



UNIVERSITI PUTRA MALAYSIA

INVESTIGATION ON INDIGENOUS BACILLUS ISOLATES WITH BIOREMEDIATION PROPERTIES FOR IMPROVING WATER QUALITY AND SHRIMP HEALTH IN MALAYSIAN AQUACULTURE

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By

DEVARAJA T.N.

Thesis Submitted to the School of Graduate Studies, Universiti Putra Malaysia in Fulfilment of the Requirement for the Degree of Doctor of Philosophy

February 2002



Dedicated to my parents, who supported me to become what I want



Abstract of thesis presented to the Senate of Universiti Putra Malaysia in fulfilment of the requirement for the degree of Doctor of Philosophy

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February 2002

Chairperson: Professor Fatimah Md. Yusoff, Ph.D.

Faculty: Science and Environmental Studies

Indigenous marine bacteria of the genus *Bacillus* were selected to study their properties as potential use for bioremediation owing to their inherent versatility. Bacteria were isolated from water and sediment samples collected along the west coast of Peninsular Malaysia in brackishwater environment. Selected isolates were identified to species level using biochemical and API CH kit and three suitable isolates, Bacillus pumilus AB58, B. subtilis AB65 and B. licheniformis AB69 were selected for the study. Optimum growth requirements of temperature, NaCl and pH were 30°C, 1.5% and 7.5 respectively, determined for the isolates by measuring the optical density and corresponding cell number. The growth curves of the isolates were plotted and all of them reached maximum cell number during a 16-20 h incubation. The cell density in overnight cultures of B. pumilus AB58, B. subtilis AB65 and B. licheniformis AB69 were 5.7×10^9 (± 0.8), 3.7×10^8 (± 0.6), 5.0×10^9 (± 0.6) cfu/ml respectively. They had the ability to tolerate ammonia levels of up to 20 mg/l without a considerable change in cell numbers for 48 h. However, the growth was suppressed completely at 25 mg/l of ammonia. At 40 ppt salinity, all the isolates survived for 4 days without significant change in initial cell numbers (10⁸ cfu/ml). The selected isolates were found to secrete extracellular enzymes viz., protease, gelatinase, amylase and



lipase as detected by clear zone formation on substrate based agar plates. Bacillus pumilus AB58 and B. subtilis AB65 produced significantly (P < 0.05) bigger protease clear zones $(19.0 \pm 2.0 \text{ and } 23.0 \pm 4.0 \text{ diameter in mm respectively})$ than B. licheniformis AB69. However, B. subtilis AB65 secreted significantly (P < 0.05) more amylase (31.0 \pm 5.0 diameter in mm) than the other two isolates. All the isolates were sensitive to most of the antibiotics tested on MHA plates. These isolates were compatible with each other in mixed culture conditions. They inhibited as well as excluded all the pathogenic vibrios (Vibrio alginolyticus M11, V. alginolyticus M12, V. parahaemolyticus M1, V. parahaemolyticus M3, V. parahaemolyticus M6, V. alginolyticus T, V. parahaemolyticus T, V. harveyi I and V. parahaemolyticus I) tested by diffusion disc, streak plate and common broth methods. Synergistic effect of isolates had significantly higher (P < 0.05) inhibition of all vibrios than the individual isolates. The isolates were confirmed for their non-pathogenicity to shrimp postlarvae (PL 29). All three isolates were tested for their effect on ammonia in simulated pond conditions. All non-aerated treatment tanks had significantly lower ammonia levels (P < 0.05) than the non-aerated control tanks, which were not treated with bacterial isolates both in case of single and combination treatments. Synergistic effect of isolates reduced the ammonia levels at a faster rate than the treatments with single isolate. Sediment properties were not significantly different between treated and control groups except for the total and available phosphorous levels, which were significantly higher in tanks treated with B. licheniformis AB69 (P < 0.05) compared to the others. The selected Bacillus isolates satisfied the criteria to qualify them for bioremediation in aquaculture.



