STUDIES ON ULCERATIVE HEAD SYNDROME AMONG OREOCHROMIS NILOTICUS FISH

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ABSTRACT

Vibrio splendidus, Aeromonas hydrophila, Trichodina centrostigeata and T. magna were isolated from Oreochromis niloticus which suffered from ulcer in head region. Various characteristics of V. splendidus have been studied. In vitro antibiotic sensitivity, pattern of isolated V. splendidus was tested against different chemotherapeutic agents. I/P inoculation of V. splendidus broth culture also mixed broth cultures of V. splendidus and A. hydrophila resulted in 100% mortality, while the challenge by immersion gave no mortalities.

INTRODUCTION

Vibrionaceae includes two genera, Vibrio Aeromonas which are distributed world wide and together constitute the most important species of fish bacterial pathogens (Inglis et al., 1993). Vibriosis is one of the major secondary bacterial diseases (Ruangpan and Kitao, 1991) and considered as enzootic disease of fish all over the world affecting marine, brackish and occasionally freshwater fishes (Schaperclaus et al., 1992) such disease may be represented in the form of focal haemorrhagic ulcer on the mouth or skin surface (Inglis et al., 1993). V. splendidus has recently been described as new bacteria causing vibriosis in juvenile and adult trubot (Lupiani et al.,1989). Pathogenicity of V. splendidus was confirmed by experimental infection (Toranzo and Barja, 1990). However, most if not all the virulence mechanism of these fish pathogens are still unknown (Woo and Brno, 1999). Genus Aeromonas are capable of causing disease in fish. Aeromonas hydrophila have been well documented as fish pathogen, and usually as opportunistic or secondary invaders rather than primary pathogens (Inglis et al., 1993). Ectoparasitic ciliates include different species that are most common parasites of fishes especially Trichodina species. Trichodinia Spp. is a frequent problem (Lom and Dykova, 1992) and never occurs in large amount on a healthy fish but when the host is weakened by an any stress Trichodiniasis reproduces in massive proportion and start to exert



their pathogenicity (Lam, 1970; VanAs and Bassan, 1988 and Duran et al., 1991). Trichodiniasis have been implicated in high moralities of fishes causing severe economic losses in various parts of the world (VanAs and Bassan, 1987) and were also responsible for growth inhibition and weight loss in cultured fishes (Duran et al., 1991).

The present study was designed for studying the relationship between these fish pathogens for inducing the ulcerative head phenomena and showing the pathogenicity of *V. splendidus* for the first time in *O. niloticus*.

MATERIAL AND METHODS

1. Fish:

- (A) Five fish out of 40 live *O. niloticus* suffered from ulcer formation in head region were collected randomly from River Nile at Moneab Giza Governorate during August, 2000 with an average body weight 100 ± 5 g. Fish were examined bacteriologically and parasitologically in Fish Diseases Department, Animal Health Research Institute.
- (B) A total number of 72 apparently normal *O. niloticus* were collected alive from Nawa farm at Kalubia Governorate, they were kept in glass aquaria supplied with dechlorinated tap water (Innes, 1966). Bacteriological mycotic and parasitological examinations were carried out on random 5 fish to ensure that experimental fish were free from the risk of natural infection.

2. Clinical examination:

The naturally infected fish were examined for any abnormalities according to Plumb and Bowser, (1982) and Lom and Dykova, (1992).

3. Bacteriological examination:

The natural infected fish were subjected to bacteriological examination according to the methods described by **Inglis** *et al.*, (1993) where aseptically samples from ulcers of such fish were streaked into trypticase soy broth, incubated at 25 C^o for 24 hr then loopfulls, from the broth were streaked onto trypticase soy agar (Difco) blood agar as well as some selective media such as Aeromonas base medium (Oxoid) nutrient agar with 2%,4% and 6.5%Na Cl.

4. Biochemical reaction:

According to Baumann and Schubert, (1965) and Krieg and Holt, (1984).

