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Clinical spectrum of SIX3-associated mutations in holoprosencephaly: correlation between genotype, phenotype and function

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ABSTRACT ► Additional data are published online only at http://jmg.bmj.

Background: Holoprosencephaly (HPE) is the most common structural malformation of the human forebrain. There are several important HPE mutational target genes, including the transcription factor SIX3, which encodes an early regulator of Shh, Wnt, Bmp and Nodal signalling expressed in the developing forebrain and eyes of all vertebrates

Objective: To characterise genetic and clinical findings in patients with SIX3 mutations.

Methods: Patients with HPE and their family members were tested for mutations in HPE-associated genes and the genetic and clinical findings, including those for additional cases found in the literature, were analysed. The results were correlated with a mutation-specific functional assav in zebrafish.

Results: In a cohort of patients (n = 800) with HPE, SIX3 mutations were found in 4.7% of probands and additional cases were found through testing of relatives. In total, 138 cases of HPE were identified, 59 of whom had not previously been clinically presented. Mutations in SIX3 result in more severe HPE than in other cases of nonchromosomal, non-syndromic HPE. An over-representation of severe HPE was found in patients whose mutations confer greater loss of function, as measured by the functional zebrafish assay. The gender ratio in this combined set of patients was 1.5:1 (F:M) and maternal inheritance was almost twice as common as paternal. About 14% of SIX3 mutations in probands occur de novo. There is a wide intrafamilial clinical range of features and classical penetrance is estimated to be at least 62%. **Conclusions:** Our data suggest that SIX3 mutations result in relatively severe HPE and that there is a genotype-phenotype correlation, as shown by functional studies using animal models.

Holoprosencephaly (HPE) is the most common structural malformation of the human forebrain and occurs after failed or abbreviated midline cleavage of the developing brain (cortex and subcortical structures) during the third and fourth weeks of gestation. HPE occurs in up to 1 in 250

gestations, but only 1 in 8000 live births.12 Classically, three degrees of severity, defined by the extent of brain malformation, have been described. In the most severe form, alobar HPE, there is a single ventricle and no interhemispheric fissure. The olfactory bulbs and tracts and the corpus callosum are typically absent and dark-grey nuclei are not separated. In semilobar HPE, the most common type of HPE in neonates who survive to clinical examination, findings include partial cortical separation with absent or hypoplastic olfactory structures and corpus callosum. In lobar HPE, findings include separate ventricles but incomplete frontal cortical separation. Recently, an additional milder form, called middle interhemispheric variant (MIHV) has been delineated, in which the posterior frontal and parietal lobes are incompletely separated and the corpus callosum may be hypoplastic.³⁻⁶ Of patients with HPE who survived the neonatal period, alobar, semilobar and lobar HPE occurred in 21%, 60% and 19%, respectively.7 A separate study showed that of patients with non-chromosomal, non-syndromic HPE (including both living children and deceased fetuses), alobar, semilobar and lobar HPE occurred in 22%, 45% and 33%, respectively.8

Clinical features may be qualitatively predicted by the specific neuroanatomical abnormalities present. These features can include characteristic craniofacial anomalies, ophthalmological abnormalities such as colobomata or microphthalmia, severe mental retardation or developmental delay, pituitary dysfunction including diabetes insipidus, oromotor dysfunction, dysautonomia and seizures. Severely affected patients do not typically survive beyond early infancy; however, less severely affected patients may have normal life-spans. Though there are exceptions, more severe brain anomalies correlate with more severe clinical sequelae and shorter life-spans.7-10

Craniofacial findings tend to correlate with the type and severity of brain anomalies. In HPE caused by single-gene mutations, facial findings may

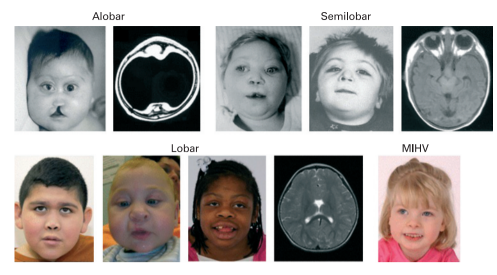


Figure 1 Patients with mutations in *SIX3*, arranged by HPE type (neuroimaging does not correspond with patients). All patients had point mutations in *SIX3* with the exception of the patient shown in the lower row, second from left, who had a complex cytogenetic rearrangement including deletion of 2p21. MRI on patients in row 2 courtesy of the Carter Centers.^{19 48} MIHV, middle interhemispheric variant.

additionally correlate with the causative gene. Patients with alobar HPE may be found to have cyclopia or synophthalmia (fusion of the optic vesicles and incomplete development of separate eyes), proboscis (a tubular nasal appendage appearing above the fused eyes), severe microcephaly and bilateral cleft lip and palate. Features in less severely affected patients may include microcephaly, hypotelorism, a flat nasal bridge and cleft lip or palate. Findings at the least severe end of the phenotypically recognisable spectrum (sometimes termed a "microform" of HPE. seen in patients without the cardinal central nervous system findings of HPE) may include solitary maxillary central incisor, hypotelorism and microcephaly. However, exceptions are often seen; a patient with severe HPE may have relatively subtle facial dysmorphisms and can have macrocephaly (as opposed to the more common microcephaly) due to hydrocephalus.^{4 9} Overall, facial anomalies have been reported in approximately 80% of patients with HPE and often lead to the diagnosis.8

HPE is aetiologically associated with teratogens such as maternal diabetes mellitus and alcohol, and has been reported in cases of prenatal exposure to pharmaceutical agents such as retinoic acid and the statin class of drugs, and to infections including cytomegalovirus, toxoplasmosis, and rubella.^{9 11 12} Up to half of patients with HPE have a numerical or structural chromosomal abnormality, whereas up to a quarter have HPE as part of a recognisable syndrome.^{13–15} HPE may also be due to single-gene mutations; < 25% of cases of HPE result from single-gene mutations in the currently commercially tested HPE-associated genes *SHH*, *ZIC2*, *SIX3* and *TGIF*.⁸

Sonic Hedgehog (*SHH*) was the first causative gene identified in human HPE.¹⁶ Since then, single mutations in at least 10 other genes have been purported to cause HPE.⁹ ^{17–24} In families in which these mutations were found, HPE seems to be inherited in an autosomal dominant fashion. Large kindreds segregating HPE-associated mutations show both incomplete penetrance and highly variable expressivity, suggesting additional environmental or genetic influences superimposed on the haploinsufficent state.²³ According to this "multiple-hit" model, a mutation in a HPE-associated gene is necessary but not sufficient for HPE, and other genes or environmental factors are required for complete phenotypic feature.²⁵