Abstrak tesis dikemukakan kepada Senat Universiti Putra Malaysia sebagai memenuhi syarat untuk mendapat ijazah Doktor Falsafah

PENYIASATAN KE ATAS ISOLAT-ISOLAT *BACILLUS* TEMPATAN BERCIRIKAN BIOREMEDIASI BAGI MENINGKATKAN KUALITI AIR DAN KESIHATAN UDANG UNTUK AKUAKULTUR DI MALAYSIA

Oleh

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Februari 2002

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Dalam kajian ini, spesies *Bacillus* marin tempatan telah dipilih untuk bioremediasi kerana sifat serba boleh yang semulajadi. Bakteria daripada sampel air dan sedimen daripada persekitaran air payau telah di kutip dari sepanjang pantai barat Semenanjung Malaysia. Isolat-isolat yang terpilih telah dikenalpasti ke peringkat spesies dengan menggunakan kaedah biokimia dan kit tersedia API CH dan tiga isolat terbaik, *Bacillus pumilus* AB58, *B. subtilis* AB65 dan *B. licheniformis* AB69 telah dipilih untuk kajian ini. Paras optimum keperluan-keperluan pertumbuhan asas; suhu, garam dan pH masing-masing adalah 30°C, 1.5% dan 7.5, yang mana telah ditentukan untuk isolat-isolat tersebut dengan mengukur densiti optik dan bilangan sel sejajar. Lengkuk pertumbuhan isolat-isolat telah diplot dan didapati bahawa semua isolat mencapai bilangan sel maksimum pada 16-20 jam (j) inkubasi. Densiti sel di dalam kultur semalaman *B. pumilus* AB58, *B. subtilis* AB65 dan *B. licheniformis* AB69 adalah $5.7 \times 10^9 (\pm 0.8)$, $3.7 \times 10^8 (\pm 0.6)$, $5.0 \times 10^9 (\pm 0.6)$ cfu/ml masing-masing. Mereka berupaya menahan paras ammonia sehingga ke 20 mg/l tanpa perubahan pada bilangan sel selama 48 j. Walau bagaimanapun, pertumbuhan terencat sepenuhnya apabila paras ammonia mencapai

tahap 25 mg/l. Pada saliniti 40 ppt, semua isolat berjaya hidup selama 4 hari tanpa perubahan yang signifikan pada bilangan sel permulaan (10⁸cfu/ml). Isolat-isolat terpilih di dapati mengeluarkan enzim-enzim luar sel, iaitu protease, gelatinase, amilase dan lipase berdasarkan zon yang terang terhasil di atas piring agar. Bacillus pumilus AB58 dan B. subtilis AB65 lebih signifikan (P < 0.05) di dalam menghasilkan zon yang lebih terang (19.0 \pm 2.0, dan 23.0 \pm 4.0 diameter dalam mm masing-masing) berbanding dengan B. licheniformis AB69. Walau bagaimanapun, B. subtilis AB65 lebih signifikan di dalam merembes amilase (P < 0.05), iaitu (31.0 \pm 5.0 diameter dalam mm) berbanding dengan kedua-dua isolat yang lain. Kesemua isolat adalah sensitif kepada kebanyakan antibiotik yang diuji di atas piring MHA. Isolat-isolat ini adalah serasi di antara satu sama lain dalam keadaan kultur campuran. Mereka merencat dan menyingkirkan semua vibrios patogenik (Vibrio alginolyticus M11, V. alginolyticus M12, V. parahaemolyticus M1, V. parahaemolyticus M3, V. parahaemolyticus M6, V. alginolyticus T, V. parahaemolyticus T, V. harveyi I and V. parahaemolyticus I) apabila diuji dengan cakera resapan, piring coretan dan medium biasa. Kesan sinergistik oleh isolat-isolat adalah secara signifikannya lebih tinggi (P < 0.05) ke atas perencatan semua vibrios berbanding dengan isolat tunggal. Isolat-isolat juga telah disahkan tidak patogenik terhadap pasca larval udang (PL 29). Ketiga-tiga isolat telah diuji kesannya terhadap ammonia dalam kolam simulasi. Semua tangki rawatan yang tidak mengandungi pengudaraan menunjukkan paras ammonia yang secara signifikannya lebih rendah (P < 0.05) berbanding dengan tangki kawalan yang tidak mengandungi pengudaraan, samada secara berasingan atau campuran. Kesan sinergistik isolat-isolat dapat mengurangkan paras ammonia pada kadar yang lebih cepat berbanding dengan