5. The in vitro antibiotic sensitivity:

The sensitivity of the isolated V. splendidus against antibiotic was carried out in Mueller and Hinton media (Difco) using disc diffusion method



according to Treagon and Pulliam, (1982). The interpretation of the result was undertaken according to Acar and Goldstein, (1986).

6. Parasitological examination:

Smears were taken from ulcers of the naturally infected O. niloticus fish they were air dried, then impregnated with 2% aqueous solutions of silver nitrate for 8 minutes followed by rinsing in distilled water, the slide was then placed in white clean dish covered with distilled water and exposed to UV (diffused day light) for about 2 hours, then slides were dried and examined microscopically Lom and Dykova, (1992). Terminology and method of measurement of the components of the adhesive disc followed the uniform specific characteristic system proposed by Lom, (1958); Wellborn, (1967) and Arthur and Lam, (1984). Detailed description of the denticles were present in accordance with the method proposed by VanAs and Bassan, (1992). Also body diameter was measured.

7. Experimental infection designs:

Experiment (1): 24 *O.niloticus* fish were divided into 4 groups (6 for each), 1^{st} group was injected I/P with 0.3ml of 24 hr old *V. splendidus* broth culture contained $(1x10^4)$ cell/ml according to Myhr *et al.*, (1991) 2^{nd} group was injected S/C with 0.5ml of the same culture while the 3^{rd} group immersed in $(1x10^4)$ cell/ml of *V. splendidus* broth culture for 1 hr while the 4^{th} group was kept as a control.

Experiment (2): 24 *O. niloticus* were divided into 4 groups (6 of each) 1^{st} group was injected I/P with 0.3 ml of 24 hr old *A. hydrophila* broth culture contained (1×10^4) cell/ml, while 2^{nd} group injected S/C with 0.5 ml of the same culture and 3^{rd} group was immersed in broth culture of *A. hydrophila* (1×10^4) cell/ml for one hour, 4^{th} group was kept as control.

Experiment (3): 24 *O. niloticus* were divided into 4 equal groups, 1^{SI} group was injected I/P with 0.3 ml of mixed broth culture prepared from equal amount (5ml) of *V. splendidus* broth culture and *A.hydrophila* broth culture, each contained $(1x10^4)$ cell/ml, while the 2^{nd} group was injected S/C by 0.5 ml of the same mixed culture and 3^{rd} were immersed in the same mixed culture for 1hr, 4^{th} group kept as a control.

Experimentally infected fish were observed for any clinical abnormalities, P/M lesions and mortality rate were recorded during 10 days (the period of experiment).

RESULTS

Result of clinical examination showed that fish suffered from whitish gray superficial ulcer on head region. (Fig.1). The bacteriological examination revealed isolation of 2 types of bacterial isolates, the first one gave identical biochemical reaction to these useful for preliminary identification of V.

splendidus (Table 1) It is Gram negative and motile. The colonies were circular, regular medium sized, shiny and translucent The organism is coccoid rods or short bacilli, sensitive to 0/129 (Fig.2), while the second isolate was identified biochemically (Table 2) as *A. hydrophila* whish gave green circular colony with dark center on *Aeromonas* base media.

In vitro Antibiogram of the used *V. splendidus* isolate to a variety of antibiotics is to be seen in (Table 3) from this table, it is clear to see that the *V. splendidus* isolate is highly sensitive to oxytetracycline, tetracyclin, garamycin, naldixic acid and nitrofurantoin, sensitive to chloramphenicol and less sensitive to Colistin sulphate, while it resistant to lincomycin, amoxycillin and ampicillin.

The parasitological examination revealed isolation of two species of Trichodinids, *Trichodina centrestigeata*, Bassan et al., (1983) and *Trichdinia magna*, VanAS and Bassan, (1989).

