

suggest that at least three quarters of patients with MR are undiagnosed by clinical dysmorphic features and karyotyping.

In the past two decades, a number of rapidly developed cytogenetic and molecular approaches have been applied to the screening or diagnosis of various congenital disorders including MR, congenital anomalies, recurrent abortion and cancer pathogenesis. Among them, array-based comparative genome hybridization (aCGH) is used to detect copy-number changes rapidly in a genome-wide manner and with high resolution. The target and resolution of aCGH depend on the type and/or design of mounted probes, and many types of microarray have been used for the screening of patients with MR and other congenital disorders: bacterial artificial chromosome (BAC)-based arrays covering whole genomes,^{9,10} BAC arrays covering chromosome X,^{11,12} a BAC array covering all subtelomeric regions,¹³ oligonucleotide arrays covering whole genomes,^{14,15} an oligonucleotide array for clinical diagnosis¹⁶ and a single nucleotide polymorphism array covering the whole genome.¹⁷ Because genome-wide aCGH has led to an appreciation of widespread copy-number variants (CNVs) not only in affected patients but also in healthy populations,^{18–20} clinical cytogeneticists need to discriminate between CNVs likely to be pathogenic (pathogenic CNVs, pCNVs) and CNVs less likely to be relevant to a patient's clinical phenotypes (benign CNVs, bCNVs).²¹ The detection of more CNVs along with higher-resolution microarrays needs more chances to assess detected CNVs, resulting in more confusion in a clinical setting.

We have applied aCGH to the diagnosis and investigation of patients with multiple congenital anomalies and MR (MCA/MR) of unknown etiology. We constructed a consortium with 23 medical institutes and hospitals in Japan, and recruited 536 clinically uncharacterized patients with a normal karyotype in conventional cytogenetic tests. Two-stage screening of copy-number changes was performed using two types of BAC-based microarray. The first screening was performed by a targeted array and the second screening was performed by an array covering the whole genome. In this study, we diagnosed well-known genomic disorders effectively in the first screening, assessed the pathogenicity of detected CNVs to investigate an etiology in the second screening and discussed the clinical significance of aCGH in the screening of congenital disorders.

MATERIALS AND METHODS

Subjects

We constructed a consortium of 23 medical institutes and hospitals in Japan, and recruited 536 Japanese patients with MCA/MR of unknown etiology from July

2005 to January 2010. All the patients were physically examined by an expert in medical genetics or a dysmorphologist. All showed a normal karyotype by conventional approximately 400–550 bands-level G-banding karyotyping. Genomic DNA and metaphase chromosomes were prepared from peripheral blood lymphocytes using standard methods. Genomic DNA from a lymphoblastoid cell line of one healthy man and one healthy woman were used as a normal control for male and female cases, respectively. All samples were obtained with prior written informed consent from the parents and approval by the local ethics committee and all the institutions involved in this project. For subjects in whom CNV was detected in the first or second screening, we tried to analyze their parents as many as possible using aCGH or fluorescence *in situ* hybridization (FISH).

Array-CGH analysis

Among our recently constructed in-house BAC-based arrays,²² we used two arrays for this two-stage survey. In the first screening we applied a targeting array, 'MCG Genome Disorder Array' (GDA). Initially GDA version 2, which contains 550 BACs corresponding to subtelomeric regions of all chromosomes except 13p, 14p, 15p, 21p and 22p and causative regions of about 30 diseases already reported, was applied for 396 cases and then GDA version 3, which contains 660 BACs corresponding to those of GDA version 2 and pericentromeric regions of all chromosomes, was applied for 140 cases. This means that a CNV detected by GDA is certainly relevant to the patient's phenotypes. Subsequently in the second screening we applied 'MCG Whole Genome Array-4500' (WGA-4500) that covers all 24 human chromosomes with 4523 BACs at intervals of approximately 0.7 Mb to analyze subjects in whom no CNV was detected in the first screening. WGA-4500 contains no BACs spotted on GDA. If necessary, we also used 'MCG X-tiling array' (X-array) containing 1001 BAC/PACs throughout X chromosome other than pseudoautosomal regions.¹² The array-CGH analysis was performed as previously described.^{12,23}

For several subjects we applied an oligonucleotide array (Agilent Human Genome CGH Microarray 244K; Agilent Technologies, Santa Clara, CA, USA) to confirm the boundaries of CNV identified by our in-house BAC arrays. DNA labeling, hybridization and washing of the array were performed according to the directions provided by the manufacturer. The hybridized arrays were scanned using an Agilent scanner (G2565BA), and the CGH Analytics program version 3.4.40 (Agilent Technologies) was used to analyze copy-number alterations after data extraction, filtering and normalization by Feature Extraction software (Agilent Technologies).

Fluorescence *in situ* hybridization

Fluorescence *in situ* hybridization was performed as described elsewhere²³ using BACs located around the region of interest as probes.

RESULTS

CNVs detected in the first screening

In the first screening, of 536 cases subjected to our GDA analysis, 54 (10.1%) were determined to have CNV (Figure 1; Tables 1 and 2).

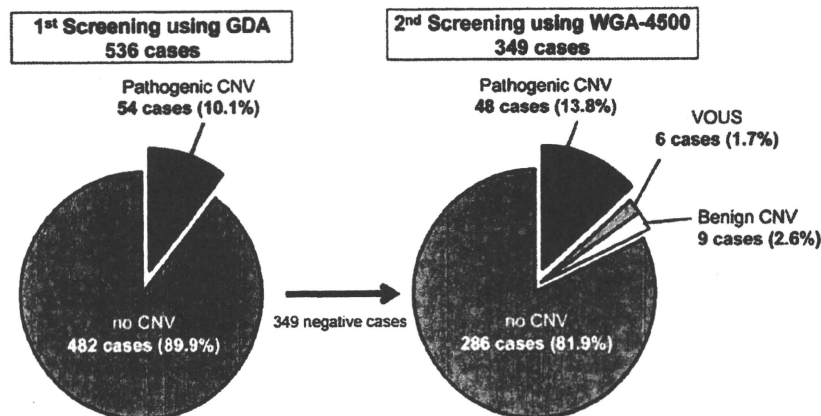


Figure 1 Percentages of each screening in the current study.

Table 1 A total of 40 cases with CNV at subtelomeric region(s) among 54 positive cases in the first screening

Gender	Position where CNV detected		Corresponding disorder ^a	OMIM or citation	Parental analysis ^b
	Loss	Gain			
M	1p36.33		Chromosome 1p36 deletion syndrome	#607872	
M	1p36.33p36.32		Chromosome 1p36 deletion syndrome	#607872	
M	1p36.33p36.32		Chromosome 1p36 deletion syndrome	#607872	
M	1p36.33p36.32		Chromosome 1p36 deletion syndrome	#607872	
M	1q44		Chromosome 1q43-q44 deletion syndrome	#612337	
F	2q37.3		2q37 monosomy ^c	Shrimpton <i>et al.</i> ²⁴	
F	2q37.3		2q37 monosomy ^c	Shrimpton <i>et al.</i> ²⁴	
M	3q29		Chromosome 3q29 deletion syndrome	#609425	
F	5p15.33p15.32		Cri-du-chat syndrome	#123450	
M	5q35.2q35.3		Chromosome 5q subtelomeric deletion syndrome	Rauch <i>et al.</i> ²⁵	
F	6p25.3		Chromosome 6pter-p24 deletion syndrome	#612582	
M	7q36.3		7q36 deletion syndrome ^d	Horn <i>et al.</i> ²⁶	
F	7q36.3		7q36 deletion syndrome ^d	Horn <i>et al.</i> ²⁶	
M	9p24.3p24.2		Chromosome 9p deletion syndrome	#158170	
F	9q34.3		Kleefstra syndrome	#610253	
F	10q26.3		Chromosome 10q26 deletion syndrome	#609625	
F	16p13.3		Chromosome 16p13.3 deletion syndrome	#610543	
F	22q13.31		Chromosome 22q13 deletion syndrome	#606232	
M	22q13.31q13.33		Chromosome 22q13 deletion syndrome	#606232	
M		15q26.3	15q overgrowth syndrome ^e	Tatton-Brown <i>et al.</i> ²⁷	
F		15q26.3	15q overgrowth syndrome ^e	Tatton-Brown <i>et al.</i> ²⁷	
M		21q22.13q22.3	Down's syndrome (partial trisomy 21)	#190685	
M		Xp22.33	A few cases have been reported; e.g. V5-130 in Lu <i>et al.</i> ²⁸		
M		Xq28	Chromosome Xq28 duplication syndrome	#300815	
F	1q44		Chromosome 1q43-q44 deletion syndrome	#612337	
		8p23.2p23.3			
M	3p26.3		3p deletion syndrome ^d	Fernandez <i>et al.</i> ²⁹	
		12p13.33p11.22			
F	3p26.3		3p deletion syndrome ^d	Fernandez <i>et al.</i> ²⁹	
		16p13.3	Chromosome 16p13.3 duplication syndrome	#613458	
F	4q35.2		4q- syndrome ^d	Jones <i>et al.</i> ³⁰	
		7q36.3			
M	5p15.33		Cri-du-chat syndrome	#123450	
		20p13			
M	5p15.33p15.32		Cri-du-chat syndrome	#123450	
		2p25.3			
F	6q27		6q terminal deletion syndrome ^d	Striano <i>et al.</i> ³¹	
		11q25			
F	6q27		6q terminal deletion syndrome ^d	Striano <i>et al.</i> ³¹	
		8q24.3			
M	7q36.3		7q36 deletion syndrome ^d	Horn <i>et al.</i> ²⁶	<i>dn</i>
		1q44			
M	9p24.3p24.2		Chromosome 9p deletion syndrome	#158170	
		7q36.3			
F	10p15.3p15.2		Chromosome 10p terminal deletion ^d	Lindstrand <i>et al.</i> ³²	<i>pat</i>
		7p22.3p22.2			
M	10p15.3		Chromosome 10p terminal deletion ^d	Lindstrand <i>et al.</i> ³²	
		2p25.3			
M	10q26.3		Chromosome 10q26 deletion syndrome	#609625	
		2q37.3	Distal trisomy 2q ^d	Elbracht <i>et al.</i> ³³	
M	18q23		Chromosome 18q deletion syndrome	#601808	
		7q36.3			
F	22q13.31q13.33		Chromosome 22q13.3 deletion syndrome	#606232	<i>pat</i>
		17q25.3	One case was reported	Lukusa <i>et al.</i> ³⁴	
M	Xp22.33/yp11.32		Contiguous gene-deletion syndrome on Xp22.3 ^d	Fukami <i>et al.</i> ³⁵	
		Xq27.3q28	Chromosome Xq28 duplication syndrome	#300815	

Abbreviations: F, female; CNV, copy-number variant; M, male; OMIM, Online Mendelian Inheritance in Man; *dn*, *de novo* CNV observed in neither of the parents.

^aThe name of disorder is based on entry names of OMIM, except for entry names in DECIPHER and description in each cited article.

^b*pat*, father had a balanced translocation involved in corresponding subtelomeric regions.

^cEntry names in DECIPHER.

^dDescription in each cited article.

All the CNVs detected in the first screening were confirmed by FISH. Among the positive cases, in 24 cases one CNV was detected. All the CNVs corresponded to well-established syndromes or already described disorders (Table 1). In 16 cases two CNVs, one deletion and one duplication, were detected at two subtelomeric regions, indicating that one of parents might be a carrier with reciprocal translocation involved in corresponding subtelomeric regions, and at least either of the two CNVs corresponded to the disorders. We also performed parental analysis by FISH for three cases whose parental samples were available, and confirmed that in two cases the subtelomeric aberrations were inherited from paternal balanced translocation and in one case the subtelomeric aberrations were *de novo* (Table 1). In the other 14 cases, CNVs (25.9%) were detected in regions corresponding to known disorders (Table 2).

CNVs detected in the second screening and assessment of the CNVs
Cases were subject to the second screening in the order of subjects detected no CNV in the first screening, and until now we have analyzed 349 of 482 negative cases in the first screening. In advance, we excluded highly frequent CNVs observed in healthy individuals and/or in multiple patients showing disparate phenotypes from the present results based on an internal database, which contained all results of aCGH analysis we have performed using WGA-4500, or other available online databases; for example, Database of Genomic Variant (<http://projects.tcag.ca/variation/>). As a result, we detected 66 CNVs in 63 cases (Figure 1; Table 3). Among them, three patients (cases 36, 42 and 44) showed two CNVs. All the CNVs detected in the second screening were confirmed by other cytogenetic methods including FISH and/or X-array. For 60 cases, we performed FISH for confirmation and to determine the size of each CNV. For five cases, cases 13, 36, 48, 57 and 63, with CNVs on the X chromosome, we used the X-array instead of FISH. For cases 4, 6, 16–19 and 34, we also used Agilent Human Genome CGH Microarray 244K to determine the refined sizes of CNVs. The maximum and minimum sizes of each CNV determined by these analyses are described in Table 3.