SIX3 has been reported as the third most common cause of HPE due to single-gene mutations.⁸ The vertebrate Six genes encode a family of related transcription factors that are orthologues of the sine oculis ("without eyes") gene in Drosophila, and are expressed in the developing fly's visual system, suggesting partially conserved phylogenetic roles. Indeed, vertebrate Six3 has been shown to be involved in midline forebrain and eye formation in several organisms including mouse, chick, Xenopus and zebrafish.²⁶⁻²⁸ Proteins encoded by the Six genes characteristically contain a DNAbinding homeobox domain, and an upstream SIX domain that can recruit additional factors to accomplish transcriptional activation or repression.^{29 30} Known biological properties of vertebrate Six3 include transcriptional repression of BMP, Wnt and Nodal targets through complex(s) formed with the general co-repressor Groucho.³¹⁻³³ It also forms a complex with a different factor, geminin, which can influence a cell's fate towards differentiation rather than proliferation in the early expanding forebrain territory.³⁴ Presumably through a distinct set of co-factors, the Six3 transcription factor can instead activate lens specification genes during eye formation.³⁵ Finally, Six3 acts as a direct regulator of Sonic hedgehog expression in the ventral forebrain.^{36 37}

SIX3 was first identified as a candidate gene because of patients with HPE who had cytogenetic anomalies involving 2p21. SIX3, which has the appropriate spatial and temporal expression pattern to result in HPE, was the most attractive candidate gene in the interval.^{19 38}

We report our findings in 59 new patients with *SIX3* mutations and an additional 79 cases collected from the literature, and correlate the clinical phenotypes with *SIX3* genotypes and functional studies in the zebrafish.^{19 38-51} The clinical findings echo the incomplete penetrance and highly variable expressivity seen in HPE due to mutations in other genes. Recent work in animal models specifically supports the multiple-hit model in mammalian *SIX3*-associated HPE.³⁶ It has been suggested that *SIX3* mutations result in a more severe holoprosencephaly phenotype than do mutations in the other HPE-related genes although a mechanism that would explain this is not yet clear.^{4 9 46}

Table 1	Patient	characteristics,	mutations	and	functional	data
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Patient*	HPE type	DNA alteration (location) in <i>SIX3</i>	Predicted protein alteration in SIX3	Functional studies†47	Gender	Proband/ relative	Ref.
	Not specified	c.109G \rightarrow T (NTD)	p.G37C	0.5–0.9	Female	Proband	NA
\$	Lobar	c.109G→T (NTD)	p.G37C	0.5-0.9	Male	Proband	42
a‡	MIHV	c.109G→T (NTD)	p.G37C	0.5-0.9	Female	Proband	NA
o‡	None	c.109G→T (NTD)	p.G37C	0.5-0.9	Female	Mother	NA
а‡	Alobar	c.206G→A (NTD)	c.G69D	>0.9	Female	Proband	48
- .		c.406dupGC (SD)	FS	<0.5			
b‡	None	c.206G \rightarrow A (NTD)	c.G69D	>0.9	Male	Father	48
0+	None	c.406dupGC (SD)	FS	<0.5	TVIC	ratio	48
-	Mana		c.G69D	<0.5 >0.9	Famala	Deternel	48
C	None	c.206G \rightarrow A (NTD)	C.009D	<i>≥</i> 0.9	Female	Paternal grandmother	
		c.406dupGC (SD)	FS	<0.5	Female		48
a‡	Alobar	c. 214G→C (NTD)	p.A72P	<0.5	Female	Proband	48
, ,	None	c. 214G→C (NTD)	p.A72P	<0.5	Female	Mother	48
·+	Alobar	c.235A→G (NTD)	p.M79V	Unknown	Not specified	Proband	47
+	Alobar	Not specified (SD)	p.V92G	<0.5	Male	Proband	45
1‡			•				45
o‡	Alobar	Not specified (SD)	p.V92G	<0.5	Male	Brother	45
;	Alobar	Not specified (SD)	p.V92G	<0.5	Male	Brother	45
1‡	Microform	Not specified (SD)	p.V92G	<0.5	Female	Mother	
÷	Semilobar	c.275T→G (SD)	p.V92G	<0.5	Female	Proband	43
t	Semilobar	c.278C→A (SD) (<i>PTCH1</i> : c.1165G→A)	p.A93D (PTCH1: p.A393T)	<0.5	Female	Proband	22 42
Da‡	Alobar	c.311A→G (SD)	p.D104G	0.5-0.9	Male	Proband	48
)b‡	None	c.311A→G (SD)	p.D104G	0.5-0.9	Male	Father	48
I‡	Not specified	c.313A→G (SD)	p.1105V	>0.9	Not specified	Proband	43
2‡	Lobar	c.338G→A (SD)	p.W113X	<0.5	Male	Proband	42
-+ Ba‡	Not specified	c.339G→A (SD)	p.W113X	<0.5	Male	Proband	NA
•			•				
3b‡	Not specified	Not tested	Not tested	Not tested	Male	Brother	NA
Bc‡	Microform	c.339G→A (SD)	p.W113X	<0.5	Female	Mother	NA 42
la‡	Alobar	c.339G→T (SD)	p.W113C	<0.5	Female	Proband	
4b‡	Microform	c.339G→T (SD)	p.W113C	<0.5	Female	Mother	42
lc‡	Microform	c.339G→T (SD)	p.W113C	<0.5	Female	Maternal aunt	42
4d‡	Microform	c.339G→T (SD)	p.W113C	<0.5	Male	Maternal uncle	42
le‡	None	c.339G→T (SD)	p.W113C	<0.5	Female	Maternal aunt	42
4f‡	Microform	c.339G \rightarrow T (SD)	p.W113C	<0.5	Female	Maternal grandmother	42
4g‡	Microform	Linkage only	Linkage only	Linkage only	Male	Maternal great uncle	42
4h‡	None	Linkage only	Linkage only	Linkage only	Female	Maternal first cousin	42
4i‡	Not specified	Linkage only	Linkage only	Linkage only	Female	Maternal half- sister	42
4j	Not specified	Not tested	Not tested	Not tested	Female	Maternal first cousin	42
4k‡	Not specified	Not tested	Not tested	Not tested	Male	Maternal first cousin	42
41‡	Not specified	Not tested	Not tested	Not tested	Female	Maternal first cousin	42
4m	Not specified	Not tested	Not tested	Not tested	Female	Maternal first cousin, once removed	42
1n	Not specified	Not tested	Not tested	Not tested	Male	Maternal great uncle	42
40	Not specified	Not tested	Not tested	Not tested	Not specified	Maternal great- grandparent	42
5‡	Semilobar	c.341C→T (SD)	p.S114L	<0.5	Female	Proband	42
ba‡	Lobar	c.385G→T (SD)	p.E129X	<0.5	Male	Proband	42
6b‡	Alobar	c.385G→T (SD)	p.E129X	<0.5	Female	Sister	42
Bc‡	Microform	c.385G→T (SD)	p.E129X	<0.5	Male	Father	42
7 7	Not specified	c.389C \rightarrow A (SD)	p.S130X	<0.5 <0.5	Male	Proband	42
	•						48
Ba‡	Alobar	c.404G→C (SD)	p.R135P	<0.5	Male	Proband	-

