ARTICLE

16p11.2–p12.2 duplication syndrome; a genomic condition differentiated from euchromatic variation of 16p11.2

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Chromosome 16 contains multiple copy number variations (CNVs) that predispose to genomic disorders. Here, we differentiate pathogenic duplications of 16p11.2-p12.2 from microscopically similar euchromatic variants of 16p11.2. Patient 1 was a girl of 18 with autism, moderate intellectual disability, behavioural difficulties, dysmorphic features and a 7.71-Mb (megabase pair) duplication (16:21 521 005–29 233 146). Patient 2 had a 7.81-Mb duplication (16:21 382 561–29 191 527), speech delay and obsessional behaviour as a boy and, as an adult, short stature, macrocephaly and mild dysmorphism. The duplications contain 65 coding genes of which Polo-like kinase 1 (PLK1) has the highest likelihood of being haploinsufficient and, by implication, a triplosensitive gene. An additional 1.11-Mb CNV of 10q11.21 in Patient 1 was a possible modifier containing the G-protein-regulated inducer of neurite growth 2 (GPRIN2) gene. In contrast, the euchromatic variants in Patients 3 and 4 were amplifications from a 945-kb region containing non-functional immunoglobulin heavy chain (IGHV), hect domain pseudogene (HERC2P4) and TP53-inducible target gene 3 (TP53TG3) loci in proximal 16p11.2 (16:31953353-32898635). Paralogous pyrosequencing gave a total copy number of 3–8 in controls and 8 to >10 in Patients 3 and 4. The 16p11.2– p12.2 duplication syndrome is a recurrent genomic disorder with a variable phenotype including developmental delay. dysmorphic features, mild to severe intellectual disability, autism, obsessive or stereotyped behaviour, short stature and anomalies of the hands and fingers. It is important to differentiate pathogenic 16p11.2-p12.2 duplications from harmless, microscopically similar euchromatic variants of proximal 16p11.2, especially at prenatal diagnosis. European Journal of Human Genetics (2013) 21, 182–189; doi:10.1038/ejhg.2012.144; published online 25 July 2012

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INTRODUCTION

The proximal short arm of chromosome 16 contains a number of copy number variation (CNV) hotspots¹ that predispose to at least four reciprocal deletions and duplications within bands 16p11.2-16p12.2 (Figure 1). From centromere to telomere, these include first, the common microdeletions/duplications of ~ 600 kb in 16p11.2 associated with neurocognitive difficulties and obesity,²⁻⁴ second, the microscopically visible duplications of 8-9 Mb from 16p11.2 to 16p12.1/2 associated with developmental delay and autism⁵⁻¹¹ and the reciprocal microdeletions associated with developmental delay, intellectual disability and subtle dysmorphic features;^{12,13} within these are, third, the distal (formerly atypical) microdeletions of \sim 220 kb in 16p11.2 associated with a phenotype that includes developmental delay or obesity but extends into the normal range¹⁴⁻¹⁶ and, fourth, the microdeletions/duplications of \sim 520 kb in 16p12.1 associated with developmental delay.^{1,17} Here, we report two new patients with cytogenetically visible duplications of 16p11.2-16p12.2 analysed using oligonucleotide array comparative

genomic hybridisation (oaCGH) and compared with 10 previous postnatal patients. $^{5\mathrm{-11}}$

CNV is common with over 66000 examples recorded in the Database of Genomic Variants (DGV) and an estimated 1% of the human population having a $CNV > 1 Mb.^{1}$ When copy number is high enough, rare CNVs of 8p23.1, 9p12, 9qh/q12, 9q13, 15q11.2 and 16p11.2 become visible in the light microscope and have been described as euchromatic variants.18 The euchromatic variants of 16p11.2 were originally defined using semi-quantitative cosmid fluorescence in situ hybridisation (FISH) and believed to result from constitutional cytogenetic amplification of paralogous sequences including non-functional IGVH, creatine transporter (SLC6A10P) and TP53TG3 genes.¹⁸⁻²⁰ Euchromatic variants of 16p11.2 can be mistaken for pathogenic duplications resulting in a termination of pregnancy in at least one documented case.²⁰ Here, we report two new cases in which bacterial artificial chromosome (BAC) FISH was used to assign the amplicons to an \sim 1-Mb region of proximal 16p11.2. Pyrosequencing²¹ was used to quantify amplicon

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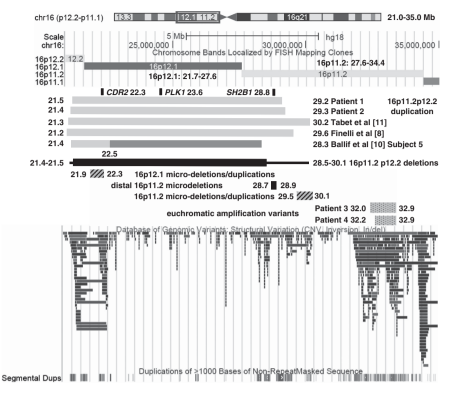