Well-documented pCNVs emerged in the second screening

CNVs identified for recently established syndromes. We assessed the pathogenicity of the detected CNVs in several aspects (Figure 2).^{21,37,38} First, in nine cases, we identified well-documented pCNVs, which are responsible for syndromes recently established. A heterozygous deletion at 1q41–q42.11 in case 2 was identical to patients in the first report of 1q41q42 microdeletion syndrome.³⁹ Likewise a CNV in case 3 was identical to chromosome 1q43–q44 deletion syndrome (OMIM: #612337),⁴⁰ a CNV in case 4 was identical to 2q23.1 microdeletion syndrome,⁴¹ a CNV in case 5 was identical to 14q12 microdeletion syndrome⁴² and a CNV in case 6 was identical to chromosome 15q26–qter deletion syndrome (Drayer's syndrome) (OMIM: #612626).⁴³ Cases 7, 8 and 9 involved CNVs of different sizes at 16p12.1–p11.2, the region responsible for 16p11.2–p12.2 microdeletion syndrome.^{44,45} Although an interstitial deletion at 1p36.23–p36.22 observed in case 1 partially overlapped with a causative region of chromosome 1p36 deletion syndrome (OMIM: #607872), the region deleted was identical to a proximal interstitial 1p36 deletion that was recently reported.⁴⁶ Because patients with the proximal 1p36 deletion including case 1 demonstrated different clinical characteristics from cases of typical chromosome 1p36 deletion syndrome, in the near term their clinical features should be redefined as an independent syndrome.⁴⁶

CNVs containing pathogenic gene(s). In four cases we identified pCNVs that contained a gene(s) probably responsible for phenotypes. In case 10, the CNV had a deletion harboring *GLI3* (OMIM: *165240)

Table 2 Other cases among 54 positive cases in the first screening

Gender	Position where CNV detected		Corresponding disorder	OMIM
	Gain	Loss		
F		4p16.3 4q35.2	Ring chromosome	
M		3q22.323	BPES	#110100
M		2q22.3	ZFX1B region	*605802
M		4q22.1	Synuclein (SNCA) region	*163890
F		7p21.1	Craniosynostosis, type 1	#123100
F		7q11.23	Williams syndrome	#194050
F		8q23.3q24.11	Langer–Giedion syndrome	#150230
M	15q11.2q13.1		Prader–Willi/Angelman	#176270/ #105830
F		17p11.2	Smith–Magenis syndrome	#182290
M		17q11.2	Neurofibromatosis, type I	+162200
M	22q11.21		DiGeorge syndrome	#188400
F		22q11.21	DiGeorge syndrome	#188400
F	Xp22.31		Kallmann syndrome 1	+308700
F	Whole X		Mosaicism	

Abbreviations: CNV, copy-number variant; F, female; M, male; OMIM, Online Mendelian Inheritance in Man.

accounting for Greig cephalopolysyndactyly syndrome (GCS; OMIM: 175700).⁴⁷ Although phenotypes of the patient, for example, pre-axial polydactyly of the hands and feet, were consistent with GCS, his severe and atypical features of GCS, for example, MR or microcephaly, might be affected by other contiguous genes contained in the deletion.⁴⁸ Heterozygous deletions of *BMP4* (OMIM: *112262) in case 11 and *CASK* (OMIM: *300172) in case 13 have been reported previously.^{49,50} In case 12, the CNV contained *YWHAE* (OMIM: *605066) whose haploinsufficiency would be involved in MR and mild CNS dysmorphism of the patient because a previous report demonstrated that haploinsufficiency of *ywhae* caused a defect of neuronal migration in mice⁵¹ and a recent report also described a microdeletion of *YWHAE* in a patient with brain malformation.⁵²

Recurrent CNVs in the same regions. We also considered recurrent CNVs in the same region as pathogenic; three pairs of patients had overlapping CNVs, which have never been reported previously. Case 16 had a 3.3-Mb heterozygous deletion at 10q24.31–q25.1 and case 17 had a 2.0-Mb deletion at 10q24.32–q25.1. The clinical and genetic information will be reported elsewhere. Likewise, cases 14 and 15 also had an overlapping CNV at 6q12–q14.1 and 6q14.1, and cases 18 and 19 had an overlapping CNV at 10p12.1–p11.23. Hereafter, more additional cases with the recurrent CNV would assist in defining new syndromes.

CNVs reported as pathogenic in previous studies. Five cases were applicable to these criteria. A deletion at 3p21.2 in case 20 overlapped with that in one case recently reported.⁵³ The following four cases had CNVs reported as pathogenic in recent studies: a CNV at 7p22.1 in case 21 overlapped with that of patient 6545 in a study by Friedman *et al.*,¹⁴ a CNV at 14q11.2 in case 22 overlapped with those of patients 8326 and 5566 in Friedman *et al.*,¹⁴ a CNV at 17q24.1–q24.2 in case 23 overlapped with that in patient 99 in Buysse *et al.*⁵⁴ and a CNV at 19p13.2 in case 24 overlapped with case P11 in Fan *et al.*⁵⁵

Large or gene-rich CNVs, or CNVs containing morbid OMIM genes. In cases inapplicable to the above criteria, we assessed CNVs

Table 3 Sixty-three cases with CNV in the 2nd screening

Case	Gender	Clinical diagnosis	Remarkable clinical features	CNV Position	WGA-4500 ^b	Base position and size of the identified CNV ^a						Protein-CNV Parental coding analysis genes ^c	Corresponding assess- or candidate men ^d genes(s)		
						Start (max)	Start (min)	End (min)	End (max)	Size (min)	Size (max)				
1	M	MCA/MR		del 1p36.23p36.22	arr cgh 1p36.23p36.22 (RP11-81J7 → RP11-19901)x1	del 1p36.23p36.22	8585127	8890860	10561097	11143717	1670237	2558590	dn	32	P
2	M	MCA/MR		del 1q41q42.11	arr cgh 1q41 (RP11-135J2 → RP11-239E10)x1	del 1q41q42.11	215986492	216532600	221534398	222467931	5001798	6481439	dn	35	P
3	F	MCA/MR	Epilepsy	del 1q44	arr cgh 1q44 (RP11-156E8)x1	del 1q44	241996973	243177632	243251660	244141010	74028	2144037		11	P
4	F	MCA/MR		del 2q22	arr cgh 2q23.1 (RP11-72H23)x1	del 2q22	147651472	147688255	149855826	149879891	2167571	2228419		7	P
5	F	MCA/MR		del 14q12q13.2	arr cgh 14q12q13.2 (RP11-36909 → RP11-26M6)x1	del 14q12q13.2	28768137	29297829	34689412	35489337	5391583	6721200		25	P
6	M	MCA/MR	CHD	del 15q26.2	arr cgh 15q26.2q26.3 (RP11-79C10 → RP11-80F4)x1	del 15q26.2	93199415	93214053	96928421	96942334	3714368	3742919		6	P
7	M	MCA/MR	CHD	del 16p12.1p11.2	arr cgh 16p12.1p11.2 (RP11-309114 → RP11-150K5)x1	del 16p12.1p11.2	25795340	27008538	29825404	31443492	2816866	5648152	dn	138	P
8	M	MCA/MR	CHD	del 16p11.2	arr cgh 16p12.1p11.2 (RP11-360L15 → RP11-150K5)x1	del 16p11.2	27184508	28873631	29825404	31443492	951773	4258984	dn	134	P
9	F	MCA/MR		del 16p11.2	arr cgh 16p11.2 (RP11-368N21 → RP11-499D5)x1	del 16p11.2	28873841	29408698	32773200	34476095	3364502	5602254		125	P
10	M	MCA/MR		del 7p14.2p13	arr cgh 7p14.2p13 (RP11-138E20 → RP11-52M17)x1	del 7p14.2p13	35621006	36470190	44657334	45508196	8187144	9887190	dn	70	P
11	F	MCA/MR	Corneal opacity	del 14q22.1q22.3	arr cgh 14q22.1q22.3 (RP11-122A4 → RP11-172G1)x1	del 14q22.1q22.3	51964774	51983834	54730496	55054754	2746662	3089980	dn	18	P
12	M	MCA/MR	Idiopathic leukodystrophy	del 17q13.3	arr cgh 17p13.3 (RP11-294J5 → RP11-35707)x1	del 17q13.3	1008128	1146211	2077151	2026967	930940	1018839	dn	22	P
13	M	MCA/MR		del Xp11.4p11.3	arr cgh Xp11.3p11.4 (RP11-1069J5 → RP11-245M24)x1	del Xp11.4p11.3	41392291	41385453	45419624	45495709	4034171	4103418	dn	9	P

Table 3 Continued

Case	Gender	Clinical diagnosis	Remarkable clinical features	CNV Position	WGA-4500 ^b	FISH ^b	Base position and size of the identified CNV ^a				Parental analysis	Protein-CNV coding genes ^c	Corresponding assess-ment ^d or candidate gene(s)			
							Start (max)	Start (min)	End (max)	End (min)				Size (min)	Size (max)	
14	M	MCA/MR		del 6q12q14.1	arr cgh 6q12q14.2(RP11-502L6→ RP11-232L4)x1 arr cgh 6q14.1 (RP11-343P23→ RP11-217L13)x1	ish del(6)(q13) (RP11-28P18)-dn ish del(6)(q14.1) (RP11-5N7-RP11-990K4-RP11-1I6+) ish del(10) (p12.1p11.23) (RP11-164A7, RP11-110B21-) ish del(10)(p11.23) (RP11-15H10-)	69029871	69731888	83926178	85101718	14194290	16071847	dn	56	P	
15	M	ZLS		del 6q14.1	arr cgh 6q14.1 (RP11-343P23→ RP11-217L13)x1	ish del(6)(q14.1) (RP11-5N7-RP11-990K4-RP11-1I6+)	75484004	76145436	79474428	79851528	3328992	4367524			10	P
16	F	MCA/MR	CHD	del 10p12.1p11.23	arr cgh 10p12.1p11.23 (RP11-89D1→ 91A23)x1	ish del(10) (p12.1p11.23) (RP11-164A7, RP11-110B21-) ish del(10)(p11.23) (RP11-15H10-)	27045285	27054002	29057401	29088950	2003399	2043665			18	P
17	M	MCA/MR		del 10p12.1p11.23	arr cgh 10p12.1p11.23 (RP11-218D6→ RP11-RP11-181I11)x1	ish del(10)(p11.23) (RP11-15H10-)	28121596	28131608	30559024	30577807	2427416	2456211			12	P
18	M	MCA/MR	CHD	del 10q24.31q25.1	arr cgh 10q24.31q25.1 (RP11-108L7→ RP11-108L7)x1	ish del(10)(q24.33) (RP11-416N2)-dn	102560783	102568462	105914057	105929608	3345595	3368825	dn		66	P
19	M	MCA/MR		del 10q24.32q25.1	arr cgh 10q24.32q25.1 (RP11-21N23→ RP11-99N20)x1	ish del(10)(q24.33) (RP11-416N2)-dn	103917900	103928189	106005827	106011522	2077638	2093622	dn		41	P
20	F	MCA/MR		del 3p21.31p21.2	arr cgh 3p21.31p21.2 (RP11-24F11→ RP11-89F17)x1	ish del(3)(p21.31) (RP11-387-)	46150261	46359965	51390597	52571544	5030632	6421283			175	P
21	M	MCA/MR		del 7p22.1	arr cgh 7p22.1 (RP11-90J23→ RP11-2K20)x1	ish del(7)(p22.1) (RP11-2K20)-dn	3185609	5892225	6233987	6409277	341762	3223668	dn		28	P
22	F	MCA/MR	Corneal opacity, CHD	dup 14q11.2	arr cgh 14q11.2 (RP11-152G22→ RP11-84D12)x3	ish dup(14)(q11.2) (RP11-152G22++)	20070731	20306624	20534929	21264945	228305	1194214			>30	P
23	M	MCA/MR		del 17q24.1q24.2	arr cgh 17q24.1q24.2 (RP11-89L7→ RP11-79K13)x1	ish del(17) (q24.1q24.2) (RP11-93E5, RP11-89L7, RP11-79K13-) ish del(19)(p13.2) (91021-)	60576365	60936391	64592701	64587782	3656310	4011417			29	P
24	M	SMS susp.		del 19p13.2	arr cgh 19p13.2 (RP11-197O4→ RP11-164D24)x1	ish del(19)(p13.2) (91021-)	9248377	10248853	11968772	12553279	1719919	3304902	dn			P
25	M	MCA/MR	Epilepsy	dup 2q11.2q13	arr cgh 2q11.2q13 (RP11-90G13→ RP11-79K7)x3	ish dup(2)(q11.2) (RP11-542D13++)	88273220	91695986	109869691	112714666	18172705	24441446			>30	P
26	M	MCA/MR	CHD	dup 4p16.1	arr cgh 4p16.1 (RP11-17I9)x3	ish dup(4)(p16.1) (RP11-301J10++)	8202790	8520479	9793705	10638054	1273226	2435264			17	P

