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1	Clinical and	molecular	genetic	finding	gs in a	autosomal	don	ninant OPA	43-related	optic
2	neuropathy									
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- 27
- 28 Word count: 17<u>53</u>08
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- 30

31 ABSTRACT

32

33 Leber hereditary optic neuropathy and autosomal dominant optic atrophy are the two most 34 common inherited optic neuropathies. The latter has been associated with mutations in the 35 OPA1 and OPA3 genes. To date, only six families with OPA3-associated dominant optic 36 atrophy have been reported. In order to identify additional families we performed Sanger 37 sequencing of the OPA3 gene in 75 unrelated optic neuropathy patients. Affected individuals 38 from two families were found to harbour the c.313C>G, p.(Gln105Glu) change in 39 heterozygous state; this genetic defect has been previously reported in four dominant optic 40 atrophy families. Intra- and inter-familial variability in age of onset and presenting symptoms 41 was observed. Although dominant OPA3 mutations are typically associated with optic 42 atrophy and cataracts, the former can be observed in isolation; we report a case with no lens 43 opacities at age 38. Conversely, it is important to consider OPA3-related disease in 44 individuals with bilateral infantile onset cataracts and to assess optic nerve health in those 45 whose vision fail to improve following lens surgery. The papillomacular bundle is primarily 46 affected and vision is typically worse than 20/40. Notably, we describe one subject who 47 retained normal acuities into the fifth decade of life. The condition can be associated with 48 extraocular clinical features: two affected individuals in the present study had sensorineural 49 hearing loss. The clinical heterogeneity observed in the individuals reported here (all having 50 the same genetic defect in OPA3) suggests that the molecular pathology of the disorder is 51 likely to be complex.

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55 KEYWORDS

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57 OPA3; optic atrophy; inherited optic neuropathy; 3-methylglutaconic aciduria type III; 58 congenital cataract; genetic ophthalmology

60 INTRODUCTION

61

62 Inherited optic neuropathies are a clinically and genetically heterogenous group of disorders 63 associated with selective loss of retinal ganglion cells. Clinically, they are characterised by 64 colour vision deficits, visual field defects and, typically, bilateral, symmetrical and 65 irreversible visual loss [1]. Mitochondrial dysfunction appears to play a central role in the 66 pathophysiology of these disorders and inheritance can be mitochondrial [MIM #535000] or 67 monogenic; autosomal dominant (for example [MIM #165500] and [MIM #165300]), 68 autosomal recessive (for example [MIM #612989]) and X-linked [MIM %311050] subtypes 69 have been described. Notably, certain types of inherited optic neuropathy form part of clinical 70 syndromes that include additional ocular or non-ocular features [2].

71

72 Defects in the OPA3 gene [MIM *606580] have been previously associated with both 73 recessive and dominant optic neuropathy. Biallelic OPA3 mutations cause 3-74 methylglutaconic aciduria type III [MIM #258501], a recessive neuro-ophthalmological 75 syndrome, most prevalent amongst individuals of Iraqi-Jewish origin and classically 76 characterized by the following triad: (i) bilateral optic atrophy diagnosed in the first decade of 77 life; (ii) a movement disorder (ataxia or extrapyramidal dysfunction) of variable severity 78 beginning in the first or second decade of life; (iii) increased urinary excretion of 3-79 methylglutaconic acid [3-8]. Disease-causing OPA3 variants inherited in an autosomal 80 dominant fashion can also cause optic neuropathy. This is often associated with 81 congenital/infantile lenticular opacities; hearing loss and neurological symptoms can also be 82 features of the disorder. Autosomal dominant OPA3-related disease is less common than the 83 recessive form and only six families have been identified to date. The following dominant 84 disease-associated variants have been reported [NCBI Reference Sequence: NM_025136.3]: 85 c.277G>A, p.(Gly93Ser); c.313C>G, p.(Gln105Glu) (recurrent mutation); 86 c.10_11insCGCCCG, p.(Val3_Gly4insAlaPro) [9,10].