Figure 1 UCSC browser screenshot annotated with extent of the duplications, triplication, deletions and euchromatic variants of proximal 16p in the context of the UCSC browser bands and data on genomic variants and segmental duplications (NCBI 36/hg18): included are the 16p11.2–p21.2 duplications in Patients 1 and 2 (light grey bars), the 16p11.2–p21.2 duplications of Finelli *et al*⁸ (light grey bar), the 16p11.2–p21.2 duplications (AcK grey bar) and duplication (light grey bar) of Ballif *et al*, ¹⁰ subject 5, the duplications of Tabet *et al*,¹¹ the reciprocal 16p11.2–p21.2 'micro'deletions (dark grey bar with narrow bars at either end indicating variable extent), the 16p12.1 microdeletions and duplications (diagonally hatched light and dark grey bar), the distal (formerly atypical) 16p11.2 microdeletions (dark grey bar), the common 16p11.2 microdeletions/duplications (diagonally hatched light and dark grey bars) and the euchromatic amplification variants in Patients 3 and 4 (dotted bars on a light grey background). The approximate start and end points of bands, imbalances, variants and candidate genes (*CDR2*, *PLK1* and *SH2B1*) are given in Mb.

copy number and corroborate estimates made using semi-quantitative FISH in euchromatic variant carriers and controls.

MATERIALS AND METHODS

Conventional G-banded chromosome analysis was carried out in lymphocytes (Patients 1 and 2) and amniotic fluid cells (Patients 3 and 4) using standard techniques. FISH was carried out with non-functional *IGVH* (cos11, 13, 33, 97 and 98) and creatine transporter (c329B6) cosmids as before.¹⁹ Dual colour BAC FISH and oaCGH with a customised 4×44 K Agilent oligonucleotide array (Santa Clara, CA, USA) were performed as described previously.²² Base pair coordinates correspond to NCBI release 36.1 (hg18). Copy number in euchromatic variant carriers and normal controls was determined by coamplifying paralogous sequences at 16p11.2 and Xq28²³ and quantifying the paralogous sequence mismatches (PSM) between them using pyrosequencing according to the manufacturer's instructions. The resultant PSM allele frequencies were calculated using the Pyrosequencing AQ software (Biotage AB, Uppsala, Sweden) and reflect the relative frequency of the paralogous sequence on each chromosome.

Clinical reports

Patient 1. This 18-year-old girl (Figures 2a–e) was the first child of healthy unrelated Caucasian parents. She had a younger brother aged 15 with normal development. The pregnancy was complicated by hyperemesis. She was born at 36 weeks gestation by spontaneous vaginal delivery weighing 2920 g (90th centile). Discharge was delayed by 5 days because of neonatal jaundice. She was a poor feeder owing to poor suck and also needed waking for feeds. She had poor weight gain and lactose intolerance. Weaning to solid food was slow and achieved by 3 years of age. Developmentally she sat at 8 months and walked at

13 months. By 18 months she had four words with meaning. During infancy, she was asthmatic and prone to respiratory tract infections but had outgrown this by the age of 9. She had otitis media and grommets inserted at the age of 3. In childhood she was a poor sleeper but this improved after the age of 12. She had vacant staring episodes but no seizures and her EEG was normal. She was toilet trained aged 7 years. She was small for her age until the age of 12 but has since caught up. She had onset of menarche aged 14 years. She has had orthodontic treatment since the age of 15 years. A Ruth Griffiths assessment at chronological age 51 months showed mental age results ranging from 28 to 40 months (sub quotient 55–78). From the age of 5 years she attended a special school for moderate learning difficulties. Since the age of 16 years she has been attending a transition college and doing a life skills course. She can dress herself, but needs help with small buttons.

From early on her behaviour was described as fixed and repetitive. She did not like plans to be changed. She would prepare well in advance for any event out of her normal routine and become fixated on it. She fixated on certain ideas or phrases and would repeat a phrase over and over to herself. She generally enjoyed meeting new people and talked very quickly to them without accounting for their lack of prior knowledge. She could be inappropriately familiar and affectionate with strangers. At other times she would happily entertain herself and not interact with others.

Clinical evaluation at the age of 15 revealed a weight of 48 kg (<25th centile), height of 158.2 cm (<25th centile) and an OFC of 55 cm (<50th centile). She had a prominent glabella, heavy eyebrows, synophrys, slight ptosis, left convergent squint, broad nasal bridge, an upturned nose with a slightly bulbous tip, bow shaped upper lip, full everted lower lip, prominent upper incisors, dental crowding and overfolded helices (Figures 2a–c). She had tapering, hypermobile fingers with brachydactyly, squared off finger tips (Figure 2d) and prominent foetal finger pads. She has had problems with painful flat feet and had bilateral medial bony

183

184

protuberances near the ankles for which she has worn orthotics (Figure 2e). Her skin was soft and velvety with wrinkled skin on the palms and soles. Her hands were clammy. She had a postural kyphosis. Neurological examination revealed poor balance and coordination.