Table 3 Continued

Case	Gender	Clinical diagnosis	Remarkable clinical features	CNV Position	WGA-4500 ^b	FISH ^b	Base position and size of the identified CNV ^a				Size (max)	Size (min)	Protein-CNV analysis genes ^c	Corresponding assess- ment ^d gene(s)	
							Start (max)	Start (min)	End (min)	End (max)					
27	F	MCA/MR		del 7q22.1q22.2	arr cgh 7q22.1q22.2 (RP11-1008 → RP11-72124)x1	ish del(7)(q22.1q22.2) (RP11-124G15+,RP11- 188E1-,RP11-95P19-)	97 314 215	98 261 079	105 604 920	106 451 506	7 343 884	9 137 291	135	P	
28	F	MCA/MR	Epilepsy	del 12q13.13	arr cgh 12q13.13 (RP11-7418 → RP11-624J6)x1	ish dup(16)(q22.3) (RP11-115E3+, RP11-90L19+, RP11-89K4)x3	50 987 232	51 016 427	51 956 291	52 180 088	93 986 4	1 192 856	44	P	
29	M	MCA/MR		dup 16q22.3	arr cgh 16q22.3 (RP11-90L19 → RP11-89K4)x3	ish dup(16)(q24.1) (RP11-770B4+, RP11-140K16 → RP11-44201)x3	70 355 260	70 848 592	72 328 913	73 785 124	1 480 321	3 429 864	25	P	
30	M	RTS susp.		dup 16q24.1	arr cgh 16q24.1 (RP11-140K16 → RP11-44201)x3	ish dup(16)(q24.1) (RP11-770B4+, RP11-140K16+, ish del(2)(q24.2) (RP11-638N12-)	82 699 729	82 797 548	83 749 375	84 123 857	95 182 7	1 424 128	16	P	
31	M	MCA/MR	Epilepsy	del 2q24.2q24.3	arr cgh 2q24.2 (RP11-89L13 → RP11-79L13)x1	ish del(3)(p26.2) (RP11-32F23-)	160 407 234	161 072 815	162 883 584	166 923 475	1 810 769	6 516 241	28	P	TBR1
32	M	MCA/MR		del 3p26.2	arr cgh 3p26.2 (RP11-32F23)x1	ish del(3)(p26.2) (RP11-32F23-)	3 943 353	4 016 797	4 198 468	4 329 970	18 167 1	3 866 617	2	P	SUMF1
33	M	MCA/MR	IgA deficiency	del 7q21.11	arr cgh 7q21.11 (RP11-22M18)x1	ish del(7)(q21.11) (RP11-115M2+, RP11-353O4-, RP11-22M18-)	83 597 839	83 601 541	84 549 609	84 788 160	94 806 8	1 190 321	3	P	SEMA3A
34	M	MCA/MR		dup 14q32.2	arr cgh 14q32.2 (RP11-128L1)x3	ish dup(14)(q32.2) (RP11-177F8+)	99 330 486	99 337 358	99 841 558	99 845 472	504 200	5 14 986	7	P	EML1, YY1
35	M	MCA/MR	Epilepsy	dup 16p13.3	arr cgh 16p13.3 (RP11-349I11)x3	ish dup(16)(p13.3) (RP11-349I11+)	4 851 459	5 678 447	5 906 909	6 165 923	228 462	1 314 464	9	P	AZBP1
36	M	MCA/MR		dup Xp22.2p22.13	arr cgh Xp22.2p22.13 (RP11-2K15 → RP11-115I10)x3	not performed (X-tiling array)	168 74 735	16 952 121	17 596 600	17 638 351	644 479	763 616	2	P	
37	F	MCA/MR		dup Xp21.3	arr cgh Xp21.3 (RP11-438J7)x3	not performed (X-tiling array)	28 704 076	28 704 076	28 868 075	28 868 075	163 999	163 999	1	P	IL1RAPL1
38	M	MCA/MR	Hyper IgE	del 1p34.3	arr cgh 1p34.3 (RP11-89N10 → RP11-166F21-1dn arr cgh 1q25.2 (RP11-177A2 → RP11-152A16)x3	ish del(1)(p34.2) (RP11-195A8+, RP11-166F21-1dn ish dup(1)(q25.2) (RP11-177A2+, RP11-152A16+)	37 830 131	38 338 265	39 466 349	39 583 645	1 128 084	1 753 514	7	P	
39	M	MCA/MR		del 2p24.1p23.3	arr cgh 2p24.1p23.3 (RP11-80H16 → RP11-88F6)x1	ish del(2)(p23.3) (RP11-88F6-, RP11-373D23+1dn	1 770 888 480	1 771 968 858	1 775 535 659	1 778 559 828	338 801	771 348	9	P	
40	F	MCA/MR	CHD	del 3p26.1p25.3	arr cgh 3p26.1p25.3 (RP11-128A5 → RP11-402P11)x1	ish del(3)(p26.1p25.3) (RP11-936E1-, RP11-402P11-, RP11-1079H21+) dn	8 190 557	8 497 949	9 930 973	10 026 217	1 433 024	1 835 660	18	P	

Table 3 Continued

Case	Gender	Clinical diagnosis	Remarkable clinical features	CNV Position	WGA-4500 ^b	FISH ^b	Base position and size of the identified CNV ^a				Parental CNV analysis		Protein-coding genes ^c	Corresponding assess-ment ^d	Candidate gene(s)
							Start (max)	Start (min)	End (min)	End (max)	Size (min)	Size (max)			
41	M	MCA/MR		del 3p22.1p21.31	arr cgh 3p22.1p21.31 (RP11-241P3→, RP11-888B)x1	ish del(3)(p22.1) (RP11-61H16+, RP11-241P3, RP11-78010+)dn ish del(3)(p14.2) (RP11-7919, RP11-230A22+)mat	41365663	42284365	48177538	49198542	5893173	7832879	dn	123	P
42	M	MCA/MR	Corneal opacity	del 3p14.3p14.2	arr cgh 3p14.3p14.2 (RP11-80H18→, RP11-7919)x1	ish del(8) (q21.11q21.13) (RP11-225J6→, RP11-214E11)x1	57370434	58149199	58742633	58887574	593434	1517140	mat	11	B
43	M	MCA/MR		del 3q26.31q26.33	arr cgh 3q26.31-q26.33 (RP11-292L5→, RP11-355N16)x1	ish del(3)(q26.32) (RP11-300L9+, RP11-105L6-)dn ish del(13)(q13.2) (RP11-142E9+, RP11-381E21, RP11-98D3+)dn	175650310	176531688	180613203	181653281	4081515	6002971	dn	12	P
44	M	MCA/MR	CHD	del 13q13.2q13.3	arr cgh 13q13.2 (RP11-269G10→, 90F5)x1	ish del(22)(q11.21) (RP11-155F20→, 54C2)x1	33451136	33895560	34813379	34909905	917819	1458769	dn	1	P
45	F	aRS		del 22q11.21	arr cgh 22q11.21 (RP11-155F20→, 54C2)x1	ish del(18)(q21.2) (RP11-159D14+, RP11-186B13, RP11-111C17-)dn	19310307	19310307	19590642	19590642	280335	280335	pat	15	B
46	M	MCA/MR		dup 19p13.3	arr cgh 19p13.3 (RP11-49M3→, RP11-268O21)x3	ish del(18)(q21.2) (RP11-159D14+, RP11-186B13, RP11-111C17-)dn	1095485	2418857	3499581	4460252	1080724	3364767	dn	113	P
47	F	MCA/MR	Autism	del 19p13.3	arr cgh 19p13.3 (RP11-30F17→, RP11-330I7)x1	ish del(19)(p13.3) (RP11-330I7-)dn	4844383	6043505	6859584	6881792	816079	2037409	dn	23	P
48	M	MCA/MR		del Xp11.3	arr cgh Xp11.3 (RP11-151G3→, RP11-48J14)x0	ish del(X)(p11.3) (RP11-203D16-)mat	44403077	44433162	46795584	46795588	2362422	2392511	mat	18	P
49	M	MCA/MR		dup 3p26.3	arr cgh 3p26.3 (RP11-63O1+)x3	ish dup(3)(p26.3) (RP11-63O1+)pat	2377366	2443357	2619407	2628216	176050	250850	pat	1	B
50	M	MCA/MR		dup 5p14.3	arr cgh 5p14.3 (RP11-91A5+)x3	ish dup(5)(p14.3) (RP11-91A5+)pat	19046234	19485530	19656108	20798445	170578	1752211	pat	1	B
51	M	MCA/MR		dup 5q13.3	arr cgh 5q13.3 (RP11-40N8→, RP11-91C10)x3	ish dup(5)(q13.1) (RP11-105A11++)mat	66417271	66481371	67501700	67838977	1020329	1421706	mat	3	B

Table 3 Continued

Case	Gender	Clinical diagnosis	Remarkable clinical features	CNV Position	WGA-4500 ^b	FISH ^b	Base position and size of the identified CNV ^a				Parental CNV analysis genes ^c		Protein-coding genes ^d	Corresponding assess- ment ^d	Candidate gene(s)
							Start (max)	Start (min)	End (min)	End (max)	Size (min)	Size (max)			
52	M	MCA/MR	dup 7p22.3	dup 7p22.3	arr cgh 7p22.3 (RP11-23D23)x3	ish dup(7)(p22.3) (RP11-23D23)++ RP11-1133D5+mat	1	954016	954584	1101944	568	1101943	mat	12	B
53	F	MCA/MR	dup 8p23.2	dup 8p23.2	arr cgh 8p23.2 (RP11-79I19+ RP11-89I12)x3	ish dup(8)(p23.2) (RP11-89I19)++ RP11-89I12+pat	3324954	3726061	4564671	5973493	838610	2648539	pat	1	B
54	M	MCA/MR	dup 9q33.1	dup 9q33.1	arr cgh 9q33.1 (RP11-150L1)x3	ish dup(9)(q33.1) (RP11-150L1)++pat	118980752	119452372	119614984	120011559	162612	1030807	pat	2	B
55	F	MCA/MR	dup 10q22.3	dup 10q22.3	arr cgh 10q22.3 (RP11-79M9)x3	ish dup(10)(q22.3) (RP11-79M9)++mat	77356915	77718484	77879148	78230039	154664	873124	mat	1	B
56	M	MCA/MR	ELBW, hepato- blastoma	dup 12q21.31	arr cgh 12q21.31 (RP11-91C4)x3	ish dup(12)(q21.31) (RP11-91C4)++ RP11-142L2+pat	80924954	82678148	82830190	85768388	152042	4843434	pat	3	B
57	M	GS	del Xp11.23	del Xp11.23	arr cgh Xp11.23 (RP11-876B24) x0 mat	not performed (X-tilling array)	47752808	47747918	47852109	47868412	104191	115604	mat	3	B
58	M	MCA/MR	dup 8q11.23	dup 8q11.23	arr cgh 8q11.23 (RP11-22I17)x3	ish dup(8)(q11.23) (RP11-22I17)++ RP11-26P22++	53665974	53717675	54235229	54576654	517554	910680		3	VOUS
59	F	MCA/MR	Micro- cephaly	dup 10q11.21	arr cgh 10q11.21 (RP11-178A10)x3	ish dup(10)(q11.21) (RP11-178A10)++	41986946	42197693	42320775	43603027	123082	1616081		15	VOUS
60	M	MCA/MR	dup 11p14.2p14.1	dup 11p14.2p14.1	arr cgh 11p14.2p14.1 (RP11-11I2)x3	ish dup(11) (p14.2p14.1) (RP11-11I2)++	26723462	27033270	27213374	27445504	180104	722042		4	VOUS
61	F	MCA/MR	dup 12p11.1	dup 12p11.1	arr cgh 12p11.1 (RP11-88P4)x3	ish dup(12)(p11.1) (RP11-472A10)++	33333493	33359944	33572956	33572956	213012	239463		2	VOUS
62	F	aRS	dup 12q21.31	dup 12q21.31	arr cgh 12q21.31 (RP11-91I24+ RP11-91C4)x3	ish dup(12)(q21.31) (RP11-91I24)++ RP11-142L2++	79949648	82172368	83968319	85768388	1795951	5818740		12	VOUS
63	F	MR	Congenital myopathy	dup Xq12	arr cgh Xq12 (RP11-90P17+ RP11-383C12)x3	Not performed (X-tilling array)	66212661	66216353	66921699	66948538	705346	735877		1	VOUS

Abbreviations: aRS, atypical Rett syndrome; B, benign; CNV, copy-number variant; *dn*: *de novo* CNV observed in neither of the parents; ELBW, extremely low birth weight; FISH, fluorescence *in situ* hybridization; GS, Gillespie syndrome; mat, CNV identified also in mother; P, pathogenic; pat, CNV identified also in father; RTS, Rubinstein-Taybi syndrome; SMS, Smith-Magenis syndrome; VOUS, variant of uncertain clinical significance; ZLS, Zimmermann-Laband syndrome.
^aThe sizes were estimated by WGA-4500, X-array, FISH or Agilent Human Genome CGH microarray 244K.
^bThe notation systems is based on ISCN2005.
^cThe number of protein-coding genes contained in the respective CNVs.
^dThe result of CNV assessment.

