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Inclusion body myositis : a nationwide study

Badrising, U.A.

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APPENDIX B

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List of abbreviations

APPENDIX C

LIST OF ABBREVIATIONS

A β	amyloid β
AIDs	autoimmune disorders
CI	confidence interval
CK	creatine kinase
CMAPs	compound muscle action potentials
EM	electron microscopy
ENMC	European Neuromuscular Center
HLA	human leucocyte antigen
IBM	inclusion body myositis
LEMS	Lambert-Eaton myasthenic syndrome
LM	light microscopy
MHC	major histocompatibility complex
MMT	manual muscle testing
MND	motor neuron disease
MRC	British Medical Research Council
MTX	methotrexate
MUAP	motor unit action potential
NMJ	neuromuscular junction
RNS	repetitive nerve stimulation
sCK	serum creatine kinase
SFEMG	single fiber electromyography
QMT	quantitative muscle testing

APPENDIX C

Curriculum Vitae

APPENDIX D

CURRICULUM VITAE

Umesh Arvind Badrising was born on July 10, 1969 in Paramaribo, Suriname. He finished his secondary school at the Rembrandt Scholen Gemeenschap in Leiden in 1987. He then started his medical study at the Leiden University. During his study he tutored courses in anatomy and participated in research towards the relationship between class-specific rheuma factors and age at the Department of Rheumatology and research concerning swallowing dysfunction in Parkinson's disease at the Department of Neurology. He passed his "doctoraal examen" (M.Sc.) in 1992 and obtained his medical degree (M.D.) in 1994. He then started as a resident and from 1995 as a research fellow at the Department of Neurology of the Leiden University Medical Center (Prof. Dr. R.A.C. Roos, Prof. Dr. A.R. Wintzen). Since 2004 he works as a neurologist at the van Weel Bethesda hospital in Dirksland, South Holland.

Umesh Arvind Badrising werd geboren op 10 juli 1969 te Paramaribo, Suriname. In 1987 behaalde hij zijn VWO diploma aan de Rembrandt Scholen Gemeenschap in Leiden. Aansluitend begon hij zijn studie geneeskunde aan de Universiteit Leiden. Gedurende de studie begeleidde hij als student-assistent practica anatomie en werkte hij mee aan onderzoek naar de relatie tussen klasse-specifieke reumafactoren en leeftijd op de afdeling reumatologie en naar slikstoornissen bij de ziekte van Parkinson op de afdeling neurologie. Het doctoraal examen behaalde hij in 1992 en het artsexamen in 1994. Hierna was hij als assistent geneeskundige niet in opleiding (1994) en vanaf 1995 als assistent geneeskundige in opleiding tot klinisch onderzoeker (AGIKO) werkzaam op de afdeling neurologie van het Leids Universitair Medisch Centrum (opleiders Prof. Dr. R.A.C. Roos, Prof. Dr. A.R. Wintzen). Sinds 2004 is hij werkzaam als neuroloog in het van Weel Bethesda ziekenhuis in Dirksland (Z.H.).

APPENDIX D

List of publications

APPENDIX E

LIST OF PUBLICATIONS

U.A. Badrising, M.L.C. Maat-Schieman, J.C. van Houwelingen, P.A. van Doorn, S.G. van Duinen, B.G.M. van Engelen, C.G. Faber, J.E. Hoogendijk, A.E. de Jager, P.J. Koehler, M. de Visser, J.J.G.M. Verschuuren, and A.R. Wintzen. Inclusion body myositis-Clinical features and clinical course of the disease in 64 patients. *J Neurol* 2005; 252:1448-1454.

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A.R. Wintzen, U.A. Badrising, R.A.C. Roos, J. Vielvoye, L. Liauw, and E.K.J. Pauwels. Dysphagia in ambulant patients with Parkinson's disease: common, not dangerous. *Can J Neurol Sci* 1994; 21(1):53-56.

APPENDIX E

A.R. Wintzen, U.A. Badrising, R.A.C. Roos, J. Vielvoye, L. Liauw, and E.K.J. Pauwels. Influence of bolus volume on hyoid movements in normal individuals with Parkinson's disease. *Can J Neurol Sci* 1994;21(1):57-59.

Acknowledgement / Nawoord

APPENDIX F

ACKNOWLEDGEMENT / NAWOORD

The result of this thesis could only be achieved thanks to the cooperation of many persons. I wish to express my extraordinary appreciation and thanks to all patients who have actively contributed to the study and to their supporting partners. And of course I wish to thank the referring neurologists for their indispensable cooperation.

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