



UNIVERSITI PUTRA MALAYSIA

**EPIDEMIOLOGY AND DIAGNOSIS OF HUMAN LEPTOSPIROSIS IN
MALAYSIA**

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**EPIDEMIOLOGY AND DIAGNOSIS OF HUMAN LEPTOSPIROSIS IN
MALAYSIA**

By

ISAM MOHAMED ALI MOHAMED EL-JALII

**Thesis Submitted in Fulfilment of the Requirements for the Degree of
Doctor of Philosophy in the Institute of Bioscience
Universiti Putra Malaysia**

September 2000



DEDICATION

**TO THE MEMORY OF MY MOTHER, FATHER AND NEPHEW
KHATAB**

TO MY WIFE EMTENAN AND DAUGHTER RAYAN

TO MY BROTHERS AND SISTERS

TO ALL OF THEM WITH LOVE AND GRATITUDE

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

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Chairman: Prof. Dr. Abdul Rani Bahaman

Institute of Bioscience

Retrospective study of human leptospirosis in Malaysia based on microscopic agglutination test (MAT) showed 13% overall prevalence of infection for the period 1983-1998. Results indicated that the prevalence was decreasing in the last five years (1994-1998). The prevalence was highest among Indians (16.67%) followed by Malays (11.48%) and the least among Chinese (5.88%). The 20-29 year-old group showed the highest prevalence of infection (17.13%). Less than 10 year-old group showed the least prevalence of infection (5.66%). Generally, many of the cases occurred between the ages of 20-50 years. Serological survey based on enzyme-linked immunosorbent assay (ELISA) showed a high overall prevalence (12.56%) of leptospiral infection. Kuala Lumpur showed the highest prevalence (19%) whilst Penang recorded the lowest prevalence (6.67%). No significant differences in the prevalence between the other states was noted.



A comparative study of three serological tests, namely enzyme-linked immunosorbent assay (ELISA), microscopic agglutination test (MAT) and indirect hemagglutination (IHA), test was carried out to evaluate these tests in the diagnosis of human leptospirosis. A total of 3000 serum samples from three groups of people were examined. In Group I, IgM and IgG-ELISA were able to detect a number of cases in the first sampling before MAT titres were detectable. In the second sampling, all samples positive for MAT were also positive for IgM-ELISA. IHA test gave positive reactions with only 38% of the samples while all samples were positive for ELISA. In Group II, ELISA detected IgM and IgG to leptospires in the samples which were negative to MAT. These were samples from patients with clinical signs of leptospirosis.

The polymerase chain reaction (PCR) was evaluated as a tool for diagnosis of leptospirosis and differentiation of leptospiral strains. Urine samples with as little as 10 serovar *hardjo* cells per ml of urine were positive on PCR indicating high sensitivity of the test. Detection of small number of leptospiral cells in urine by PCR was an advantage over culture. Random amplification polymorphic DNA (RAPD) fingerprinting was applied to differentiate leptospiral strains. Two primers were tested for their abilities to generate individual RAPD fingerprints. The DNA profiles obtained with each primer were distinct and reproducible. The fingerprint obtained could be useful for distinguishing the serovars up to the strain level. Profiles obtained revealed genetic heterogeneity between serovars belong to one serogroup.

Analysis of leptospiral DNA with restriction enzymes, *Hind* III, *BamH* I and *EcoR* I revealed a high heterogeneity between the serovars examined. This high heterogeneity may be due to the large genome of the genus *Leptospira*. No relationship was found between the restriction patterns and the species from which the isolate was isolated. Similarities were observed among isolates of the same serovar. The 10 leptospiral field isolates were assayed for presence of plasmid DNA. Only two isolates were found to harbour plasmid DNA. The plasmid profiling obtained is of limited epidemiological value for differentiation of leptospiral isolates.

It appears that human leptospirosis is an endemic infection in Malaysia. The findings showed that ELISA was a suitable serological test for diagnosis of leptospirosis compared to MAT and IHA tests. On the other hand, the application of PCR and REA in the diagnosis of leptospirosis would be useful.