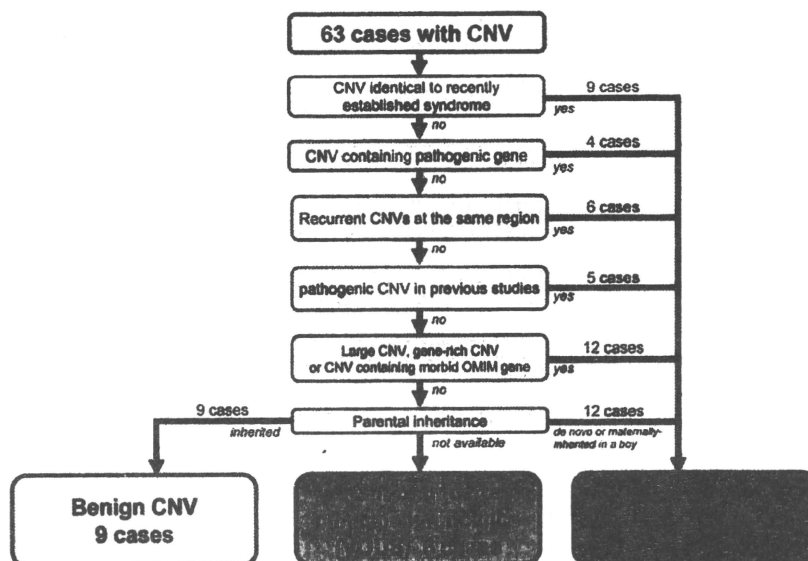


Figure 2 A flowchart of the assessment of CNVs detected in the second screening.

from several aspects. A CNV that contains abundant genes or is large (> 3 Mb) has a high possibility to be pathogenic.²¹ The CNVs in cases 25–30 probably correspond to such CNVs. Also, we judged a CNV containing a morbid OMIM gene as pathogenic.²¹ *TBR1* (OMIM: *604616) in case 31,⁵⁶ *SUMF1* (OMIM: *607939) in case 32,^{57,58} *SEMA3A* (OMIM: *603961) in case 33,⁵⁹ *EML1* (OMIM: *602033) and/or *YY1* (OMIM: *600013) in case 34,^{60,61} *A2BP1* (OMIM: *605104) in case 35⁶² and *ILIRAPL1* (OMIM: *300206) in case 36.⁶³ Several previous reports suggest that these genes are likely to be pathogenic, although at present no evidence of a direct association between these genes and phenotypes exists.

CNVs de novo or X maternally inherited. Among the remaining 27 cases, 12 cases had CNVs considered pathogenic as their CNVs were *de novo* (cases 37–47) or inherited del(X)(p11.3) from the mother (case 48). In the second screening we performed FISH for 36 CNVs of the 34 cases whose parental samples were available to confirm that 24 cases had *de novo* CNVs, which were probably pathogenic. A CNV in case 48, a boy with a nullizygous deletion at Xp11.3 inherited from his mother, was also probably relevant to his phenotype (Tables 3 and 4). Meanwhile, although case 57 was a boy with a deletion at Xp11.23 inherited from his mother, he was clinically diagnosed with Gillespie syndrome (OMIM: #206700) that was reported to show an autosomal dominant or recessive pattern,⁶⁴ thus we judged that the deletion was not relevant to his phenotype. As a result, cases 49–57 had only CNVs inherited from one of their parents which are likely to be unrelated to the phenotypes; that is, bCNV (Table 4).

As a result, we estimated that 48 cases among 349 analyzed (13.8%) had pCNV(s) in the second screening (Table 3; Figure 2). The CNVs of the remaining six cases, cases 58–63, were not associated with previously reported pathogenicity and their inheritance could not be evaluated, thus we estimated they were variants of uncertain clinical significance (VOUS).³⁸

DISCUSSION

Because aCGH is a high-throughput technique to detect CNVs rapidly and comprehensively, this technique has been commonly used for

analyses of patients with MCA and/or MR.^{38,65–68} However, recent studies of human genomic variation have uncovered surprising properties of CNV, which covers 3.5–12% of the human genome even in healthy populations.^{18–20,69} Thus analyses of patients with uncertain clinical phenotypes need to assess whether the CNV is pathogenic or unrelated to phenotypes.²¹ However, such an assessment may diminish the rapidness or convenience of aCGH.

In this study, we evaluated whether our in-house GDA can work well as a diagnostic tool to detect CNVs responsible for well-established syndromes or those involved in subtelomeric aberrations in a clinical setting, and then explored candidate pCNVs in cases without any CNV in the first GDA screening. We recruited 536 cases that had been undiagnosed clinically and studied them in a two-stage screening using aCGH. In the first screening we detected CNVs in 54 cases (10.1%). Among them, 40 cases had CNV(s) at subtelomeric region(s) corresponding to the well-established syndromes or the already described disorders and the other 14 cases had CNVs in the regions corresponding to known disorders. Thus about three quarters of cases had genomic aberrations involved in subtelomeric regions. All the subtelomeric deletions and a part of the subtelomeric duplications corresponded to the disorders, indicating that especially subtelomeric deletions had more clinical significance compared to subtelomeric duplications, although the duplication might result in milder phenotypes and/or function as a modifier of phenotypes.⁷⁰ Moreover, parental analysis in three cases with two subtelomeric aberrations revealed that two of them were derived from the parental balanced translocations, indicating that such subtelomeric aberrations were potentially recurrent and parental analyses were worth performing. Recently several similar studies analyzed patients with MCA/MR or developmental delay using a targeted array for subtelomeric regions and/or known genomic disorders and detected clinically relevant CNVs in 4.4–17.1% of the patients.^{28,65,70,71} Our detection rate in the first screening was equivalent to these reports. Although such detection rates depend on the type of microarray, patient selection criteria and/or number of subjects, these results suggest that at least 10% of cases with undiagnosed MCA/MR and a normal karyotype would be detectable by targeted array.

Table 4 Parental analysis of 34 cases in the second screening

Case	Gender	Clinical diagnosis	CNV		Size of CNV (bp)		Protein-coding genes	Parental analysis	Pathogenicity
			del/dup	Position	Min.	Max.			
1	M	MCA/MR	del	1p36.23p36.22	1670237	2558590	32	<i>de novo</i>	P
2	M	MCA/MR	del	1q41q42.11	5001798	6481439	35	<i>de novo</i>	P
7	M	MCA/MR	del	16p12.1p11.2	2816866	5648152	138	<i>de novo</i>	P
8	M	MCA/MR with CHD	del	16p11.2	951773	4258984	134	<i>de novo</i>	P
10	M	MCA/MR	del	7p14.2p13	8516513	9421233	70	<i>de novo</i>	P
11	F	MCA/MR	del	14q22.1q22.3	2746662	3089980	18	<i>de novo</i>	P
12	M	MCA/MR	del	17q13.3	930940	1018839	22	<i>de novo</i>	P
13	M	MCA/MR	del	Xp11.4p11.3	4034171	4103418	9	<i>de novo</i>	P
14	M	MCA/MR	del	6q12q14.1	14194290	16071847	56	<i>de novo</i>	P
18	M	MCA/MR	del	10q24.31q25.1	3345595	3368825	66	<i>de novo</i>	P
19	M	MCA/MR	del	10q24.32q25.1	2077638	2093622	41	<i>de novo</i>	P
21	M	MCA/MR	del	7p22.1	341762	3223668	28	<i>de novo</i>	P
24	M	SMS susp.	del	19p13.2	1719919	3304902	23	<i>de novo</i>	P
37	F	MCA/MR	del	1p34.3	1128084	1753514	7	<i>de novo</i>	P
38	M	MCA/MR	dup	1q25.2	338801	771348	9	<i>de novo</i>	P
39	M	MCA/MR	del	2p24.1p23.3	3721550	8376636	86	<i>de novo</i>	P
40	F	MCA/MR	del	3p26.1p25.3	1433024	1835660	18	<i>de novo</i>	P
41	M	MCA/MR	del	3p22.1p21.31	5893173	7832879	123	<i>de novo</i>	P
42 ^a	M	MCA/MR	del	8q21.11q21.13	5289394	5770485	12	<i>de novo</i>	P
42 ^a	M	MCA/MR	del	3p14.3p14.2	593434	1517140	11	Maternal	B
43	M	MCA/MR	del	3q26.31q26.33	4081515	6002971	12	<i>de novo</i>	P
44 ^b	M	MCA/MR	del	13q13.2q13.3	917819	1458769	1	<i>de novo</i>	P
44 ^b	M	MCA/MR	del	22q11.21	917819	1458769	15	Paternal	B
45	F	Rett syndrome	del	18q21.2	2121913	3642522	9	<i>de novo</i>	P
46	M	MCA/MR	dup	19p13.3	2041395	2404096	113	<i>de novo</i>	P
47	F	MCA/MR	del	19p13.3	816079	2037409	23	<i>de novo</i>	P
48 ^c	M	MCA/MR	del	Xp11.3	2362422	2392511	18	Maternal	P
49	M	MCA/MR	dup	3p26.3	176050	250850	1	Paternal	B
50	M	MCA/MR	dup	5p14.3	170578	1752211	1	Paternal	B
51	M	MCA/MR	dup	5q13.3	1020329	1421706	3	Maternal	B
52	M	MCA/MR	dup	7p22.3	568	1101943	12	Maternal	B
53	F	MCA/MR	dup	8p23.2	838610	2648539	1	Paternal	B
54	M	MCA/MR	dup	9q33.1	162612	1030807	2	Paternal	B
55	F	MCA/MR	dup	10q22.3	154664	873124	1	Maternal	B
56	M	MCA/MR	dup	12q21.31	152042	4843434	3	Paternal	B
57	M	Gillespie syndrome	del	Xp11.23	104191	115604	3	Maternal	B

Abbreviations: B, benign; CNV, copy-number variant; F, female; M, male; MCA/MR, multiple congenital anomalies and mental retardation; P, pathogenic.

^aTwo CNVs were detected in case 42.

^bTwo CNVs were detected in case 44.

^cNullizygous deletion inherited from his mother probably affected the phenotype.

Another interesting observation in the first screening was that subtelomeric rearrangements frequently occurred even in patients with MCA/MR of uncertain whose karyotype had been diagnosed as normal. This result may be consistent with a property of subtelomeric regions whose rearrangements can be missed in conventional karyotyping,⁷² and in fact other techniques involving subtelomeric FISH or MLPA also identified subtelomeric abnormalities in a number of patients with MCA and/or MR in previous reports.^{70,73,74} Our result may support the availability of prompt screening of subtelomeric regions for cases with uncertain congenital disorders.

In the second screening we applied WGA-4500 to 349 cases to detect 66 candidate pCNVs in 63 cases (18.1%), and subsequently assessed the pathogenicity of these CNVs. The pCNVs included nine

CNVs overlapping identical regions of recently recognized syndromes (cases 1–9; deletion at 1p36.23–p36.22, 1q41–q42.11, 1q43–q44, 2q23.1, 14q12, 15q26–qter and 16p11.2–p12.2, respectively), four CNVs containing disease-associated genes (cases 10–13; *GLI3*, *BMP4*, *YWHAE* and *CASK*, respectively), three pairs of CNVs of recurrent deletions (cases 14, 15: at 6q12–q14.1 and 6q14.1; case 16, 17: at 10p12.1–p11.23 and case 18, 19: at 10q24.31–q25.1 and 10q24.32–q25.1), five CNVs identical to pCNVs in previous studies (cases 20–24), six large and/or gene-rich CNVs (cases 25–30) and six CNVs containing a morbid OMIM gene (cases 31–36). For the remaining cases, we estimated the pathogenicity of the CNVs from a parental analysis (Table 4). We judged the 11 *de novo* CNVs (cases 37–47) and 1 CNV on chromosome Xp11.3 inherited from

the mother (case 48) as probably pathogenic. And nine inherited CNVs (cases 49–57) were probably benign. The clinical significance of CNVs in the other six cases, cases 58–63, remains uncertain (VOUS). As a result we estimated CNVs as pathogenic in 48 cases among 349 cases (13.8%) analyzed in the second screening. None of the pCNVs corresponded to loci of well-established syndromes. This may suggest that our two-stage screening achieved a good balance between rapid screening of known syndromes and investigation of CNV of uncertain pathogenicity.

Table 5 Summary of parental analyses

	Average size (bp)		The average number of protein-coding genes	
	Min.	Max.		
Pathogenic CNVs^a				
del	23	3 309 267	4 597 689	43
dup	2	1 190 098	1 587 722	61
Total	25	3 139 733	4 356 892	44
Benign CNVs^b				
del	3	538 481	1 030 504	10
dup	8	334 432	1 740 327	3
Total	11	390 082	1 546 739	5

Abbreviation: CNV, copy-number variant.
^aTwenty-four *de novo* CNVs and case 48.
^bEleven inherited CNVs other than case 48.

