

**ISOLATION AND CHARACTERIZATION OF
BACTERIOPHAGE FOR VIBRIOSIS THERAPY IN
FISH**

MOHAMMAD TAMRIN BIN MOHAMAD LAL

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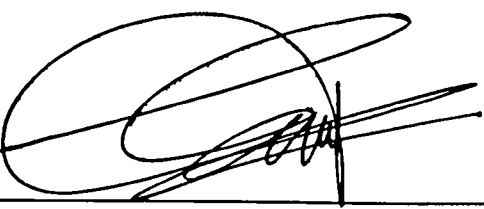
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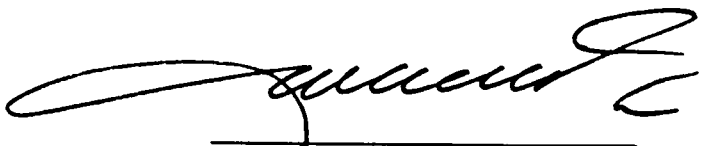
NAME : **MOHAMMAD TAMRIN BIN MOHAMAD LAL**
MATRIC NO. : **PY1211001T**
TITLE : **ISOLATION AND CHARACTERIZATION OF
BACTERIOPHAGE FOR VIBRIOSIS THERAPY IN FISH**
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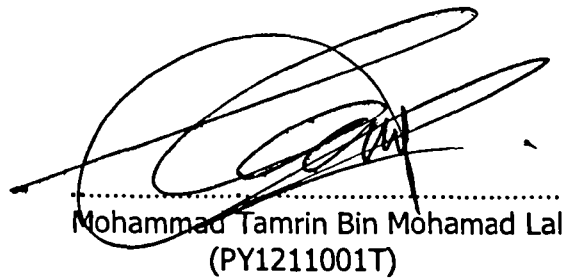
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I hereby declare that the material in this thesis is my own except for quotations, excepts, equations, summaries and references, which have been duly acknowledged.

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ABSTRACT

Bacterial infections in aquaculture are commonly treated using antibiotics. However, due to health concern and environmental issues, new control strategies for bacterial diseases are needed. Therefore, this study was conducted to isolate and characterize bacteriophage that are potentially be used as therapy for fish bacterial diseases. Four species of bacterial pathogens (*Vibrio alginolyticus*, *V. harveyi*, *V. parahaemolyticus* and *Photobacterium damselae*) were targeted for bacteriophage isolation. Each bacteriophage isolate was spotted onto different bacterial pathogens (*V. alginolyticus*, *V. harveyi*, *V. parahaemolyticus* and *Ph. damselae*) lawns. The bacteriophage morphology was determined using TEM and the whole genome sequence of bacteriophage was achieved using Illumina sequencing and *de novo* assembly. The stability of the bacteriophage was evaluated on different levels of pH, temperatures and bile concentrations. The bactericidal effect of the bacteriophage was evaluated using the *in vitro* co-culture method. In addition, the toxicity of the bacteriophage was evaluated against brine shrimp (*Artemia sp.*) and Asian seabass (*Lates calcarifer*) juveniles. This study has successfully isolated bacteriophage which were effective against *V. alginolyticus*, *V. harveyi* and *V. parahaemolyticus*. The bacteriophage isolates exhibited high specificity to its host with exception to *V. harveyi* phage that was also capable of infecting *V. parahaemolyticus* ATCC 17802. All phage isolates were classified under the double stranded DNA phage. The TEM analysis revealed that the *V. alginolyticus* phage, *V. harveyi* phage and *V. parahaemolyticus* phage were belong to the Family of *Myoviridae*, *Myoviridae* and *Siphoviridae*, respectively. The complete genome of *V. alginolyticus* phage was estimated at 248,088 bp and has high homology to *Vibrio* phage VH7D. Meanwhile, *V. parahaemolyticus* phage genome was 56,637 bp and hypothetically novel. Interestingly, all the phages possess methylated genome. The bioinformatics analyses revealed that the phage genomes have low significant homologies to vibrio virulent genes and toxin related proteins. All phage isolates were stable at 50 °C but completely deactivated at temperatures higher than 60 °C. The phage also stable at wide range of pH (4-9?) and high bile concentrations. Further analysis showed that the *V. parahaemolyticus* phage required high level of multiplicity of infection (MOI 100) to suppress the growth of its host but *V. harveyi* and *V. alginolyticus* phages required low MOI (0.01) to achieve similar effect. The findings of this study showed that the characteristics of the bacteriophage complied with the phage therapy requirement whereby all phages exhibited bactericidal effect and highly specific. The methylated genome allows the bacteriophage to survive from the defence mechanisms of the host bacteria. Lack of virulence genes prohibits the phage from contributing virulence to host bacteria through horizontal gene transfer. Furthermore, the phages were stable in both acidic and alkaline conditions which make them withstand the extreme condition of the gastrointestinal environment during therapy through oral administration. Most importantly, the bacteriophage were not toxic to the target animals. With these characteristics, the isolated phages seem beneficial for therapeutic use against vibriosis in aquaculture.