T. centrestigeata Bassan et al., (1983) is a medium sized with a very high body, surrounded by a finely striated border membrane. The center of the adhesive disc has a characteristic center ridges reaching from 12 to 15 in number of denticles ranged from 27-30. Blade is angular, truncate and slanting backward. The junction of the blade with the central part is narrow. In the same species, tips of the blades are tanget to the border of the adhesive disc. Rays (thorns) are straight or sometimes slightly curved posteriorly, thick at the base and dapering gradually to sharp rounded point central part conical shaped (Fig. 3, Table 4). T. magna VanAS and Bassan, (1989), is the largest Trichodinid with disc shaped or saucer-like body surrounded by finely striated border membrane. The center of the adhesive disc is dark and limely granulated. Massive denticles and provided with strongly falcated blades and wedge-like rays. The blade is broad with a round apex. The central part of the denticle is broad at the base and tapers to a rounded point in close association with the preceding denticle rays are long and slightly curved anteriorly They taper from thick bases towards sharp points (Fig. 4, Table 4) Some stages were obtained during division of T. magna by binary fission (Fig. 5, Table 4).

I/P inoculation of *V. splendidus* in *O. niloticus* revealed 100% mortality (Table 5) with 48 hr post infection where the fish was suffered from loss of scales, haemorrhage in all body surface (Fig. 6) inflamed vent with small amount of yellowish exudate in body cavity.

In case of S/C inoculation, there was erected scales at lateral sides of fish, white discoloration with large amount of yellowish red exudate in body cavity, haemorrhage in all internal organs with inflammed vent (Fig. 7), with severe ulceration at the site of inoculation which surrounded with haemorrhagic area, while there was no mortality in fish group challenge by immersion, fish was suffered from slight haemorrhage at the body surface which disappeared 96 hr after immersion .I/P inoculation with mixed culture of A. hydrophila and V. splendidus showed clear respiratory manifestations

that fish try to gasp atmospheric air, white discoloration in body surface, haemorrhage in head region at the base of pectoral fins with protruded vent (Fig.8). The mortality rate was 100%.

While in S/C inoculation of mixed culture, there was haemorrhage at the pectoral and pelvic fin, ulcer formation with loss of musculature at the site of inoculation which surrounded by haemorrhagic area.

During P/M examination there was severe haemorrhage in all organs including gonads moderate amount of yellowish brown exudate in body cavity, the mortality rate was 66.6% (Table 6).

There was no change in fish group which infected by immersion method.

In case of fish group which injected I/P with *A. hydrophila*, the mortality rate was 100% with signs of general septicaemia in liver, spleen, and mesentery), while in S/C inoculation the mortality rate was 50% after 66hr post infection (Table 7) with haemorrhage of internal organs. There were no clinical signs in fish group which challenged by immersion.

DISCUSSION

In most cases where ectoparasite infestation occurs on fish, there is a possibility that secondary infection by bacteria and fungi may take place (VanAS and Bassan, 1988), in the present, study we isolated 2 species of *Trichodina* and 2 types of bacteria from naturally infected *O. niloticus* were isolated.

Isolation of *V. splendidus* from ulcer of naturally infected *O. niloticus* for the first time in Egypt agree with **Inglis** et al., (1993) who recorded that more than one genus of Vibrio have been isolated in outbreaks of Vibriosis in fish and shellfish (*V. anguillarium*, *V. vulnificus*, *V. splendidus* and *V. pelagius*) also with **Lupiani** et al., (1989) who isolated *V. splendidus* from cultured turbot.

Examination of Gram stained smear from suspected colonies of *V. splendidus* isolates, showed it was Gram -ve, short bacilli (Myhr et al., 1991and Inglis et al., 1993).

Biochemical reactions and other growth characters of *V. splendidus* were summarized in (Table 1), the result basically agrees with that reported by (Lupiani *et al.*, 1989) except gelatin hydrolysis and VP reaction which agree with Myhr *et al.*, (1991) who reported that VP was 4/36 and Baumann and Schubert, (1965) who reported that *V. splendidus* hydrolysed gelatin.