rawatan menggunakan isolat tunggal. Kandungan sedimen adalah tidak signifikan di antara kumpulan yang dirawat dan kawalan kecuali untuk jumlah dan tahap tersedia fosforus, yang mana secara signifikannya lebih tinggi di dalam tangki yang dirawat dengan *B. licheniformis* AB69 (P < 0.05) berbanding dengan yang lain. Isolat-isolat *Bacillus* yang terpilih memenuhi kriteria untuk melayakkannya sebagai agen bioremediasi di dalam akuakultur.



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I certifiy that an Examination Committee met on 1st February 2002 to conduct the final examination of Devaraja T.N. on his Doctor of Philosophy thesis entitled "Investigation on Indigenous *Bacillus* Isolates with Bioremediation Properties for Improving Water Quality and Shrimp Health in Malaysian Aquaculture" in accordance with Universiti Pertanian Malaysia (Higher Degree) Act 1980 and Universiti Pertanian Malaysia (Higher Degree) Regulations 1981. The Committee recommends that the candidate be awarded the relevant degree. Members of the Examination Committee are as follows:

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DECLARATION

I hereby declare that the thesis is based on my original work except for quotations and citations, which have been duly acknowledged. I also declare that it has not been previously or concurrently submitted for any other degree at UPM or other institutions.

Devaraja TA

Date: 25.03.2002





TABLE OF CONTENTS

| DEDICATION | ii |
|-----------------------|------|
| ABSTRACT | iii |
| ABSTRAK | v |
| ACKNOWLEDGEMENTS | viii |
| APPROVAL SHEETS | x |
| DECLARATION | xii |
| LIST OF TABLES | xvii |
| LIST OF FIGURES | xix |
| LIST OF PLATES | XX |
| LIST OF ABBREVIATIONS | xxi |

CHAPTER

| 1 | GENI | ERAL INTRODUCTION | 1 |
|---|------|---|----|
| | 1.1 | Background and Scope of the Study | 1 |
| | 1.2 | Statement of the Problem | 8 |
| 2 | LITE | RATURE REVIEW | 12 |
| | 2.1 | Microorganisms and Nutrient Cycles in Aquatic System | 12 |
| | | 2.1.1 Nitrogen Cycle | 13 |
| | | 2.1.2 Sulphur Cycle | 15 |
| | | 2.1.3 Carbon Cycle | 16 |
| | | 2.1.4 Phosphorus Cycle | 18 |
| | 2.2 | Organic Matter in Aquatic System | 19 |
| | | 2.2.1 Production of Organic Matter | 19 |
| | | 2.2.2 Degradation of Organic Matter | 20 |
| | | 2.2.3 Organic Matter in Aquaculture Pond | 23 |
| | 2.3 | Need for Water and Soil Quality Management in Aquaculture | 25 |
| | | 2.3.1 Water Quality Management | 25 |
| | | 2.3.2 Soil Quality Management | 26 |
| | 2.4 | Toxicity of Ammonia, Nitrite and Sulphide to Shrimp | 29 |
| | | 2.4.1 Ammonia | 30 |
| | | 2.4.2 Nitrite | 31 |
| | | 2.4.3 Hydrogen Sulphide | 31 |
| | 2.5 | Bioremediation | 33 |
| | | 2.5.1 Bioremediation in Waste Water Management | 36 |
| | | 2.5.2 Bioremediation of Toxic Chemical Polluted Environment | 37 |
| | | 2.5.3 Biotransformation of Metals and Metallic Compounds | 38 |
| | | 2.5.4 Oil Bioremediation | 39 |
| | 2.6 | Microorganisms and Microbial Products in Aquaculture | 40 |
| | | 2.6.1 Extracellular Enzymes in Aquatic Systems | 40 |
| | | 2.6.2 Manipulation of Microbial Communities in | |
| | | Aquaculture Ponds | 42 |