ABSTRAK

PEMENCILAN DAN PENGENALPASTIAN BAKTERIOFAJ UNTUK TERAPI VIBRIOSIS IKAN

Jangkitan bakteria di akuakultur pada umumnya dirawat menggunakan antibiotik. Namun, penggunaanya yang boleh menyebabkan masalah kesihatan dan menjelaskan alam sekitar memerlukan strategi kawalan jangkitan bakteria yang baru. Oleh itu, kajian ini dilakukan untuk memencil dan mengenalpasti bakteriofaj yang berpotensi untuk digunakan bagi tujuan terapi. Empat spesies bakteria pathogen (*Vibrio alginolyticus*, *V. harveyi*, *V. parahaemolyticus* dan *Photobacterium damselae*) digunakan untuk tujuan pemencilan bakteriofaj. Setiap isolat bakteriofaj diuji ke atas hamparan bakteria pathogen. Morfologi bakteriofaj tersebut ditentukan menggunakan TEM dan penjurukan keseluruhan genom bakteriofaj dihasilkan menggunakan penjurukan Illumina dan pemasangan genome de novo. Tahap kestabilan bakteriofaj dikaji pada tahap pH, suhu dan kepekatan hempedu yang berbeza. Kesan bakterisidal bakteriofaj ditentukan menggunakan ujian ko-kultur secara *in vitro*. Kesan toksik bakteriofaj pula ditentukan menggunakan ujian toksik terhadap anak udang (*Artemia sp.*) dan ikan siakap (*Lates calcarifer*). Kajian ini berjaya memmenculkan bakteriofaj yang berkesan melawan *V. alginolyticus*, *V. harveyi* dan *V. parahaemolyticus*. Isolat bakteriofaj tersebut amat spesifik terhadap perumahnya kecuali pada isolat bakteriofaj *V. harveyi* yang boleh menjangkiti *V. parahaemolyticus* ATCC 17802. Semua isolate faj (*V. alginolyticus*, *V. harveyi* and *V. parahaemolyticus*) adalah faj DNA dwibebenang. Analisis TEM menunjukkan faj *V. alginolyticus*, faj *V. harveyi* and faj *V. parahaemolyticus* masing-masing berada pada Famili Myoviridae, Myoviridae and Siphoviridae. Jujukan genom lengkap bagi faj *V. alginolyticus* dianggarkan pada 248,088 bp dan homolog kepada *Vibrio* phage VH7D. Manakala, genom *V. parahaemolyticus* adalah 56,637 bp dan berkemungkinan novel. Menariknya, kebanyakkan faj tersebut memiliki genom bermetil. Analisa bioinfomatik menunjukkan genom-genom tersebut memiliki homolg yang rendah terhadap gen virulen dan protein toksin vibrio. Semua faj adalah stabil pada suhu 50 °C tetapi tidak aktif pada suhu lebih tinggi dari 60 °C. Faj tersebut stabil pada julat pH yang besar (4-9) dan boleh bertoleransi pada tahap kepekatan hempedu yang tinggi. Analisis lanjut menunjukkan bahawa Faj *V. parahaemolyticus* memerlukan MOI yang tinggi (MOI 100) untuk membantutkan pertumbuhan perumahnya, namun, faj *V. harveyi* dan *V. alginolyticus* boleh membantutkan pertumbuhan perumahnya pada MOI yang rendah (MOI 0.01). Dapatkan kajian ini menunjukkan bakteriofaj tersebut menepati kriteria-kriteia untuk calon terapi dimana ia menunjukkan aktiviti bakterisidal yang tinggi dan amat spesifik. Genom bermetil juga membolehkan bakteriofaj bermandiri dari mekanisma pertahanan perumah. Gen virulen yang tidak dikesan menghadkan peningkatan tahap virulen bakteria melalui perpindahan gen. Selain itu, faj tersebut stabil dalam keadaan berasid dan beralkali membolehkan mereka bertoleransi dengan keadaan ekstrem gastrousus ikan selepas pemberian secara oral. Seterusnya, faj tersebut tidak toksik pada haiwan sasaran. Kesimpulan dari sifat-sifat yang dinyatakan, faj yang dipenculkan dalam kajian ini mungkin berfaedah untuk kegunaan terapeutik menentang vibriosis di akuakultur.