The biochemical reactions of A. hydrophila were shown in (Table 2) agree with Krieg and Holt, (1984).

Sensitivity of *V. splendidus* to some chemotherapeutic agents were shown in (Table 3) and this may agree with **Myhr** et al., (1991) who reported that *V. splendidus* is senstive to oxytetracycline and resistant to oxolinic acid



while of Inglis et al., (1993), reported that V. splendidus is resistant to ampicillin, oxytetracycline and streptomycine.

In our present work *T. centrostigeata* was isolated from naturally infected *O. niloticus* as reported by Nativided et al., (1986) who isolated *T. centrostigeata* from *O. niloticus* in philippines, *T. centrostigeata* was originally described from cichlid in South Africa (Bassan et al., 1983) it has been recorded from Eastern caprivi (VanAs and Bassan, 1992), wild and cultured freshwater fishes in Taiwan (Bassan and VanA, s 1994). The body dimensions of the present specimen fall within the range of those previously reported from Taiwan as well as the type population from South Africa. From the previous data it can be confirmed suggestions that *T. centostigeata* is a parasite endimic to Africa which at some stage was translocated to Taiwan via the fish introduction from the African contiment.

T. magna seems to be widely distributed in Africa. This parasite was first recorded in South Africa by VanAs and Bassan, (1989), then by the same author in the Zambezi River system (1992) and Ali, (1992) our specimens recovered from O. niloticus nearly agree well with the descriptions of VanAs and Bassan, (1989) VanAs and Bassan, (1992) and Ali, (1992).

During the division by binary fission, the first sign is a thickening of radial pins of the mother individuals (Fig. 5) Lom and Dykova, (1992). The occurrence of stressors such as parasitism enabled Vibrio to produce infection in form of haemorrhagic ulcers on the mouth or skin surface (Inglis et al., 1993).

For the best of our knowledge's the pathogenicity of *V. splendidus* in *O. niloticus* has not been studied yet the experimental infection of *O. niloticus* with *V. splendidus* by S/C route gave 66.6 % mortality with signs of anorexia, beginning of ulcer formation in head region, severe ulceration at the site of inoculation Fig. (6), these this may nearly agree with **Inglis** *et al.*, (1993) who reported that *V. anguillarum* infection may be subdermal also **Abd El G iber** *et al.*, (1997) who studied the pathogenicity of some *Vibrio Spp in O. niloticus as V. anguillarum*, *V. ordalli*, *V. damselea and V. vulnificus*.

The P/M examination showed that intestine was swollen filled with mucoid liquid, large amount of reddish liquid in the peritoneal cavity with inflammed vent (Luipiane et al., 1989).

Also mortality rate was 66.6% in fish group which injected S/C by mixed culture of *A. hydrophila* and *V. splendidus*).

In case of intraperitoneal inoculation of *V. splendidus* or mixed culture of *A. hydrophila* and *V. splendidus* the mortality rate was 100% within 48hr post infection (Fig.8) there were no changes in the internal organs with absence of reddish exudate.

The virulence mechanism of $V_{\cdot\cdot}$ splendidus is still unknown (Woo and Bruno, 1999) In case of fish group which infected by immersion in $V_{\cdot\cdot}$ splendidus broth culture, the mortality rate was zero, no gross lesion except

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slight haemorrhage on body surface which disappeared after 96hr post immersion.

This result may indicate that *V. splendidus* need predisposing factor as reported by **Inglis** *et al.*, (1993) who reported that the occurrence of stresses such as ectoparasitism enabled Vibrio to produce infection in the form of haemorrhagic ulcer on the mouth or skin surface.

In such cases the mortality of fish may occur due to secondary infection (Van As and Bassan, 1988).