| | | 2.6.3 | | |
|---|------|----------|--|-----|
| | | | Applications | 45 |
| | | 2.6.4 | The Significance of the Use of Microbial Products in | |
| | | | Aquaculture | 53 |
| | | 2.6.5 | Bioremediation in Aquaculture and Bacillus Species | 55 |
| 3 | ISOL | ATION | AND IDENTIFICATION OF | |
| | BAC | ILLUS IS | SOLATES | 61 |
| | 3.1 | | | 61 |
| | 3.2 | | ials and Methods | 62 |
| | | | Sample Collection | 62 |
| | | | Isolation and Identification | 62 |
| | | 3.2.3 | Colony Characteristics and Cell Morphology | 65 |
| | 3.3 | Result | S | 66 |
| | | 3.3.1 | Isolation and Identification | 66 |
| | | 3.3.2 | Colony Characteristics and Cell Morphology | 68 |
| | 3.4 | Discus | ssion | 70 |
| 4 | PRO | PERTIES | S OF SELECTED BACILLUS ISOLATES | 76 |
| | 4.1 | Introd | uction | 76 |
| | 4.2 | Mater | ials and Methods | 77 |
| | | 4.2.1 | Measurement of Optical Density in Overnight | |
| | | | Bacterial Culture | 77 |
| | | | Cell Enumeration in Overnight Bacterial Culture | 77 |
| | | | Optimum Growth Requirements | 78 |
| | | | Growth Curves of Bacillus Isolates | 79 |
| | | | Effect of Ammonia on the Growth of Bacillus Isolates | 80 |
| | | 4.2.6 | | 81 |
| | | 4.2.7 | Detection of Extracellular Enzymes | 83 |
| | | 4.2.8 | | 84 |
| | | 4.2.9 | | 85 |
| | | 4.2.10 | Interaction Effect of Bacillus Isolates Using | |
| | | | Diffusion Disc Method | 88 |
| | | | Statistical Analysis | 88 |
| | 4.3 | Result | | 89 |
| | | 4.3.1 | Optimum Growth Requirements | 89 |
| | | 4.3.2 | Growth Curves of <i>Bacillus</i> Isolates | 93 |
| | | 4.3.3 | | 95 |
| | | 4.3.4 | | 97 |
| | | 4.3.5 | 5 | 99 |
| | | 4.3.6 | Antibiotic Sensitivity of <i>Bacillus</i> Isolates | 105 |
| | | 4.3.7 | | 106 |
| | A A | 4.3.8 | Interaction Effect | 106 |
| | 4.4 | Discus | | 107 |
| | | 4.4.1 | Growth Requirements and Growth Curves | 107 |
| | | 4.4.2 | Effect of Ammonia and Salinity on the Growth | 100 |
| | | | of Bacillus Isolates | 109 |