Continued

Table 1 Continued

Patient*	HPE type	DNA alteration (location) in <i>SIX3</i>	Predicted protein alteration in SIX3	Functional studies†47	Gender	Proband/ relative	Ref.
18b‡	None	c.404G→C (SD)	p.R135P	<0.5	Female	Mother	48
19	Not specified	c.404_407dupGCGC (SD)	FS	<0.5	Female	Proband	42
20‡	Semilobar	c.405_409CGCCG (SD)	FS	<0.5	Male	Proband	42
21a	Not specified	c.463_465delCAC (SD)	p.H155del	Unknown	Not specified	Proband	47
21b	Not specified	c.463 465delCAC (SD)	p.H155del	Unknown	Female	Mother	47
22‡	Not specified		p.F157l	<0.5	Male	Proband	42
23a‡	Alobar	c.507delG (SD)	FS	<0.5	Female	Proband	48
23b‡	None	c.507delG (SD)	FS	<0.5	Male	Father	48
24a‡	Lobar	c.515C→T (SD)	p.A172V	0.5-0.9	Male	Proband	42
24b‡	Semilobar	Not tested	Not tested	Not tested	Female	Sister	42
24c‡	Not specified	Not tested	Not tested	Not tested	Female	Mother	42
25a	Alobar	c.518A→C (SD)	p.H173P	0.5-0.9	Female	Proband	45
	None				Female		45
25b	Semilobar	c.518A \rightarrow C (SD)	p.H173P	0.5-0.9		Mother	43
26a		c.518A \rightarrow C (SD)	p.H173P	0.5-0.9	Female	Proband	43
26b	None	c.518A \rightarrow C (SD)	p.H173P	0.5-0.9	Female	Mother	43
26c	None	c.518A→C (SD)	p.H173P	0.5–0.9	Female	Maternal aunt	43
26d	Not specified	Not tested	Not tested	Not tested	Not specified	Maternal first cousin	
26e	Not specified	Not tested	Not tested	Not tested	Not specified	Maternal first cousin	43
26f	Not specified	Not tested	Not tested	Not tested	Not specified	Maternal first cousin	43
27‡	Not specified	c.520T→C (SD)	p.Y174H	0.5–0.9	Female	Proband	42
28a‡	Alobar	c.542 576dup (SD)	FS	Unknown	Female	Proband	43 45
28b‡	None	c.542 576dup (SD)	FS	Unknown	Male	Father	43 45
29‡	Semilobar	c.551delC (SD)	FS	<0.5	Female	Proband	42
30a‡	Alobar	c.556 557dupGG (SD)	FS	<0.5	Female	Proband	44 45
30b‡	Alobar	c.556 557dupGG (SD)	FS	<0.5	Male	Brother	44 45
30c‡	Alobar	-	FS	<0.5 <0.5	Female	Sister	44 45
-		c.556_557dupGG (SD)	FS	<0.5 <0.5		Sister	44 45
30d‡	Not specified	c.556_557dupGG (SD)	FS	<0.5 <0.5	Female		44 45
30e‡	Microform	c.556_557dupGG (SD)			Male	Father	42
31a‡	Alobar	Not tested	Not tested	Not tested	Female	Proband	42
31b‡	Semilobar	c.582dup C (SD)	FS	<0.5	Female	Sister	42
31c‡	Microform	c.582dup C (SD)	FS	<0.5	Male	Father	43 45
32a‡	Semilobar	c.605C→T (SD)	p.T202I	<0.5	Female	Proband	
32b‡	None	c.605C→T (SD)	p.T202l	<0.5	Female	Mother	43 45
33a‡	Semilobar	c.619G→T (HD)	p.E207X	<0.5	Male	Proband	42
33b‡	Alobar	c.619G→T (HD)	p.E207X	<0.5	Female	Sister	42
34	Not specified	c.637T→G (HD)	p.F213V	<0.5	Male	Proband	42
35‡	Lobar	c.652C→T (HD)	p.R218W	0.5-0.9	Female	Proband	42
36‡	Not specified	c. 653G→C (HD)	p.R218P	<0.5	Female	Proband	42
37‡	Not specified	c.676C→G (HD)	p.L226V	>0.9	Female	Proband	19 52
38	Not specified	c.680A→C (HD)	p.Q227P	0.5–0.9	Not specified	Proband	42
39‡	Microform	c.686C→T (HD)	p.P229L	Unknown	Male	Proband	46
40‡	Alobar	c.692C→G (HD)	p.P231R	<0.5	Female	Proband	45
41‡	Alobar	c.698 706del (HD)	p.N233_S235del	<0.5	Female	Proband	19 39
42	Not specified	c.GC718_719AA (HD)	p.A240K	<0.5	Not specified	Proband	42
42 43a‡	Not specified	c.721C→T (HD)	p.0241X	⊂0.5 Unknown	Male	Proband	42
-	Not specified	. ,	Not tested		Not specified		42
43b 42o*		Not tested		Unknown		Sibling	42
43c‡	None	c.721C→T (HD)	p.0241X	Unknown	Female	Mother	42
44	Not specified	c.730G→T (HD)	p.G244C	<0.5	Not specified	Proband	42
45	Not specified	c.736delA (HD)	FS	Unknown	Not specified	Proband	
46	Not specified	c.743_745delCAG (HD) (<i>TGIF</i> : c.478C→T)	p.248K delQ (TGIF: p.160S)	Unknown	Female	Proband	NA
47a‡	Microform	c.749T→C (HD)	p.V250A	0.5–0.9	Male	Proband	19
47b‡	None	c.749T→C (HD)	p.V250A	0.5-0.9	Female	Mother	19
47c‡	None	c.749T→C (HD)	p.V250A	0.5-0.9	Female	Maternal aunt	19
							19