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Table (1): Biochemical reactions of V. splendidus

Test	Result	Test	Result	Test	Result
Gram reaction	- ve	Indole	- ve	Lactose	+ ve
Oxidase	+ ve	MR	+ ve	Xylose	+ ve
Catalase	+ve	VP	- ve	Arabinose	- ve
Motility	+ ve	Phenylalanine	- ve	Mannitol	- ve
H2S on TSI	-ve	Gelatin	+ ve	Growth in NaCl	
Citrate	+ ve	Anaerobic growth	- ve	at 2% at 4%	+ve + ve
Urea	- ve	Sugar fermer	ntation	at 6.5%	- ve
Hydrolysis of Tween-80	+ ve	Glucose	+ ve	Sensitivity to	· + ve
Nitrate	+ ve	Sorbitol	+ ve	0/129	, VC

Table (2): Biochemical reactions of A. hydrophila.

· Test	Result	Test	Result	Test	Result
Gram reaction	- ve	Indole	+ ve	Glucose	+ ve
Oxidase	+ ve	Urea	- ve	Sorbitol	- ve
Catalase	+ ve	VP	+ ve	Mannitol	+ ve
Motility	+ ve	Gelatin	+ ve	Sucrose	+ ve
Growth at 37C°	+ ve	H2S production	- ve	Arabinose	+ ve
Simmon's Citrate	+ ve		L	l	

Table (3): Invitro antibiotic sensitivity listing for V. splendidus isolate

	Justing 101 7. spicitulans isolate
Antibiotic disc	Sensitivity
Oxytetracycline OT (30µg)	+++
Tetracyclin TE (30μg)	+++
Garamycin GM (30µg)	+++
Nitrofurantoin F(300μg)	+++
Chloramephenicoll C(30µg)	++
Naldxic acid NA (30µg)	+++
Colistin Sulphate CT (10µg)	+
Ampicillin AML(10µg)	-
Lincomycin. MY (2µg)	-
TY* 1 1	

Highly sensitive

Intermediate sensitive ++

Less sensitive

+

Resistance

Table (4): Morphological data (in μm) of *T. centrostigeata* and *T. magna* from ulcer of naturally infected *O. niloticus* fish.

	T. centrostigeata Bassan et al.,1983(n=25)	T. magna VanAS and Bassan 1989(n=25)
Diameter of Body Adhesive disc Denticular ring Number of Denticles Radialpins/denticle	45.7 (44-48) 37.3 (36-40) 21.2 (17-36) 28(27-30) 6 - 8	61.2 (50-68) 52.6 (45-59) 32.2 (27-36) 24 (24-29) 9 (8-11)
Dimension of a denticle Blade Central part Ray Length Span Width of the border membrane Central of adhesive disc	6 (4-8) 2 (1-3) 4.2 (4-6) 4.6 (4-6) 13.7 (12-15)	7.2 (6-8) 2.4 (2-3) 10.5 (9-12) 9.8 (9-11) 19.8 (19-21)
Adoral spiral	With central ridges 400°	Dark and finely granulated 405°

Table (5): Virulence of V. splendidus in O. niloticus.

Group	No. of fish	Route of inoculation	Bacterial culture	Dose cell/ml	Days post infection 0 24hr 48hr	Mortality %
1	6	I/P	V.splendidus	0.3mI (1×10^4)	6/6	100
2	6	S/C	V.splendidus	0.5ml (1×10^4)	4/6	66.6
3	6	Immersion	V.splendidus	1x10 ⁴ cel	0/6	0
4	6	I/P	Saline	0.3	0/6	0

Table (6): Mortality rate in O. niloticus experimentally infected with mixed culture (V. splendidus and A. hydrophila)

Group	No. of fish	Route of inoculation	Type of culture	Dose cell/ml	Days post infection 0 24hr 48hr	Mortality %
1	6	I/P	Mixed culture	0.3ml (1x10 ⁴)	6/6	100
2	6	S/C	of A. hydrophila and V.	0.5ml (1×10^4)	2/6 4/6	66.6
3	6	Immersion	splendidus	1x10 ⁴ cel	0/6	0
4	6	1/P	Saline	0.3	0/6	0

Table (7): Virulence of A. hydrophila in O. niloticus.

