

Contents

Review

- 361** Hereditary diffuse gastric cancer: updated clinical guidelines with an emphasis on germline *CDH1* mutation carriers *R S van der Post, I P Vogelaar, F Carneiro, P Guilford, D Huntsman, N Hoogerbrugge, C Caldas, K E C Schreiber, R H Hardwick, M G E M Ausems, L Bardram, P R Benusiglio, T M Bisseling, V Blair, E Bleiker, A Boussioutas, A Cats, D Coit, L DeGregorio, J Figueiredo, J M Ford, E Heijkoop, R Hermens, B Humar, P Kaurah, G Keller, J Lai, M J L Ligtenberg, M O'Donovan, C Oliveira, H Pinheiro, K Ragunath, E Rasenberg, S Richardson, F Roviello, H Schackert, R Seruca, A Taylor, A ter Huurne, M Tischkowitz, S T A Joe, B van Dijk, N C T van Grieken, R van Hillegersberg, J W van Sandick, R Vehof, J H van Krieken, R C Fitzgerald*

Complex traits

- 375** Progressive influence of body mass index-associated genetic markers in rural Gambians *A J Fulford, K K Ong, C E Elks, A M Prentice, B J Hennig*

Developmental defects

- 381** Nonsense mutation in the *WDR73* gene is associated with Galloway-Mowat syndrome *T Ben-Omran, S Fahiminiya, N Sorfazlian, M Almurieki, Z Nawaz, J Nadaf, K A Khadija, S Zaineddin, H Kamel, J Majewski, V Tropepe*
- 391** Minichromosome maintenance complex component 8 (MCM8) gene mutations result in primary gonadal failure *Y Tenenbaum-Rakover, A Weinberg-Shukron, P Renbaum, O Lobel, H Eideh, S Gulsuner, D Dahary, A Abu-Rayyan, M Kanaan, E Levy-Lahad, D Bercovich, D Zangen*

New loci

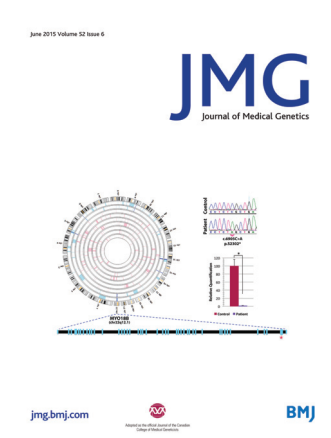
- 400** A novel syndrome of Klippel-Feil anomaly, myopathy, and characteristic facies is linked to a null mutation in *MYO18B* *A M Alazami, A Y Kentab, E Fageih, J Y Mohamed, H Alkhalidi, H Hijazi, F S Alkuraya*

Genotype-phenotype correlations

- 405** Familial periventricular nodular heterotopia, epilepsy and Melnick-Needles Syndrome caused by a single *FLNA* mutation with combined gain-of-function and loss-of-function effects *E Parrini, D Mei, M A Pisanti, S Catarzi, D Pucatti, C Bianchini, M Mascalchi, E Bertini, A Morrone, M L Cavaliere, R Guerrini*
- 413** Rare variants in *SOS2* and *LZTR1* are associated with Noonan syndrome *G L Yamamoto, M Aguen, M Gos, C Hung, J Pilch, S Fahiminiya, A Abramowicz, I Cristian, M Buscarilli, M S Naslavsky, A C Malaquias, M Zatz, O Bodamer, J Majewski, A A L Jorge, A C Pereira, C A Kim, M R Passos-Bueno, D R Bertola*

Cancer genetics

- 422** Bilateral vestibular schwannomas in older patients: NF2 or chance? *D G Evans, S Freeman, C Gokhale, A Wallace, S K Lloyd, P Axon, C L Ward, S Rutherford, A King, S M Huson, R T Ramsden, on Behalf of the Manchester NF2 service*
- 426** A germline mutation in *PBRM1* predisposes to renal cell carcinoma *P R Benusiglio, S Couvé, B Gilbert-Dussardier, S Deveau, H Le Jeune, M Da Costa, G Fromont, F Memeteau, M Yacoub, I Coupier, D Leroux, A Méjean, B Escudier, S Giraud, A-P Gimenez-Roqueplo, C Blondel, E Frouin, B T Teh, S Ferlicot, B Bressac-de Paillerets, S Richard, S Gad*



Cover credit: Homozygosity mapping of the disease locus to *MYO18B* from Alazami *et al*, pg 400.



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