| | | 4.4.3 | Extracellular Enzymes, Antibiogram, Species Differentia | |
|---|------|-----------------|---|-----|
| | | | and Interaction | 111 |
| 5 | BIOO | CONTRO | DL OF <i>VIBRIO</i> SPECIES BY <i>BACILLUS</i> ISOLATES | 114 |
| | 5.1 | Introdu | uction | 114 |
| | 5.2 | Materi | als and Methods | 116 |
| | | 5.2.1 | Selected Bacillus Isolates | 116 |
| | | 5.2.2 | Pathogenic Vibrios | 116 |
| | | 5.2.3 | Diffusion Disc Method | 117 |
| | | 5.2.4 | Perpendicular Streak Plate Method | 118 |
| | | 5.2.5 | Competitive Exclusion of Vibrios by Bacillus Isolates | 118 |
| | 5.3 | Result | S | 120 |
| | | 5.3.1 | Inhibition by Diffusion Disc Method | 120 |
| | | 5.3.2 | Inhibition by Perpendicular Streak Plate Method | 121 |
| | | 5.3.3 | Competitive Exclusion by Common Broth Method | 121 |
| | 5.4 | Discus | sion | 124 |
| 6 | ΡΑΤ | HOGENI | ICITY OF BACILLUS ISOLATES ON PENAEUS | |
| Ũ | | | POSTLARVAE | 129 |
| | 6.1 | Introdu | | 129 |
| | 6.2 | | als and Methods | 130 |
| | | 6.2.1 | Experimental Animals | 130 |
| | | 6.2.2 | Selected Isolates | 130 |
| | | 6.2.3 | Experimental Set-up | 131 |
| | | 6.2.4 | Examination With Light Microscope | 132 |
| | | 6.2.6 | Survival Percentages of P. monodon Postlarvae | 132 |
| | 6.3 | Result | S | 133 |
| | | 6.3.1 | General Parameters | 133 |
| | | 6.3.2 | External Observation With Light Microscope | 134 |
| | 6.4 | Discus | sion | 135 |
| 7 | EFFE | ECT OF <i>E</i> | <i>ACILLUS</i> ISOLATES | |
| | | | IA CONCENTRATIONS | 140 |
| | 7.1 | Introdu | uction | 140 |
| | 7.2 | Materi | als and Methods | 142 |
| | | 7.2.1 | Selected Isolates | 142 |
| | | 7.2.2 | Measurement of Ammonia Concentrations | 143 |
| | | 7.2.3 | Preliminary Experiments | 144 |
| | | 7.2.4 | Effect of Individual Bacillus Isolate on | |
| | | | Ammonia Concentration and Sediment Quality | 145 |
| | | 7.2.5 | , , | |
| | | | on Ammonia Concentration | 146 |
| | | 7.2.6 | Statistical Analysis | 147 |
| | 7.3 | Result | | 147 |
| | | | Preliminary Experiments | 147 |
| | | 7.3.2 | Effect of Individual Bacillus Isolate on | |
| | | | Ammonia Concentration and Sediment Quality | 148 |

| | 7.3.3 Synergistic Effect of Two <i>Bacillus</i> Isolates on Ammonia Concentration 7.4 Discussion | 156 161 |
|---|---|-------------------|
| 8 | GENERAL DISCUSSION AND RECOMMENDATIONS | 170 |
| | BIBLIOGRAPHY APPENDICES VITA | 179 201 211 |



