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Genetic and clinical aspects of inherited syndromes associated with adenomatous polyposis

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Citation

Ghorban Oghli, Z. (2023, May 25). *Genetic and clinical aspects of inherited syndromes associated with adenomatous polyposis*. Retrieved from <https://hdl.handle.net/1887/3618966>

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Note: To cite this publication please use the final published version (if applicable).

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.

ACKNOWLEDGMENTS

Many have contributed in different ways to this thesis. Patients, supervisors, friends and family. This thesis would not have been possible without the help and support of all of these individuals. I am deeply thankful to each and every one of them. Some I would like to thank especially.

First and foremost, I would like to express my deep gratitude to my primary advisor, Prof. dr. Hans Vasen for his guidance and encouragement. Without your unwavering support, this thesis would not have been completed. For this, I am eternally grateful.

Second, I would like to thank my co-advisors Prof. James Hardwick and Dr. Alexandra Langers for their encouragement and support. I would also like to express my gratitude to Dr. Juul Wijnen, for early morning meetings for the SNP project and Prof. Jeanine Houwing-Duistermaat for the complicated statistical discussions. I am also grateful to the members of 'Care for CMMRD' (C4CMMRD) group for their collaboration on the CMMRD projects.

I would also like to thank the ladies at the Stichting Opsporing Erfelijke Tumoren (StOET) for their unprecedented commitment to families with hereditary cancer syndromes. I am especially grateful to Magdalena for helping me with many aspects of living and surviving in the Netherlands. Also especial thanks to Mary and Ingrid for helping with polyposis database.

I am grateful to my friends for their constant encouragement and support, especially during the times when I felt overwhelmed or discouraged. I am particularly thankful to Isaura, a friend and colleague who I could talk and de-stress about PhD life. I am also grateful for my friends Setareh, Elham, Shima, Zary, Zohreh (Zahedi) and Zeinab (Neshati) for our lunch meetings in LUMC. Thank you all for your help and friendship.

I am deeply appreciative to my dearest sister, Zahra, for her support and encouragements. I am also grateful to my dear brother and my sister-in-law. Dear Sajjad and Zahra, you gave me the best gift of all by making me an aunt to

perfect little Ali. It makes me extremely proud to watch him grow into the kind, polite, funny and intelligent person he is becoming.

Lastly, but most importantly, I would like to thank my parents for their love and sacrifice. Their belief in me and their willingness to support me in any way possible have made all the difference. I am forever grateful for your guidance and support throughout my life.

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.