Abstrak tesis yang dikemukakan kepada Senat Universiti Putra Malaysia sebagai memenuhi keperluan untuk ijazah Doktor Falsafah

EPIDEMIOLOGI DAN DIAGNOSIS LEPTOSPIROSIS DI MALAYSIA

Oleh

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Kajian retrospektif terhadap leptospirosis manusia di Malaysia berdasarkan ujian pengaglutinatan mikroskopi (MAT) menunjukkan prevalens jangkitan keseluruhan 13% untuk tempoh 1983 - 1998. Hasil kajian menunjukkan prevalens ini semakin kurang dalam tempoh lima tahun terakhir (1994-1998). Prevalens ini paling tinggi di kalangan kaum India (16.67%) diikuti kaum Melayu (11.48%) dan paling sedikit di kalangan kaum Cina (5.88%). Kumpulan umur 20-29 menunjukkan prevalens jangkitan paling tinggi (17.13%). Kumpulan umur kurang daripada 10 tahun menunjuk prevalens paling rendah (5.66%). Umumnya, banyak daripada kes ini berlaku pada kumpulan umur 20 - 25 tahun. Tinjauan serologi berdasarkan asai imunoserap terangkai enzim (ELISA) menunjukkan prevalens jangkitan leptospira keseluruhan tinggi (12.56%). Kuala Lumpur menunjukkan prevalens paling tinggi (19%), manakala Pulau Pinang merekodkan prevalens paling rendah (6.67%). Tiada kelainan tererti wujud di antara negeri lain.

Suatu kajian perbandingan terhadap tiga ujian serologi, iaitu asai imunoerap terangkai enzim (ELISA), ujian pengaglutinatan mikroskopi (MAT), dan ujian penghemagglutinatan tak langsung (IHA) telah dijalankan untuk menilai ujian ini dalam diagnosis leptospirosis manusia. Sejumlah 3000 sampel serum manusia telah diperiksa. Dalam kumpulan I, IgM- dan IgG-ELISA dapat mengesan beberapa kes dalam pensampelan pertama, sebelum titer MAT dapat dikesan. Dalam pensampelan kedua, kesemua sampel yang positif dengan MAT juga positif dengan IgM-ELISA. Ujian IHA memberikan tindak balas positif untuk 38% daripada sampel sahaja, sambil kesemua sampel positif dengan ELISA. Dalam kumpulan II, ELISA mengesan IgM dan IgG terhadap leptospira dalam sampel yang negatif dengan MAT. Ini ialah daripada pesakit yang menunjukkan petanda klinikal untuk leptospira.

Tindak balas berangkai polimerase (PCR) telah dinilai sebagai alat untuk diagnosis leptospirosis dan untuk pembezaan strain leptospira. Sampel urin yang mengandungi hanya 10 *hardjo* sel per ml urin masih positif dalam ujian PCR, menunjukkan tingginya kepekaan ujian tersebut. Pengesan sejumlah sel yang begitu kecil dalam urin melalui PCR memberi ujian ini kelebihan berbanding pengkulturan. Penyidikjarian DNA polimorfus penguatan rawak (RAPD) telah diguna untuk membezakan strain leptospira. Dua primer telah diuji untuk keupayaannya menjana sidikjari RAPD individu. Profil DNA yang diperolehi daripada setiap primer ini adalah nyata dan boleh dihasil semula. Sidikjari yang diperolehi mungkin

berguna untuk membeza serovar sehingga pada aras strain. Profil diperolehi menunjukkan keheterogenan genetik di antara serovar daripada satu serokumpulan.

Analisis DNA leptospira dengan enzim pengehadan, *Hind III*, *BamH I* dan *EcoR I* menunjukkan keheterogenan tinggi di antara serovar yang diperiksa. Keheterogenan tinggi ini mungkin disebabkan oleh genom besar pada genus *Leptospira*. Tiada perkaitan yang ditemui di antara pola pengehadan dan spesies daripada mana pencilan itu diperolehi. Persamaan telah dicerap di kalangan pencilan yang serovarnya sama. Pencilan 10 leptospira liar telah diasaiakan untuk kewujudan DNA plasmid. Hanya dua pencilan telah ditemui mengandungi DNA plasmid. Pemprofilan plasmid yang diperolehi itu mempunyai nilai epidemiologi terhad dalam pembezaan pencilan leptospira.

