ONLINE MUTATION REPORT

A novel mutation in the Connexin 46 gene causes autosomal dominant congenital cataract with incomplete penetrance

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ongenital or paediatric cataract is a phenotypically and genetically heterogeneous disorder consisting of lens opacities in early life. Thirteen genes have been described for autosomal dominant congenital cataract (ADCC). These include genes for seven members of the crystallin family,^{1 2} which are responsible for the refractive index and transparency of the lens, two connexin genes^{3 4} and major intrinsic protein of the lens (MIP)⁵ which are involved in the transport directly between cells of small metabolites and water, respectively, the cytoskeletal protein beaded filament structural protein-2 (BFSP2),⁶ and transcription factors paired-like homeodomain transcription factor-3 (PITX3)⁷ and heat shock factor-4 (HSF4).⁸ Five additional loci have been described on chromosomes 1pter-p36.1,⁹ 15q21q22,¹⁰ 17p13,¹¹ 17q24,¹² and 20p12-q12.¹³

We used a linkage approach to investigate these 13 genes and five loci in a large pedigree from Victoria, Australia, with zonular pulverulent cataract with the aim of identifying the causative mutation.

METHODS

Ethics approval for this study was obtained from the Human Research Ethics Committees of the Royal Children's Hospital, Melbourne, Australia, the Royal Victorian Eye and Ear Hospital, Melbourne, Australia, and the University of Tasmania, Hobart, Australia.

Patient ascertainment and collection of genetic material

The pedigree crch13 was identified through a database maintained by the Royal Children's Hospital, Melbourne, Australia and the Royal Victorian Eye and Ear Hospital, Melbourne, comprising paediatric cataract patients from south-eastern Australia with any type of lens opacity.14 Written informed consent was obtained from all participating individuals or their guardians. When possible, family members were examined by one or more ophthalmologists (MGW, DAM, JEE, JEC, or IR-E). Due to the rural location of most family members, affection status was determined from medical records when direct examination was not feasible. In many cases pre-operative visual acuity was not available. Buccal mucosal swabs were either collected during examination or by mailed kits and DNA extracted using the PureGene DNA Isolation Kit (Gentra Systems). Unaffected controls were ascertained from nursing homes for the elderly in Launceston, Tasmania, Australia, and were found to be free of ophthalmic disorders, including any form of cataract. Blood was collected from control individuals and DNA extracted with the Nucleon BACC3 kit (Amersham Pharmacia Biotech).

Linkage analysis

All individuals were genotyped at microsatellite markers representing known cataract genes and loci by the analysis of

Key points

- Congenital or paediatric cataract is a highly heterogeneous disorder with 13 known genes and at least five additional loci identified.
- A linkage approach was used to investigate a large pedigree from south-eastern Australia with a faint lamellar opacity surrounding a nuclear pulverulent cataract, with occasional fine needle-like cortical riders.
- Linkage to the Connexin46 locus was identified.
- An R76H mutation was identified in all affected individuals and not found in 100 control chromosomes.
- The mutation was also identified in six unaffected carriers, indicating reduced penetrance of the mutation.

fluorescently-tagged PCR products on an ABI PRISM 310 Genetic Analyzer (Applied Biosystems). Two-point LOD scores were calculated using the FASTLINK package.¹⁵ Allele frequencies were obtained from the population control samples. Penetrance was set at 0.95 in heterozygotes and homozygous variants. Disease gene frequency was set to 0.0001.

Sequencing

The coding region of *CX46* was sequenced using three overlapping PCR fragments. Primer sequences were: 1F: 5'-CGGTGTTCATGAGCATTTTC-3', 1R: 5'-GACGTAGGT-CCGCAGCAG-3', 2F: 5'-GCAGGACAATCCCTCGTC-3', 2R: 5'-GGTCAGGGCTAGCAGTTTGA-3', 3F: 5'-TCGGGTTCCC-ACCCTACTAT-3', 3R: 5'-TGCACTTTGGTTTTGGTTTC-3'. PCR products were cycle sequenced with Big Dye Terminator Ready Reaction Mix (Applied Biosystems) and analysed on an ABI PRISM 310 Genetic Analyzer (Applied Biosystems).