Continued

Table 1 Continued

Patient*	HPE type	DNA alteration (location) in <i>SIX3</i>	Predicted protein alteration in SIX3	Functional studies†47	Gender	Proband/ relative	Ref.
47e‡	Alobar	c.749T→C (HD)	p.V250A	0.5–0.9	Not specified	Sibling	19
17f	Alobar	c.749T→C (HD)	p.V250A	0.5-0.9	Not specified	Sibling	19
8	Not specified	c.762T→A (HD)	p.F254L	0.5-0.9	Male	Proband	42
19‡	Not specified	c.769C→T (HD)	p.R257W	<0.5	Female	Proband	42
50a	Semilobar	c. 769C→T (HD)	p.R257W	<0.5	Female	Proband	43 45
50b	Alobar	Not tested	Not tested	Not tested	Male	Brother	43 45
50c‡	None	c. 769C→T (HD)	p.R257W	<0.5	Male	Father	43 45
i1a	Lobar	c. 769C→G (HD)	p.R257G	Unknown	Not specified	Proband	47
i1b	Not specified	c. 769C→G (HD)	p.R257G	Unknown	Male	Father	47
2‡	Semilobar	c.770G→C (HD)	p.R257P	>0.9	Female	Proband	19 52
3	Not specified	c.773G→C (HD)	p.R258L	<0.5	Not specified	Proband	42
4	Not specified	c.785G→A (HD)	p.R262H	0.5-0.9	Not specified	Proband	42
5a	Not specified	Not tested	Not tested	Unknown	Not specified	Proband	47
i5b	None	Not tested	Not tested	Unknown	Not specified	Obligate carrier	47
5c	Not specified	c.806G→C (CTD)	p.R269T	Unknown	Male	Grandfather	47
6a‡	Semilobar	c.806G→T (CTD)	p.R269M	Unknown	Female	Proband	47 51
6b‡	Alobar	Not tested	Not tested	Unknown	Male	Sibling	47
6c‡	Lobar	c.806G→T (CTD)	p.R269M	Unknown	Male	Sibling	47
6d‡	Microform	c.806G→T (CTD)	p.R269M	Unknown	Female	Mother	47 51
7	Semilobar	c.806+1G→T (CTD)	Splice	Unknown	Not specified	Proband	47
8‡	Not specified	c.G807C (CTD)	p.R269S	0.5-0.9	Male	Proband	42
9	Not specified	c.820_832delinsCTGGACCT (CTD)	p.A274X	Unknown	Male	Proband	NA
0	Semilobar	c.850G→C (CTD) (<i>ZIC2</i> : c.910T→C)	p.A284P (ZIC2: p.W304R)	Unknown	Female	Proband	NA
1‡	Semilobar	c.890C→T (CTD)	p.P297L	0.5-0.9	Female	Proband	42
2‡	Semilobar	c.944C→T (CTD)	p.T315l	Unknown	Female	Proband	48
3	Not specified	Deletion	NA	Unknown	Not specified	Proband	47
4	Not specified	Microdeletion by qPCR	NA	Unknown	Not specified	Proband	40 41
5	Not specified	Microdeletion by qPCR	NA	Unknown	Not specified	Proband	40 41
6	Not specified	Microdeletion by qPCR	NA	Unknown	Not specified	Proband	41
7	Not specified	Microdeletion by qPCR	NA	Unknown	Female	Proband	41
8‡	Semilobar	del(2)(p2101p2109)	NA	Unknown	Female	Proband	38 49
9‡	None	del(2)(p21)	NA	Unknown	Male	Proband	38
0‡	Semilobar	del(2)(p21p23)	NA	Unknown	Male	Proband	38
1‡	Semilobar	del(2)(p16p22)	NA	Unknown	Female	Proband	38
2	Lobar	del(2)(p21p22.2)	NA	Unknown	Female	Proband	38 50
3‡	Alobar	del(2)(p21p22.1)	NA	Unknown	Male	Proband	38
4‡	Semilobar	inv ins(2)(p21q24q13)	NA	Unknown	Female	Proband	38
/5‡	Semilobar	t(1;2)(p21;p21)	NA	Unknown	Male	Proband	38
6	Alobar	t(1;2)(p22.3;p21)	NA	Unknown	Male	Proband	38
17‡	Lobar	del(2)(p16.3p21)§	NA	Unknown	Male	Proband	NA

CTD, C-terminal domain; HD, homeodomain; HPE, holoprosencephaly; MIHV, middle interhemispheric variant; NA, not applicable (unpublished when seeming in the Reference column); NTD, N-terminal domain; qPCR, quantitative PCR: SD, SIX domain.

*Each kindred is numbered separately; within a kindred, individual members have separate letter identifier.

†Protein Activity Index.

‡Clinical information available.

\$Complex rearrangement additionally resulting in chromosome 2 inversion, translocation involving chromosomes 7, 13 and 18 and deletions of regions of chromosomes 7 and 18.

METHODS

The study protocol was approved by the National Human Genome Research Institute, and informed consent was obtained from all participants or their guardians.

Of the 65 cases for whom inheritance was known, 57.8% had maternal inheritance, 26.6% had paternal inheritance, 13.8% were de novo and the condition in 2 maternal half-siblings in one family seems to be due either an undetected germline mutation or parental mosaicism (the mother had negative mutation testing on peripheral blood analysis). We found no significant association between HPE severity and the parent of origin.

Of the 113 patients for whom gender was known, 40.7% were male and 59.3% were female, giving male:female ratio of 1:1.5. By χ^2 analysis, there was a statistically significant difference in the increased prevalence of affected females overall ($\chi^2 = 3.903$, p = 0.0482). Of the 61 kindreds in which the gender of the proband was known, 59.0% were female and 41.0% were male, giving a male:female ratio of 1:1.4. There was no significant difference in the increased prevalence of affected female probands ($\chi^2 = 1.984$, p = 0.159).