Nampaknya leptospirosis manusia itu adalah suatu jangkitan endemik di Malaysia. Penemuan ini menunjukkan ELISA itu merupakan ujian serologi yang sesuai untuk diagnosis leptospirosis berbanding ujian MAT dan IHA. Disebaliknya, penggunaan PCR dan REA dalam diagnosis leptospirosis adalah memuaskan.

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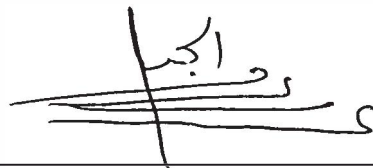
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I hereby declare that the thesis is based on my original work except for quotations and citations, which have been duly acknowledged. I also declare that it has not been previously or currently submitted for any other degree at UPM or other institutions.



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TABLE OF CONTENTS

| | Page |
|---|-------------|
| DEDICATION..... | 2 |
| ABSTRACT..... | 3 |
| ABSTRAK..... | 6 |
| ACKNOWLEDGEMENTS..... | 9 |
| APPROVAL SHEETS..... | 12 |
| DECLARATION FORM..... | 18 |
| LIST OF TABLES..... | 18 |
| LIST OF FIGURES..... | 20 |
| LIST OF ABBREViations..... | 22 |
| CHAPTER | |
| I INTRODUCTION..... | 24 |
| II LITERATURE REVIEW..... | 29 |
| Leptospires..... | 29 |
| Epidemiology of Human Leptospirosis..... | 33 |
| Transmission of Leptospirosis..... | 35 |
| Pathogenesis of Leptospirosis..... | 35 |
| Immunity to Leptospirosis..... | 37 |
| Clinical Signs and Symptoms of Leptospirosis..... | 39 |
| Laboratory Diagnosis of Leptospirosis..... | 41 |
| Microscopic Examination..... | 42 |
| Histological Staining Techniques..... | 43 |
| Culture..... | 44 |
| Microscopic Agglutination Test (MAT)..... | 45 |
| Enzyme-Linked Immunosorbent Assay (ELISA)..... | 46 |
| Indirect Haemagglutination (IHA) Test | 49 |
| Complement Fixation Test (CFT)..... | 51 |
| DNA Based Methods..... | 51 |
| Leptospiral Plasmids..... | 63 |
| Human Leptospirosis in Malaysia..... | 64 |
| Animal Leptospirosis in Malaysia..... | 72 |
| III EPIDEMIOLOGY OF HUMAN LEPTOSPIROSIS IN MALAYSIA..... | 73 |
| Introduction..... | 73 |
| Materials and Methods..... | 76 |
| Retrospective Study..... | 76 |
| Serological Survey..... | 77 |
| Statistical Analysis..... | 83 |
| Results..... | 83 |