Computational methods

ClustalW¹⁶ (http://www.ebi.ac.uk/clustalw/) was used to align CX46 protein sequences from five mammalian species represented in GenBank. The likely structure of the CX46 protein was determined using HMMTOP^{17 18} (http://www. enzim.hu/hmmtop/index.html) and MEMSAT from the PSIPRED server^{19 20} (http://bioinf.cs.ucl.ac.uk/psipred/).

Abbreviations: ADCC, autosomal dominant congenital cataract; BFSP2, beaded filament structural protein-2; HSF4, heat shock factor-4; MIP, major intrinsic protein; PITX3, paired-like homeodomain transcription factor-3

RESULTS

The phenotype in this pedigree is a faint lamellar nuclear opacity surrounding pulverulent nuclear opacities (fig 1), some with fine gold dots or haze and some with needle-like peripheral riders. The median age of diagnosis was 5 years (range 0–73 years), however the 10 patients in the two most recent generations (generations VI and VII) were diagnosed at 6 months to 2 years. Of 42 eyes with cataracts, 20 had not had surgery and have good or minimally (6/9) reduced vision. The median age of surgery was 17–26 years (range 10–67 years). No other systemic or ocular abnormalities, such as nystagmus, strabismus, or iris abnormalities, were noted. One older female diagnosed at age 5 years and operated on at age 67 years, developed pseudophakic glaucoma.

A LOD score of 2.96 was obtained at $\theta = 0.04$ from D13S1236, the marker included to detect linkage to the *CX46* gene. Other candidate loci implicated in ADCC (*CX50, CRYGD/GC, CRYBA1*, 1pter-p36.1, 15q21-q22, 17p13, 17q24) were excluded by linkage analysis (data not shown). An equivocal LOD score of 1.36 at $\theta = 0.08$ from marker D16S496 representing *HSF4* was the only other positive LOD score. Once the significant result at D13S1236 was obtained, additional ADCC loci (*BFSP2, PITX3, CRYAB, MIP, CRYAA, CRYBB2*, and 20p12-q12) were not investigated for this monogenic disorder.

CX46 was sequenced in affected individuals and the variant 226G>A (GenBank reference NM_021954) causing an R76H substitution was identified in all 21 affected individuals but not in 100 control chromosomes by direct sequencing. Six unaffected relatives (IV:4, V:7, V:8, V:9, V:27, and VI:15) were also found to carry the mutation (fig 2). This residue is conserved across species represented in GenBank (fig 3). The residue is predicted by MEMSAT to be at the boundary between extracellular loop 1 and transmembrane loop 2 while HMMTOP predicts that it is within the second transmembrane domain.

DISCUSSION

The investigation of this large Australian cataract pedigree has revealed a novel mutation, R76H, in the *CX46* gene. The R76H mutation is likely to be causative as it segregates with affected status amongst reasonably distant branches of the pedigree with the same phenotype and was not detected in unaffected, unrelated controls. The inheritance in this pedigree is clearly autosomal dominant, although not fully penetrant.

Connexin proteins form hexamers known as connexons in the cell membranes. Connexons in neighbouring cells dock to

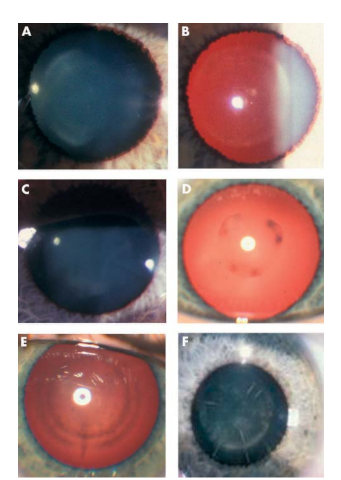


Figure 1 A range of pulverulent cataract phenotypes of pedigree crch13. (A) Mild pulverulent phenotype of the nucleus of individual VI:8, directly illuminated and (B) retroilluminated. (C) Mild pulverulent phenotype of the nucleus with a cortical lamella opacity of individual VII:2, directly illuminated and (D) retroilluminated. (E) Mild pulverulent phenotype of the nucleus with a cortical lamella opacity of individual VII:1, retroilluminated. (F) Pulverulent phenotype in the nucleus with cortical riders of individual VI:3, directly illuminated.

form gap junctions which allow the transport of small metabolites directly between cells.¹ Two connexins, *CX46* and *CX50*, are expressed in lens fibre cells. Previously reported mutations of these genes associated with congenital cataracts