Of those kindreds with molecularly identified specific mutations in the proband, 40.3% had multiple affected family members identified, though familial testing or clinical description

Table 2Mutations (n = 63)

	n (%)
Location	
N-terminal domain	6 (9.5)
SIX domain	27 (42.9)
Homeodomain	22 (34.9)
C-terminal domain	8 (12.7)
Exon 1	57 (90.5)
Exon 2	6 (9.5)
Mutation type	
Missense	44 (69.8)
Nonsense	6 (9.5)
$Duplication \to FS$	6 (9.5)
Deletion→FS	3 (4.8)
Splice mutation	1 (1.6)
9 bp deletion \rightarrow 3 aa deletion	1 (1.6)
3 bp deletion \rightarrow 1 aa deletion	1 (1.6)
Deletion/insertion \rightarrow nonsense	1 (1.6)

aa, Amino acid; FS, frameshift; MIHV, middle interhemispheric variant.

was not available in all cases. Of the cases with described phenotypes, the penetrance of recognised phenotypic effect of mutations in *SIX3* was estimated to be at least 82% (n = 111). Within kindreds with multiple affected members and where clinical description was available, penetrance was estimated at 73% (n = 61). However, testing was often performed only on individuals with the phenotype and their parents. Many others were ascertained only because of the presence of a relative with severe HPE and only on later examination were considered to have microform HPE (eg, hypotelorism). Considering these patients to be "unaffected" results in a penetrance estimate of 62% (n = 61) (fig 1, table 1).

Mutations

Of those with molecularly identified mutations, the 62 kindreds encompassed 63 mutations in *SIX3* (kindred 4 had two mutations in *SIX3*), of which 93.7% were unique. Three unrelated kindreds had the same mutation in the N-terminal domain, two had the same mutation in the SIX domain and two had the same mutation in the homeodomain. Three patients had mutations in

Figure 2 Results of mutation studies.

two HPE genes: one (patient 9) in both *SIX3* and *PTCH*, one (patient 46) in both *SIX3* and *TGIF* and one (patient 60) in both *SIX3* and *ZIC2* (tables 1, 2; figs 2, 3).

Clinical features

Patients with mutations in *SIX3* (not including cytogenetic cases) had a significantly different distribution of HPE types than previously published cases of non-chromosomal, non-syndromic HPE ($\chi^2 = 24.179$, p<1 \cdot 10⁻⁴).⁸ Overall, patients with *SIX3* mutations had a higher proportion of severe HPE (table 3).

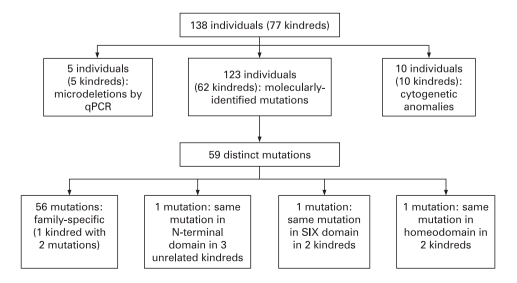
The quality of patient data was highly variable, so it is difficult to accurately calculate the prevalence of specific phenotypic features. However, we present the most commonly reported craniofacial anomalies (table 4). Severe craniofacial findings such as cyclopia and proboscis were only reported with alobar HPE. Overall, the severity of facial dysmorphisms seemed to correlate with the degree of brain anomalies—for example, the degree of hypotelorism was more pronounced in patients with alobar HPE.

In terms of clinical features other than craniofacial anomalies, the most commonly reported findings, in decreasing order of prevalence are: mental retardation or developmental delay, seizure disorder and diabetes insipidus (supplemental tables 1a,b online).

Functional studies

Using the functional analysis developed by Domené *et al*,⁴² 46 mutations (representing 99 patients with HPE who had neuroimaging performed or who were neurologically normal with *SIX3* mutations) had functional studies performed using a zebrafish assay.⁴² Functional study results were divided into three categories, with protein activity described relative to a normal control (value of 1.0): (1) protein activity <0.5 (alleles with the least functional activity), (2) protein activity 0.5–0.9 (alleles with moderate functional activity) and (3) protein activity >0.9 (alleles with near-normal functional activity). One kindred (kindred 4) had two separate mutations in *SIX3* and was categorised as belonging to the functional group with the more severe impairment of the two.

Including those patients who had alobar, semilobar and lobar HPE types with a functional protein index <0.5, there was an



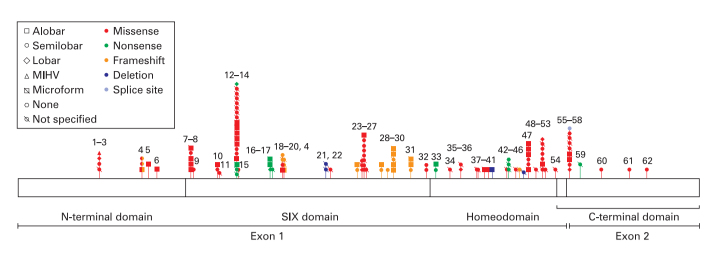


Figure 3 Known mutations in *SIX3*, showing holoprosencephaly and mutation type. MIHV, middle interhemispheric variant. Numbers refer to kindreds.

unequal distribution of HPE types ($\chi^2 = 12.071$, p = 0.0039), with more severe HPE over-represented. Of those with functional protein index <0.5 and HPE on neuroimaging, 60.7% had alobar, 32.1% had semilobar and 7.1% had lobar HPE. Within the group of patients whose mutations had a functional protein index of 0.5–0.9, a significantly unequal distribution of HPE types was not found ($\chi^2 = 0.727$, p = 0.8054). (fig 4, table 5).

Including those with the most severe types of HPE (alobar, semilobar and lobar) and functional protein index either <0.5 and 0.5–0.9, we performed the linear trend alternative to independence test,⁵² using SAS V9.1 software. As there was only one person with functional protein index >0.9 whose HPE type was known, this category was not included in the statistical analysis. Our results showed that, within the group of patients whose mutations had a functional protein index <0.9 there was a significant correlation between HPE types and the functional protein index ($\chi^2 = 12.99$, p = 0.0003). That is, the functional protein index is an explanatory variable with ordered categories for traditional HPE types.

DISCUSSION

We present 138 cases of HPE with mutations in *SIX3*, many of whom have not been previously clinically described; this is the largest comprehensive evaluation of a cohort of patients with mutations in a gene involved in HPE. Analysis of this group allows several conclusions to be made.