She had an autism diagnostic interview-revised (ADI-R) assessment and observations by an occupational therapist, an art therapist and a consultant psychiatrist at the age of 16 years and 9 months and the results confirmed the diagnosis of autism. The assessment showed difficulties with non-verbal regulation of social interaction, difficulty in developing peer relationships and a lack of shared enjoyment and socio-emotional reciprocity. She had difficulty initiating and sustaining conversation in a reciprocal way, some stereotyped and repetitive speech and did not have the social awareness to avoid asking inappropriate things at inappropriate times. The assessment showed that she had more difficulties with minor changes in routine or environment when she was younger compared with when she was assessed. Her parents had made a great deal of effort to avoid any changes of her routine to prevent distress. She has become more aware of her difficulties with time

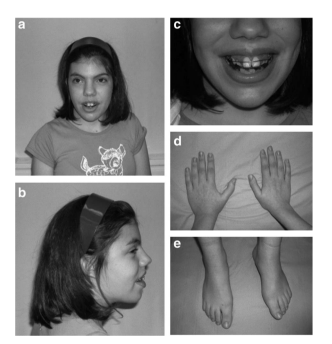


Figure 2 (a–e) The phenotype of Patient 1 aged 15 years: (a) Facial features showing heavy eyebrows, synophrys, slight ptosis, left convergent squint, broad nasal bridge, bow shaped upper lip and full lower lip; (b) lateral facial features showing overfolded helix, prominent glabella, upturned nose with a slightly bulbous tip and full everted lower lip; (c) oral features showing prominent upper incisors and dental crowding; (d) hands of the proband showing brachydactyly and squared off finger tips; and (e) feet of the proband showing bilateral medial protuberances near her ankles.

and she gets frustrated at not being able to do things alone. As a result, she has been verbally and physically aggressive towards her brother and peers at times. At college, one to one support is being put in place to try and help her behaviour. She enjoys reading and writing.

Patient 2 (Database of Chromosomal Imbalance and Phenotype in Humans Using Ensembl Resources (DECIPHER) 2131). This male patient was referred at the age of 45 so that his elderly parents could plan for his future. He had been a full-term normal delivery with a good birth weight and no very early health concerns. He was a little late to walk but his speech was markedly delayed and a diagnosis of glue ear was made. He attended special school from the age of 6 after an extra year at nursery school. He was a generally good natured boy with some obsessional behaviour. He had an abnormal fixation on buses and trucks, which involved large amounts of time engaged in spinning the wheels, but is not regarded as having had autism. He can now read well but cannot write more than a poor rendition of his name and is not good with numbers. He was unable to pass any exams but does have a number of skills such as excellent map-reading ability. He was generally healthy as a child but did have one episode of unexplained hypercalcemia. Since leaving school he has had jobs in sheltered employment. As an adult he has been well other than recurrent rectal prolapse. On examination, he is short at 158 cm (0.4th centile), has an OFC of 59 cm (98th centile) and is of average build. He has a wide mouth, a large furrowed tongue and broad alveolar margins. His hands are broad with a generalised brachydactyly and his digits are untapered. He has one large lipoma on his back and freckling on the palms and soles. A diagnosis of the Bannavan-Rilev-Ruvalcaba syndrome was considered but molecular analysis of the phosphatase and tensin homologue (PTEN) gene was negative.

Patient 3. This healthy 43-year-old woman was in her third pregnancy and delivered a boy with normal measurements at the fortieth week of pregnancy (weight 3160 g, length 54 cm, OFC 34 cm and APGAR scores 10/10/10). The healthy boy is now 6 years old and will start school in the autumn of this year.

Patient 4. This patient was referred for prenatal diagnosis after two miscarriages. No further clinical details were available.

RESULTS

Patient 1

An extra G-dark band was present in the proximal short arm of chromosome 16 (Figure 3a). FISH with non-functional *IGVH* (cos33 and 98) and creatine transporter cosmids (c329B6) from proximal 16p11.2¹⁹ was normal but signals with BAC CTB2515A14 from 16p12.1 were duplicated (data not shown) (Table 1). OaCGH confirmed a duplication of 16p12.2–16p11.2 with a minimum size of 7.71 Mb and a maximum of 8.07 Mb (Figure 4e) (Table 1). An additional duplication CNV of band 10q11.21 had a minimum size of 1.11 Mb and a maximum of 2.47 Mb. Parental chromosomes were normal. The karyotype of the proband was: 46,XX, dup (16) (p11.2–p12.2)dn.ish dup (16) (p11.2–p12.2)(CTB2515 A14 + +,CTB74E23 + RP11-165M2 +).arr 16p12.2p11.2(21 507 188

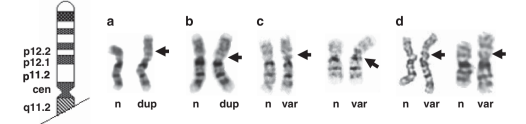


Figure 3 (a-d) G-banded partial karyotypes from Patients 1, 2, 3 and 4 with the normal chromosome (n) on the left and the duplicated (dup) or euchromatic variant (var) chromosome 16 on the right: (a) Patient 1 and (b) Patient 2 with the duplicated region of chromosome 16 indicated by the large black arrows; (c) Patient 3 and (d) Patient 4 with the euchromatic variant region of chromosome 16 indicated by the large black arrows. Note the similarity between the G-banded duplications in the Patients 1 and 2 (a, b) and the euchromatic variants in Patients 3 and 4 (c, d).