Group	No of fish	Route of inoculation	Type of culture	Dose cell/ml	Days post infection 0 24- 48- 96hr	Mortality %
1	19 6	I/P	a	0.3ml (1×10^4)	2/6 –6/6	100
2	6	S/C	A. hydrophila	0.5ml (1×10^4)	1/6 2/6 3/6	50
3	6	Immersion	V	1x10 ⁴ cel 1 for 1 hr	0/6	0
4	6	I/P	Saline	0.3	0/6	0



Fig (1): O. niloticus suffered from ulcer formation on head region (Naturally infected)

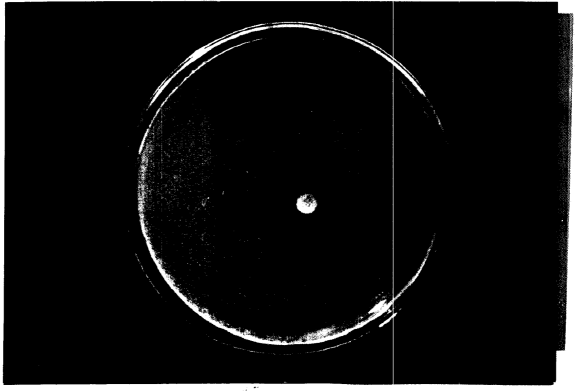


Fig (2): Sensitivity of *V. splendidus* to vibriostate 0/129.

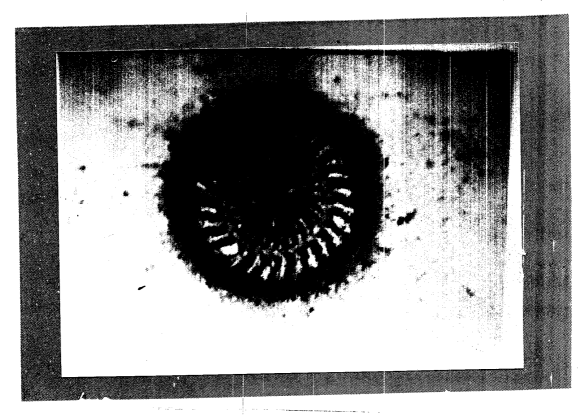


Fig.(3): T. centrestigeata



Fig.(4): Mature old specimen of *T. magna*

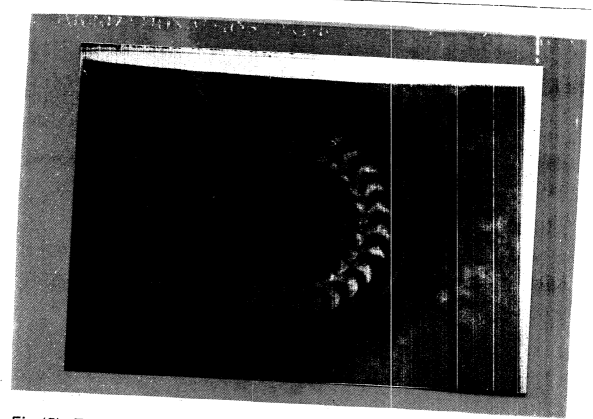


Fig.(5): *T. magna* prior to binary fission as appearing from radial pins arrows

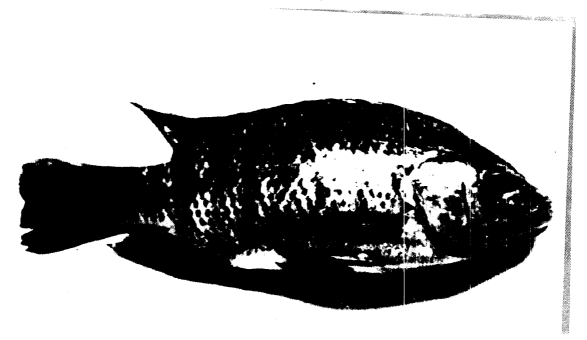


Fig. (6): O. niloticus injected I/P with V. splendidus after48 hr of infection. While in S/C inoculation of mixed culture, there was haemorrhage at the pectoral and pelvic fin, ulcer formation with loss of musculature at the site of inoculation which surrounded by Haemorrhagic area.