Table 3Distribution of HPE types (n = 138)

HPE type	n (% of total) (% of cases with known HPE type)	
Alobar	27 (19.6, 29.3)	
Semilobar	22 (15.9, 23.9)	
Lobar	9 (6.5, 9.8)	
MIHV	1 (0.7, 1.1)	
Microform	13 (9.4, 14.1)	
None*	20 (14.5, 21.7)	
Not specified	46 (33.3)	

MIHV, middle interhemispheric variant.

*No evidence for penetrance.

First, as has been previously posited, our study shows that mutations in *SIX3* correlate with more severe HPE than with non-chromosomal, non-syndromic HPE overall.^{4 9 48} Despite the overall high severity, the clinical features are highly variable. Within large kindreds, people with *SIX3* mutations include both phenotypically normal people and people with severe holoprosencephaly incompatible with life. The basis of this variable expressivity is largely unknown. One potential explanation is a multi-hit mechanism, examples of which are the three cases where mutations were found in *SIX3* and in one other HPE-associated gene. In these patients, mutations in *SIX3* may be necessary but not sufficient for HPE. Another insult, either genetic (eg, changes in at least one other HPE-associated gene) or environmental (eg, gestational diabetes mellitus) must also occur.

Second, mutations in *SIX3* result in holoprosencephaly through a loss-of-function mechanism. Of patients whose mutations result in the greatest functional impairment in the zebrafish assay, protein activity correlates with human HPE severity.⁴² This analysis allows for a more refined genotype-phenotype discussion. Instead of correlating the gene involved or the location or type of the mutation with severity of phenotype, we can begin to predict features based on functional analyses.

Third, these patients do confirm the idea that in HPE, "the face predicts the brain". That is, more severe facial dysmorphisms (such as cyclopia) tend to correlate with more severe HPE.

Table	4	Reported*	phenotypic	findings	(n = 91)
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Finding	Prevalence (%)
Craniofacial anomalies	
Hypotelorism	44.0
Microcephaly	36.3
Cleft lip and/or palate	35.2
Flat nasal bridge/absent nasal septum	17.6
Philtral agenesis/hypoplasia	13.2
Coloboma	9.9
Solitary maxillary central incisor	8.8
Cyclopia	6.6

*It is likely that many of these findings, such as hypotelorism, occur more often than was reported.

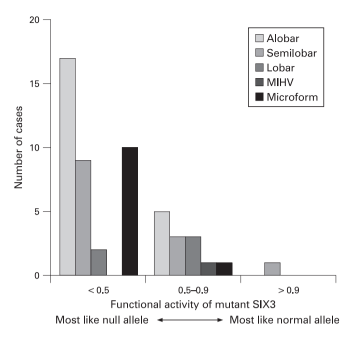


Figure 4 Functional results and HPE types, showing overrepresentation of severe HPE of whose mutation resulted in the greatest functional impairment by zebrafish assay.⁴² MIHV, middle interhemispheric variant.

Although this adage does not apply to all cases of HPE, it does seem to hold true in HPE due to *SIX3* mutations.

Fourth, we estimated penetrance at 82% of all cases, 73% of cases from well-described families with multiple mutation carriers and 62% of those diagnosed on clinical grounds alone. In the practice of clinical genetics, this lowest estimate could be considered the most accurate. However, this latter penetrance estimate may be inaccurate for at least two reasons. First, the quality of clinical data was variable and subtle signs of midline defects may have been missed, resulting in an under-estimate of penetrance. Second, as full testing was not available on family members in many cases, there may have been many cases with *SIX3* mutations who were not ascertained. This would result in an overestimate of penetrance. If more members of families could be tested, it would be possible to better calculate the penetrance, quantify expressivity and calculate the rate of sporadic mutations.

Fifth, it is interesting that more female patients have been reported with *SIX3* mutations. The female predominance has been reported in some, but not all studies of HPE.^{4 8} The fact that the difference does reach significance when all mutation

Table 5 Functional result vs. HPE type (n = 99)

	<0.5	0.5–0.9	>0.9	Total
Alobar	17	5	0	22
Semilobar	9	3	1	13
.obar	2	3	0	5
VIIIV	0	1	0	1
/licroform	10	1	0	11
one	9	7	0	16
lot specified	19	10	2	31
otal	66	30	3	99

MIHV, middle interhemispheric variant.

carriers are considered lends credence to the idea that being female is somehow protective. One explanation is that the *SIX3* mutations in males may be more likely to be embryonic-lethal. However, in patients who survived long enough to have the type of HPE identified, the correlation between gender and severity of HPE was not significant.

One shortcoming of this report is that most of the patients discussed here were not seen in person, although we did perform the laboratory analysis to identify the SIX3 mutation in approximately half the cases. Details of the data available from referring clinicians varied greatly; in some cases, extensive medical records were sent, whereas in other cases, relatively little was available. For this reason, it is difficult to make certain genotype-phenotype correlations. However, it can be noted that in cases for which more clinical information was available. the severity of features often seemed more impressive. Thus, these data may under-represent the severity of the clinical features of SIX3 mutations. On the other hand, many people who had either no or very subtle features were ascertained only because they were related to a person with the same mutation but much more severe presenting signs. Following this logic, these data may over-represent severity.

Despite the challenges interpreting the large and varied data, the number of patients and families described here greatly enriches our understanding of the spectrum of features in patients with mutations in *SIX3*. These considerations argue for the importance of a combined and comprehensive approach to clinical and genetic studies of complex genetic disorders such as HPE.

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NODAL signaling strength via mutation of several pathway members including FOXH1 is linked to human heart defects and holoprosencephaly. Am J Hum Genet 2008-83-18-29

