted from the column using 25 ml of hexane. However, the unsaturated aromatic fraction (F2) was eluted with 60 ml of hexane and dichloromethane (80:20; V/V). Both of

F1 and F2 fractions were concentrated using a gentle stream of pure nitrogen to about 0.2 ml, before being injected into GC/FID (UNEP/ IOC/ IAEA, 1992).

Gas chromatography

All samples were analyzed by a Hewlett Packard 5890 series II GC gas chromatograph equipped with a flame ionization detector (FID). The instrument was operated in split less mode (3 μ l split less injection) with the injection port maintained at 290^o C and the detector maintained at 300^oC. Samples were analyzed on a fused silica capillary column HP-1; 100% dimethyl polysiloxane (30 m length, 0.32 mm i.d, 0.17 μ m film thickness). The temperature was programmed from 60 - 290^oC, changing at a rate of 3^oC /min and maintained at 290^oC for 25 min. The carrier gas was nitrogen flowing at 1.2 ml/ min (UNEP/ IOC/ IAEA, 1992).

Quantification and mixture preparation

Aliphatic standard mixture (100 μ g/ml) brought from MERCK was used for F1 analysis, this standard mixture containing C11, C12, C13, C15, C17, pristane, C18, phytane, C19, C20, C21, C22, C23, C24 and C30. Chlorofluorobenzene (CFB); 20 μ g/ml was used as

Table 5. The microbiological examinations of *M. cephalus* after 45 days of rearing.

Treatments	Bacterial count CFU × 10 ² /100 ml			
	Muscle	Skin	internal organs	Mean*
Control	2	0.8	2	1.6 ^d
B&F	1.3	3.8	3.8	2.97 ^c
O&F&B100	2.2	3.6	2.3	2.7 ^c
O&F&B300	2.5	6	4	4.16 ^b
O&F&B500	1.5	4.7	5	3.73 ^b
Chem300	4.8	10	8.8	7.87 ^a
Mean*	2.38 ^c	4.81 ^a	4.31 ^b	

Mean values in the same column or in the same row which have the same letter are insignificantly different at $P < 0.01$ while the mean values with different letters are significantly different at $P < 0.01$.

internal standard for aliphatic fraction. In addition, a stock solution containing the following PAHs was used for quantification of hydrocarbons: naphthalene, acenaphthylene, acenaphthene, fluorene, phenanthrene, anthracene, fluoranthene, benzo(a) anthracene, chrysene, benzo(b)fluoranthene, benzo(k)fluoranthene, pyrene, benzo(a)pyrene, dibenzo(a,h)anthracene, benzo(ghi) perylene and indeno(1,2,3-cd) pyrene by dilution to create a series of calibration standards of PAHs at 0.1, 0.25, 0.5, 0.75, 1.0, 2.0, 5.0 and 10 µg/ml. The detection limit was 0.01 µg/ml for each PAH. For analytical reliability and recovery efficiency of the results, 6 analyses were conducted on PAH reference materials, HS-5 and 2974 (provided by EIMP-IAEA). The laboratory results showed recovery efficiency ranging from 92 - 111% with coefficient of variation (cv) 8 - 14% for all studied pollutants (16 PAHs fractions).

All solvents were pesticide grade purchased from Merck and appropriate blanks (1000 fold concentrates) were analyzed.

Statistics

The statistical analyses of the data were carried out in triplicates using ANOVA test and the least significant difference L.S.D. according to Steel and Torrie (1980).

RESULTS AND DISCUSSION

The isolation of the used bacterial strain was carried out from a bottom water sample collected from Sidi Kerir, West Alexandria, where many industrial oil companies pour their wastes. The selection process was based on its high ability to degrade crude oil through enriched water samples with oil. The identification process was carried out at Mubarak City for Scientific Research and Technology Applications using the molecular technique. The results indicated the isolation of a new bacterial strain identified as *P. aeruginosa strain elnaggar1* with an

associated number of GQ505252 in the data base of the GenBank library.

However, the growth performance and the survival percentage of *M. cephalus* were estimated in the examined oily contaminated seawater after 30 and 45 days of the rearing process (Table 2). The obtained data showed in general that the bacterial treatment led to slight impacts on the growth performance of the *M. cephalus* compared to the untreated fish (control), while the chemical treatment showed serious impacts on both the growth performance and the survival percentage of *M. cephalus* especially on testing the 500 ppm oil degradation process (Chem₅₀₀) compared to the control.