× 2 21 521 005-29 233 146 × 3 29 581 455 × 2), 10q11.21 (45 480 521 × 2 46 371 243-47 485 737 × 3 47 946 216 × 2).

Patient 2 (DECIPHER 2131)

An extra G-dark band was present in the proximal short arm of chromosome 16 (Figure 3b) and duplicated signals were found using FISH with BAC RP11-142A12 from distal 16p12.1 (Figure 4a) (Table 1). OaCGH confirmed a duplication of 16p12.2-16p11.2 (Figure 4a) with a minimum of 7.81 Mb and a maximum of 8.37 Mb (Table 1). An additional duplication CNV of 1.52 Mb from Xp22.31 was found with a minimum size of 1.52 Mb. Parental chromosomes were normal and the karvotype of the proband was: 46,XY,dup (16) (p11.2-p12.2) dn.ish dup (16) (p11.2-12.2) (RP11-142A12,RP11-165M2 +).arr 16p12.1p11.2 (21 209 438 × 2 21 382 561-29 191 527 × 3 29 581 455 × 2), Xp22.31(6 467 403-7 992 061) × 3.

Patient 3

An extra GTG-positive band in proximal 16p was found during chromosome analysis of amniotic fluid cultures and lymphocytes from the mother (Figure 3c). The extra band was CBG negative but enhanced (enh) signals were seen using FISH with IGVH cosmids (cos11, 13, 33, 97 and 98) (data not shown)¹⁹ Enh signals were also seen with BACs RP11-408D2 (Figure 4b) and RP11-378C4. The results with BAC RP5-1044J9 were less convincing than those in Patient 4 implying an amplicon of 692 kb (Table 1). The duplicated signals with BAC RP11-410P5 are likely to represent independent variation as they are non-contiguous. The karyotype was 46,XX, var (16) (p11.2p11.2) mat.ish var (16) (RP11-410P5++,RP11-196G11+,RP5-1044J9?enh,RP11-408D2enh,RP11-378C4enh,RP11-488I20+,RP11-80F22+,RP11-65B9+). Pyrosequencing gave a total diploid copy number of 8 (Supplementary Figure 1c) and, assuming that the normal chromosome had the median copy number of 2, the euchromatic variant chromosome would have at least six copies of the amplicon spanning 4.2 Mb.

Table 1 FISH and oaCGH results

Patient 4

An extra GTG-positive band was found during chromosome analysis of amniotic fluid cultures (Figure 3d). No information on the karyotypes of the parents was available. Enh signals were seen using FISH with IGVH cosmids (cos11, 33 and 98; data not shown)¹⁹ and an increased interval between the 16p BAC RP11-67I10 and a 16q11.2-microdissection probe (Figure 4c). Enh signals were also found with BACs RP5-1044J9, RP11-408D2 and RP11- 378C4 (Figure 4d) (Table 1) implying an amplicon of 945 kb. Enh signals from non-contiguous BACs RP11-410P5 and RP11-80F22 (Table 1) were thought to be independent variation especially as RP11-80F22 is one of the commonest CNVs found using diagnostic oaCGH.²⁴ The formal karyotype was: 46,XY,var(16) (p11.2p11.2). ish var (16) (RP11-410P5enh,RP11-196G11+,RP5-1044J9enh,RP11-408D2enh,RP11-378C4enh,RP11-488I20 + ,RP11-80F22enh,RP11-65B9 +). Pyrosequencing gave a total diploid copy number of > 10 (Supplementary Figure 1d) and, if the normal chromosome had the median copy number of 2, the euchromatic variant chromosome would have at least eight copies of the amplicon spanning 7.6 Mb.

Copy number in controls

Paralogous pyrosequencing gave total copy numbers of between three and eight in 46 controls (20 males and 26 females) with a median value of four (Supplementary Figures 1a and b).

DISCUSSION

Our results indicate how similar large single-copy duplications of 16p11.2-p12.2 and amplification variants of 16p11.2 appear under the light microscope (Figures 3a-d) and how distinct they are at the molecular level (Figure 1). The importance of differentiating duplications from variants has been illustrated by the termination of a pregnancy when a euchromatic variant was misinterpreted as a duplication.²⁰ The presence or absence of the pathogenic duplication can be tested using FISH, qPCR or microarrays (Table 1). The same methods