Among the cases with parental analyses, the 25 pCNVs had larger sizes and contained more protein-coding genes (average size, 3.1 Mb at minimum to 4.4 Mb at maximum; average number of genes, 44) as compared with the 11 inherited bCNVs that were probably unrelated to phenotypes (average size, 0.39 Mb at minimum to 1.5 Mb at maximum; average number of genes, 5) (Table 5). Although all of the 25 pCNVs except 2 were deletions, about three quarters (8 of 11 cases) of the inherited bCNVs were duplications (Table 5). These findings are consistent with previously reported features of pCNVs and bCNVs.^{21,38}

We also compared our current study with recent aCGH studies meeting the following conditions: (1) a microarray targeted to whole genome was applied; (2) patients with MCA and/or MR of uncertain etiology, normal karyotype and the criteria for patients selection were clearly described; (3) pathogenicity of identified CNVs were assessed. On the basis of the above criteria, among studies reported in the past 5 years, we summarized 13 studies (Table 6).^{10,14,15,17,54,55,75–81} Diagnostic yield of pCNVs in each study was 6.3–16.4%, and our current diagnostic yield of the second screening was 13.8%. Though cases with subtelomeric aberration detected in the first screening had been excluded, our diagnostic yield was comparable to those of the reported studies. It is not so important to make a simple comparison between diagnostic yields in different studies as they would depend on the conditions of each study, for example, sample size or array resolution,^{38,82} however it seems interesting that the higher resolution of a microarray does not ensure an increase in the rate of detection of pCNVs. One recent study showed data that may explain the discrepancy between the resolution of microarray and diagnostic yield.^{54,83} The authors analyzed 1001 patients with MCA and/or MR using one

Table 6 Previous studies of analyzing patients with MCA and/or MR using aCGH targeted to whole genome

Author (year)	Applied array			Patients		Pathogenic CNV	
	Type	Number ^a	Distribution ^b	Number	Type of disorders	Number	%
Schoumans <i>et al.</i> ⁷⁵	BAC	2600	1.0 Mb*	41	MCA and MR	4	9.8
de Vries <i>et al.</i> ⁷⁶	BAC	32 477	Tiling	100	MCA and/or MR	10	10.0
Rosenberg <i>et al.</i> ⁷⁷	BAC	3500	1.0 Mb*	81	MCA and MR	13	16.0
Krepischi-Santos <i>et al.</i> ⁷⁸	BAC	3500	1.0 Mb*	95	MCA and/or MR	15	15.8
Friedman <i>et al.</i> ¹⁴	SNP	Affymetrix 100K	23.6 kb**	100	MR	11	11.0
Thuresson <i>et al.</i> ⁷⁹	BAC		1.0 Mb*	48	MCA and MR	3	6.3
Wagenstaller <i>et al.</i> ⁸⁰	SNP	Affymetrix 100K	23.6 kb**	67	MR	11	16.4
Fan <i>et al.</i> ⁵⁵	Oligo	Agilent 44K	24 kb–43 kb**	100 ^c	MCA and MR, Autism	15 ^d	15.0
Xiang <i>et al.</i> ¹⁵	Oligo	Agilent 44K	24 kb–43 kb**	40 ^e	MR, DD and autism	3	7.5
Pickering <i>et al.</i> ¹⁰	BAC	2600	1 Mb*	354 ^f	MCA and/or MR	36 ^g	10.2
McMullan <i>et al.</i> ¹⁷	SNP	Affymetrix 500K	2.5 kb–5.8 kb**	120	MCA and/or MR	18	15.0
Bruno <i>et al.</i> ⁸¹	SNP	Affymetrix 250K	2.5 kb–5.8 kb**	117	MCA and/or MR	18	15.4
Buysse <i>et al.</i> ⁵⁴	BAC	3431	1 Mb*	298	MCA and/or MR	26	8.7
	Oligo	Agilent 44K	24 kb–43 kb**	703	MCA and/or MR	74	10.5
Our current study	BAC	4523	0.7 Mb	349	MCA and MR	48	13.8
Total				2613		305	11.7

Abbreviations: BAC, bacterial artificial chromosome; CNV, copy-number variant; DD, developmental delay; MCA, multiple congenital anomalies; MR, mental retardation; SNP, single nucleotide polymorphism.

^aThe number of clones or name of array is described.

^bEach distribution referred to each article (*) or manual of each manufacturer (**).

^cAll cases were analyzed by both a targeted array and a genome-wide array.

^dIn five cases, CNVs were also identified by a targeted array.

^eTen cases with an abnormal karyotype were excluded.

^fOnly cases studied with an array throughout the genome are described. Ninety-eight cases were also analyzed by a targeted array.

^gSeventeen cases with an abnormal karyotype were excluded.

of two types of microarray, BAC array and oligonucleotide array. The BAC array was applied for 298 patients to detect 58 CNVs in 47 patients, and among them 26 CNVs (8.7%) were determined to be causal (pathogenic). Conversely, the oligonucleotide arrays were applied for 703 patients to detect 1538 CNVs in 603 patients, and among them 74 CNVs (10.5%) were determined to be pathogenic. These results may lead to the following idea: a lower-resolution microarray detects a limited number of CNVs likely to be pathogenic, because such CNVs tend to be large, and a higher-resolution microarray detects an increasing number of bCNVs or VOUS.³⁸ Indeed, in studies using a high-resolution microarray, most of the CNVs detected were smaller than 500 kb but almost all pCNVs were relatively large.^{54,81,83} Most of the small CNVs were judged not to be pathogenic, and the percentage of pCNVs stabilized at around 10%. This percentage may suggest a frequency of patients with MCA/MR caused by CNV affecting one or more genes, other than known syndromes and subtelomeric aberrations. The other patients may be affected by another cause undetectable by genomic microarray; for example a point mutation or microdeletion/duplication of a single gene, aberration of microRNA, aberration of methylation states, epigenetic aberration or partial uniparental disomy.

As recently hypothesized secondary insult, which is potentially another CNV, a mutation in a phenotypically related gene or an environmental event influencing the phenotype, may result in clinical manifestation.⁸⁴ Especially, in two-hit CNVs, two models have been hypothesized: (1) the additive model of two co-occurring CNVs affecting independent functional modules and (2) the epistatic model of two CNVs affecting the same functional module.⁸⁵ It also suggests difficulty in selecting an optimal platform in the clinical screening. Nevertheless, information on both pCNVs and bCNVs detected through studies using several types of microarrays is unambiguously significant because an accumulation of the CNVs will create a map of genotype-phenotype correlation that would determine the clinical significance of each CNV, illuminate gene function or establish a new syndrome.

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- 1 Roeleveld, N., Zielhuis, G. A. & Gabreëls, F. The prevalence of mental retardation: a critical review of recent literature. *Dev. Med. Child Neurol.* **39**, 125–132 (1997).
- 2 Hunter, A. G. Outcome of the routine assessment of patients with mental retardation in a genetics clinic. *Am. J. Med. Genet.* **90**, 60–68 (2000).
- 3 Smith, D. W. & Bostian, K. E. Congenital anomalies associated with idiopathic mental retardation. *J. Pediatr.* **65**, 189–196 (1964).
- 4 Gustavson, K. H., Hagberg, B., Hagberg, G. & Sars, K. Severe mental retardation in a Swedish county. II. Etiologic and pathogenetic aspects of children born 1959–1970. *Neuropadiatrie* **8**, 293–304 (1977).
- 5 Fryns, J. P., Kleczkowska, A., Kubief, E. & Van den Berghe, H. Cytogenetic findings in moderate and severe mental retardation. A study of an institutionalized population of 1991 patients. *Acta. Paediatr. Scand. Suppl.* **313**, 1–23 (1984).

- 6 Gustavson, K. H., Holmgren, G. & Blomquist, H. K. Chromosomal aberrations in mildly mentally retarded children in a northern Swedish county. *Ups. J. Med. Sci. Suppl.* **44**, 165–168 (1987).
- 7 Schrepers-Tijdink, G. A., Curfs, L. M., Wieggers, A., Kleczkowska, A. & Fryns, J. P. A systematic cytogenetic study of a population of 1170 mentally retarded and/or behaviourally disturbed patients including fragile X-screening. The Hondsberg experience. *J. Genet. Hum.* **36**, 425–446 (1988).
- 8 van Karnebeek, C. D., Koevoets, C., Sluijter, S., Bijlsma, E. K., Smeets, D. F., Redeker, E. J. et al. Prospective screening for subtelomeric rearrangements in children with mental retardation of unknown aetiology: the Amsterdam experience. *J. Med. Genet.* **39**, 546–553 (2002).
- 9 Vissers, L. E., de Vries, B. B., Osoegawa, K., Janssen, I. M., Feuth, T., Choy, C. O. et al. Array-based comparative genomic hybridization for the genomewide detection of submicroscopic chromosomal abnormalities. *Am. J. Hum. Genet.* **73**, 1261–1270 (2003).
- 10 Pickering, D. L., Eudy, J. D., Olney, A. H., Dave, B. J., Golden, D., Stevens, J. et al. Array-based comparative genomic hybridization analysis of 1176 consecutive clinical genetics investigations. *Genet. Med.* **10**, 262–266 (2008).
- 11 Bauters, M., Van Esch, H., Marynen, P. & Fryns, G. X chromosome array-CGH for the identification of novel X-linked mental retardation genes. *Eur. J. Med. Genet.* **48**, 263–275 (2005).
- 12 Hayashi, S., Honda, S., Minaguchi, M., Makita, Y., Okamoto, N., Kosaki, R. et al. Construction of a high-density and high-resolution human chromosome X array for comparative genomic hybridization analysis. *J. Hum. Genet.* **52**, 397–405 (2007).
- 13 Kok, K., Dijkhuizen, T., Swart, Y. E., Zorgdrager, H., van der Vlies, P., Fehrmann, R. et al. Application of a comprehensive subtelomere array in clinical diagnosis of mental retardation. *Eur. J. Med. Genet.* **48**, 250–262 (2005).
- 14 Friedman, J. M., Baross, A., Delaney, A. D., Ally, A., Arbour, L., Armstrong, L. et al. Oligonucleotide microarray analysis of genomic imbalance in children with mental retardation. *Am. J. Hum. Genet.* **79**, 500–513 (2006).
- 15 Xiang, B., Li, A., Valentin, D., Nowak, N. J., Zhao, H. & Li, P. Analytical and clinical utility of whole-genome oligonucleotide array comparative genomic hybridization for pediatric patients with mental retardation and developmental delay. *Am. J. Med. Genet.* **146A**, 1942–1954 (2008).
- 16 Shen, Y., Irons, M., Miller, D. T., Cheung, S. W., Lip, V., Sheng, X. et al. Development of a focused oligonucleotide-array comparative genomic hybridization chip for clinical diagnosis of genomic imbalance. *Clin. Chem.* **53**, 2051–2059 (2007).
- 17 McMullan, D. J., Bonin, M., Hehir-Kwa, J. Y., de Vries, B. B., Dufke, A., Rattenberry, E. et al. Molecular karyotyping of patients with unexplained mental retardation by SNP arrays: a multicenter study. *Hum. Mutat.* **30**, 1082–1092 (2009).
- 18 Iafate, A. J., Feuk, L., Rivera, M. N., Listewnik, M. L., Donahoe, P. K., Qi, Y. et al. Detection of large-scale variation in the human genome. *Nat. Genet.* **36**, 949–951 (2004).
- 19 Sebat, J., Lakshmi, B., Troge, J., Alexander, J., Young, J., Lundin, P. et al. Large-scale copy number polymorphism in the human genome. *Science* **305**, 525–528 (2004).
- 20 Redon, R., Ishikawa, S., Fitch, K. R., Feuk, L., Perry, G. H., Andrews, T. D. et al. Global variation in copy number in the human genome. *Nature* **444**, 444–454 (2006).
- 21 Lee, C., Iafate, A. J. & Brothman, A. R. Copy number variations and clinical cytogenetic diagnosis of constitutional disorders. *Nat. Genet.* **39**, S48–S54 (2007).
- 22 Inazawa, J., Inoue, J. & Imoto, I. Comparative genomic hybridization (CGH)-arrays pave the way for identification of novel cancer-related genes. *Cancer Sci.* **95**, 559–563 (2004).
- 23 Hayashi, S., Kurosawa, K., Imoto, I., Mizutani, S. & Inazawa, J. Detection of cryptic chromosome aberrations in a patient with a balanced t(1;9)(p34.2;p24) by array-based comparative genomic hybridization. *Am. J. Med. Genet.* **139**, 32–36 (2005).
- 24 Shrimpton, A. E., Braddock, B. R., Thomson, L. L., Stein, C. K. & Hoo, J. J. Molecular delineation of deletions on 2q37.3 in three cases with an Albright hereditary osteodystrophy-like phenotype. *Clin. Genet.* **66**, 537–544 (2004).
- 25 Rauch, A. & Dörr, H. G. Chromosome 5q subtelomeric deletion syndrome. *Am. J. Med. Genet. C* **145C**, 372–376 (2007).
- 26 Horn, D., Tönnies, H., Neitzel, H., Wahl, D., Hinkel, G. K., von Moers, A. et al. Minimal clinical expression of the holoprosencephaly spectrum and of Currarino syndrome due to different cytogenetic rearrangements deleting the Sonic Hedgehog gene and the HLXB9 gene at 7q36.3. *Am. J. Med. Genet. A* **128A**, 85–92 (2004).
- 27 Tatton-Brown, K., Pilz, D. T., Orstavik, K. H., Patton, M., Barber, J. C., Collinson, M. N. et al. 15q overgrowth syndrome: a newly recognized phenotype associated with overgrowth, learning difficulties, characteristic facial appearance, renal anomalies and increased dosage of distal chromosome 15q. *Am. J. Med. Genet. A* **149A**, 147–154 (2009).
- 28 Lu, X., Shaw, C. A., Patel, A., Li, J., Cooper, M. L., Wells, W. R. et al. Clinical implementation of chromosomal microarray analysis: summary of 2513 postnatal cases. *PLoS One* **2**, e327 (2007).
- 29 Fernandez, T. V., García-González, I. J., Mason, C. E., Hernández-Zaragoza, G., Ledezma-Rodríguez, V. C., Anguiano-Alvarez, V. M. et al. Molecular characterization of a patient with 3p deletion syndrome and a review of the literature. *Am. J. Med. Genet. A* **146A**, 2746–2752 (2008).
- 30 Jones, K. L. *Smith's Recognizable Patterns of Human Malformation*, 6th edn. (Elsevier Saunders, Philadelphia, 2006).
- 31 Striano, P., Malacarne, M., Cavani, S., Pierluigi, M., Rinaldi, R., Cavaliere, M. L. et al. Clinical phenotype and molecular characterization of 6q terminal deletion syndrome: five new cases. *Am. J. Med. Genet. A* **140**, 1944–1949 (2006).