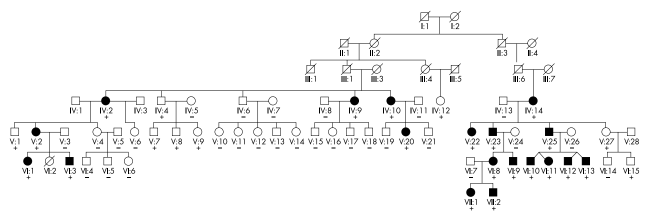


Figure 2 Pedigree diagram of crch13 indicating the presence of Connexin46 R76H mutation. Shaded symbols indicate the presence of an ophthalmologist-confirmed cataract. Squares indicate males, circles females, "+" heterozygote, and "-" wild type. Individuals with no "+" or "-" symbol have not been typed.

Mouse	CENVCYDRAFPISHIRFWALQIIFVSTPTLIYLGHVLHIVRMEEKKKEREEELLRRDNPQ
Rat	CENVCYDRAFPISHIRFWALQIIFVSTPTLIYLGHVLHIVRMEEKKKEREEELLRRDNPQ
Human	CENVCYDRAFPISHIRFWALQIIFVSTPTLIYLGHVLHIVRMEEKKKEREEEEQLK
Cow	CENVCYDRAFPISHVRFWVLQIIFVSTPTLIYLGHVLHLVRMEEKKKEREEE
Danio	CENVCYDEAFPISHIRFWVLQIIFVSTPTLIYLGHVLHIVRMEEKRKEREEELRKASRLQ

Figure 3 Cross species alignment of Connexin46. ClustalW alignment of residues 61–120 of human Cx46 with mouse, rat, cow, and zebra fish (Danio rerio) orthologs. R76 is highlighted in grey.

are shown in table 1. All seven mutations have been linked to zonular pulverulent congenital cataracts which are "pulverised" dust-like or punctate opacities in developmental zones of the lens. The phenotype in the Australian pedigree in the present study resembles those previously reported. All mutations of connexin genes described previously appear fully penetrant in the pedigrees in which they were detected.

The R76 residue of CX46 is conserved between species (fig 3), indicating that the arginine is likely to be functionally important and that the mutation may therefore have a detrimental physiological effect. It is not clear whether R76 is located in the first extracellular loop or in the second transmembrane domain. Other cataract mutations have been detected in both these domains (table 1). Mutations in the extracellular domains may affect connexon docking if the conformation of the loop is changed. It is unclear what the affect of variation within the transmembrane domain would be, however, other mutations of transmembrane domains of connexin genes have been reported. The wild type arginine has a positive charge while histidine can be either positively charged or neutral, depending on the microenvironment. This may help explain the incomplete penetrance observed if cataract formation is dependant on the ionisation of this residue.

A lack of gross effects on protein structure is implied by both the range of ages at which surgery was performed, indicating a phenotype of variable severity, and the incomplete penetrance observed in the pedigree. Individuals V:27 and VI:15 have been examined thoroughly by several of the investigators and are clearly unaffected. Carrier individuals IV:4, V:7, V:8, and V:9 have not been examined by a member of our research group and, therefore, it is possible that they may be subtly affected. The only other report of a monogenic paediatric cataract mutation with incomplete penetrance is a 5 bp insertion in the γ -crystallin gene also causing a variable zonular pulverulent phenotype.²¹ The variability was suggested to be due to environmental factors or modifying genes. Unaffected individuals IV:4 and V:27 have passed on the R76H mutation to their offspring, who also remain unaffected. Environmental factors are unlikely to show this type of pattern, unless there are significant household effects, suggesting the possibility of a second modifying or protective gene.

Investigations of animal models may help elucidate the nature of modifying genes. Mice with a disrupted *cx46* gene

develop a nuclear lens cataract. The cataract phenotype and presence of cleaved γ -crystallin in the lens was variable, depending on the genetic background of the mouse, indicating the presence of modifier genes involved in the development of the phenotype.²² The *Lop10* mutation in mice may also provide some insight. This phenotype is caused by the G22R variant of murine *cx50*,²³ however, the phenotype is variable and dependent on the genetic background of the mouse.²⁴ These examples provide evidence for modifier genes in the development of congenital cataract and support the hypothesis of modifier genes acting in the pedigree described here.

