


Contents



Review

- 81**  Guidelines for the diagnosis and management of individuals with neurofibromatosis 1 R E Ferner, S M Huson, N Thomas, C Moss, H Willshaw, D G Evans, M Upadhyaya, R Towers, M Gleeson, C Steiger, A Kirby

Original articles



- 89** A molecular and clinical study of Larsen syndrome caused by mutations in *FLNB* L S Bicknell, C Farrington-Rock, Y Shafeghati, P Rump, Y Alanay, Y Alembik, N Al-Madani, H Firth, M H Karimi-Nejad, C A Kim, K Leask, M Maisenbacher, E Moran, J G Pappas, P Prontera, T de Ravel, J-P Fryns, E Sweeney, A Fryer, S Unger, L C Wilson, R S Lachman, D L Rimoin, D H Cohn, D Krakow, S P Robertson
- 99** Features associated with germline *CDKN2A* mutations: a GenoMEL study of melanoma-prone families from three continents A M Goldstein, M Chan, M Harland, N K Hayward, F Demenais, D T Bishop, E Azizi, W Bergman, G Bianchi-Scarra, W Bruno, D Calista, L A Cannon Albright, V Chaudru, A Chompret, F Cuellar, D E Elder, P Ghiorzo, E M Gillanders, N A Gruis, J Hansson, D Hogg, E A Holland, P A Kanetsky, R F Kefford, M T Landi, J Lang, S A Leachman, R M MacKie, V Magnusson, G J Mann, J Newton Bishop, J M Palmer, S Puig, J A Puig-Butlle, M Stark, H Tsao, M A Tucker, L Whitaker, E Yakobson, The Lund Melanoma Study Group, and the Melanoma Genetics Consortium (GenoMEL)
- 107** Evaluation of *BRCA1* and *BRCA2* mutation prevalence, risk prediction models and a multistep testing approach in French-Canadian families with high risk of breast and ovarian cancer J Simard, M Dumont, A-M Moisan, V Gaborieau, H Vézina, F Durocher, J Chiquette, M Plante, D Avar, P Bessette, C Brousseau, M Dorval, B Godard, L Houde, Y Joly, M-A Lajoie, G Leblanc, J Lépine, B Lespérance, H Malouin, J Parboosingh, R Pichette, L Provencher, J Rhéaume, D Sinnett, C Samson, J-C Simard, M Tranchant, P Voyer, INHERIT BRCAs, D Easton, S V Tavtigian, B-M Knoppers, R Laframboise, P Bridge, D Goldgar

- 122** Schimke immuno-osseous dysplasia: a clinicopathological correlation J M Clewing, B C Antalfy, T Lücke, B Najafian, K M Marwedel, A Hori, R M Powel, A F S Do, L Najera, K SantaCruz, M J Hicks, D L Armstrong, C F Boerkoel

Short report

- 131** Expansion of the genotypic and phenotypic spectrum in patients with *KRAS* germline mutations M Zenker, K Lehmann, A L Schulz, H Barth, D Hansmann, R Koenig, R Korinthenberg, M Kreiss-Nachtsheim, P Meinecke, S Morlot, S Mundlos, A S Quante, S Raskin, D Schnabel, L-E Wehner, C P Kratz, D Horn, K Kutsche

Letters to JMG

- 136** An atypical deletion of the Williams-Beuren syndrome interval implicates genes associated with defective visuospatial processing and autism L Edelmann, A Prosnitz, S Pardo, J Bhatt, N Cohen, T Lauriat, L Ouchanov, P J González, E R Manghi, P Bondy, M Esquivel, S Monge, M F Delgado, A Splendore, U Francke, B K Burton, L A McInnes
- 144** Methylation analysis of *KvDMR1* in human oocytes E Geuns, P Hilven, A Van Steirteghem, I Liebaers, M De Rycke
- 148**  Correlation between clinical severity in patients with Rett syndrome with a p.R168X or p.T158M *MECP2* mutation, and the direction and degree of skewing of X-chromosome inactivation H Archer, J Evans, H Leonard, L Colvin, D Ravine, J Christodoulou, S Williamson, T Charman, M E S Bailey, J Sampson, N de Klerk, A Clarke
- 153**  Development of a genotyping microarray for Usher syndrome F P M Cremers, W J Kimberling, M Külm, A P de Brouwer, E van Wijk, H te Brinke, C W R J Cremers, L H Hoefsloot, S Banfi, F Simonelli, J C Fleischhauer, W Berger, P M Kelley, E Haralambous, Z Bitner-Glindzicz, A R Webster, Z Saihan, E De Baere, B P Leroy, G Silvestri, G J McKay, R K Koenekoop, J M Millan, T Rosenberg, T Joensuu, E-M Sankila, D Weil, M D Weston, B Wissinger, H Kremer

JMG Unlocked

Articles carrying the



logo are available free via the journal's website: www.jmedgenet.com. Funding for this open access experiment is provided by the UK's Joint Information Systems Committee (JISC); see <http://jmg.bmj.com/cgi/content/full/42/2/97> for further information.

contd...

Contents ...contd

Miscellaneous

147 Correction

160 Correction

Electronic letters

e65 A novel X-linked recessive form of Mendelian susceptibility to mycobacterial disease *J Bustamante, C Picard, C Fieschi, O Filipe-Santos, J Feinberg, C Perronne, A Chapgier, L de Beaucoudrey, G Vogt, D Sanlaville, A Lemainque, J-F Emile, L Abel, J-L Casanova*

e66 Genetic variants in brain-derived neurotrophic factor associated with Alzheimer's disease *R Huang, J Huang, H Cathcart, S Smith, S E Poduslo*

Online mutation report

e67 A truncation in the RYR1 gene associated with central core lesions in skeletal muscle fibres *D Rossi, P De Smet, A Lyfenko, L Galli, S Lorenzini, D Franci, F Petrioli, A Orrico, C Angelini, V Tegazzin, R Dirksen, V Sorrentino*