Band	BAC/oligonucleotide	Position (Mb from telomere, hg18)	Patient 1	Patient 2	Patient 3	Patient 4
16p12.2 A_14_P201422		21 209 438		Normal		
16p12.2	A_14_P125957	21 382 561		Dup		
16p12.2			Normal	Dup	NT	NT
16p12.2	A_14_P110982	21 521 005	Dup	Dup	NT	NT
16p12.1	CTD-2515A14	24 637 484-24 856 494	Dup	NT	NT	NT
16p11.2	RP11-410P5	28082641-28266523	NT	NT	Dup	Enh
16p11.2	A_14_P114769	29 191 527	Dup	Dup	NT	NT
16p11.2	A_14_P107156	29 233 146	Dup	?Dup	NT	NT
16p11.2	A_14_P200817	29 581 455	Normal	Normal	NT	NT
16p11.2	RP11-74E23	29 550 782-29 724 963	Normal	NT	NT	NT
16p11.2	RP11-196G11	30862944-31045174	NT	NT	Normal	Normal
16p11.2	RP5-1044J9	31 953 353-32 079 506	NT	NT	?Enh	Enh
16p11.2	RP11-408D2	32206388-32345224	NT	NT	Enh	Enh
16p11.2	RP11-378C4	32722382-32898635	NT	NT	Enh	Enh
16p11.1/2	RP11-488I20	34 289 152-34 490 212	NT	NT	Normal	Normal
16p11.1	RP11-80F22	34476095-34627143	NT	NT	Normal	Enha
16p11.1	RP11-65B9	34 982 278-35 143 302	NT	NT	Normal	Normal
16cen	D16Z2		NT	NT	Normal	Normal
16qh	D16Z3		NT	NT	Normal	Normal
16q12.2	RP11-165M2	54 425 480–54 559 961	Normal	NT	NT	NT

Abbreviations: CNV, copy number variation; dup, duplication; enh, enhanced signal; FISH, fluorescence in situ hybridisation; NT, not tested; oaCGH, oligonucleotide array comparative genomic hybridisation. The Ensembl Genome browser G-dark band assignments are in bold. ^aCommon benign CNV,²⁴ grey shading highlights the regions involved in pathogenic duplication or euchromatic variation.

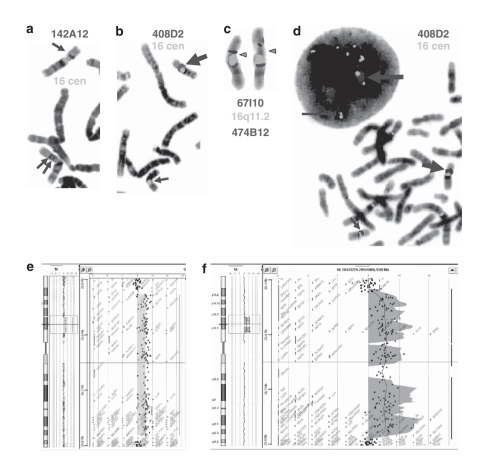


Figure 4 (**a**–**f**) Dual colour BAC FISH results and oaCGH results: (**a**) Distinct pairs of signals from RP11-142A12 (red arrows) indicating the duplication in Patient 2; (**b**) enh (large red arrow) and normal signals (small red arrow) with RP11-408D2 (red) consistent with amplification in a metaphase cell from Patient 3; (**c**) increased interval (blue arrows) between BAC RP11-67110 (red) and the 16q11.2 microdissection probe (green) in a pair of chromosomes 16 from Patient 4; (**d**) enh (large red arrows) and normal signals (small red arrows) with RP11-408D2 (red) consistent with amplification in interphase and metaphase cells from Patient 4 (the centromeres of chromosomes 1, 5 and 19 have also been labelled in green in these images); (**e**) and (**f**) oaCGH results from Patients 1 (**e**) and 2 (**f**) analysed with the Agilent Analytics software. The vertical coloured bars and background represent the extent of the duplications adjacent to idiograms of chromosome 16 in the left hand panels, magnified in the right hand panels, with black dots representing oligonucleotides with normal copy number and red dots oligonucleotides with increased copy number. The colour reproduction of this figure is available at the *European Journal of Human Genetics* journal online.

can be used to obtain results consistent with a euchromatic variant (Figures 4a–d), but care is needed in the interpretation of results from this copy number variable region, qPCR can be sensitive to template DNA quality and targeted arrays may lack adequate coverage and require careful choice of controls. Accurate determination of copy numbers higher than 10 remains problematic even with the pyrosequencing assay used here. The less likely possibility of an unbalanced insertion would require a combination of techniques. Both pathogenic duplications of 16p11.2–p12.1 and euchromatic variants of 16p11.2 have been directly transmitted from parents to children but one of the parents has been affected, to some extent, in all the duplications.⁷

16p11.2-p21.2 duplication phenotype

Table 2 summarises the differing phenotypic consequences in Patients 1 and 2 together with seven individuals with duplications of 16p11-16p13,^{5–9} monozygotic twins with a slightly larger duplication¹¹ and a patient with a triplication and duplication (Ballif *et al*,¹⁰ subject 5). A prenatal case was excluded as no sonographic anomalies were found before termination of the pregnancy.²⁵ Only developmental and/or psychomotor delay was present in all patients. In one patient and the twins, normal

development was followed by a marked decline after the age of 2. Intellectual disability in nine patients varied from severe in four to mild or moderate in others. Autism or autistic behaviour was recorded in eight patients but only Patient 1, the twins and one other patient had a formal ADI-R or DSM IV diagnosis. Patient 2 was not thought to have autism as a child. Obsessive and/or stereotyped behaviour was found in seven patients, social problems in six, ADHD or hyperactivity in four and echolalia in three. MRI/CT scan anomalies were noted in four patients and epilepsy, seizures and/or EEG anomalies in three.