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LIST OF ABBREVIATIONS

Acc. No.	Accession Number
ATCC	American Type Culture Collection
BLAST	Basic Local Alignment Search Tools
bp	Base pair
CaCl₂	Calcium chloride
CDS	Coding sequences
DNA	Deoxyribonucleic acid
dTMP	Thymidine monophosphate
dUMP	Deoxyuridine monophosphate
dUTP	Deoxyuridine triphosphate
et al.	And others
FDA	Food and Drug Administration
GC	Guanine-cytosine
HCl	Hydrochloric acid
ICTV	International Committee on Taxonomy of Viruses
i.e.	That is
JCVI	J. Craig Venter Institute
LPS	Lipopolysaccharide
MOI	Multiplicity of infection
NaCl	Sodium chloride
NCBI	National Centre for Biotechnology Information
ORF	Open reading frame
PBS	Phosphate Buffered Saline
PEG	Polyethylene glycol
PES	Polyethersulfone
RAPD	Random amplified polymorphic DNA
RFLP	Restriction fragment length polymorphism
RNA	Ribonucleic acid
SDS	Sodium dodecyl sulfate
SDS-PAGE	Sodium dodecyl sulphate-polyacrylamide gel electrophoresis
TCBS	Thiosulphate citrate bile sucrose agar
TEM	Transmission Electron Microscopy
TEMED	Tetramethylethylenediamine
TSA	Tryptic soy agar
TSB	Tryptic soy broth
VALATCC	<i>V. alginolyticus</i> ATCC® 17749™
VANATCC	<i>V. anguillarum</i> ATCC
VHATCC	<i>V. harveyi</i> ATCC® 35084™
VPATCC	<i>V. parahaemolyticus</i> ATCC® 17802™

LIST OF SYMBOLS

$^{\circ}\text{C}$	Degree Celcius
μl	microliter
OD_{600}	Optical Density at 600 nm
%	Percent
mM	Milimolar
ml	Milliliter
l	Liter
$\times\text{g}$	Times gravity
M	Molar
$\text{ng } \mu\text{l}^{-1}$	Nanogram per microliter
nm	Nanometer
φ	Bacteriophage
Φ	Bacteriophage
Φ	Bacteriophage
Ψ	Bacteriophage
cfu ml^{-1}	Colony forming unit per milliliter
pfu ml^{-1}	Plaque forming unit per milliliter

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CHAPTER 1

GENERAL INTRODUCTION

1.1 Common Bacterial Pathogens in Marine Fish Aquaculture

Fish in captivity as well as in the natural habitat are exposed to many kinds of bacterial diseases including as vibriosis, streptococcosis and bacterial kidney disease (BKD) (Toranzo *et al.*, 2005). Vibriosis is one of the bacterial diseases which often occurs in aquaculture. Vibriosis may cause by various *Vibrios* such as *V. anguillarum*, *V. ordalii*, *V. salmonicida*, *V. vulnificus*, *V. harveyi*, *V. alginolyticus*, *V. cholerae*, *V. fischeri*, *V. furnisii*, *V. ichthyoenteri*, *V. logei*, *V. pelagius*, *V. splendidus*, *V. tapetis* or *V. wodanis* (Toranzo *et al.*, 2005; Won and Park, 2008; Austin and Austin, 2007). The outbreak of vibriosis has been reported to occur worldwide involving many marine organisms (Austin and Austin, 2007) and also freshwater fishes (Geng *et al.*, 2014). Fish affected by this disease generally shows typical signs of haemorrhage on the base of fins, exophthalmia, corneal opacity and skin lesions. Meanwhile, the moribund fish will experience severe anemia which manifested by pale gills (Toranzo *et al.*, 2005). Study by Ransangan and Mustafa (2009) showed that *V. harveyi* is responsible for mortality in Asian seabass (*Lates calcarifer*) cultured in Sabah, Malaysia.

Streptococcosis had been reported both in freshwater and marine fish aquaculture. Although it can be caused by many *Streptococcus* species, most of the infections in marine aquaculture are due to *Streptococcus iniae* (Musa *et al.*, 2007). Infected fish normally showed meningoencephalitis, panophthalmitis, skin lesion, necrosis, corneal opacity and hemorrhage (Musa *et al.*, 2007). Streptococcosis can easily be transmitted through contact with infected fish or contaminated feeds (Musa *et al.*, 2007).

*Photobacterium damsela*e is a marine bacterium that causes infection in a variety of marine fish (Rivas *et al.*, 2013). Fish species which are reported to be affected by this pathogen include rainbow trout (Pedersen *et al.*, 2009), seabass

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