- 25 Ming JE, Muenke M. Multiple hits during early embryonic development: digenic diseases and holoprosencephaly. Am J Hum Genet 2002;71:1017-32
- 26 Gestri G, Carl M, Appolloni I, Wilson SW, Barsacchi G, Andreazzoli M. Six3 functions in anterior neural plate specification by promoting cell proliferation and inhibiting Bmp4 expression. Development 2005;132:2401-13.
- 27. Kobayashi M, Toyama R, Takeda H, Dawid IB, Kawakami K. Overexpression of the forebrain-specific homeobox gene six3 induces rostral forebrain enlargement in zebrafish. Development 1998:125:2973-82.
- 28. Oliver G, Mailhos A, Wehr R, Copeland NG, Jenkins NA, Gruss P. Six3, a murine homologue of the sine oculis gene, demarcates the most anterior border of the developing neural plate and is expressed during eye development. Development 1995 121 4045-55
- Kawakami K, Ohto H, Takizawa T, Saito T. Identification and expression of six family 29 genes in mouse retina. FEBS Lett 1996;393:259-63.
- Granadino B, Gallardo ME, Lopez-Rios J, Sanz R, Ramos C, Ayuso C, Bovolenta P, 30. Rodriguez de Cordoba S. Genomic cloning, structure, expression pattern, and chromosomal location of the human SIX3 gene. Genomics 1999;55:100-5.
- Inbal A, Kim SH, Shin J, Solnica-Krezel L. Six3 represses nodal activity to establish 31 early brain asymmetry in zebrafish. Neuron 2007:55:407-15.
- Kobayashi M, Nishikawa K, Suzuki T, Yamamoto M. The homeobox protein Six3 32 interacts with the Groucho corepressor and acts as a transcriptional repressor in eye and forebrain formation. Dev Biol 2001;232:315-26.
- 33. Lagutin OV, Zhu CC, Kobayashi D, Topczewski J, Shimamura K, Puelles L, Russell HR, McKinnon PJ, Solnica-Krezel L, Oliver G. Six3 repression of Wnt signaling in the anterior neuroectoderm is essential for vertebrate forebrain development. Genes Dev 2003;17:368-79.
- 34 Del Bene F. Tessmar-Raible K. Wittbrodt J. Direct interaction of geminin and Six3 in eye development. Nature 2004;427:745-9.
- 35 Liu W, Lagutin OV, Mende M, Streit A, Oliver G. Six3 activation of Pax6 expression is essential for mammalian lens induction and specification. EMBO J 2006;25:5383-95
- 36. Geng X, Speirs C, Lagutin O, Inbal A, Liu W, Solnica-Krezel L, Jeong Y, Epstein DJ, Oliver G. Haploinsufficiency of Six3 fails to activate Sonic hedgehog expression in the ventral forebrain and causes holoprosencephaly. Dev Cell 2008;15:236-47.
- Jeong Y, Leskow FC, El-Jaick K, Roessler E, Muenke M, Yocum A, Dubourg C, Li X, 37 Geng X, Oliver G, Epstein DJ. Regulation of a remote Shh forebrain enhancer by the Six3 homeoprotein. Nat Genet 2008;40:1348-53.
- Schell U, Wienberg J, Köhler A, Bray-Ward P, Ward DE, Wilson WG, Allen WP, Lebel 38. RR, Sawyer JR, Campbell PL, Aughton DJ, Punnett HH, Lammer EJ, Kao FT, Ward DC, Muenke M. Molecular characterization of breakpoints in patients with holoprosencephaly and definition of the HPE2 critical region 2p21. Hum Mol Genet 1996:5:223-9.
- 39 Nanni L, Croen LA, Lammer EJ, Muenke M. Holoprosencephaly: molecular study of a California population. Am J Med Genet 2000;90:315-19.
- 40 Bendavid C, Dubourg C, Gicquel I, Pasquier L, Saugier-Veber P, Durou MR, Jaillard S, Frebourg T, Haddad BR, Henry C, Odent S, David V. Molecular evaluation of foetuses with holoprosencephaly shows high incidence of microdeletions in the HPE genes. Hum Genet 2006;119:1-8.
- 41. Bendavid C, Haddad BR, Griffin A, Huizing M, Dubourg C, Gicquel I, Cavalli LR, Pasquier L, Shanske AL, Long R, Ouspenskaia M, Odent S, Lacbawan F, David V, Muenke M. Multicolour FISH and quantitative PCR can detect submicroscopic deletions in holoprosencephaly patients with a normal karyotype. J Med Genet 2006;43:496-500.
- Domené S, Roessler E, El-Jaick K, Boorech J, Vélez JI, Bale S, Lacbawan F, Muenke 42. M, Feldman B. Mutations in the human SIX3 gene in holoprosencephaly are loss-offunction. Hum Mol Genet 2008;17:3919-28.
- 43. Dubourg C, Lazaro L, Pasquier L, Bendavid C, Blayau M, Le Duff F, Durou MR, Odent S, David V. Molecular screening of SHH, ZIC2, SIX3, and TGIF genes in patients with features of holoprosencephaly spectrum: Mutation review and genotype- phenotype correlations. Hum Mutat 2004;24:43-51.
- Pasquier L, Dubourg C, Blayau M, Lazaro L, Le Marec B, David V, Odent S. A new 44. mutation in the six-domain of SIX3 gene causes holoprosencephaly. Eur J Hum Genet 2000;8:797-800.
- Pasquier L, Dubourg C, Gonzales M, Lazaro L, David V, Odent S, Encha-Razavi F. 45 First occurrence of aprosencephaly/atelencephaly and holoprosencephaly in a family with a SIX3 gene mutation and phenotype/genotype correlation in our series of SIX3 mutations. J Med Genet 2005;42:e4.
- El-Jaick KB, Fonseca RF, Moreira MA, Ribeiro MG, Bolognese AM, Dias SO, 46. Pereira ET, Castilla EE, Orioli IM. Single median maxillary central incisor: new data and mutation review. Birth Defects Res A Clin Mol Teratol 2007;79:573-80
- 47. Paulussen A, Tserpelis D, Spierts S, Smeets H, Herbergs J. Holoprosencephaly mutations in the Dutch population. Poster presented at the European Society of Human Genetics, Barcelona, Spain, 31 May to 3 June, 2008 48
- Ribeiro LA, El-Jaick KB, Muenke M, Richieri-Costa A. SIX3 mutations with holoprosencephaly. Am J Med Genet 2006;140:2577-83.
- 49 Wilson WG, Shanks DE, Sudduth KW, Couper KA, McIlhenny J. Holoprosencephaly and interstitial deletion of 2(p2101p2109). Am J Med Genet 1989;34:252-4.

