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Genetic and clinical aspects of inherited syndromes associated with adenomatous polyposis

Ghorban Oghli, Z.

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LIST OF PUBLICATIONS

High yield of surveillance in patients diagnosed with constitutional mismatch repair deficiency.

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Journal of Medical Genetics 2014; 51:283–93.

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CURRICULUM VITAE

Zeinab Ghorban Oghli is born on 18th February 1983 in Tehran, Iran. She completed Grade 11 at Tazkiyeh High school and then moved to South Africa and matriculated at Pretoria High School for Girls in 2000. She then started studying Medicine at the University of Witwatersrand (Johannesburg, South Africa) in 2001 and graduated in December 2006 with Bachelor of Medicine and Surgery (MBBCh). Thereafter, she moved back to Iran and attended 18 months of medical internship at Iran University of Medical Sciences (Tehran, Iran) and was then licensed to practice as a medical doctor by the Islamic Republic of Iran Medical Council.

Zeinab moved to the Netherlands in 2011 when she decided to pursue her education by applying for Master of biomedical science but she was advised to start her PhD. She then started her research in the field of hereditary cancers at Leiden University Medical School (LUMC) and Dutch Foundation for Hereditary Tumours (StOET) under supervision of Prof. Hans Vasen. During her PhD study, she presented the results of her research in international conferences and also the meetings of the European Consortium 'Care for CMMRD' (C4CMMRD). During her PhD study, Zeinab also helped to establish the Middle East Network on Hereditary Colorectal Cancer (<https://www.hccn-me.com>) with the goal to improve care for these high-risk groups in the Middle East and Eastern Mediterranean and North African Countries which also resulted in two publications.

