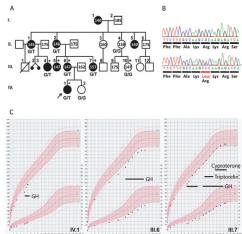


December 2013 Volume 50 Issue 12

JMG
Journal of Medical Genetics



BMJ

jmg.bmj.com

Cover credit: Genealogy and anthropometry of *CDKN1C* mutations causing Russell-Silver Syndrome, from Brioude *et al*, page 823.

Receive regular table of contents by email. Register using this QR code.



Contents

Review

- 785** Criteria and prediction models for mismatch repair gene mutations: a review *A K Win, R J MacInnis, J G Dowty, M A Jenkins*

Genome-wide studies

- 794** Genome-wide association study of sex hormones, gonadotropins and sex hormone-binding protein in Chinese men *Z Chen, S Tao, Y Gao, J Zhang, Y Hu, L Mo, S-T Kim, X Yang, A Tan, H Zhang, X Qin, L Li, Y Wu, S Zhang, S L Zheng, J Xu, Z Mo, J Sun*

Cognitive and behavioural genetics

- 802** Identification of pathogenic gene variants in small families with intellectually disabled siblings by exome sequencing *J H M Schuurs-Hoeijmakers, A T Vulto-van Silfhout, L E L M Vissers, I I G M van de Vondervoort, B W M van Bon, J de Ligt, C Gilissen, J Y Hehir-Kwa, K Neveling, M del Rosario, G Hira, S Reitano, A Vitello, P Failla, D Greco, M Fichera, O Galesi, T Kleefstra, M T Greally, C W Ockelo, M H Willemsen, E M H F Bongers, I M Janssen, R Pfundt, J A Veltman, C Romano, M A Willemsen, H van Bokhoven, H G Brunner, B B A de Vries, A P M de Brouwer*

New loci

- 812** Association analyses identifying two common susceptibility loci shared by psoriasis and systemic lupus erythematosus in the Chinese Han population *Y Li, H Cheng, X-b Zuo, Y-j Sheng, F-s Zhou, X-fa Tang, H-y Tang, J-p Gao, Z Zhang, S-m He, Y-m Lv, K-j Zhu, D-y Hu, B Liang, J Zhu, X-d Zheng, L-d Sun, S Yang, Y Cui, J-j Liu, X-j Zhang*

December 2013 Volume 50 Issue 12

- 819** A novel mutation in *KIAA0196*: identification of a gene involved in Ritscher-Schinzel/3C syndrome in a First Nations cohort *A M Elliott, L R Simard, G Coghlan, A E Chudley, B N Chodirker, C R Greenberg, T Burch, V Ly, G M Hatch, T Zelinski*

Developmental defects

- 823** *CDKN1C* mutation affecting the PCNA-binding domain as a cause of familial Russell Silver syndrome *F Brioude, I Oliver-Petit, A Blaise, F Praz, S Rossignol, M L Jule, N Thibaud, A-M Faussat, M Tauber, Y L Bouc, I Neichine*


Genotype-phenotype correlations


- 831** Genetic variants in *CHI3L1* influencing YKL-40 levels: resequencing 900 individuals and genotyping 9000 individuals from the general population *A D Kjaergaard, J S Johansen, B G Nordestgaard, S E Bojesen*

- 838** A new face of Borjeson-Forsman-Lehmann syndrome? De novo mutations in *PHF6* in seven females with a distinct phenotype *C Zweier, C Kraus, L Brueton, T Cole, F Degenhardt, H Engels, G Gillessen-Kaesbach, L Graul-Neumann, D Horn, J Hoyer, W Just, A Rauch, A Reis, B Wollnik, M Zeschneigle, H-J Lüdecke, D Wiczorek*

Vision science

- 848** A novel heterozygous *OPA3* mutation located in the mitochondrial target sequence results in altered steady-state levels and fragmented mitochondrial network *T Grau, L F Burbulla, G Engl, C Delettre, B Delprat, K Oexle, B Leo-Kottler, T Roscioli, R Krüger, D Rapaport, B Wissinger, S Schimpf-Linzenbold*

 This article has been chosen by the Editor to be of special interest or importance and is freely available online.

 This article has been made freely available online under the BMJ Journals Open Access scheme. See <http://jmg.bmj.com/site/about/guidelines.xhtml#open>

C O P E COMMITTEE ON PUBLICATION ETHICS

This journal is a member of and subscribes to the principles of the Committee on Publication Ethics
www.publicationethics.org.uk

 equator network

 recycle
When you have finished with this please recycle it