Dysmorphism was seen in 10 patients but absent in one mother and daughter diad.⁷ Common dysmorphisms included a depressed, broad or large nasal bridge, upslanting or narrow palpebral features, hypertelorism and a long or tented philtrum. Short stature was found in five patients and microcephaly in four but normal OFC was found in Patient 1 and macrocephaly in Patient 2. Tapering, long, short and/or hypermobile fingers were found in eight patients. Patient 1 also had the prominent fingertip pads noted before (Ballif *et al*,¹⁰ subject 5) as well as brachydactyly and medial bony protuberances on both feet. Skin syndactyly of the feet was seen in three patients. Recurrent infections and nystagmus or strabismus were found in four

					þ			(01		
	11-p13	16p11-p13 16p11-p13 16p11.2-p1	16p11.2-p12	16p11.2-p12.1	16p11.2-p12.1	16p11.2-p12.2	16p11.2-p13.1	16p11.2-p12.2	16p11.2-p12.2	16p11.2-p12.2	16p11.2-p12.2
	L (- = L	- = L	00	00 - -	00 - -	- = L	6 trp;1 dup	8.95	7.71-8.07	7.81-8.37
SJ	35	Full term	Full term		Full term		Full term			30	Full term
Birth Weight (g) Sev (M/F)	2200	2030	3/00	ц	4050 (>p9/)	(50) UCCZ	DG 77	ц	(p4) (p4) M	2920 (p90) E	W
ace (W/17) Age at examination (vears)			~	35. 39 and 40	- גר	3, 25	1.5.4	- []	17	15	45
Height (cm)			I	158 (< D3)	103 (p3)	151 (0.03)	NR	-3 to -4 SDS	173 (mean)	158 (< p25)	p0.4
Weight (kg)				.	16.3 (p50)	42 (<p3)< td=""><td>NR</td><td>-2 SDS</td><td>49 (-2 SDS)</td><td>48 (< p25)</td><td>_</td></p3)<>	NR	-2 SDS	49 (-2 SDS)	48 (< p25)	_
	Micro			52.5 (<p3)< td=""><td>49 (p3)</td><td>54 (p25)</td><td>Micro</td><td>-1SDS</td><td>55 (-1 SDS)</td><td>55 (< p50)</td><td>Macro</td></p3)<>	49 (p3)	54 (p25)	Micro	-1SDS	55 (-1 SDS)	55 (< p50)	Macro
Dev/psychomotor delay	+	+	+	-+ TD	After 24/12	+	+	+++	After 26/12	+	+
Autism			Behaviour	DSM-IV	Partial	+	+	I	DSM-IV	ADI-R	I
Obesessive behaviours				+		+		+		+	+
Stereotyped behaviours				+	I	+			+	+	
Echolalia				+	+					+	
Social problems				+	+	+			+:	+	
				+				+,	Ŧ,		
Intellectual disability				Mild	Mild	Severe	+	Severe	Severe	Moderate	+
				00	00	AC < − CZ	-	47	-		
			_	I		+ -	÷		÷		
E E di Intranes E ni lonev/enizurae	I	_	+ -	I	I	+ -				I	
Lpriepay acizurea Ataxia	I	F	F	I	I	+ +				-	I
Hvnotonia	+	I	I	I	I	- 1	+			-	
Dysmorphism	+	+	+	None	I	+	- +	+	+	+	+
Depressed/large/broad nasal bridge	+	+	+	I	I	+	+	+	+	+	
	Upslant		Narrow	I	Upslant		Upslant	Narrow	Upslant	Ι	Ι
Hypertelorism		+	+	Ι	I	+	+	+	+		
Long philtrum	+	+	+	I	I	I	+	+		Ι	
Round face		I	+	I	I	I	+	+		I	
B		I		Ι	I	+	+			+	
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Uther single reatures				AU; EF; CU		5	CPC; HI	НА	EV; MH; IVV; VSU		FI; L; MSI

Table 2 Phenotypic features found in patients with postnatal duplications of proximal 16p

centile etc; SD, sternum deformity, SDS, SD, VSD, ventral septal defect (in twin P2 that spontaneously closed); +, present; -, absent. Greater than 10: chromosome z =1.28% (two copies), chromosome 16 = 87.2% (> 10 copies). ^bDsta added from Finelli *et a*,1¹¹ Patients 2 and 3.

