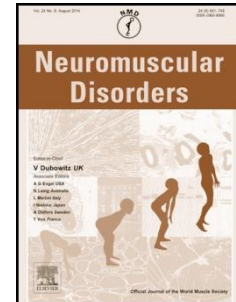


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Author: György Máté Milley, Edina Timea Varga, Zoltán Grosz, Csilla Nemes, Zsuzsanna Arányi, Judit Boczán, Péter Diószeghy, Mária Judit Molnár, Anikó Gál



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Genotypic and phenotypic spectrum of the most common causative genes of Charcot-Marie-Tooth disease in Hungarian patients

György Máté Milley M.D.¹, Edina Timea Varga M.D. Ph.D.^{1,2}, Zoltán Grosz M.D.¹, Csilla Nemes M.Sc. Ph.D.¹, Zsuzsanna Arányi M.D. Ph.D.³, Judit Boczán M.D. Ph.D.⁴, Péter Diószeghy M.D. Ph.D.⁵, Mária Judit Molnár M.D. D.Sc.^{1*} and Anikó Gál M.Sc. Ph.D.^{1*}

1. *Institute of Genomic Medicine and Rare Disorders, Semmelweis University, Budapest, Hungary*

2. *Department of Neurology, University of Szeged, Szeged, Hungary*

3. *MTA-SE NAP B Peripheral Nervous System Research Group, Dept. of Neurology, Semmelweis University, Budapest, Hungary*

4. *Department of Neurology, Medical Center, University of Debrecen, Debrecen, Hungary*

5. *Department of Neurology, Andras Josa Teaching Hospital, Nyiregyhaza, Hungary*

*These authors contributed equally to the manuscript.

Corresponding author:

Maria Judit Molnar M.D., D.Sc.

Institute of Genomic Medicine and Rare Disorders, Semmelweis University

H-1083, Budapest Tömő str. 25-29., Hungary

Tel: +36-20-6632513 Fax: +36-1-459-14-92

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