| | |
|--|-----|
| Retrospective Study..... | 83 |
| Serological Survey..... | 92 |
| Discussion..... | 95 |
| IV COMPARISION BETWEEN ENZYME-LINKED IMMUNOSORBENT ASSAY (ELISA) , MICROSCOPIC AGGLUTINATION TEST (MAT) AND INDIRECT HAEMAGGLUTINATION (IHA) TEST FOR DIAGNOSIS OF HUMAN LEPTOSPIROSIS..... | 104 |
| Introduction..... | 104 |
| Materials and Methods..... | 108 |
| Human Test Sera..... | 108 |
| Microscopic Agglutination Test (MAT)..... | 110 |
| Enzyme-linked Immunosorbent Assay (ELISA)..... | 111 |
| Indirect Haemagglutination (IHA) Test..... | 112 |
| Cross Reactivity..... | 113 |
| Statistical Analysis..... | 114 |
| Results..... | 114 |
| Group I Samples..... | 114 |
| Group II Samples..... | 120 |
| Group III Samples..... | 122 |
| Overall IgM and IgG-ELISA for the Different Groups..... | 123 |
| ELISA IgM Titres to Leptospires for Group I amples..... | 124 |
| ELISA IgG Titres to Leptospires for Group I Samples..... | 125 |
| Discussion..... | 127 |
| V POLYMERASE CHAIN REACTION FOR DETECTION OF LEPTOSPIRES IN URINE AND DIFFERENTIATION OF LEPTOSPIRAL SEROVARS..... | 136 |
| Introduction..... | 136 |
| Materials and Methods..... | 140 |
| Leptospiral Strains..... | 140 |
| Preparation of Leptospiral DNA..... | 141 |
| Determination of DNA Concentration and Purity..... | 143 |
| Urine Samples..... | 143 |
| Preparation of Urine Samples for Specific PCR..... | 143 |
| Specific PCR Primers..... | 144 |
| RAPD-PCR Primers..... | 145 |
| Specific PCR..... | 146 |
| RAPD-PCR..... | 146 |
| Gel Electrophoresis of Amplification Products..... | 147 |
| Results..... | 147 |
| Specific PCR..... | 147 |
| RAPD-PCR..... | 148 |
| Discussion..... | 160 |

| | | |
|------------|--|-----|
| VI | APPLICATION OF RESTRICTION ENDONUCLEASES ANALYSIS IN THE IDENTIFICATION OF LEPTOSPIRAL ISOLATES | 165 |
| | Introduction..... | 165 |
| | Materials and Methods..... | 167 |
| | Leptospiral Isolates..... | 167 |
| | Preparation of Chromosomal Leptospiral DNA..... | 168 |
| | Determination of DNA Concentration and Purity..... | 168 |
| | DNA Digestion with Restriction Endonucleases..... | 169 |
| | Agarose Gel Electrophoresis and Photography of Digested Products..... | 169 |
| | Molecular Size Estimation of DNA Fragments..... | 170 |
| | Purification of Plasmid DNA..... | 170 |
| | Agarose Gel Electrophoreses of Plasmid DNA..... | 171 |
| | Results..... | 172 |
| | Digestion with Restriction Enzymes..... | 172 |
| | Plasmid DNA..... | 179 |
| | Discussion..... | 184 |
| VII | GENERAL DISCUSSION AND CONCLUSION..... | 188 |
| | BIBLIOGRAPHY..... | 200 |
| | APPENDICES..... | 223 |
| | Appendix A: Buffers Used for ELISA..... | 224 |
| | Appendix B: Preparation of Johnson and Seiter (JS) Medium... | 227 |
| | Appendix C: Reagents Supplied for IHA Kit..... | 231 |
| | Appendix D: Buffers and Reagents for Molecular Biology | 233 |
| | BIODATA Of AUTHOR..... | 237 |

LIST OF TABLES

| Table | | Page |
|--|--|-------------|
| 2.1 Leptospiral serovars that have been isolated from man and animals in Malaysia..... | | 31 |
| 3.1 Prevalence of leptospirosis according to different states (1990-1998)..... | | 85 |
| 3.2 Distribution of positive cases according to races..... | | 87 |
| 3.3 Prevalence of leptospirosis amongst different races..... | | 88 |
| 3.4 Distribution of positive cases of leptospiral infection (1988-1998) according to age groups..... | | 89 |
| 3.5 Clinical manifestations of positive cases of leptospiral infection (1988-1998)..... | | 91 |
| 3.6 Distribution of IgM, IgG and overall ELISA positive samples amongst different states..... | | 93 |
| 3.7 IgM and IgG titres to leptospires for ELISA positive samples..... | | 94 |
| 4.1 Distribution of sera amongst different groups..... | | 109 |
| 4.2 Infecting serovars amongst MAT positive patients of Group I..... | | 115 |
| 4.3 Distribution of MAT titres during first and second samplings..... | | 116 |
| 4.4 IgM-ELISA during first and second samplings for Group I samples..... | | 117 |
| 4.5 IgG-ELISA during first and second samplings for Group I samples..... | | 118 |
| 4.6 Comparison between IgM, IgG-ELISA and IHA Test..... | | 119 |
| 4.7 IgM-ELISA during first and second samplings for Group II samples..... | | 121 |
| 4.8 IgG-ELISA during first and second samplings for Group II samples..... | | 121 |