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REFERENCES

- Matsunaga E, Shiota K. Holoprosencephaly in human embryos: epidemiologic 1. studies of 150 cases. Teratology 1977;16:261-72.
- 2 Leoncini E, Baranello G, Orioli IM, Annerén G, Bakker M, Bianchi F, Bower C, Canfield MA, Castilla EE, Cocchi G, Correa A, De Vigan C, Doray B, Feldkamp ML, Gatt M, Irgens LM, Lowry RB, Maraschini A, Mc Donnell R, Morgan M, Mutchinick O, Poetzsch S, Riley M, Ritvanen A, Gnansia ER, Scarano G, Sipek A, Tenconi R, Mastroiacovo P. Frequency of holoprosencephaly in the International Clearinghouse Birth Defects Surveillance Systems: Searching for population variations. Birth Defects Res A Clin Mol Teratol 2008:82:585-91.
- 3 Muenke M, Beachy PA. Genetics of ventral forebrain development and holoprosencephaly. Curr Opin Genet Dev 2000;10:262-9.
- 4 Cohen MM Jr. Holoprosencephaly: clinical, anatomic, and molecular dimensions. Birth Defects Res A Clin Mol Teratol 2006;76:658-73.
- 5 Barkovich AJ, Simon EM, Clegg NJ, Kinsman SL, Hahn JS. Analysis of the cerebral cortex in holoprosencephaly with attention to the sylvian fissures. AJNR Am J Neuroradiol 2002;23:143-50.
- Barkovich AJ, Quint DJ. Middle interhemispheric fusion: an unusual variant of 6. holoprosencephaly. AJNR Am J Neuroradiol 1993;14:431-40.
- 7 Hahn JS, Barkovich AJ, Stashinko EE, Kinsman SL, Delgado MR, Clegg NJ. Factor analysis of neuroanatomical and clinical characteristics of holoprosencephaly. Brain Dev 2006;28:413-9.
- Lazaro L, Dubourg C, Pasquier L, Le Duff F, Blayau M, Durou MR, de la Pintière AT, 8 Aguilella C, David V, Odent S. Phenotypic and molecular variability of the holoprosencephalic spectrum. Am J Med Genet 2004;129A:21-4.
- 9. Dubourg C, Bendavid C, Pasquier L, Henry C, Odent S, David V. Holoprosencephaly. Orphanet J Rare Dis 2007;2:8.
- Plawner LL, Delgado MR, Miller VS, Levey EB, Kinsman SL, Barkovich AJ, Simon 10 EM, Clegg NJ, Sweet VT, Stashinko EE, Hahn JS. Neuroanatomy of holoprosencephaly as predictor of function: beyond the face predicting the brain. Neurology 2002;59:1058-66.
- Edison RJ, Muenke M. Mechanistic and epidemiologic considerations in the 11. evaluation of adverse birth outcomes following gestational exposure to statins. Am J Med Genet 2004;131:287-98.
- Edison RJ, Muenke M. Central nervous system and limb anomalies in case reports 12. of first-trimester statin exposure. N Engl J Med 2004;350:1579-82. Erratum in: N Engl J Med 2005:352:2759.
- 13. Cohen MM Jr. Perspectives on holoprosencephaly: Part I. Epidemiology, genetics, and syndromology. Teratology 1989;40:211-35.
- 14 Cohen MM Jr, Sulik KK. Perspectives on holoprosencephaly: Part II. Central nervous system, craniofacial anatomy, syndrome commentary, diagnostic approach, and experimental studies. J Craniofac Genet Dev Biol 1992;12:196-244.
- 15. Croen LA, Shaw GM, Lammer EJ. Holoprosencephaly: epidemiologic and clinical characteristics of a California population. Am J Med Genet 1996;64:465-72.
- 16. Roessler E, Belloni E, Gaudenz K, Jay P, Berta P, Scherer SW, Tsui LC, Muenke M. Mutations in the human Sonic Hedgehog gene cause holoprosencephaly. Nat Genet 1996:14:357-60.
- 17. Brown SA, Warburton D, Brown LY, Yu CY, Roeder ER, Stengel-Rutkowski S, Hennekam RC, Muenke M. Holoprosencephaly due to mutations in ZIC2, a homologue of Drosophila odd-paired. Nat Genet 1998:20:180-3.
- 18. Wallis D, Muenke M. Mutations in holoprosencephaly. Hum Mutat 2000;16: 99-108
- Wallis DE, Roessler E, Hehr U, Nanni L, Wiltshire T, Richieri-Costa A, Gillessen-19. Kaesbach G, Zackai EH, Rommens J, Muenke M. Mutations in the homeodomain of the human SIX3 gene cause holoprosencephaly. Nat Genet 1999; 22: 196-8.
- Gripp KW, Wotton D, Edwards MC, Roessler E, Ades L, Meinecke P, Richieri- Costa 20 A, Zackai EH, Massagué J, Muenke M, Elledge SJ. Mutations in TGIF cause holoprosencephaly and link NODAL signalling to human neural axis determination. Nat Genet 2000;25:205-8.
- de la Cruz JM, Bamford RN, Burdine RD, Roessler E, Barkovich AJ, Donnai D, Schier 21. AF, Muenke M. A loss-of-function mutation in the CFC domain of TDGF1 is associated with human forebrain defects. Hum Genet 2002;110:422-8.
- 22. Ming JE, Kaupas ME, Roessler E, Brunner HG, Golabi M, Tekin M, Stratton RF, Sujansky E. Bale SJ, Muenke M, Mutations in PATCHED-1, the receptor for SONIC HEDGEHOG, are associated with holoprosencephaly. Hum Genet 2002:110:297-301
- Roessler E, Du YZ, Mullor JL, Casas E, Allen WP, Gillessen-Kaesbach G, Roeder ER, 23. Ming JE, Ruiz i Altaba A, Muenke M. Loss-of-function mutations in the human GLI2 gene are associated with pituitary anomalies and holoprosencephaly-like features. Proc Natl Acad Sci USA 2003;100:13424-9. Roessler E, Ouspenskaia MV, Karkera JD, Vélez JI, Kantipong A, Lacbawan F, 24.

- Sawyer JR, Jones E, Hawks FF, Quirk JG Jr, Cunniff C. Duplication and deletion of chromosome band 2(p21p22) resulting from a familial interstitial insertion (2;11)(p21;p15). *Am J Med Genet* 1994;49:422–7.
- Zwetsloot CP, Brouwer OF, Maaswinkel-Mooy PD. Holoprosencephaly: variation of expression in face and brain in three sibs. J Med Genet 1989;26:274–6.
- Laflamme C, Filion C, Labelle Y. Functional characterization of SIX3 homeodomain mutations in holoprosencephaly: interaction with the nuclear receptor NR4A3/NOR1. *Hum Mutat* 2004;24:502–8.
- 53. Agresti A. An introduction to categorical data analysis. New York: John Wiley and Sons, 1996.

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