In summary, a novel mutation of the human *CX46* gene has been found to segregate with a pulverulent phenotype. The mutation is only the second reported congenital cataract mutation with incomplete penetrance in the literature and, as such, provides an opportunity for the investigation of modifying genes and their interactions.

ACKNOWLEDGEMENTS

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Conflict of interest: none declared.

Gene	Mutation	Location	Reference
CX46	1137insC	C-terminal cytoplasmic tail	Mackay <i>et al</i> , 1999⁴
	N63S	First extracellular loop	Mackay et al, 1999⁴
	P187L	Second extracellular loop	Rees et al, 2000 ²⁵
	F32L	First transmembrane domain	Jiang et al, 2003 ²⁶
CX50	E48K	First extracellular loop	Berry <i>et al</i> , 1999 ²⁷
	1247M	C-terminal cytoplasmic tail	Polyakov et al, 2001 ²⁸
	P88S	Second transmembrane domain	Shiels et al, 1998 ³

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REFERENCES

- 1 Francis PJ, Berry V, Moore AT, Bhattacharya S. Lens biology: development
- and human cataractogenesis. Trends Genet 1999;15:191–6.
 Berry V, Francis P, Reddy MA, Collyer D, Vithana E, MacKay I, Dawson G, Carey AH, Moore A, Bhattacharya SS, Quinlan RA. Alpha-B crystallin gene (CRYAB) mutation causes dominant congenital posterior polar cataract in humans. Am J Hum Genet 2001;**69**:1141–5.
- 3 Shiels A, Mackay D, Ionides A, Berry V, Moore A, Bhattacharya S. A missense mutation in the human connexin50 gene (GJA8) underlies autosomal dominant "zonular pulverulent" cataract, on chromosome 1q. Am J Hum Genet 1998;**62**:526-32
- 4 Mackay D, Ionides A, Kibar Z, Roulenu G, Berry V, Moore A, Shiels A, Bhattacharya S. Connexin46 mutations in autosomal dominant congenital cataract. Am J Hum Genet 1999;64:1357-64.
- 5 Berry V, Francis P, Kaushal S, Moore A, Bhattacharya S. Missense mutations in MIP underlie autosomal dominant 'polymorphic' and lamellar cataracts linked to 12q. Nat Genet 2000;25:15-7
- 6 Conley TP, Erturk D, Keverline A, Mah TS, Keravala A, Barnes LR, Bruchis A, Hess JF, Fitzgerold PG, Weeks DE, Ferrell RE, Gorin MB. A Juvenile-onset, progressive cataract locus on chromosome 3q21-q22 is associated with a missense mutation in the beaded filament structural protein-2. Am J Hum Genet 2000;66:1426-31.
- 7 Semina EV, Ferrell RE, Mintz-Hittner HA, Bitoun P, Alward WL, Reiter RS, Funkhauser C, Daack-Hirsch S, Murray JC. A novel homeobox gene PITX3 is mutated in families with autosomal-dominant cataracts and ASMD. Nat Genet 1998;19:167-70.
- Bu L, Jin Y, Shi Y, Chu R, Ban A, Eiberg H, Andres L, Jiang H, Zheng G, Qian M, Cui B, Xia Y, Liu J, Hu L, Zhao G, Hayden MR, Kong X. Mutant DNA-Mariner cataract. Nat Genet 2002;31:276–8.
- **Eiberg H**, Lund AM, Warburg M, Rosenberg T. Assignment of congenital cataract Volkmann type (CCV) to chromosome 1p36. *Hum Genet* 1995:96:33-8
- 10 Vanita SJR, Sarhadi VK, Singh D, Reis A, Rueschendorf F, Becker-Follmann J, Jung M, Sperling K. A novel form of "central pouchlike" cataract, with sutural opacities, maps to chromosome 15q21-22. Am J Hum Genet 2001:68:509-14.
- Berry V, Ionides AC, Moore AT, Plant C, Bhattacharya SS, Shiels A. A locus 11 for autosomal dominant anterior polar cataract on chromosome 17p. Hum Mol Genet 1996;5:415-9.