16p11.2-p12.2 duplications and euchromatic variants JCK Barber et al

Table 3 UCSC genes within the 945-kb euchromatic CNV region of 16p11.2 (16:31953353-32898635) (hg18)

No	Start (bp)	End (bp)	Size (bp)	UCSC gene ID	UCSC description
1	31 957 501	31 985 174	27 673	IGVH (uc010cat.1)	Immunoglobulin A heavy chain mRNA IgA5741-27
2	31970984	31971143	159	IGHV3-30 (uc002ecu.1)	Immunoglobulin gamma heavy chain variable region mRNA for <i>IGHV3-30</i> gene, clone MGgis13b
3	31978049	31978136	87	IGH (uc002ecv.1)	IGHV VHDJ region mRNA, clone:H35
4	31984750	31 984 975	5	IGH (uc002ecw.1)	IGHV VHDJ region mRNA, clone:H155
5		32071375	1265	HERC2P4 (uc002ecx.1)	hect domain and RLD 2 pseudogene 4 mRNA (cDNA clone IMAGE:5416019)
6	32075067	32 085 501	10434	X69637 (uc010cau.1)	mRNA sequence
7		32174741	2596	TP53TG3b (uc002ecy.2)	TP53TG3b mRNA
8	32172145	32174741	2596	TP53TG3b (uc010cav.1)	TP53-inducible gene 3b mRNA
9			2438	BC042588 (uc002eda.2)	cDNA clone IMAGE:4826738
10	32208370	32 229 286	20916	<i>BC041879</i> (uc002edb.2)	Similar to protein phosphatase 2A 48 kDa regulatory subunit isoform 1; serine/threonine protein phosphatase 2A, 48 kDa regulatory subunit; PP2A, subunit B, PR48 isoform; PP2A B subunit PR48; NY-REN-8 antigen, mRNA (cDNA clone IMAGE:5272051).
11	32236530	32 236 557	27	DQ571479 (uc002edc.1)	DQ571479
12	32 592 342	32 596 379	4037	TP53TG3 (uc002edd.2)	TP53TG3
13	32 592 342	32 596 379	4037	TP53TG3b (uc002ede.2)	TP53TG3b mRNA
14	32688520	32 688 546	26	DQ574674 (uc002edg.1)	DQ574674
15	32693751	32695017	1266	BC038215 (uc010caw.1)	Hect domain and RLD 2 pseudogene 4 mRNA (cDNA clone IMAGE:5416019).
16	32679767	32690418	10651	LOC440366 (uc002edf.1)	Hect domain and RLD 2 pseudogene (LOC440366) non-coding RNA.
17	32796298	32 799 423	3125	SLC6A10P (uc002edh.1)	Solute carrier family 6 (neurotransmitter transporter, creatine), member 10
18	32796298	32 803 964	7666	SLC6A10P (uc002edi.1)	Pseudogene; cDNA FLJ43855 fis, clone TESTI4007163, highly similar to Sodium- and chloride-dependent creatine transporter 2.
19	32822426	32822513	87	IGHV (uc002edj.1)	IGHV variable region mRNA, clone N85.

Abbreviations: CNV, copy number variant; IGHV, immunoglobulin heavy chain; TP53TG3, TP53-inducible target gene 3.

Note: the graded background shading highlights loci that imply an inverted repeat structure; HGNC genes in bold.

patients each. Only one of the twins had a congenital heart defect (VSD) that resolved spontaneously.

Candidate genes

A minimum of 65 known coding genes are common to the duplication intervals in Patients 1 and 2 of which 11 are OMIM Morbid genes. Of these 11, many are autosomal recessive and only 3 have an estimated haploinsufficiency likelihood score (HLS), with implied dosage sensitivity, of <10%.²⁶ These include the sarcoplasmic/endoplasmic reticulum calcium ATPase 1 (ATP2A1) gene (8.8%), associated with autosomal recessive Brody myopathy, and the Amiloride-sensitive sodium channel subunits β and γ genes SCNN1B (9.2%) and SCNN1G (8.8%) associated with the autosomal dominant hypertensive disorder Liddle syndrome. Of the other 54 genes, the Polo-like kinase 1 (PLK1) gene has the lowest HLS of 0.4% and no other gene has an HLS of <10%. PLK1 is a protein kinase superfamily member and the human form of Drosophila polo, which is an important cell cycle regulator implicated in mammals as both oncogene and tumour suppressor. Polo mutants can switch development from neurons to additional neuroblasts in Drosophila,27 which suggests that dosage of PLK1 might conceivably have an effect on neuronal development and microcephaly in humans.

Despite higher HLS scores, the *cerebellar degeneration-related* protein 2 (CDR2) gene (46.9%) is a candidate for the 16p12.1 microdeletion syndrome²⁸ and might contribute to the developmental or behavioural problems when duplicated. Similarly, duplication of the Ataxin-2-like protein (ATXNL2) gene (31.5%) might be related to the ataxia seen in Patient 1 and Case 1 of Cohen *et al.*⁵ The SH2B adapter protein 1 (SH2B1) gene (19.2%) is a candidate for the developmental delay and obesity associated with the 200-kb microdeletions of 16p11.2. This gene lies within the minimum extent of the present and published 16p11.2–p12.2 duplications (Figure 1)¹⁵ but not within the triplication that has a more severe but similar phenotype to the duplications reported here (Ballif *et al*, ¹⁰ subject 5).