- 32 Lindstrand, A., Malmgren, H., Verri, A., Benetti, E., Eriksson, M., Nordgren, A. et al. Molecular and clinical characterization of patients with overlapping 10p deletions. *Am. J. Med. Genet. A* **152A**, 1233–1243 (2010).
- 33 Elbracht, M., Roos, A., Schönerr, N., Busse, S., Damen, R., Zerres, K. et al. Pure distal trisomy 2q: a rare chromosomal abnormality with recognizable phenotype. *Am. J. Med. Genet. A* **149A**, 2547–2550 (2009).
- 34 Lukusa, T. & Fryns, J. P. Pure *de novo* 17q25.3 micro duplication characterized by micro array CGH in a dysmorphic infant with growth retardation, developmental delay and distal arthrogryposis. *Genet. Couns.* **21**, 25–34 (2010).
- 35 Fukami, M., Kirsch, S., Schiller, S., Richter, A., Benes, V., Franco, B. et al. A member of a gene family on Xp22.3, VCXA, is deleted in patients with X-linked nonspecific mental retardation. *Am. J. Hum. Genet.* **67**, 563–573 (2000).
- 36 Shaffer, L. G. & Tommerup, N. *An International System for Human Cytogenetic Nomenclature* (2005) (Karger, Basel, 2005).
- 37 Koolen, D. A., Pfundt, R., de Leeuw, N., Hehir-Kwa, J. Y., Nillesen, W. M., Neefs, I. et al. Genomic microarrays in mental retardation: a practical workflow for diagnostic applications. *Hum. Mutat.* **30**, 283–292 (2009).
- 38 Miller, D. T., Adam, M. P., Aradhya, S., Biesecker, L. G., Brothman, A. R., Carter, N. P. et al. Consensus statement: chromosomal microarray is a first-tier clinical diagnostic test for individuals with developmental disabilities or congenital anomalies. *Am. J. Hum. Genet.* **86**, 749–764 (2010).
- 39 Shaffer, L. G., Theisen, A., Bejjani, B. A., Ballif, B. C., Aylsworth, A. S., Lim, C. et al. The discovery of microdeletion syndromes in the post-genomic era: review of the methodology and characterization of a new 1q41q42 microdeletion syndrome. *Genet. Med.* **9**, 607–616 (2007).
- 40 van Bon, B. W., Koolen, D. A., Borgatti, R., Magee, A., Garcia-Minaur, S., Rooms, L. et al. Clinical and molecular characteristics of 1qter microdeletion syndrome: delineating a critical region for corpus callosum agenesis/hypogenesis. *J. Med. Genet.* **45**, 346–354 (2008).
- 41 van Bon, B. W., Koolen, D. A., Brueton, L., McMullan, D., Lichtenbelt, K. D., Adès, L. C. et al. The 2q23.1 microdeletion syndrome: clinical and behavioural phenotype. *Eur. J. Hum. Genet.* **18**, 163–170 (2010).
- 42 Mencarelli, M. A., Kleefstra, T., Katzaki, E., Papa, F. T., Cohen, M., Pfundt, R. et al. 14q12 microdeletion syndrome and congenital variant of Rett syndrome. *Eur. J. Med. Genet.* **52**, 148–152 (2009).
- 43 Rump, P., Dijkhuizen, T., Sikkema-Raddatz, B., Lemmink, H. H., Vos, Y. J., Verheij, J. B. et al. Drayer's syndrome of mental retardation, microcephaly, short stature and absent phalanges is caused by a recurrent deletion of chromosome 15(q26.2→qter). *Clin. Genet.* **74**, 455–462 (2008).
- 44 Ballif, B. C., Hornor, S. A., Jenkins, E., Madan-Khetarpal, S., Surti, U., Jackson, K. E. et al. Discovery of a previously unrecognized microdeletion syndrome of 16p11.2-p12.2. *Nat. Genet.* **39**, 1071–1073 (2007).
- 45 Shinawi, M., Liu, P., Kang, S. H., Shen, J., Belmont, J. W., Scott, D. A. et al. Recurrent reciprocal 16p11.2 rearrangements associated with global developmental delay, behavioral problems, dysmorphism, epilepsy, and abnormal head size. *J. Med. Genet.* **47**, 332–341 (2010).
- 46 Kang, S. H., Scheffer, A., Ou, Z., Li, J., Scaglia, F., Belmont, J. et al. Identification of proximal 1p36 deletions using array-CGH: a possible new syndrome. *Clin. Genet.* **72**, 329–338 (2007).
- 47 Johnston, J. J., Olivos-Glander, I., Killoran, C., Elson, E., Turner, J. T., Peters, K. F. et al. Molecular and clinical analyses of Greig cephalopolysyndactyly and Pallister-Hall syndromes: robust phenotype prediction from the type and position of GLI3 mutations. *Am. J. Hum. Genet.* **76**, 609–622 (2005).
- 48 Johnston, J. J., Olivos-Glander, I., Turner, J., Aleck, K., Bird, L. M., Mehta, L. et al. Clinical and molecular delineation of the Greig cephalopolysyndactyly contiguous gene deletion syndrome and its distinction from acrocallosal syndrome. *Am. J. Med. Genet. A* **123A**, 236–242 (2003).
- 49 Hayashi, S., Okamoto, N., Makita, Y., Hata, A., Imoto, I. & Inazawa, J. Heterozygous deletion at 14q22.1-q22.3 including the BMP4 gene in a patient with psychomotor retardation, congenital corneal opacity and feet polysyndactyly. *Am. J. Med. Genet. A* **146A**, 2905–2910 (2008).
- 50 Hayashi, S., Mizuno, S., Migita, O., Okuyama, T., Makita, Y., Hata, A. et al. The CASK gene harbored in a deletion detected by array-CGH as a potential candidate for a gene causative of X-linked dominant mental retardation. *Am. J. Med. Genet. A* **146A**, 2145–2151 (2008).
- 51 Toyooka, K., Shionoya, A., Gambello, M. J., Cardoso, C., Leventer, R., Ward, H. L. et al. 14-3-3epsilon is important for neuronal migration by binding to NUDEL: a molecular explanation for Miller-Dieker syndrome. *Nat. Genet.* **34**, 274–285 (2003).
- 52 Mignon-Ravix, C., Cacciagli, P., El-Waly, B., Moncia, A., Milh, M., Girard, N. et al. Deletion of YWHAE in a patient with periventricular heterotopias and marked corpus callosum hypoplasia. *J. Med. Genet.* **47**, 132–136 (2010).
- 53 Haldeman-Englert, C. R., Gai, X., Perin, J. C., Ciano, M., Halbach, S. S., Geiger, E. A. et al. A 3.1-Mb microdeletion of 3p21.31 associated with cortical blindness, cleft lip, CNS abnormalities, and developmental delay. *Eur. J. Med. Genet.* **52**, 265–268 (2009).
- 54 Buysse, K., Delle Chiaie, B., Van Coster, R., Loeys, B., De Paep, A., Mortier, G. et al. Challenges for CNV interpretation in clinical molecular karyotyping: lessons learned from a 1001 sample experience. *Eur. J. Med. Genet.* **52**, 398–403 (2009).
- 55 Fan, Y. S., Jayakar, P., Zhu, H., Barbouth, D., Sacharow, S., Morales, A. et al. Detection of pathogenic gene copy number variations in patients with mental retardation by genomewide oligonucleotide array comparative genomic hybridization. *Hum. Mutat.* **28**, 1124–1132 (2007).
- 56 Hevner, R. F., Shi, L., Justice, N., Hsueh, Y., Sheng, M., Smiga, S. et al. Tbr1 regulates differentiation of the preplate and layer 6. *Neuron* **29**, 353–366 (2001).
- 57 Cosma, M. P., Pepe, S., Annunziata, I., Newbold, R. F., Grompe, M., Parenti, G. et al. The multiple sulfatase deficiency gene encodes an essential and limiting factor for the activity of sulfatases. *Cell* **113**, 445–456 (2003).
- 58 Dierks, T., Schmidt, B., Borissenko, L. V., Peng, J., Preusser, A., Mariappan, M. et al. Multiple sulfatase deficiency is caused by mutations in the gene encoding the human C(alpha)-formylglycine generating enzyme. *Cell* **113**, 435–444 (2003).
- 59 Behar, O., Golden, J. A., Mashimo, H., Schoen, F. J. & Fishman, M. C. Semaphorin III is needed for normal patterning and growth of nerves, bones and heart. *Nature* **383**, 525–528 (1996).
- 60 Eudy, J. D., Ma-Edmonds, M., Yao, S. F., Talmadge, C. B., Kelley, P. M., Weston, M. D. et al. Isolation of a novel human homologue of the gene coding for echinoderm microtubule-associated protein (EMAP) from the Usher syndrome type 1a locus at 14q32. *Genomics* **43**, 104–106 (1997).
- 61 He, Y. & Casaccia-Bonelli, P. The Yin and Yang of YY1 in the nervous system. *J. Neurochem.* **106**, 1493–1502 (2008).
- 62 Martin, C. L., Duvall, J. A., Ilkin, Y., Simon, J. S., Arreaza, M. G., Wilkes, K. et al. Cytogenetic and molecular characterization of A2BP1/FOX1 as a candidate gene for autism. *Am. J. Med. Genet.* **144B**, 869–876 (2007).
- 63 Tabolacci, E., Pomponi, M. G., Pietrobono, R., Terracciano, A., Chiurazzi, P. & Neri, G. A truncating mutation in the IL1RAPL1 gene is responsible for X-linked mental retardation in the MRX21 family. *Am. J. Med. Genet.* **140**, 482–487 (2006).
- 64 Nelson, J., Flaherty, M. & Grattan-Smith, P. Gillespie syndrome: a report of two further cases. *Am. J. Med. Genet.* **71**, 134–138 (1997).
- 65 Shaffer, L. G. & Bejjani, B. A. Medical applications of array CGH and the transformation of clinical cytogenetics. *Cytogenet. Genome Res.* **115**, 303–309 (2006).
- 66 Shaffer, L. G., Bejjani, B. A., Torchia, B., Kirkpatrick, S., Coppinger, J. & Ballif, B. C. The identification of microdeletion syndromes and other chromosome abnormalities: cytogenetic methods of the past, new technologies for the future. *Am. J. Med. Genet. C Semin. Med. Genet.* **145C**, 335–345 (2007).
- 67 Bejjani, B. A. & Shaffer, L. G. Clinical utility of contemporary molecular cytogenetics. *Annu. Rev. Genomics Hum. Genet.* **9**, 71–86 (2008).
- 68 Edelman, L. & Hirschhorn, K. Clinical utility of array CGH for the detection of chromosomal imbalances associated with mental retardation and multiple congenital anomalies. *Ann. NY Acad. Sci.* **1151**, 157–166 (2009).
- 69 de Ståhl, T. D., Sandgren, J., Piotrowski, A., Nord, H., Andersson, R., Menzel, U. et al. Profiling of copy number variations (CNVs) in healthy individuals from three ethnic groups using a human genome 32K BAC-clone-based array. *Hum. Mutat.* **29**, 398–408 (2008).
- 70 Shao, L., Shaw, C. A., Lu, X. Y., Sahoo, T., Bacino, C. A., Lalani, S. R. et al. Identification of chromosome abnormalities in subtelomeric regions by microarray analysis: a study of 5,380 cases. *Am. J. Med. Genet. A* **146A**, 2242–2251 (2008).
- 71 Lu, X., Phung, M. T., Shaw, C. A., Pham, K., Neil, S. E., Patel, A. et al. Genomic imbalances in neonates with birth defects: high detection rates by using chromosomal microarray analysis. *Pediatrics* **122**, 1310–1318 (2008).
- 72 Xu, J. & Chen, Z. Advances in molecular cytogenetics for the evaluation of mental retardation. *Am. J. Med. Genet. C Semin. Med. Genet.* **117C**, 15–24 (2003).
- 73 Ravnan, J. B., Tepperberg, J. H., Papenhausen, P., Lamb, A. N., Hedrick, J., Eash, D. et al. Subtelomere FISH analysis of 11 688 cases: an evaluation of the frequency and pattern of subtelomere rearrangements in individuals with developmental disabilities. *J. Med. Genet.* **43**, 478–489 (2006).
- 74 Ahn, J. W., Ogilvie, C. M., Welch, A., Thomas, H., Madula, R., Hills, A. et al. Detection of subtelomere imbalance using MLPA: validation, development of an analysis protocol, and application in a diagnostic centre. *BMC Med. Genet.* **8**, 9 (2007).
- 75 Schoumans, J., Ruivenkamp, C., Holmberg, E., Kyllerman, M., Anderlid, B. M. & Nordenskjöld, M. Detection of chromosomal imbalances in children with idiopathic mental retardation by array based comparative genomic hybridisation (array-CGH). *J. Med. Genet.* **42**, 699–705 (2005).
- 76 de Vries, B. B., Pfundt, R., Leisink, M., Koolen, D. A., Vissers, L. E., Janssen, I. M. et al. Diagnostic genome profiling in mental retardation. *Am. J. Hum. Genet.* **77**, 606–616 (2005).
- 77 Rosenberg, C., Knijnenburg, J., Bakker, E., Vienna-Morgante, A. M., Sloos, W., Otto, P. A. et al. Array-CGH detection of micro rearrangements in mentally retarded individuals: clinical significance of imbalances present both in affected children and normal parents. *J. Med. Genet.* **43**, 180–186 (2006).
- 78 Krepisch-Santos, A. C., Vienna-Morgante, A. M., Jehewe, F. S., Passos-Bueno, M. R., Knijnenburg, J., Suzhik, K. et al. Whole-genome array-CGH screening in undiagnosed syndromic patients: old syndromes revisited and new alterations. *Cytogenet. Genome Res.* **115**, 254–261 (2006).
- 79 Thuresson, A. C., Bondeson, M. L., Edeby, C., Ellis, P., Langford, C., Dumanski, J. P. et al. Whole-genome array-CGH for detection of submicroscopic chromosomal imbalances in children with mental retardation. *Cytogenet. Genome Res.* **118**, 1–7 (2007).
- 80 Wagenstaller, J., Spranger, S., Lorenz-Depiereux, B., Kazmierczak, B., Nathrath, M., Wahl, D. et al. Copy-number variations measured by single-nucleotide-polymorphism oligonucleotide arrays in patients with mental retardation. *Am. J. Hum. Genet.* **81**, 768–779 (2007).