- 12 Armitage MM, Kivlin JD, Ferrell RE. A progressive early onset cataract gene maps to human chromosome 17q24. Nat Genet 1995;9:37-40.
- 13 Yamada K, Tomita H, Yoshiura K, Kondo S, Wakni K, Fukushima Y, Ikegawa S, Nakamura Y, Amemiya T, Niikawa N. An autosomal dominant posterior polar cataract locus maps to human chromosome 20p12-q12. Eur J Hum Genet 2000;8:535-9.
- 14 Wirth MG, Russell-Eggitt IM, Craig JE, Elder JE, Mackey DA. Aetiology of congenital and paediatric cataract in an Australian population. Br J Ophthalmol 2002;**86**:782-6.
- Cottingham RW Jr, Idury RM, Schaffer AA. Faster sequential genetic linkage computations. Am J Hum Genet 1993;53:252–63. 15
- Higgins DG, Sharp PM. CLUSTAL: a package for performing multiple sequence alignment on a microcomputer. Gene 1988;73:237–44. 16
- Tusnady GE, Simon I. The HMMTOP transmembrane topology prediction 17 server. Bioinformatics 2001;17:849-50.
- Tusnady GE, Simon I. Principles governing amino acid composition of integral membrane proteins: application to topology prediction. J Mol Biol 1998;283:489-506.
- Jones DT. Do transmembrane protein superfolds exist? FEBS Lett 19 1998;423:281-5.
- Jones DT, Taylor WR, Thornton JM. A model recognition approach to the prediction of all-helical membrane protein structure and topology. 20 Biochemistry 1994;33:3038-49.
- Ren Z, Li A, Shastry BS, Padma T, Ayyagari R, Scott MH, Parks MM, Kaiser-Kupfer MI, Heitmancik JF. A 5-base insertion in the gammaC-crystallin gene is associated with autosomal dominant variable zonular pulverulent cataract. Hum Genet 2000;**106**:531–7
- 22 Gong X, Agopian K, Kumar NM, Gilula NB. Genetic factors influence cataract formation in alpha 3 connexin knockout mice. Dev Genet 1999;24:27-32.
- 23 Chang B, Wang X, Hawes NL, Ojakian R, Davisson MT, Lo WK, Gong X. A Gja8 (Cx50) point mutation causes an alteration of alpha 3 connexin (Cx46) in semi-dominant cataracts of Lop10 mice. *Hum Mol Genet* 2002;**11**:507–13. **Runge PE**, Hawes NL, Heckenlively JR, Langley SH, Roderick TH. Autosomal
- dominant mouse cataract (Lop-10). Consistent differences of expression in heterozygotes. *Invest Ophthalmol Vis Sci* 1992;**33**:3202–8.
- Rees MI, Watts P, Fenton I, Clarke A, Snell RG, Owen MJ, Gray J. Further evidence of autosomal dominant congenital zonular pulverulent cataracts linked to 13q11 (CZP3) and a novel mutation in connexin 46 (GJA3). Hum Genet 2000;106:206-9.
- 26 Jiang H, Jin Y, Bu L, Zhang W, Liu J, Cui B, Kong X, Hu L. A novel mutation in GJA3 (connexin46) for autosomal dominant congenital nuclear pulverulent cataract. Mol Vis 2003;**9**:579–83.
- Berry V, Mackay D, Khaliq S, Francis PJ, Hameed A, Anwar K, Mehdi SQ, Newbold RJ, Ionides A, Shiels A, Moore T, Bhattacharya SS. Connexin 50 mutation in a family with congenital "zonular nuclear" pulverulent cataract of Pakistani origin. *Hum Genet* 1999;105:168–70. 27
- Polyakov AV, Shagina IA, Khlebnikova OV, Evgrafov OV. Mutation in the connexin 50 gene (GJA8) in a Russian family with zonular pulverulent 28 cataract. Clin Genet 2001;60:476-8.

PostScript

BOOK REVIEW

Hereditary Hearing Loss and its Syndromes, 2nd edn

Edited by H V Toriello, W Reardon, R J Gorlin. Oxford University Press, 2004, £110, pp 502. ISBN 0-19-513849-X

Although formally classified as the second edition, this is essentially the third version of groundbreaking book by the Bruce Konigsmark and Bob Gorlin entitled Genetic and Metabolic Deafness as originally published in 1976. Subsequent recognition of the pressing need to incorporate the rapid expansion in knowledge of new syndromes prompted the appearance of the first editon of Hereditary Hearing Loss and its Syndromes in 1995, as a sister publication to Syndromes of the Head and Neck. This new edition represents the coming of age of the marriage between molecular biology and conventional clinical genetics and provides an excellent state of the art synthesis of contemporary knowledge.