Moreover, it was observed that after 30 days of the rearing period the addition of 100, 300 and 500 ppm crude oil in the tested aquaria O&F₁₀₀, O&F₃₀₀ and O&F₅₀₀, respectively, led to severe effects on the growth rates of *M. cephalus* fries. The daily weight gain was reduced by 71.5, 78.6 and 78.6% compared to the untreated fish (control), respectively and the daily length gain was reduced by 46.7, 46.7 and 53.3%, respectively. While, the chemical treatment of 300 ppm oil (Chem₃₀₀) led to a reduction of 78.6 and 46.7% in the daily weight gain and the daily length gain, respectively, compared to the control. Moreover, the chemical treatment of 500 ppm crude oil in the aquarium Chem₅₀₀ led to a complete death of the tested *M. cephalus* fries within two days of the treatment.

On the other hand, the *P. aeruginosa strain elnaggar1* in contact with these fish fries (F&B) led to 10.7 and 6.7%, reduction percentage in the daily weight gain and the daily length gain respectively, compared to the control. In addition, the microbial treatment which carried out in the aquaria O&F&B₁₀₀, O&F&B₃₀₀ and O&F&B₅₀₀, it led to a reduction of 7.1, 10.7 and 39.3% in the daily weight gain, respectively and a reduction of 6.7, 20 and 33.3% in the daily length gain compared to the control.

However, after 45 days of the rearing process these estimated parameters showed to be highly affected by both the presence of the crude oil and the use of the oil

dispenser in the treatment process. The results showed that the presence of the crude oil in concentrations ranged from 100 to 500 ppm led to a reduction in the daily weight gain and the daily length gain of the *M. cephalus* fries, it ranged from 4.4 mg/fish/day to 7.8 mg/fish/day and from 0.24 mm/fish/day to 0.39 mm/fish/day, respectively compared to the untreated fish (45.7 mg/fish/day and 0.94 mm/fish/day) While the use of the oil disperser led to a daily weight gain of 12.89 mg/fish/day and a daily length gain of 0.56 mm/fish/day. Moreover, the data presented in Tables 2 and 3 showed also the impact of the bacterial treatment (*P. aeruginosa*) on the growth performance and the survival rate of *M. cephalus* after 45 days of rearing, which including two weeks for adaptation, two weeks for treatment and two weeks for washing. It was observed that the microbial treatment of 100, 300 and 500 ppm crude oil had the slight impacts on the rearing process of *M. cephalus* fries compared to the untreated fish. The daily weight gain and the daily length gain were fish/day and from 0.8 to 0.89 mm/fish/day, ranged from 35.11 to 43.56 mg/fish/day and the most effective microbial O&F&B₁₀₀ using the marine *P. aeruginosa*, the degradation percent were 36, 29 and 21.2%, respectively, compared to the original crude oil added to these aquaria.

On the other hand, the chemical treatment of 300 and 500 ppm crude oil using a commercial disperser led to degradation percentages of 8.4 and 16.6%, respectively, compared to the original crude oil added to these aquaria. This great effectiveness explained by several authors they mentioned this bacterial species able to produce the bio-surfactant of rhamnolipid in the presence of the crude oil which facilitate the biodegradation process (Balba et al., 2002; Tang et al., 2007; Urum et al., 2003). Moreover, the production of this biosurfactant in the surrounding medium of the bacterial cells showed to extract the lipopolysaccharides from the cellular envelope which increase the cell hydrophobicity and it subsequently stimulates the uptake process of the hydrocarbon droplets via the direct contact with these bacterial cells (Guo-liang et al., 2005).

Moreover similar results were obtained by Norman et al. (2002), they studied the effect of extracting the lipopolysaccharides (LPS) from the cell surfaces of *P. aeruginosa* strains on the crude oil degradation (n-alkane). It was found that the cell hydrophobicity increased and the degradation of n-alkane increased. On the other hand, Guo-liang et al. (2005) observed that neither cell growth nor rhamnolipid production were obtained when *P. aeruginosa* grown on crude oil as a sole carbon source.