87

The OPA3 gene is composed of at least 3 exons that are alternatively spliced to produce two major transcripts: *OPA3A* (exon 1 plus exon 2a; encodes the 179 amino acid isoform b; NCBI Reference Sequence: NM_025136.3) and *OPA3B* (exon 1 plus exon2b; encodes the 180 amino acid isoform a; NCBI Reference Sequence: NM_001017989.2). Although cDNA studies indicate ubiquitous expression of both transcripts, *OPA3A* is much more strongly expressed in most tissues, including the brain [11]. Notably, the OPA3A amino acid sequence 94 appears to be more conserved in evolution and yet no human disease has been associated with 95 mutations in the *OPA3B*-specific exon 2b [7,11]. At the subcellular level, OPA3 localises 96 predominantly to the mitochondrial inner membrane and although its function remains 97 unclear previous studies have suggested involvement in the regulation of mitochondrial 98 morphology [10,12].

99

100 In the present study we report clinical and genetic findings in two families with dominant 101 *OPA3*-related optic neuropathy. The phenotypic spectrum of the disorder is broadened and 102 intra- and inter-familial variability is discussed.

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- 104

105 MATERIALS & METHODS

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107 Initially, 74 unrelated individuals with a presumed diagnosis of inherited optic neuropathy 108 (mean age 43 years; range 24 to 66 years) were tested for OPA3 mutations using Sanger 109 sequencing of the three exons and flanking intron-exon boundaries of the OPA3 gene 110 (primers and conditions available on request). All these subjects were previously screened 111 and excluded for: (i) defects in the OPA1 gene ([MIM *605290]; the major cause of 112 dominant optic neuropathy [2,13]) and (ii) the three primary mitochondrial mutations 113 associated with Leber hereditary optic neuropathy ([MIM #535000]; m.11778G>A, m.14484T>C and m.3460G>A). Subsequently, members of an additional family with 114 115 suspected dominant optic atrophy and cataracts were recruited and tested for OPA3 mutations in a similar fashion. 116

117

118 Clinical assessment of individuals with OPA3-related disease included detailed history, best 119 corrected Logarithm of the Minimum Angle of Resolution (logMAR) visual acuity, dilated 120 fundus examination and optic disc imaging. Optical coherence tomographs of the optic nerve 121 head were obtained with the spectral-domain Cirrus platform (Carl Zeiss Meditec, Dublin, 122 CA, USA) in two cases. An audiogram was performed in three patients. Informed consent 123 was obtained from all participants and all investigations were conducted in accordance with 124 the principles of the Declaration of Helsinki. Institutional Review Board (IRB)/Ethics 125 Committee approval was obtained from the Multicentre Research Ethics Committee 126 (MREC).

128

129

130 **RESULTS**

131

132 Three *OPA3* coding variants that have not been previously reported in publicly available 133 databases (1000 genomes project database, National Heart, Lung, and Blood Institute Exome 134 Sequencing Project or NHLBI ESP, dbSNP Build 139, accessed Jun 2014) were identified in 135 the 180 amino acid isoform a (OPA3B; [NCBI Reference Sequence: NM_001017989.2]: 136 c.227C>T, p.(Ala76Val); c.379G>A, p.(Gly127Ser) and c.389G>A, p.(Gly130Glu) 137 (Supplementary Table S1). Each of these changes was detected in the heterozygous state in a 138 simplex sporadic case and it was not possible to perform segregation analysis to support their 139 pathogenicity. Also, none of these alters the amino acid sequence of isoform b (OPA3A) and 140 \underline{w} we therefore, consider them to be variants of unknown significance.

141

142 In addition to these variants, a heterozygous c.313C>G, p.(Gln105Glu) change (only 143 affecting OPA3A, the transcript encoding the 179 amino acid isoform b; [NCBI Reference 144 Sequence: NM_025136.3]) was identified in an individual with a diagnosis of dominant optic 145 atrophy (subject C1; Figure 1A). This sequence alteration is the most common OPA3 146 mutation associated with dominant disease as it was previously identified in four families 147 segregating optic atrophy [9,10]; therefore no further evidence was needed to confirm that 148 this is a functional variant. The proband as well as his affected sister (subject C2) and 149 daughter (subject C3) presented in the first years of life with nystagmus and abnormal optic 150 disc appearance. The clinical findings are detailed in Table 1.

151

The same disease-associated variant, c.313C>G, p.(Gln105Glu), was detected in all three affected members of a family (Figure 1B) that were recruited and tested for *OPA3* mutations at a later time. Colour disc images and clinical findings are presented in Figure 1C and Table 1 respectively.