| | | |
|------|--|-----|
| 4.9 | IgM and IgG-ELISA for Group III samples..... | 122 |
| 4.10 | Overall IgM and IgG-ELISA for Group 1, 11 and 111 samples.... | 123 |
| 4.11 | IgM titres to leptospires for Group I samples during first and second samplings..... | 124 |
| 4.12 | IgG titres to leptospires for Group I samples during first and second samplings..... | 126 |
| 5.1 | Leptospiral reference strains used in the study..... | 140 |
| 5.2 | Leptospiral field isolates used in the study..... | 141 |
| 5.3 | Sequences of 20 oligonucleotides screened for RAPD-PCR analysis..... | 145 |
| 5.4 | Distribution of fingerprinting profiles of reference strains obtained by RAPD-PCR using primers OPA 3 and OPA 20..... | 159 |
| 5.5 | Distribution of fingerprinting profiles of field isolates obtained by RAPD-PCR using primers OPA 3 and OPA 20..... | 159 |
| 6.1 | Leptospiral field isolates and reference strains used in the study | 168 |
| 6.2 | Molecular size of leptospiral DNA samples estimated from the fragments generated by digestion with Hind III restriction endonuclease enzyme..... | 174 |
| 6.3 | Molecular size of leptospiral DNA samples estimated from the fragments generated by digestion with BamH I restriction endonuclease enzyme..... | 176 |
| 6.4 | Molecular size of leptospiral DNA samples estimated from the fragments generated by digestion with EcoR I restriction endonuclease enzyme..... | 178 |

LIST OF FIGURES

| Figure | Page |
|---|-------------|
| 3.1 Epidemiology of leptospirosis..... | 75 |
| 3.2 Prevalence of human leptospirosis in Malaysia (1983- 1998)..... | 84 |
| 3.3 Prevalence of leptospirosis according to different states (1990-1998)..... | 85 |
| 3.4 IgM and IgG titres to leptospires for ELISA positive samples..... | 94 |
| 4.1 Distribution of MAT titres during first and second samplings..... | 116 |
| 4.2 IgM titres to leptospires for Group1 samples during first and second samplings..... | 125 |
| 4.3 IgG titres to leptospires for Group I samples during first and second samplings..... | 126 |
| 5.1 Gel electrophoresis of PCR products obtained from pure genomic DNA of leptospiral field isolates..... | 151 |
| 5.2 Gel electrophoresis of PCR products obtained from urine samples seeded with 10, 20, 50, 100, 200, 400 <i>hardjo</i> cells per ml urine..... | 152 |
| 5.3 RAPD profiles of leptospiral reference strains obtained with primer OPA 3 and electrophoresed on 2% agarose gel..... | 153 |
| 5.4 RAPD profiles of leptospiral field isolates obtained with primer OPA 3 and electrophoresed on 2% agarose gel..... | 154 |
| 5.5 RAPD profiles of leptospiral field isolates obtained with primer OPA 3 and electrophoresed on 2% agarose gel..... | 155 |
| 5.6 RAPD profiles of leptospiral reference strains obtained with primer OPA 20 and electrophoresed on 2% agarose gel..... | 156 |
| 5.7 RAPD profiles of leptospiral reference strains obtained with primer OPA 20 and electrophoresed on 2% agarose gel..... | 157 |
| 5.8 RAPD profiles of leptospiral field isolates obtained with primer OPA 20 and electrophoresed on 2% agarose gel..... | 158 |

| | | |
|-----|---|-----|
| 6.1 | REA profiles of leptospiral field isolates after digestion with Hind III and electrophoresed on 0.7% agarose gel..... | 180 |
| 6.2 | REA profiles of leptospiral field isolates after digestion with BamH I and electrophoresed on 0.7% agarose gel..... | 181 |
| 6.3 | REA profiles of leptospiral field isolates after digestion with EcoR I and electrophoresed on 0.7% agarose gel..... | 182 |
| 6.4 | Agarose gel (0.7%) electrophoresis of plasmid DNA from leptospiral field isolates..... | 183 |