However, the impact of the microbial and the chemical treatments on the total bacterial count of the muscle, skin and the internal organs of the tested *M. cephalus* was presented in Table 5. It was observed that the most affected fish part was the skin followed by the internal organs. The lowest bacterial count was estimated in the

from 0.8 to 0.89 mm/fish/day, respectively. While the presence of *P. aeruginosa* in the aquaria F&B, O&F&B₁₀₀, O&F&B₃₀₀, and O&F&B₅₀₀ led to a survival percentage ranged from 70 to 80% compared to the control (90%). On the other hand, the survival percentage was reduced to 50, 40 and 30% on the addition of the crude oil in the tested aquaria O&F₁₀₀, O&F₃₀₀ and O&F₅₀₀, respectively. However, the data presented in Figure 2 showed that with increasing of the oil concentration from 100 to 3000 ppm in the aquaria the mean bacterial count was mg/l to 900 mg/l mainly after the presence of in contact with bacterial growth the first three days washing process of this integrated on the surrounding media since after a week of washing, it led to reduce the bacterial content to a level slightly similar to that of the control. Moreover, the statistical analysis presented in Table 4 showed that with increasing oil concentration the efficiency of the microbial treatment increase. The maximum mean degradation rate was observed after four days followed by O&F&B₅₀₀ and day of the experiment regard less to the used treatment was O&F&B treatment (1914 ppm/day) fish muscle.

The chemical treatment led to increase the fish susceptibility towards the bacterial accumulation compared to the control. It showed a high significant difference at $P < 0.01$ in accumulating the bacterial counts in these examined fish the mean bacterial count was 7.87×10^2 cfu/100 ml compared to that of the control (1.6×10^2 cfu/100 ml). However the bacterial treatments showed lower bacterial accumulation in these examined fish parts especially in case of O&F&B₁₀₀ and the O&B the estimated counts were 2.7 and 2.97×10^2 cfu/100 ml, respectively.

Moreover, the impacts of these all treatments on the morphological symptoms of the tested fish were summarized in Figure (3). The data showed a significant susceptibility of *M. cephalus* to the rot fin disease especially the caudal fin it almost degraded due to the presence of either high crude oil concentration or even the presence of the oil disperser in the surrounding media. Similarly, Douglas and Yarbrough (2006) showed that after Six to eight days of the oil exposure all the examined mullet which taken from the treated ponds had varying degrees of fin rot on one or more of their fins.

Where, the fin erosion involved primarily the caudal, pectoral and pelvic fins, the caudal fin was the most severely damaged. In addition, they observed the degree of damage was varied from a slight discoloration with no visible fraying to complete erosion of all of the fin elements. They isolated a gram negative rod tentatively identified as *Vibrio* sp. and they considered it as the primary pathogen responsible for the fin erosion. Mean while, it can be noticed this microbial treatment may kept the survival percent and the growth performance of the tested fish as partially similar as that of the control and this can be explained by Spanggaard et al. (2001), they showed the fish pathogenic bacterium *V. anguillarum* growth was most prominently inhibited by *Pseudomonas*

spp. using the well diffusion assay and the all diseaseprotecting strains were pseudomonads. Moreover, they mentioned pseudomonads are typically siderophore producers and the addition of live bacterial cultures to fish-rearing water may thus improve survival of the fish.

However, these results were also in agreement with that of Gram et al. (1999), they observed the combined probiotic treatment using *Pseudomonas fluorescens* resulted in a 46% reduction of calculated accumulated mortality; it was 25% after 7 days in the probiotic-treated fish, whereas mortality was 47% in fish not treated with the probiont.

CONCLUSION

The results indicated the use of the chemical treatment for escaped crude oil in marine environment it may led to increase fish susceptibility towards bacterial accumulation especially on the skin part compared to the untreated fish. Moreover, the bacterial treatment using the marine *P. aeruginosa* led to the degradation of the oil with low impact on the fish in the surrounding medium which more or less seemed to be similar to the untreated fish. So, it could concluded it may be useful to use such integrated microbial system for crude oil bioremediation in marine contaminated areas and in the same time it could act as a potent probiotic agent for the reared *M. cephalus* which enable them to control the fish diseases like rot fins and tails.

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Moreover, although the exposure of the *M. cephalus* to the water-soluble fractions (WSF) of crude oil increase the susceptibility to bacterial infection it was noticed that the muscle of the examined fish was the lowest infected part. This can be explained by the work done by Thomas (2006); he studied the effect of the exposure of the mullet to the water-soluble fractions (WSF) of crude oil and two fuel oils. It was obtained that these tested oils altered the ascorbic acid (AsA) content of several striped mullet, *M. cephalus* tissue. In addition, the exposure to sub-lethal concentrations of the WSFs caused a depletion of As A reserves in brain, gill, kidney and liver tissues, but not in muscles.

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