156

157

158 **DISCUSSION**

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160 Dominant *OPA3*-related disease is a clinically heterogeneous disorder (Table 1); some 161 patients present with poor visual behaviour and nystagmus from birth (for example subject 162 C1) while others remain asymptomatic until later in life (for example subject M2). The optic 163 neuropathy is characterised by primary involvement of the papillomacular bundle and has 164 similarities with that observed in individuals with *OPA1* mutations [14]. Age of onset is in 165 the first two decades of life (Table 1) and, although comprehensive longitudinal data are 166 lacking, patients typically experience a slowly progressive, symmetrical decrease in vision 167 [9,10]. Visual acuity is usually worse than 0.3 logMAR and it is of interest that one subject in 168 the present series had preserved acuity and no visual complaints at age 41 (subject M2; 0.1 169 logMAR right, 0.2 logMAR left; Table 1 and Figure 1D). Previous reports have shown that 170 peripheral visual fields are spared and that colour vision is impaired [9,10]. Nevertheless, loss 171 of colour discrimination is variable and without a systemic axis [10]. Subject M3 reported 172 dyschromatopsia only in the right eye, in keeping with findings from colour vision testing 173 with the Ishihara plates (right eye 4 of 17 plates; left eye 16 of 17 plates).

174

175 On fundus examination, temporal optic disc pallor was the most common finding in the 176 present cohort although diffuse pallor involving the whole neuroretinal rim and optic disc 177 excavation are not unusual [9,10]. Notably, subject M1 was referred to the clinic after a 178 routine eye test at age 60 revealed an enlarged cup-to-disc ratio (0.8) in the left eye (Figure 179 1C); an erroneous initial diagnosis of glaucoma was made as consideration was not given to 180 the fact that she had infantile-onset cataracts operated at age 35. Furthermore, subject M4 had 181 a healthy optic disc appearance in the right eye and only subtle pallor in the left eye at age 14. 182 In such equivocal cases measurement of retinal nerve fibre layer thickness with optical 183 coherence tomography can be helpful (for example Figure 1D). Future imaging studies in 184 patients with OPA3-related disease are expected to provide further insights.

185

186 Crystalline lens opacities (i.e. cataracts) are observed in most affected individuals [9,10]. 187 Nevertheless, a patient with a heterozygous p.(Val3 Gly4ins2) mutation and no cataracts at 188 age 52 has been reported by Grau and colleagues [10]. Only one of the seven patients 189 reported here had no clinical history of cataract: subject C3 was examined at age 38 and no 190 lens opacity was detected. It is worth highlighting that a number of patients with OPA3-191 related disease presented with bilateral lens opacities at birth or in early childhood (including 192 subject M4 who was only noted to have optic atrophy after lens extraction at age 15); 193 therefore, OPA3 mutations should be in the differential in cases of bilateral infantile-onset 194 cataract and it is important to assess optic nerve function and health before concluding about 195 the likely amblyogenicity of the lens opacities.

196

197 Ocular OPA3-related disease often occurs in parallel with extraocular clinical features. 198 Reynier and colleagues reported affected members of a family with a heterozygous 199 p.(Gly93Ser) change in OPA3 to have mild neurological signs including spasticity and 200 extrapyramidal dysfunction [9]. Subsequently, Grau and colleagues reported four individuals 201 with a heterozygous p.(Val3_Gly4ins2) change to have hearing loss. Expression analysis of 202 OPA3 in murine cochlear tissue was consistent with the notion that auditory neuropathy can 203 be an extraocular feature of OPA3-related disease [10]. Notably, two affected individuals 204 from the present cohort had sensorineural hearing loss (Table 1); hearing impairment has 205 been previously described in a patient with the same genetic defect [10]. Clinicians should be 206 vigilant to the development of such a complication and audiograms should be performed in 207 all patients with OPA3-related disease at least once.

208

209 All OPA3 families with dominant disease reported to date have affected individuals in two or 210 more generations and disease-associated variants appear to be highly penetrant (Figure 1A-B, 211 [9,10]). The p.(Gln105Glu) change identified in both families reported here has been 212 previously shown to be a recurrent mutation [10]; it was not possible to perform haplotype 213 analysis in subjects from the present cohort and thus, the possibility of a recent common 214 ancestor in the two families cannot be excluded. The p.(Gln105Glu) change affects an amino 215 acid in a predicted coiled-coil structure (Uniprot) but it is unclear how this common cause of 216 OPA3-related disease affects mitochondrial function and/or structure.