LIST OF TABLES

| Table | 1 | Page |
|-------|--|------|
| 1 | Commercial microbial products available in the market and their functions | 7 |
| 2 | Approximate feed wastage in different type of cultures | 25 |
| 3 | Water quality parameters for penaeid shrimp grow-out ponds | 27 |
| 4 | Lethal toxicity level of ammonia and nitrite to shrimp, Penaeus monodon | 32 |
| 5 | Some of the vital functions of bacteria involved in mineralisation | 46 |
| 6 | Microorganisms possessing in vitro probiotic properties | 50 |
| 7 | Probiotics and feed supplements used in aquaculture | 50 |
| 8 | Basic generic characteristics of the selected Bacillus isolates | 66 |
| 9 | Carbohydrate based tests for species identification of <i>Bacillus</i> isolates using API CH 50 kit | 67 |
| 10 | Supplementary tests in API 20 E kit | 68 |
| 11 | Cell density of <i>Bacillus</i> isolates after 18 hours of incubation under different temperature indicated by optical density (at 600 nm) and corresponding cell number (cfu/ml) respectively | 90 |
| 12 | Cell density of <i>Bacillus</i> isolates after 18 hours of incubation under different pH levels indicated by optical density (at 600 nm) and corresponding cell number (cfu/ml) respectively | 91 |
| 13 | Cell density of <i>Bacillus</i> isolates after 18 hours of incubation under different salt levels indicated by optical density (at 600 nm) and corresponding cell number (cfu/ml) respectively | 92 |
| 14 | Optical density (at 600 nm) and cell number (cfu/ml) in 18 hour culture of <i>Bacillus</i> isolates under optimum growth conditions | 93 |
| 15 | Cell density of <i>Bacillus</i> isolates after 18 hours of incubation under different levels of total ammonia-nitrogen (mg/l) | 96 |
| 16 | Cell density of <i>Bacillus</i> isolates after 18 hours of incubation under different levels of salinity (ppt) | 98 |



| 17 | Clear zone diameter (in mm) formed by extracellular enzymes from <i>Bacilla</i> isolates after 18 hours of incubation | us 99 |
|----|---|----------|
| 18 | Antibiogram of <i>Bacillus</i> isolates | 105 |
| 19 | Pathogenic vibrios used in biocontrol experiments | 117 |
| 20 | Inhibition zones (diameter in mm) formed in diffusion disc method by <i>Bacillus</i> isolates against pathogenic vibrios | 122 |
| 21 | Inhibition zones (diameter in mm) formed in diffusion disc method by combination of <i>Bacillus</i> isolates against pathogenic vibrios | 122 |
| 22 | Growth dominance of <i>Bacillus</i> isolates against the pathogenic vibrios at the point of intersection in perpendicular streak plate method | 123 |
| 23 | Viable counts of VaM11 (cfu/ml) after 48 h incubation with <i>Bacillus</i> isolates | 123 |
| 24 | Viable counts of VpM1 (cfu/ml) after 48 h incubation with <i>Bacillus</i> isolates | 124 |
| 25 | Water quality parameters, total plate count (TPC) and percentages of shrimp postlarvae survived in different treatments for 14 days | 133 |
| 26 | Water quality parameters in control and treated tanks | 148 |
| 27 | Total plate counts, TPC (cfu/ml) of water in all tanks during the experiment | 149 |
| 28 | Physical and chemical composition of sediment in control and treatment tanks at the beginning and after 14 days of experiment | 155 |
| 29 | Water quality parameters in control tanks and tanks treated with combination of <i>Bacillus</i> isolates | 157 |



LIST OF FIGURES

| Figure | | Page |
|--------|--|------|
| 1 | Water and sediment samples collection sites along the west coast of Peninsular Malaysia | 63 |
| 2 | Growth curves of selected Bacillus isolates at 30°C and pH 7.5 | 94 |
| 3 | Specific growth rate (μ) of <i>Bacillus</i> isolates at log phase | 94 |
| 4 | Cell numbers (cfu/ml) of Bacillus at 20 mg/l total ammonia-nitrogen | 97 |
| 5 | Cell numbers (cfu/ml) of Bacillus at 40 ppt salinity | 99 |
| 6 | Total ammonia-nitrogen concentrations (mg/l) in aerated tanks treated with <i>Bacillus</i> isolates | 153 |
| 7 | Total ammonia-nitrogen concentrations (mg/l) in non-aerated tanks treated with <i>Bacillus</i> isolates | 154 |
| 8 | Total ammonia-nitrogen (mg/l) in aerated tanks treated with combinations of <i>Bacillus</i> isolates | 159 |
| 9 | Total ammonia-nitrogen (mg/l) in non-aerated tanks treated with combinations of <i>Bacillus</i> isolates | 160 |