LIST OF ABBREVIATIONS

| | |
|-----------------------------------|---|
| Ab | Antibody |
| ABTS | 2,2-azino-bis (3-ethylbenzthiazoline-6-Sulfonic acid) |
| AP-PCR | Arbitrary primed polymerase chain reaction |
| Bp | Base pair |
| BSA | Bovine serum albumin |
| °C | Degree Celcius |
| CAAT | Cross agglutination absorption test |
| CFT | Complement fixation test |
| CHEF | Contour-clamped homogenous electric field |
| CSF | Cerobrospinal fluid |
| D.D | De-ionized distilled |
| dNTP | Deoxy-nucleotide triphosphate |
| DFM | Dark-field microscopy |
| DNA | Deoxyribonucleic acid |
| Dr. | Doctor |
| EDTA | Ethylene diamine tetra-acetone |
| ELISA | Enzyme-linked Immunosorbent assay |
| FAO | Food and Agriculture Organization |
| FAT | Fluorescent antibody test |
| g | Gram |
| g/l | Gram/litre |
| G+C | Guanine+Cytosine |
| H₂O₂ | Hydrogen peroxide |
| IgA | Immunoglobulin A |
| IgG | Immunoglobulin G |
| IgM | Immunoglobulin M |
| IHA | Indirect haemagglutination |
| IMR | Institute for Medical Research |
| JS | Johnson and Seiter |
| Kbp | Kilobase pairs |
| K L | Kuala Lumpur |
| L | Leptospira |
| LPS | Lipopolysaccharide |
| M | Molar |
| MAT | Microscopic agglutination test |
| Mda | Megadalton |
| mM | Milmole |
| M.S | Molecular size |
| M.W | Molecular weight |
| N.S | Negri Sembilan |
| O.D | Optical density |
| PBS | Phosphate buffer saline |
| PCR | Polymerase chain reaction |

| | |
|---------------|-----------------------------------|
| pH | Hydrogen-ion concentration |
| PLGE | Pulse field gel electrophoresis |
| PUO | Pyrexia of unknown origin |
| RAPD | Random amplified polymorphic DNA |
| REA | Restriction endonuclease analysis |
| RNA | Ribonucleic acid |
| rpm | Round per minute |
| SAT | Slide agglutination test |
| SDS | Sodium dodecyl sulphate |
| U | Unit |
| UK | United Kingdom |
| UM | Universiti Malaya |
| UPM | Universiti Putra Malaysia |
| US | United States |
| USA | United States of America |
| UV | Ultraviolet |
| μg | Microgram |
| μl | Microlitre |
| μM | Micro mole |
| V | Volt |
| VRI | Veterinary Research Institute |
| V/V | Volume/volume |
| WHO | World Health Organization |
| % | Percent |

CHAPTER I

INTRODUCTION

Leptospirosis, also known as “march fever” and “mud fever,” is an important zoonotic disease, with important veterinary and public health impact (Dikken and Kmety, 1978; Gussenhoven *et al.*, 1997; Marcos *et al.*, 1997). Leptospirosis is caused by *Leptospira interrogans*. Based on immunological tests, more than 200 leptospiral serovars have been identified. The serovars could be placed into 23 serogroups (Soltys, 1979; Woodward *et al.*, 1997; Chu *et al.*, 1998). In Malaysia, 38 serovars from 13 serogroups have been known to occur (Bahaman, 1988).

In livestock, the disease causes important economic losses. Although this disease is usually mild and often subclinical, it can lead to great losses due to abortions, stillbirths, infertility, mastitis, weak progeny, decreased milk production, and with certain leptospiral serovars, death (Songer *et al.*, 1983; Thiermann, 1984; Bey and Johnson, 1986).

Leptospiral serovars can infect mammals including man. However, the pathogenicity and clinical manifestations of the disease depend on the animal