217

218 Findings from this and other studies [9,10] suggest that dominant OPA3-related disease is 219 associated with significant intra- and inter-familial phenotypic variability. Any attempt to 220 draw genotype-phenotype correlations would therefore be highly speculative and secondary 221 genetic factors are likely to be the basis for the observed clinical heterogeneity; a similar 222 conclusion has been reached for the much commoner OPA1-associated optic neuropathy [14]. 223 Expansion of the phenotypic spectrum of OPA3-related disease has led to greater 224 understanding of the condition and the risks of developing extraocular complications. 225 However, it is still unclear what disease mechanism explains multi-system tissue involvement 226 and future studies on the pathogenesis of the disorder are expected to provide important 227 insights.

- 228
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TABLE

|--|

Subject (family ID)	t Age; VA $(\log MAR)$ Pres right/left		Presentation/ onset	Lens	Heterozygous OPA3 change identified	Other features	
C1 (GCC1)	72; M 1.30/1.30		Nystagmus in infancy	Blue-dot cataracts diagnosed at age 2	p.(Gln105Glu)	-	
C2 (GCC1)	62; F 1.00/1.00		Nystagmus in infancy	Cataracts operated at age 4	p.(Gln105Glu)	-	
C3 (GCC1)	38; F	0.80/0.80	Nystagmus in infancy	No significant lens opacity at age 38	p.(Gln105Glu)	-	
M1 (G43755)	65; F	0.72/1.04	Cataracts at age 34; re-referred as found by optometrist to have cupped left disc at age 60	Cataracts operated at age 34; bilateral aphakia	p.(Gln105Glu)	Right mild-moderate s/n hearing loss; left moderate-severe s/n hearing loss; bilateral ocular hypertension on topical treatment	
M2 (G43755)	41; M	Mild cataracts at M 0.10/0.20 age 9; presently asymptomatic		Nuclear cataracts operated at age 35	p.(Gln105Glu)	Right and left mild- moderate s/n hearing loss	
M3 (G43755)	₍₅₎ 36; F 0.60/0.30 ^I		Decrease in VA from age 19	Blue-dot cataracts operated at age 36&37	p.(Gln105Glu)	Normal audiogram; familial hypercholesterolaemia	
M4 (G43755)	13; F	0.08/0.18	Cataracts at age 13	Lamellar cataracts operated at age 15	p.(Gln105Glu)	Congenital AV malformation in frontal lobe	
III.3 (#1) [9]	38; F	0.70/0.70	Poor vision from infancy	Posterior cortical cataracts operated at age 51	p.(Gly93Ser)	Tremor of hands; extrapyramidal rigidity of upper limbs; absence of deep tendon reflexes	
IV.1 (#1) [9]	15; F	0.30/0.30	Decreased VA before age 10	Posterior cortical cataracts operated at age 47&48	p.(Gly93Ser)	Mild postural tremor & mild rigidity of upper extremities	
IV.2 (#1) [9]	57; F	0.60/0.50	Decreased VA before age 10	Anterior cortical cataracts operated at age 45&46	p.(Gly93Ser)	Postural tremor without extrapyramidal signs	
V.1 (#1) [9]	29; F	0.70/0.52	Decrease in VA from age 12	Anterior cortical cataracts operated at age 25	p.(Gly93Ser)	Normal neurological examination	
VI.1 (#1) [9]	4; F	0.15/0.15	Visual impairment investigated at age 3	Anterior & posterior cortical cataracts operated at age 4	p.(Gly93Ser)	Normal neurological examination	
IV.6 (#1) [9]	50, F	NA	Visual impairment from infancy	Cataracts	p.(Gly93Ser)	Normal neurological examination	
V.6 (#1) [9]	5; F	NA	Visual impairment from infancy	Cataracts operated at age 5	p.(Gly93Ser)	Normal neurological examination	
III.3 (#2) [9]	37; F	1.70/1.70	Decrease in VA from age 12	Posterior capsular cataract diagnosed at age 56	p.(Gln105Glu)	-	
III.7 (#2)[9]	49; F	"legally blind"	Decrease in VA from age 10	Cataracts diagnosed at age 45	p.(Gln105Glu)	-	
IV.1 (#2) [9]	33; F	1.30/1.30	Decrease in VA from age 6	Cataracts diagnosed at age 10; cerulean cataracts at age 33	p.(Gln105Glu)	Normal neurological examination	