LIST OF PLATES

| Plate | | Page |
|-------|---|-----------|
| 1 | Endospore stained cells of a) <i>Bacillus pumilus</i> AB58 – usually occur as single rods; b) <i>B. subtilis</i> AB65 – shortest rods among three isolates; and c) <i>B. licheniformis</i> AB69 - note the long rods | 71 |
| 2 | Scanning electron microphotographs showing rod shaped vegetative cells of a) <i>Bacillus pumilus</i> AB58 (mag. x7000); b) <i>B. subtilis</i> AB65 (mag. x4000); and c) <i>B. licheniformis</i> AB69 (mag. x6,500)-longest rods among three isolates | 72 |
| 3 | Clear zones formed on skim milk agar plates due to the action of protease secreted by a) <i>B. pumilus</i> AB58 showed bigger clear zone than; b) <i>B. subtilis</i> AB65; and c) <i>B. licheniformis</i> AB69 | 101 |
| 4 | Clear zones formed on nutrient gelatin agar plates due to the action of gelatinase secreted by a) <i>B. pumilus</i> AB58 showed bigger clear zone that b) <i>B. subtilis</i> AB65; and c) <i>B. licheniformis</i> AB69 | n; 102 |
| 5 | Clear zones formed on starch plates due to the action of amylase secreted by a) <i>Bacillus pumilus</i> AB58 showed bigger clear zone than; b) <i>B. subtilis</i> AB65; and c) <i>B. licheniformis</i> AB69 | 103 |
| 6 | Clear zones formed on lipid plates due to the action of lipase secreted by a) <i>Bacillus pumilus</i> AB58; b) <i>B. subtilis</i> AB65; and c) <i>B. licheniformis</i> AB69 (biggest clear zone) | 104 |
| 7 | RAPD of selected Bacillus isolates using wide range primer | 107 |
| 8 | Interaction of a) <i>Bacillus pumilus</i> AB58 (\Box) and <i>B. subtilis</i> AB65 (∇ , small inhibition zone) with <i>B. licheniformis</i> AB69; b) <i>B. pumilus</i> AB58 (\Box) and <i>B. licheniformis</i> AB69 (+,small inhibition zone) with <i>B. subtilis</i> AB65; and c) <i>B. subtilis</i> AB65 (∇) and <i>B. licheniformis</i> AB69 (+) with <i>B. pumilus</i> AB58 – note the inhibition of <i>B. pumilus</i> AB58 by the other two isolates | 108 |
| 9 | Microphotographs of <i>Penaeus monodon</i> postlarvae showing a) healthy muscle (mag. x250); b) intact telson (mag. x1000); b) undamaged rostral spine (mag. x1000); and d) undamaged c) rostrum tip (mag. x1000) | 135 |

LIST OF ABBREVIATIONS

| AAHU | – aquatic animal health unit |
|------------------|--|
| ANOVA | – analysis of variance |
| Bl | – Bacillus licheniformis AB69 |
| BOD | - biological oxygen demand |
| Вр | – Bacillus pumilus AB58 |
| bp | – base pair |
| Bs | – Bacillus subtilis AB65 |
| BWS | – bacterial white spot |
| cfu | – colony forming units |
| COD | – chemical oxygen demand |
| DDW | - double distilled water |
| DNA | – deoxyribo nucleic acid |
| dNTPs | - deoxyribonucleoside triphosphates |
| DO | dissolved oxygen |
| DW | - distilled water |
| EDTA | – ethylene dinitro tetraacetic acid |
| ERMs | - environmentally relevant microorganisms |
| FAO | - Food and Agriculture Organisation |
| GEMs | - genetically engineered microorganisms |
| h | – hour |
| kDa | – kilo dalton |
| LD ₅₀ | – lethal dose 50 |

