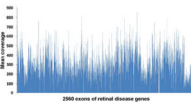
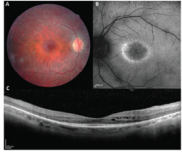


# Journal of Medical Genetics

October 2013 Volume 50 Issue 10

**JMG**  
Journal of Medical Genetics



BMJ

jmg.bmj.com

Cover credit: Exon coverage for NGS diagnosis of retinal disorders from Wang *et al*, p 674 in this issue.

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



## Contents

### Reviews

- 641** Digenic inheritance in medical genetics  
*A A Schäffer*
- 653** An overview on molecular biology of KIT/PDGFR $\alpha$  wild type (WT) gastrointestinal stromal tumours (GIST) *M Nannini, G Biasco, A Astolfi, M A Pantaleo*

### Developmental defects

- 662** Bicuspid aortic valve and aortic coarctation are linked to deletion of the X chromosome short arm in Turner syndrome *C Bondy, V K Bakalov, C Cheng, L Olivieri, D R Rosing, A E Arai*

### Complex traits

- 666** Large-scale genotyping identifies a new locus at 22q13.2 associated with female breast size  
*J Li, J N Foo, N Schoof, J S Varghese, P Fernandez-Navarro, G L Gierach, S T Quek, M Hartman, S Nord, V N Kristensen, M Pollán, J D Figueroa, D J Thompson, Y Li, C C Khor, K Humphreys, J Liu, K Czene, P Hall*

### Genotype-phenotype correlations

- 674** Comprehensive molecular diagnosis of 179 Leber congenital amaurosis and juvenile retinitis pigmentosa patients by targeted next generation sequencing *X Wang, H Wang, V Sun, H-F Tuan, V Keser, K Wang, H Ren, I Lopez, J E Zaneveld, S Siddiqui, S Bowles, A Khan, J Salvo, S G Jacobson, A Iannaccone, F Wang, D Birch, J R Heckentively, G A Fishman, E I Traboulsi, Y Li, D Wheaton, R K Koenekoop, R Chen*

## October 2013 Volume 50 Issue 10

### Cancer genetics


- 689** Confirmation of papillary thyroid cancer susceptibility loci identified by genome-wide association studies of chromosomes 14q13, 9q22, 2q35 and 8p12 in a Chinese population *Y-L Wang, S-H Feng, S-C Guo, W-J Wei, D-S Li, Y Wang, X Wang, Z-Y Wang, Y-Y Ma, L Jin, Q-H Ji, J-C Wang*


### Copy-number variation

- 696** CAG size-specific risk estimates for intermediate allele repeat instability in Huntington disease *A Semaka, C Kay, C Doty, J A Collins, E K Bijlsma, F Richards, Y P Goldberg, M R Hayden*

### Mutation report

- 704** Prevalence of rare mitochondrial DNA mutations in mitochondrial disorders  
*S Bannwarth, V Procaccio, A S Lebre, C Jardel, A Chausseot, C Hoarau, H Maoulida, N Charrier, X Gai, H M Xie, M Ferre, K Fragaki, G Hardy, B M de Camaret, S Marlin, C M Dhaenens, A Slama, C Rocher, J P Bonnefont, A Röitig, N Aoutil, M Gilleron, V Desquiret-Dumas, P Reynier, J Ceresuela, L Jonard, A Devos, C Espil-Taris, D Martinez, P Gaignard, K-H L Q Sang, P Amati-Bonneau, M J Falk, C Florentz, B Chabrol, I Durand-Zaleski, V Paquis-Fluckelinger*

 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](http://www.publicationethics.org.uk)

 equator network

 recycle  
When you have finished with this please recycle it