Modifiers or second hits

The phenotype of patients with 520-kb microdeletions of 16p12.1 was more severe when a second large CNV (gain or loss > 500 kb) was present and these microdeletions are thought to predispose to neuropsychiatric phenotypes in the presence of other large deletions or duplications.^{17,28} The genes within the remaining 7.2–7.3 Mb minimum interval in Patients 1 and 2 may therefore be sufficient to explain the cognitive phenotype in 16p11.2–p12.2 duplication syndrome but both patients had additional CNVs > 500 kb in size. In the female Patient 1 with autism, the additional 1.11-Mb duplication CNV of 10q11.2 overlaps multiple CNVs in the DGV but also contains a minimum of 9 genes. Of these, duplication of the *G-protein-regulated inducer of neurite growth 2 (GPRIN2)* gene is a candidate modifier of severity in Rett syndrome²⁹ and might conceivably interact with other candidate autism genes in 16p11.2–p12.2 duplications. In the male Patient 2 without autism, an additional duplication CNV of 1.52 Mb in Xp22.31 contains a minimum of 5 genes, including the *steroid sulfatase (STS)* gene, but there is growing evidence that duplication CNVs containing *STS* are benign population variants.^{30,31} The inheritance pattern of these two CNVs was not determined.

Breakpoints

The breakpoints mapped in the present patients with oaCGH and the previously reported patients with BAC- or M-FISH⁸⁻¹¹ correspond broadly to clusters of segmental duplications in proximal 16p (Figure 1). In both their families, Finelli et al⁸ mapped the telomeric breakpoints to BAC RP11-98D10 in 16p12.2 (16:21 354 957-21 532 536) and the centromeric breakpoints to BAC RP11-368N21 in 16p11.2 (16:29408797-29609842). These authors also suggested that the duplications are mediated by non-allelic homologous recombination (NAHR) between a low copy repeat (LCR16v) in 16p11.2 and 16p12.2 that contains a truncated paralogue of the SLCL7A5 gene from 16q24.3.8 Both the centromeric breakpoints in the present the Patients 1 and 2 lie within an interval of 390 kb in 16p11.2 (Table 1) that overlaps with BAC 368N21 and contains the SLCL7A5P1 gene. The telomeric breakpoints lie within different 173 kb and 13.8 kb intervals in 16p12.2 (Table 1) that overlap with BAC 98D10 but do not contain a copy of the SLCL7A5P1 gene or, in the case of Patient 1, correspond to any known segmental duplication. Similar variation has been recorded in the reciprocal deletion syndrome¹¹⁻¹³ (Figure 1) but further highresolution analysis of the breakpoints will be needed to establish whether all recurrent deletions and duplications of 16p11.2-16p12.2 arise by NAHR between a variety of repeats and/or via the more complex mechanisms being found in other genomic disorders.³²

Constitutional euchromatic amplification variants of 16p11.2

Including the present patients, euchromatic variants of 16p11.2 have been described in 14 families^{18,19} and have no known phenotypic consequences. The amplicons in Patients 3 and 4 have been approximately mapped with FISH (Table 1) and come from a 945kb region of proximal 16p11.2 (16:31 953 353–32 898 635) that contains 19 UCSC (University of California Santa Cruz) sequences of which only two are HUGO Gene Nomenclature Committee (HGNC) genes (Table 2). The region is flanked by non-functional *immunoglobulin heavy chain (IGHV)* genes, with *hect domain pseudogenes* (*HERC2P4*) and *TP53-inducible target gene 3 (TP53TG3)* protein loci that imply an asymmetrical palindromic structure (Table 3) that might mediate amplification.³³

A diploid copy number of 4 in normal individuals and 10–12 in 16p11.2 euchromatic variant carriers was previously estimated using semi-quantitative cosmid FISH.¹⁹ The results of paralogous pyrosequencing (previously reported in abstract³⁴) showed a wider range of copy number from 3 to 8 in controls and 8 to >10 in 16p11.2 euchromatic variant carriers. We could not quantify copy numbers greater than 10 but accurate, cost effective and high-throughput methods are needed to determine the possible effect of large CNVs on quantitative phenotypic traits in large cohorts.³⁵

CONCLUSIONS

Our results indicate that the 16p11.2–p12.2 duplication syndrome is a recurrent genomic disorder characterised by developmental delay, autistic and/or repetitive behaviour, dysmorphic features, microcephaly, short stature and tapering fingers. Duplications of 16p11.2–p12.1 can be mistaken for harmless euchromatic variants of proximal 16p11.2 in the light microscope but are distinct at the molecular level and can be distinguished using FISH or aCGH.

CONFLICT OF INTEREST

The authors declare no conflict of interest.

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WEB RESOURCES

DGV: http://projects.tcag.ca/variation/

DECIPHER v5.0: https://decipher.sanger.ac.uk/

Ensembl Genome Browser: http://www.ensembl.org/Homo_sapiens/Info/Index UCSC Genome Browser: http://genome.ucsc.edu/

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