- 81 Bruno, D. L., Ganesamoorthy, D., Schoumans, J., Bankier, A., Coman, D., Delatycki, M. *et al*. Detection of cryptic pathogenic copy number variations and constitutional loss of heterozygosity using high resolution SNP microarray analysis in 117 patients referred for cytogenetic analysis and impact on clinical practice. *J. Med. Genet.* **46**, 123–131 (2009).
- 82 Sagoo, G. S., Butterworth, A. S., Sanderson, S., Shaw-Smith, C., Higgins, J. P. & Burton, H. Array CGH in patients with learning disability (mental retardation) and congenital anomalies: updated systematic review and meta-analysis of 19 studies and 13,926 subjects. *Genet. Med.* **11**, 139–146 (2009).
- 83 Wincent, J., Anderlid, B. M., Lagerberg, M., Nordenskjöld, M. & Schoumans, J. High-resolution molecular karyotyping in patients with developmental delay and/or multiple congenital anomalies in a clinical setting. *Clin. Genet.* (e-pub ahead of print 8 May 2010).
- 84 Girirajan, S., Rosenfeld, J. A., Cooper, G. M., Antonacci, F., Siswara, P., Itsara, A. *et al*. A recurrent 16p12.1 microdeletion supports model for severe developmental delay. *Nat. Genet.* **42**, 203–209 (2010).
- 85 Veltman, J. A. & Brunner, H. G. Understanding variable expressivity in microdeletion syndromes. *Nat. Genet.* **42**, 192–193 (2010).

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Rapid improvement of life-threatening capillary leak syndrome after stem cell transplantation by bevacizumab

Hiromasa Yabe, Miharu Yabe, Takashi Koike, Takashi Shimizu, Tsuyoshi Morimoto and Shunichi Kato

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Aristoteles Giagounidis

Medizinische Klinik II, St Johannes Hospital,
Duisburg, Germany

Stefano A. Pileri

Hematopathology Unit, Policlinico S. Orsola,
University of Bologna, Bologna, Italy

Mario Cazzola

Department of Hematology Oncology, University of Pavia Medical School,
Fondazione IRCCS Policlinico San Matteo, Pavia, Italy

Andrew N. J. McKenzie

Medical Research Council Laboratory of Molecular Biology,
Cambridge, United Kingdom

James S. Wainscoat

LRF Molecular Haematology Unit, Nuffield Department of Clinical Laboratory
Sciences, John Radcliffe Hospital, Oxford, United Kingdom

Jacqueline Boulwood

LRF Molecular Haematology Unit, Nuffield Department of Clinical Laboratory
Sciences, John Radcliffe Hospital, Oxford, United Kingdom

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Correspondence: Dr Jacqueline Boulwood, University Reader at University

of Oxford, Co-Director, LRF Molecular Haematology Unit, Nuffield Department of Clinical Laboratory Sciences, John Radcliffe Hospital, Oxford OX3 9DU, United Kingdom; e-mail: jacqueline.boulwood@ndcls.ox.ac.uk.

References

1. McGowan KA, Li JZ, Park CY, et al. Ribosomal mutations cause p53-mediated dark skin and pleiotropic effects. *Nat Genet.* 2008;40(8):963-970.
2. Jones NC, Lynn ML, Gaudenz K, et al. Prevention of the neurocristopathy Treacher Collins syndrome through inhibition of p53 function. *Nat Med.* 2008;14(2):125-133.
3. Danilova N, Sakamoto KM, Lin S. Ribosomal protein S19 deficiency in zebrafish leads to developmental abnormalities and defective erythropoiesis through activation of p53 protein family. *Blood.* 2008;112(13):5228-5237.
4. Harris NL, Jaffe ES, Diebold J, et al. The World Health Organization classification of neoplastic diseases of the hematopoietic and lymphoid tissues. Report of the Clinical Advisory Committee meeting, Airie House, Virginia, November 1997. *Ann Oncol.* 1999;10(12):1419-1432.
5. Swerdlow SH, Campo E, Harris NL, et al. *WHO Classification of tumours of Haematopoietic and Lymphoid Tissues.* Lyon, France: IARC;2008.
6. Barlow JL, Drynan LF, Hewett DR, et al. A p53-dependent mechanism underlies macrocytic anemia in a mouse model of human 5q- syndrome. *Nat Med.* 2010;16(1):59-66.
7. Pellagatti A, Cazzola M, Giagounidis AA, et al. Gene expression profiles of CD34+ cells in myelodysplastic syndromes: involvement of interferon-stimulated genes and correlation to FAB subtype and karyotype. *Blood.* 2006;108(1):337-345.
8. Boulwood J, Pellagatti A, Cattan H, et al. Gene expression profiling of CD34+ cells in patients with the 5q- syndrome. *Br J Haematol.* 2007;139(4):578-589.
9. Paterson JC, Ballabio E, Mattsson G, Turner SH, Mason DY, Marafioti T. Labeling of multiple cell markers and mRNA using automated apparatus. *Appl Immunohistochem Mol Morphol.* 2008;16(4):371-381.
10. Ebert BL, Pretz J, Bosco J, et al. Identification of RPS14 as a 5q- syndrome gene by RNA interference screen. *Nature.* 2008;451(7176):335-339.

To the editor:

Rapid improvement of life-threatening capillary leak syndrome after stem cell transplantation by bevacizumab

Capillary leak syndrome (CLS) is a severe complication of allogeneic stem cell transplantation (SCT) characterized by weight gain, generalized edema, hypotension, and hypoalbuminemia.¹ The main CLS pathogenesis is injury of the capillary endothelium resulting in a loss of intravascular fluid into interstitial spaces. Treatment is limited to withdrawal of growth factors and systemic corticosteroids; however, a good response is limited and most severe CLS cases progress to fatal multiple-organ dysfunction syndrome. Vascular endothelial growth factor (VEGF) is a potent inducer of vascular permeability and may have a crucial role in the mechanism underlying CLS formation.² In the present study, we report the successful treatment of life-threatening CLS that developed after allogeneic SCT using the anti-VEGF antibody bevacizumab (Avastin; Chugai).³

A 6-year-old male with Fanconi anemia received marrow cells from a HLA-DRB1 mismatched unrelated donor as previously described.⁴ On day 22 after SCT, the patient developed posterior reversible encephalopathy syndrome with mild systemic edema, suggesting generalized injury of the vascular endothelium. Subsequently, grade 2 acute graft-versus-host disease of the skin and gastrointestinal tract ensued but was easily controlled with prednisolone. However, systemic edema accompanying consciousness disturbance, tachypnea and tachycardia developed 68 days after SCT. Computed tomography (CT) revealed massive pleural effusion (Figure 1A) and ascites, and the patient was diagnosed with CLS. Despite intensive conventional treatments, including prednisolone (1 mg/kg daily), ulinastatin (10 000 units/kg daily), and

albumin (0.8 g/kg every other day), hypotension, negative central venous pressure, and anuria developed 72 days after SCT. Because of the patient's critical condition and lack of response to other therapies, his case was discussed in the transplantation peer review group. Off-label use of bevacizumab was recommended. Written informed consent to the treatment in accordance with the Declaration of Helsinki and permission to publish results were obtained from the parents separately before the study and after the study, respectively. The publication of this study involving bevacizumab administration was approved by the institutional review board of Tokai University Hospital. Rationale and potential side effects were also discussed with the parents. Intravenous bevacizumab (5 mg/kg body weight) was administered over a 90-minute period. On the first day after treatment, urine production started to improve, and blood pressure and central venous pressure returned to the normal range. On the second day, all symptoms were ameliorated. A marked decrease in the amounts of pleural effusion was evident on the CT films obtained on the fifth day after bevacizumab administration (Figure 1B), and complete resolution of pleural effusion was revealed on the CT films taken 20 days after the treatment (Figure 1C). Plasma VEGF level before bevacizumab administration was not elevated (27 pg/mL; normal, < 115 pg/mL).

To the best of our knowledge, this is the first report on bevacizumab treatment of CLS developing after SCT. CLS after SCT has been difficult to ameliorate; however, bevacizumab was

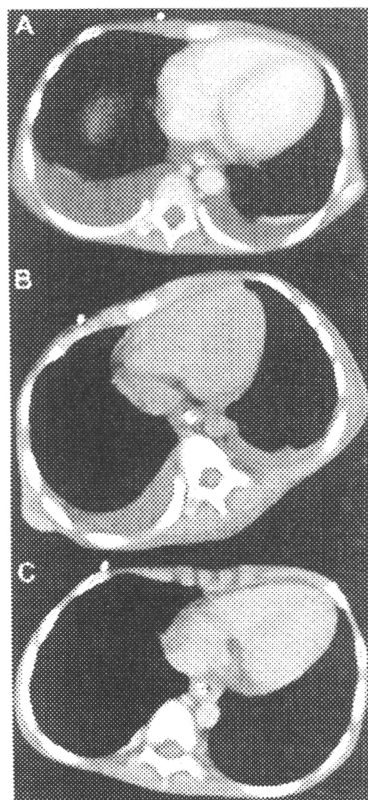


Figure 1. Administration of bevacizumab. Chest CT before (A), 5 days after (B), and 20 days after (C) treatment with bevacizumab.

shown to be highly effective against CLS in a patient even when plasma VEGF level was not increased, and may be useful under coexisting illness after SCT. Vascular endothelial damage plays a causal role in early complications of vascular origin after SCT, including hepatic veno-occlusive disease, engraftment syndrome, thrombotic microangiopathy, and idiopathic pneumonia syndrome.⁵ Bevacizumab may have a broad spectrum of efficacy against these complications.