IV.2 (#2)[9]	12; M 0.52/0.52 Decreased VA at age 12		Decreased VA at age 12	Posterior capsular cataracts operated p.(Gln105Glu) at age 19		-
II.2 (OAK1) [10]	57 1.00/1.00 Diagnosed at age 18		Monocular cataract	p.(Gln105Glu)	-	
II.6 (OAK1) [10]	54	0.40/0.50	Onset in infancy	Cataracts	p.(Gln105Glu)	Hearing loss
III.1 (OAK1) [10]	35	0.52/0.30 Onset in infancy		No significant lens opacity at age 19	p.(Gln105Glu)	-
III.2 (OAK1) [10]	31	0.40/0.52	Onset in infancy	Cataracts	p.(Gln105Glu)	-
III.3 (OAK1) [10]	17	17 0.52/0.40 Onset in infancy		Cataracts	p.(Gln105Glu)	Chiari malformation type I
I.1 (OAK61)[10]	57	1.30/1.40	Diagnosed at age 10	No information	p.(Gln105Glu)	-
II.1 (OAK61)[10]	33	0.40/0.40	Diagnosed at age 19	No significant lens opacity at age 25	p.(Gln105Glu)	-
II.4 (OAK105)[10]	84	1.15/1.30	Onset in infancy	Congenital cataracts	p.(Val3_Gly4ins 2)	Hearing loss
III.2 (OAK105)[10]	61	0.30/0.40	Onset in adolescence	No significant lens opacity at age 52	p.(Val3_Gly4ins 2)	Hearing loss
III.3 (OAK105) [10]	62	62 0.52/0.40 Diagnosed at age 19		Cataracts	p.(Val3_Gly4ins 2)	Hearing loss
III.5 (OAK105) [10]	57	1.15/1.15	Onset in adolescence	Congenital cataracts	p.(Val3_Gly4ins 2)	Hearing loss
IV.1 (OAK105) [10]	32	32 1.30/1.30 Diagnosed at age 19		Congenital cataracts	p.(Val3_Gly4ins 2)	Morbus Scheuermann
III.1 (OAK255) [10]	46	NA	Onset in infancy	Cataracts	p.(Gln105Glu)	Intestinal pseudo- obstruction
III.2 (OAK255) [10]	47	NA	Onset in infancy	Cataracts	p.(Gln105Glu)	

F, female; M, male; VA, visual acuity; logMAR, logarithm of the minimum angle of resolution (equivalent); BE, both eyes; NA, not available; s/n, sensorineural; AV, arteriovenous. The term cataracts is used to denote bilateral crystalline lens opacities. <u>Variants are annotated according to the NCBI Reference Sequence NM 025136.3 (OPA3 isoform b).</u>

- **FIGURE LEGEND**
- 288

289 **Figure 1.**

- A. Pedigree from a family segregating optic atrophy. Subjects C1 (IV:2), C2 (IV:5) and C3
- 291 (V:3) had a heterozygous *OPA3* change, c.313C>G, p.(Gln105Glu).
- **B.** Pedigree from a family segregating optic atrophy and cataracts. DNA from subjects M1
- 293 (II:6), M2 (III:3) and M3 (III:4) was available for testing; a heterozygous c.313C>G,
- p.(Gln105Glu) change in *OPA3* was identified in all there patients. See text and Table 1 forclinical findings of all tested individuals as well as subject M4 (IV:2).
- 296 C. Right and left optic disc appearance of an individual with a confirmed *OPA3* mutation
- (subject M1) showing pallor of the neuroretinal rim, which is more marked temporally. Left
 optic nerve appears more affected than the right and has optic disc excavation in addition to
 the temporal pallor.
- **D.** Left optic disc appearance and pattern of retinal nerve fibre layer (RNFL) thinning in an individual with a confirmed *OPA3* mutation (subject M2). Sparing of the nasal, superior and inferior peripapillary quadrants is observed. The RNFL profile for each eye is superimposed
- 303 on the normal distribution percentiles. The normal distribution indices are colour-coded: (i)
- 304 red <1%, (ii) yellow <5%, (iii) green <95%, and (iv) white >95%.
- 305