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Facioscapulohumeral disease

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Curriculum vitae

The author of this study was born September 16th, 1948 in Wassenaar, The Netherlands. After secondary schooling (Gymnasium B), he started the study of medicine at the State University of Leiden in 1966. The Dutch medical licence examination (arts-examen) was passed in January, 1974. He was introduced to Neurology by his father during several months of residency at the "St. Annadal" Hospital in Maastricht. From July 1974 till July 1975 the author was a rotating intern at Baylor College of Medicine, Houston, Texas, USA, and obtained the Texas Medical Licence in August 1975. Subsequently he fulfilled his military service of which 12 months were spent as resident in Neurology at the Central Military Hospital, "Dr. A. Mathijssen" in Utrecht. The residency in Neurology was continued at the Department of Neurology of the "Wilhelmina Gasthuis" in Amsterdam. In his last year of training he was a resident in Psychiatry at the Psychiatric Hospital "Endegeest" in Oegstgeest. In May 1980 he was licenced to practice Neurology. At present he is a staff-neurologist at the University Hospital in Leiden.

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