Hiromasa Yabe
Tokai University School of Medicine,
Isehara, Japan

Miharu Yabe
Tokai University School of Medicine,
Isehara, Japan

Takashi Koike
Tokai University School of Medicine,
Isehara, Japan

Takashi Shimizu
Tokai University School of Medicine,
Isehara, Japan

Tsuyoshi Morimoto
Tokai University School of Medicine,
Isehara, Japan

Shunichi Kato
Tokai University School of Medicine,
Isehara, Japan

Contribution: H.Y. designed the study and wrote the paper; M.Y. performed diagnosis and planned treatment; T.K., T.S., and T.M. were substantially involved in clinical management; and S.K. performed real-time PCR and chimerism analysis.

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Correspondence: Dr Hiromasa Yabe, Department of Cell Transplantation and Regenerative Medicine, Tokai University School of Medicine, 143 Shimokasuya, Isehara, Kanagawa, 259-1193, Japan; e-mail: yabeh@is.icc.u-tokai.ac.jp.

References

1. Nümberger W, Willers R, Burdach S, et al. Risk factor for capillary leak syndrome after bone marrow transplantation. *Ann Hematol.* 1997;74(5):221-224.
2. Grove CS, Lee YC. Vascular endothelial growth factor: the key mediator in pleural effusion formation. *Curr Opin Pulm Med.* 2002;8(4):294-301.
3. Pichelmayer O, Zielinski C, Raderer M. Response of a nonmalignant pleural effusion to bevacizumab. *N Engl J Med.* 2005;353(7):740-741.
4. Yabe H, Inoue H, Matsumoto M, et al. Allogeneic haematopoietic cell transplantation from alternative donors with a conditioning regimen of low-dose irradiation, fludarabine and cyclophosphamide in Fanconi anaemia. *Br J Haematol.* 2006;134(2):208-212.
5. Woywodt A, Haubitz M, Buchholz S et al. Counting the cost: markers of endothelial damage in hematopoietic stem cell transplantation. *Bone Marrow Transplant.* 2004;34(12):1015-1023.



病気のはなし

糖原病 II 型 (ポンペ病, ライソゾーム病)

えとうよしかつ
衛藤義勝*



ポンペ病 (Pompe disease) は 1932 年ポンペにより報告された¹⁾ ライソゾーム病の一つであり、酸性 α -グルコシダーゼ (acid alpha glucosidase, GAA) の遺伝的酵素欠損により発症する。臨床的には乳児型と小児・成人型 (遅発型) に分類される。乳児型は乳児期早期に心拡大、心不全で 2 歳までに死亡する。遅発型では筋力低下、歩行障害、呼吸障害を呈し、最後は呼吸不全で死亡する重篤な疾患である。遺伝形式は常染色体劣性遺伝形式をとる。酵素補充療法が開発され、早期治療により症状の悪化を予防できる。ただし、酵素に対する抗体産生は治療効果を減弱することから、抗体に対する治療も試みられている²⁾。



用語解説

ライソゾーム病
細胞内の小胞器官で、多くの酸性水解酵素を含む。遺伝的に各種ライソゾーム酵素が欠損することにより種々のライソゾーム病を発症する。

ポンペ病

ポンペにより発見されたのでポンペ病と名付けられた。

酵素補充療法

酵素欠損の患者に酵素を投与して治療する方法。

シャペロン療法

低分子による酵素活性化療法。

病因と病態

本症は、ライソゾームに局在する酸性 α -グルコシダーゼ (GAA) の遺伝的酵素欠損により細胞内にグリコーゲンが蓄積することで発症する³⁾。図 1⁴⁾ にみられるように、筋肉組織のライソゾームにグリコーゲンが大量に蓄積すると同時にオートファジーによる細胞内蓄積が組織障害として重要である。

筋線維はそのために断裂し、炎症細胞の浸潤が認められる。過剰なオートファジーがさらに患者の筋崩壊に拍車をかける。GAA 活性はリンパ球、皮膚線維芽細胞において酵素活性を測定すると、対照の 10% 以下の活性であり、乳児型と遅発型 (後述) では残存酵素活性は遅発型で高値を示す⁵⁾。 α -グルコシダーゼは中性 α -グルコシダーゼ、マルターゼ活性などもあり、ポンペ病で低下

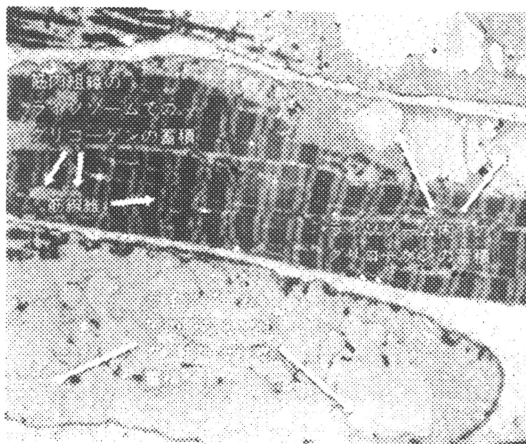


図 1 ポンペ病の筋組織像 (文献 4 から転載)
ライソゾームへのグリコーゲンの蓄積、筋線維の断裂。

* 東京慈恵会医科大学遺伝病研究講座 ☎105-8461 東京都港区西新橋 3-25-8

しているのはGAA活性である。GAA酵素をコードするGAA遺伝子は17番染色体の長腕にあり、約20 kbで20のエクソンを含む。cDNAのサイズは3.6 kbであり、952個のアミノ酸をコードする。糖鎖がつかないGAAの分子量は10,537である^{6,7)}。

ポンペ病の遺伝子変異としては欠失、挿入、重複、スプライシング異常、ノンセンス異常、ミスセンス異常など200種以上知られている。このほかの活性低下をきたさない偽性低下症(pseudo-deficiency)も知られている^{7,8)}。

欧米人で多い遺伝子変異はc-32-13TG, c525delT, c2481+102-2646+31の順で多いが現在まで多数の変異が見いだされている。日本人ではpSer. 529Val, pArg600Cys, pSer. 619Arg, pAsp645Glu, pAr4g672Glnの5種類が多い^{8,9)}。

疫学

ポンペ病の頻度は表1に示すように約40,000人に1人であるが、今後診断法が進歩し、治療も確立されたことから増える可能性がある。

臨床症状、病型および臨床経過

ポンペ病は臨床的には乳児型と遅発型に分類される。古典的乳児型はポンペによって1932年に報告された病型である¹⁾。乳児型は生後早期に心症状を呈する。それに対し、心症状が少なく、筋力低下、呼吸筋症状を主体とし、小児期発症あるいは15歳以降の成人期発症に発症するタイプを遅発型という。

残存酵素活性の程度により乳児型、非古典的乳児型、小児型、成人型に分類される(図2)。

1. 乳児型

乳児古典型の発症年齢は生後約2か月頃からで、心肥大、心不全症状が現れる(図3)¹⁰⁾。また、筋緊張は著明に低下を呈する。表1に示したとおり乳児型症例の臨床症状の頻度は約140,000人に1人である。初発症状として哺乳力障害を呈する。巨舌、左室肥大に伴う心不全、呼吸障害をきたす。筋力低下も著しい。運動発達も著明に障害されており、定頸、お座りも難しく、寝たきりの状況が続く。呼吸筋も障害を受け、横隔膜の動きも悪く、呼吸不全に陥る。1歳までには呼吸不

表1 ポンペ病の頻度

乳児型	1/138,000 (1/43,000~1/536,000)
遅発型	1/57,000 (1/27,000~1/128,000)
一般頻度*	1/40,000 (1/17,000~1/100,000)

全、心不全で90%は死亡する。

なお、非古典的乳児型は生後約4.8か月で発症、筋緊張低下が著明であり、ウェルトニッヒ・ホフマン病のような臨床型を呈する。心症状は少ない。哺乳力低下、体重増不良、発達障害、筋力低下で、1~3歳頃には呼吸不全のために人工呼吸管理が必要となる。心肥大は軽度であり、巨舌も軽度である^{5,11)}。

2. 遅発型

遅発型では16歳未満の発症を小児型、以降の発症を成人型としている。発症年齢はさまざまであり、残存酵素活性による。図4にわが国の各種ポンペ病の発症年齢を示す。Winkelらの報告では、ポンペ病225例のうち1歳未満の発症は15%、1~6歳では11%、6~18歳13%、18歳以上で62%である^{12,13)}。

1) 小児型

幼児期~学童期に筋力低下が徐々に発症する。立ち上がりが難しくなるなど、筋力低下が上肢より下肢にみられ、筋萎縮が少しずつ進行する。舌の筋力が低下するため、言語も不明瞭になりやすい。呼吸障害のため学童期~20歳頃までに人工換気が必要になる。また、横隔膜の障害のための呼吸筋障害により早朝の頭痛、日中の眠気、夜間の呼吸障害などがみられる。心肥大は通常認めないが、2~5%の患者で軽度の心肥大がみられる。

2) 成人型

発症年齢はさまざまであり、30歳くらいにピークがあるが、60~70歳で発症する患者もいる。発症年齢は残存酵素活性と相関がある。肢帯型筋ジストロフィーや先天性ミオパシーに類似した臨床症状を呈する。頸部から肢体を中心に筋萎縮が起こり、下肢の大腿筋、胸鎖乳突筋、横隔膜筋の萎縮が起こりやすい。また、翼状肩甲を示す(図5)。筋力低下のための歩行障害が生じ、転倒などしやすくなるため車いすが必要な状態とな

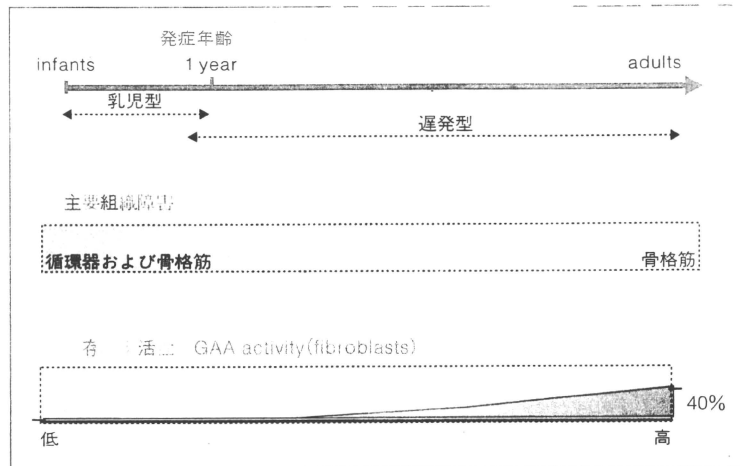


図2 ポンペ病の臨床的スペクトラム
残存酵素活性が高いほど発症年齢は遅くなる。

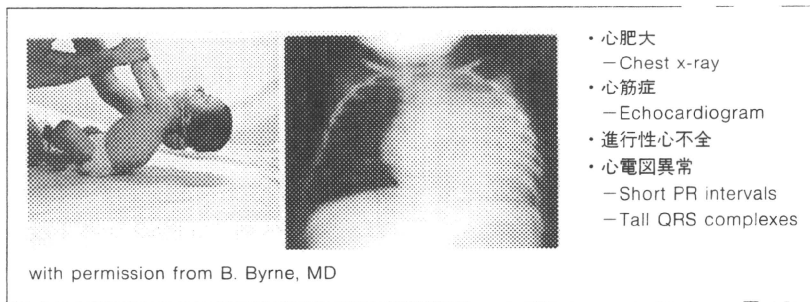


図3 乳児型ポンペ病の心症状

(文献10から転載)

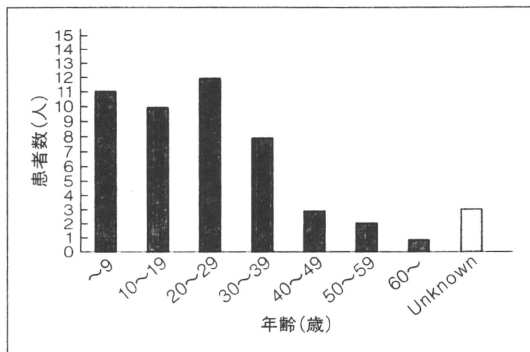


図4 日本人ポンペ病の治療患者年齢分布

る。嚥下障害も少しずつ現れ、発語も明瞭でなくなる。

30~40歳前後で呼吸筋の障害のために人工呼吸管理となる患者が多い。朝の頭痛、不眠なども病状の早期にはみられる。顔の筋も萎縮することから細い顔つきが多い。多くは気胸、肺炎などを

合併し呼吸不全で死亡する。

臨床検査所見と画像所見

1. 血液・生化学所見

通常血清CK以外は正常である。CK値は200~2,000 IU/lと正常から中等度上昇がみられる。AST, ALT, LDHも軽度上昇する。末梢血リンパ球の空胞化が乳児型でみられることもある。

2. 心電図所見

乳児型では左室肥大、PR間隔の短縮、QRSの高振幅を認める。小児型、成人型ではWPW症候群を呈する患者もいる。

3. CT, MRIなどの所見

筋CT, MRIでは小児型、成人型で大腿筋のうち大腿直筋、長内転筋の高吸収像などを認める。脳MRAでは椎骨脳底動脈での動脈瘤が遅発型で、また乳児型では頭部CT, MRIで脳白質の髄鞘化の遅れなどがみられる症例も認める。