A reviewer's task in making criticism of an outstanding and definitive textbook is not an easy one. In planning this new edition the editors have resisted the temptation to tinker with a successful format to the extent that the overall structure is virtually unchanged. The existing chapter on endocrine and metabolic disorders has been split into two and the miscellaneous chapter has disappeared, to be replaced by a chapter on cardiac syndromes. Otherwise the chapter headings are as in the previous edition with much of their text reproduced, albeit with expanded sections on "heredity" to embrace the many new discoveries of the last few years. Thus the contents can be subdivided into four introductory chapters which set the scene, followed by 12 chapters describing system associated hearing loss syndromes. In general these are excellent, with each providing detailed accounts of an exhaustive list of common and rare conditions in which hearing loss can occur. All these are lavishly illustrated with ample references for those who wish to delve further

Against this background of general excellence any possible hint of criticism might well be viewed as petty and inappropriate, so it is hoped that the editors will forgive a few personal comments. Most readers will be very familiar with the basic principles of human genetics so that on turning to the chapter on genetic counselling it was disappointing to find that this is largely limited to an explanation of traditional patterns of inheritance. The real challenge facing most clinical geneticists and genetic counsellors is how to counsel the hearing parents of a child with isolated non-syndromal hearing loss. Chapter 2 provides useful suggestions for investigation but the subsequent chapter on genetic counselling provides little in the way of assistance. True, there is a useful table (of unstated source) providing empirical risks, but with little in the way of guidance as to how these should be applied. Should these risks be modified on the basis of age of onset, laterality, asymmetry, progression, vestibular involvement, audiology or a normal Connexin 26/30 mutation analysis? Presumably they should, but how? The editors and chapter authors embrace most of the world's experts on genetic hearing loss and it is a little unfortunate that they could not expand on this crucial component of the counselling process. An overview of how genes and their products interact to facilitate the hearing process would also be useful, as would expansion of some of the sections on molecular pathogenesis in the system orientated chapters. Finally, the era when medical books can include full frontal nude photographs of children and adults must be coming to a close and one wonders how many of the stark naked adults appearing in some of the syndrome chapters gave informed consent for their publication in perpetuity.

Clearly these are minor criticisms of an excellent textbook which will provide an invaluable resource and be consulted widely. It is difficult to see how any department encountering patients with hearing loss could possibly manage without it.

I Young

CORRECTIONS

doi: 10.1136/jmg.2005.18333corr1

An error has been detected in the online mutation report by Burdon *et al* (*J Med Genet* 2004;**41**:e106). The mutation is identified in the manuscript as 226G>A in regards to the Genbank reference NM_021954. However, it should be 227G>A. The amino acid designation, R76H, is correct and this numbering error does not change any of the other results or conclusions of the article. The author apologises for this error.

doi: 10.1136/jmg.2004.013151corr1

Several errors have been detected in the electronic letter by Toyama *et al* (*J Med Genet* 2004;**41**:e74).

First, the abbreviations for Table 1 should read:

Ex, exon; (FAM)-,FAM-labelled; (HEX)-, HEX-labelled; (ROX)-,ROX-labelled; (NED)-, NED-labelled; UP, upstream; Pro, prometer; Int, intron; Fl, flanking; STR, short tandem repeat.

Second, the parenthesised section of the last sentence of the Results should read:

(7.3 \pm 1.3 mmol/l (K287I) and 7.63 \pm 1.0 mmol/l (M310I) compared to that of the wild type (3.8 \pm 0.4 mmol/l).

In addition, in Ex4 of Table 3 the "Type" should read C235 (R79W), in Figure 1 "Euro. Am" is the abbreviation for "European American," and in Table 4 the title should read "Catalytic activity of recombinant AMPD1 expressed in *E. coli*".

We apologise for these errors.

doi: 10.1136/jmg.2004.019190corr1

The authors for the paper titled Positive association of the DIO2 (deiodianase type 2) gene with mental retardation in the iodine-deficient areas of china (*J Med Genet* 2004:**41**;585–590) have identified an error within their abstract. The second line from the results section should read: Particularly with rs255012, CC genotype frequency was significantly higher in MR cases than in controls (chi squared = 9.18, p = 0.00246). The author apologises for this mistake.