| mmt | – million metric tonne |
|----------|--|
| mt | – metric tonne |
| OD | - optical density |
| PCR | – polymerase chain reaction |
| ppt | - parts per thousand |
| RAPD | – random amplified polymorphic DNA |
| 16S rRNA | - 16 subunit ribosomal ribose nucleic acid |
| S.E. | – standard error |
| TBE | – tris boric acid EDTA |
| TCBS | - thiosulphate citrate bile salt sucrose |
| TPC | – total plate count |
| TSA | – trypticase soy agar |
| TSB | - trypticase soy broth |
| UPM | – Universiti Putra Malaysia |
| USD | – Dollar of United States of America |
| VaM11 | – Vibrio alginolyticus Malaysia 11 |
| VaM12 | – Vibrio alginolyticus Malaysia 12 |
| VaT | – Vibrio algninolyticus Thailand |
| VhI | Vibrio harveyi Indonesia |
| VpI | - Vibrio paraheamolyticus Indonesia |
| VpM1 | – Vibrio parahaemolyticus Malaysia 1 |
| VpM3 | – Vibrio parahaemolyticus Malaysia 3 |
| VpM6 | – Vibrio parahaemolyticus Malaysia 6 |
| VpT | – Vibrio parahaemolyticus Thailand |



CHAPTER 1

GENERAL INTRODUCTION

1.1 Background and Scope of the Study

In recent decades aquaculture has become a major food production industry helping to meet the increasing demand for food. The world population has crossed six billion during the year 2000 (Census Bureau, 2000) causing an increase in demand for food. Therefore, food production by agriculture and capture fisheries has to be supplemented by other alternatives. Total world fish and shellfish production has reached 126.17 million metric tonnes (mmt) in 1999 (FAO, 2000) of which, 33.31 mmt come from aquaculture.

Dehadrai (1993) predicted that aquaculture has to fill the gap of 19.6 mmt by 2000, 37.5 mmt by 2010 and 62.54 mmt by 2020. Among various aquaculture practices, shrimp culture is gaining increasing importance world-wide due to the short period of culture and high profits. Even though shrimp farming has grown to become a booming export oriented industry, shrimp production and trade have undergone fluctuations during the past four years indicating uncertainty over its sustainability in days to come due to different social and environmental problems.

The shrimp industry has to develop strategies to tackle the viral disease problems, which have disrupted shrimp farming. Generally in shrimp farming, the ultimate goal is to maximise the production with high stocking density, increase feeding and increase water exchange often accompanied with heavy use of chemicals. This has caused various other problems like disease outbreaks, environmental pollution and other socio-economic fall out (Primavera, 1994). The shrimp industry is now looking for ways to rebuild itself and aims at a long-term sustainability that will not pollute the environment or minimise pollution.

The incidence and severity of several infectious diseases largely depends upon the quality of environment in which the host lives. Outbreak of diseases might be avoided by maintaining a healthy environment through suitable water quality management practices. Healthy environment can also be achieved by stocking at optimum density, providing sufficient water exchange and aeration (Wang *et al.*, 1999a). In the past, water exchange was the only solution widely practised to get rid of the accumulated wastes but recent viral disease outbreak has limited this practice. The recent practice of using closed or semi-closed system with heavy chlorination to reduce introduction of viral infection is not popular due to the extra cost involved. Reducing stocking density and feeding rate to minimise the accumulation of organic matter has also indirectly decreased the rate of production thus reducing the profit margin (Phillips *et al.*, 1993). Frequent water exchange to flush out shrimp pond effluent eutrophicates the coastal environment.

The ultimate strategy is to manipulate the pond environment by enhancing the mineralisation process in culture ponds so as to maintain a healthy environment and minimise the nutrient load to reduce pollution before disposing the effluents (Phillips *et al.*, 1993). Several management practices have been adopted for maintaining optimal water and sediment quality like the use of chemicals, physical methods like bottom