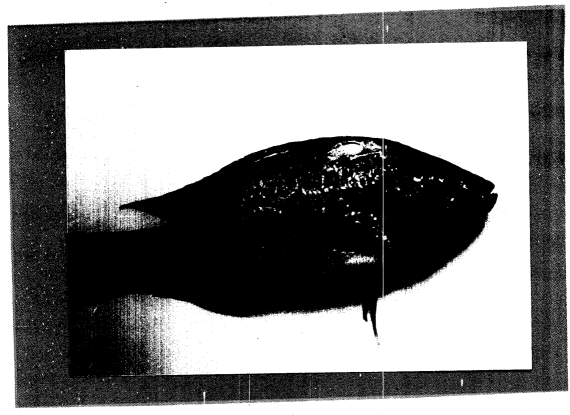


Fig.(7): O.niloticus infected S/C with V. splendidus showed ulcer formation and inflamed vent

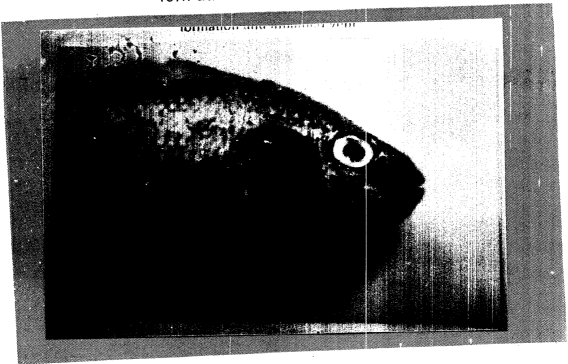


Fig.(8): O. niloticus experimentally infected I/P by mixed culture of A. hydrophila and V. splendidus.

مركز البحوث الزراعية معهد بحوث صحه الحيوان

إستمارة توفيق أوضاع للابحاث المقدمة للتقيم للحصول على درجة رئيس بحوث / أمراض أسماك البحث رقم (١)

التوقيع	نسبة	نوع البحث	عنوان البحث	القائمون بالبحث	۾
	المشاركة				
N N allus	0/00 .	منفردفي	دراسة عن ظاهرة تقرح	د.عبد المحسن حسن	١
		التخصص	الرأس في أسماك البلطي	محمد	
	%0.	وغير	الراس في المهاك البلطي	د. منی مصطفی حسین	۲
otheric		مستخلص من		د. سی سسی حسین	
		رسالة			

١١ تحديد الهدف من البحث

٢ وضع خطة البحث العام

ستجميع العينات وتجهيزها

٤ الجزء العملى من البحث

متدوين النتائج

: قام الباحث بتحديد الهدف من البحث

: قام الباحث بالأشتراك في وضع خطة البحث

: شارك الباحث في جمع العينات وتجهيزها

: قام الباحث بالجزء العملي الخاص بتخصيصة

: شارك الباحث في تدوين النتائج

7 صياغة وكتابة البحث و أعدادة للنشر: قام الباحث بالمشاركة في صياغة البحث وكتابتة

وأعدادة للنشر

%0.:

