


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- 289** Increased prevalence of imprinting defects in patients with Angelman syndrome born to subfertile couples *M Ludwig, A Katalinic, S Groß, A Sutcliffe, R Varon, B Horsthemke*


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- 292** Distinct *CDH3* mutations cause ectodermal dysplasia, ectrodactyly, macular dystrophy (EEM syndrome) *K W Kjaer, L Hansen, G C Schwabe, A P Marques-de-Faria, H Eiberg, S Mundlos, N Tommerup, T Rosenberg*
- 299** Disruption of the gene *Euchromatin Histone Methyl Transferase 1 (Eu-HMTase1)* is associated with the 9q34 subtelomeric deletion syndrome *T Kleefstra, M Smidt, M J G Banning, A R Oudakker, H Van Esch, A P M de Brouwer, W Nillesen, E A Sistermans, B C J Hamel, D de Bruijn, J-P Fryns, H G Yntema, H G Brunner, B B A de Vries, H van Bokhoven*
- 307**  Multiple mechanisms are implicated in the generation of 5q35 microdeletions in Sotos syndrome *K Tatton-Brown, J Douglas, K Coleman, G Baujat, K Chandler, A Clarke, A Collins, S Davies, F Faravelli, H Firth, C Garrett, H Hughes, B Kerr, J Liebelt, W Reardon, G B Schaefer, M Splitt, I K Temple, D Waggoner, D D Weaver, L Wilson, T Cole, V Cormier-Daire, A Irrthum, N Rahman on behalf of the Childhood Overgrowth Collaboration*

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- 318** Subset of individuals with autism spectrum disorders and extreme macrocephaly associated with germline *PTEN* tumour suppressor gene mutations *M G Butler, M J Dasouki, X-P Zhou, Z Talebizadeh, M Brown, T N Takahashi, J H Miles, C H Wang, R Stratton, R Pilarski, C Eng*

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- 328** Deletion 9q34.3 syndrome: genotype-phenotype correlations and an extended deletion in a patient with features of Opitz C trigonocephaly *S A Yatsenko, S W Cheung, D A Scott, M J M Nowaczyk, M Tarnopolsky, S Naidu, G Bibat, A Patel, J G Leroy, F Scaglia, P Stankiewicz, J R Lupski*
- 336** Comprehensive genomic analysis of *PKHD1* mutations in ARPKD cohorts *A M Sharp, L M Messiaen, G Page, C Antignac, M-C Gubler, L F Onuchic, S Somlo, G G Germino, L M Guay-Woodford*
- 350** *DHCR7* nonsense mutations and characterisation of mRNA nonsense mediated decay in Smith-Lemli-Opitz syndrome *L S Correa-Cerro, C A Wassif, J S Wayne, P A Krakowiak, D Cozma, N R Dobson, S W Levin, G Anadiotis, R D Steiner, M Krajewska-Walasek, M J M Nowaczyk, F D Porter*
- 358** Genetics of Charcot-Marie-Tooth disease type 4A: mutations, inheritance, phenotypic variability, and founder effect *R Claramunt, L Pedrola, T Sevilla, A López de Munain, J Berciano, A Cuesta, B Sánchez-Navarro, J M Millán, G M Saifi, J R Lupski, J J Vilchez, C Espinós, F Palau*
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### Online mutation reports

**e22** No *MSH6* germline mutations in breast cancer families with colorectal and/or endometrial cancer *P Vahteristo, S Ojala, A Tamminen, J Tommiska, H Sammalkorpi, S Kiuru-Kuhlefelt, H Eerola, L A Aaltonen, K Aittomäki, H Nevanlinna*

**e23** High frequency of *APOB* gene mutations causing familial hypobetalipoproteinaemia in patients of Dutch and Spanish descent *S W Fouchier, R R Sankatsing, J Peter, S Castillo, M Pocovi, R Alonso, J J P Kastelein, J C Defesche*