


## Contents

### Cancer genetics

- 355**  Diagnostic criteria for constitutional mismatch repair deficiency syndrome: suggestions of the European consortium 'Care for CMMRD' (C4CMMRD) *K Wimmer, C P Kratz, H F A Vasen, O Caron, C Colas, N Entz-Werle, A-M Gerdes, Y Goldberg, D Ilencikova, M Muleris, A Duval, N Lavoine, C Ruiz-Ponte, I Slavic, B Burkhardt, L Brugieres, on behalf of the EU-Consortium Care for CMMRD (C4CMMRD)*

- 366** *RB1* mutation spectrum in a comprehensive nationwide cohort of retinoblastoma patients *C J Dommering, B M Mol, A C Moll, M Burton, J Cloos, J C Dorsman, H Meijers-Heijboer, A H van der Hout*

### Copy-number variation

- 375** Comprehensive genotyping and clinical characterisation reveal 27 novel NKX2-1 mutations and expand the phenotypic spectrum *A Thorwarth, S Schmittner-Hübener, P Schruppf, I Müller, S Jyrch, C Dame, H Biebermann, G Kleinau, J Katchanov, M Schuelke, G Ebert, A Steinger, C Bönemann, K Brockmann, H-J Christen, P Crock, F de Zegher, M Griese, J Hewitt, S Ivarsson, C Hübner, K Kapelari, B Plecko, D Rating, I Stoeva, H-H Ropers, A Grüters, R Ullmann, H Krude*

### New loci

- 388** Mutation in *KANK2*, encoding a sequestering protein for steroid receptor coactivators, causes keratoderma and woolly hair *Y Ramot, V Molho-Pessach, T Meir, R Alper-Pinus, I Siam, S Tams, S Babay, A Zlotogorski*

### Complex traits

- 395** HLA alleles as biomarkers of high-titre neutralising antibodies to interferon- $\beta$  therapy in multiple sclerosis *C Núñez, M<sup>a</sup> C Cénit, R Alvarez-Lafuente, J Río, M Fernández-Arquero, R Arroyo, X Montalbán, O Fernández, B Oliver-Martos, L Leyva, M Comabella, E Urceley*
- 401** A meta-analysis identifies adolescent idiopathic scoliosis association with *LBX1* locus in multiple ethnic groups

*D Londono, I Kou, T A Johnson, S Sharma, Y Ogura, T Tsunoda, A Takahashi, M Matsumoto, J A Herring, T-P Lam, X Wang, E M S Tam, Y-Q Song, Y-H Fan, D Chan, K S E Cheah, X Qiu, H Jiang, D Huang, Japanese Scoliosis Clinical Research Group, TSRHC IS Clinical Group, the International Consortium for Scoliosis Genetics, P Su, P Sham, K M C Cheung, K D K Luk, D Gordon, Y Qiu, J Cheng, N Tang, S Ikegawa, C A Wise*

### Phenotypes

- 407** A familial disorder of altered DNA-methylation *A Caliebe, J Richter, O Ammerpohl, D Kanber, J Beygo, S Bens, A Haake, E Jüttner, B Korn, D J G Mackay, J I Martin-Subero, I Nagel, N J Sebire, L Seidmann, I Vater, C S von Kaisenberg, I K Temple, B Horsthemke, K Buiting, R Siebert*
- 413** Pathogenic mutations in *GLI2* cause a specific phenotype that is distinct from holoprosencephaly *K A Bear, B D Solomon, S Antonini, I J P Arnhold, M M França, E H Gerkes, D K Grange, D W Hadley, J Jääskeläinen, S S Paulo, P Rump, C A Stratakis, E M Thompson, M Willis, T L Winder, A A L Jorge, E Roessler, M Muenke*

### Methods

- 419**  A blinded international study on the reliability of genetic testing for GGGGCC-repeat expansions in *C9orf72* reveals marked differences in results among 14 laboratories *C Akimoto, A E Volk, M van Blitterswijk, M Van den Broeck, C S Leblond, S Lumbroso, W Camu, B Neitzel, O Onodera, W van Rheenen, S Pinto, M Weber, B Smith, M Proven, K Talbot, P Keagle, A Chesi, A Ratti, J van der Zee, H Alstermark, A Birve, D Calini, A Nordin, D C Tradowsky, W Just, H Daoud, S Angerbauer, M DeJesus-Hernandez, T Konno, A Lloyd-Jani, M de Carvalho, K Mouzat, J E Landers, J H Veldink, V Silani, A D Gitler, C E Shaw, G A Rouleau, L H van den Berg, C Van Broeckhoven, R Rademakers, P M Andersen, C Kubisch*

### PostScript

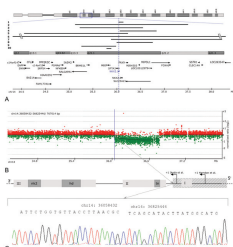
- 425** Correspondence

### Book review

- 428** The PKU Paradox: A Short History of a Genetic Disease *H L Levy*

June 2014 Volume 51 Issue 6

**JMG**  
Journal of Medical Genetics




BMJ


[jmg.bmj.com](http://jmg.bmj.com)

Cover credit: Copy-number variants involving NKX2-1, from Thorwarth *et al.* pg 375.

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



 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  
<http://publicationethics.org/>

 equator network

 recycle  
When you have finished with this please recycle it