٧ نسبة المشاركة في البنث

رنيس القسم

التوقيع

وررو الرراب مررات مركز البحوث الزراعية معهد بحوث صحة الحيوان

استمارة رقم (٣) واسف مشروع بحث جدید مقترح

	: الطب البيطرى ـ أمراض الأسماك	١ – مجال التخصص العام
حدة البكتريولوجي)	: أمراض الأسماك (وحدة الطفيليات وو	٢- القيي
	: وحده الطغيليات _ وحده البكتر	الوحدة / المعمل الفرعى
	:	المعمل المتخصص
		~~~ مبررات ودوافع البحث :
النيلي.	منطقة الرأس في بعض الأسماك البلطي	ظهور بعض التقرحات على
		٤- عنوان البحث المحدد:
		عربي :
	اهرة تقرح في أسماك البلطي النيلي.	الصورة الميكروبية لط
		انجليزى:
Microbial profile of ulce	erative syndrome in Oreochromi	s niloticus.
and purples mad and any and and and and any red run over deer had the and hed	ن البحث :	٥- الهدف والنتائج المتوقعة ،
العدوى لها.	المسببة لهذه التقرحات مع دراسة مظاهر	
en e	ك المصابة لتحديد الصورة الإكلينيكية.	١- فحص عينات الأسمال
	وبات المسببة لهذه الإصابة.	
	ساسية للميكروب المعزول.	
بة للمرض.	عية لدراسة الصورة الإكلينيكية والتشريحي	

## البحث الأول

عنوان البحث : دراسة عن ظاهرة تقرح الرأس في أسماك البلطي.

القائمون بالبحث: د. عبد المحسن حسن و د. منى مصطفى حسين

مكان النشر : مجلة طب بيطرى بنى سويف المؤتمر العلمي الثاني في أكتوبر ٢٠٠١

مجلد ۱۱عدد (۲ أ): صفحة ۳۲۳ ـ ۳۷۹ سنة ۲۰۰۱

#### الملخص العربي

تمت هذه الدراسة بقسم بحوث الأسماك بمعهد بحوث صحة الحيوان اثناء تجميع اسماك البلطى النيلى وجد ان عدد خمس سمكات من أربعون سمكة مصابة بنقرحات في منطقة الرأس وأثبت الفحص البكتيريولوجى إصابتهم بميكروب (الفييروسبلينديدس) وذلك لأول مرة في مصر و ميكروب (الإيرومونس هيدروفيلا) كما أثبت الفحص الطفيلي إصابتهم بطفيل (التيريكودينا سنتر جاتا) و (التيركودينا مجنا) وأثبت إختبار الحساسية حساسية ميكروب (الفييروسبلينديدس) العالمية الي الأوكسي نتراتسيكلين والنتراسيكلين والكلورمفينيكول ومقاومتهاإلى النيوميسين والأموكسيلين وقد أجريت العدوى الصناعية بميكروب (الفييروسبلينديدس) بالحقن بالغشاء البريتوني وأدت الى نسبة نفوق وفي حالة العدوى الصناعية بكلاً من (الفييروسبلينديدس) و العدوى بطريقة الغمس الى اي نفوق وفي حالة العدوى الصناعية بكلاً من (الفييروسبلينديدس) و (الإيرومونس هيدروفيلا) ادت الى نسبة نفوق م ١٠٠% أما الحقن تحت الجلد ادى (الإيرومونس هيدروفيلا) ادت الى نسبة نفوق الغمس الى اى نفوق او ظهور أعراض مرضية.

#### Research

(1)

#### **Published**

Studies on ulcerative head syndrome among Oreochromis niloticus fish.

Abd El-Mohsen Hassan and Mona Mostafa Husien

Second Scientific Congress Proceeings Beni-Suef Vet. Med. J. Vol. XI., No. (2) Oct. (2001): (363-379).

#### **Summary**

Vibrio splendidus, Aeromonas hydrophila, Trichodina centrostigeata and T.magna were isolated from Oreochromais niloticus which suffered from ulcer in head region .Various characteristic of V splendidus have been studied .In vitro antibiotic sensitivity,pattern of isolated V. splendidus was tested against different chemotherapeutic agent .I/P inoculation of V. splendidus broth culyure also mixed broth cultures of V. splendidus and A. hydrophila resulted in 100% mortality, while the challenge by immersion gave no mortalities .