CASE REPORT

Interstitial deletion within 7q31.1q31.3 in a woman with mild intellectual disability and schizophrenia

Keiko Akahoshi¹ Toshiyuki Yamamoto²

¹Department of Pediatrics, Tokyo Children's Rehabilitation Hospital, Tokyo, Japan; ²Institute of Medical Genetics, Tokyo Women's Medical University, Tokyo, Japan **Abstract:** We report the case of a Japanese woman with an interstitial deletion within the 7q31.1q31.3 region, she presented with mild intellectual disability since infancy, and later developed characteristic psychiatric manifestations, including abnormal behavior, delusions, and hallucinations. She was diagnosed with paranoid schizophrenia (F20.0, International Statistical Classification of Diseases and Related Health Problems 10th Revision). Array comparative genomic hybridization examination revealed the deletion involving several important genes for neurodevelopment. Particularly, *FOXP2*, *DOCK4*, *MET*, and *WNT2* in this region are suggested to be related to language impairment, autistic disorders, and cognitive disorders, via the WNT pathway. In addition, the WNT signal pathway has been suggested to be implicated in the pathogenesis of psychiatric disorders such as schizophrenia and bipolar disorder. However, there is no case report regarding schizophrenia associated with a 7q31 microdeletion. We suspect that the disruptions of these one or plural genes among the interstitial deletion of 7q31.1q31.3 may be involved in the development of schizophrenia in this woman. This is the first report on schizophrenia associated with a 7q31 microdeletion.

Keywords: chromosomal microarray, psychiatric disorder, autism spectrum disorder, ASD, Wnt pathway

Introduction

Schizophrenia is the most well-known chronic psychiatric disorder, characterized primarily by hallucinations and delusions and abnormal behaviors, predominantly developing in late puberty. Although the lifetime prevalence is approximately 0.5%–1.0% in the general population and a great amount of evidence has been accumulating on its cause, including responsible genes, it has not been clearly elucidated yet.¹

The 7q31 chromosomal region is rich in genes. Within this region, there are several genes involved in neurodevelopment. These genes affected by the Wnt pathway may participate in the development of psychiatric disorders.

In this study, we describe the case of a 28-year-old Japanese woman with mild intellectual disability associated with an interstitial deletion in 7q31 region. This patient also showed the phenotypes of schizophrenia.

Patient report

Our case involved a 29-year-old female, who was described as cheerful and peaceful. Although her father and older brother are healthy, her mother succumbed to fungal pneumonia. The patient was born at 38 weeks of gestation via vaginal delivery with early membrane rupture. Her birth weight was 2,700 g. During pregnancy, maternal edema and proteinuria were observed. There was no birth asphyxia; however, the patient showed poor sucking and muscular hypotonia. Her development was delayed

Neuropsychiatric Disease and Treatment 2018:14 1773-1778

1773

Correspondence: Keiko Akahoshi Tokyo Children's Rehabilitation Hospital, 4-10-1 Musashi-Murayama, Tokyo 208-0011, Japan Tel +81 42 561 2521 Fax +81 42 566 3753 Email fwkt4124@mb.infoweb.ne.jp



Comparison of the second second

as evidenced by gaining head control at 6 months, rolling over at 18 months, and standing at 22 months. The diagnosis of intellectual disability was made at 7 years of age, because her intelligence quotient measured by the Binet– Tanaka test was 53. A neurological examination at that time confirmed the presence of psychomotor retardation. Some minor anomalies, such as a horizontal line in her palm, higharched palate, and mandibular protrusion, were observed (Figure 1). After graduating from a special education school, she attended a workshop for the handicapped. She was active, helped her family well, and even participated in the Special Olympics.

At the age of 27 years, she suddenly became inactive and began neglecting her regular household chores. She refused to go to work and became withdrawn; she did not watch television, an activity she previously enjoyed. She was expressionless, was unkempt, and had stopped eating and bathing by herself. She suffered from insomnia, and her mood was unstable, irritable, and aggressive. She developed sluggish movements; talked to and laughed at herself; occasionally cried, shouted, and got irritated; and distressed her family.



Figure I The present patient, a 29-year-old female.

Laboratory tests, including thyroid function, revealed no abnormalities with the exception of a chromosomal abnormality. Brain magnetic resonance imaging demonstrated no remarkable changes, and electroencephalography revealed no paroxysmal activity. She was prescribed psychotropic drugs, but she did not use them. Thereafter, she demonstrated obsessive-compulsive, repetitive, ritualized behavior, and she was no longer able to eat, bath, and excrete by herself. She also experienced hallucinations and delusions. Some of her complaints included "some people whisper in my ears," "someone is chasing me," "something moves my hand," and "something touched my feet." Eight months later, her family reported that she behaved as a completely different person. She was diagnosed with paranoid schizophrenia (F20.0, International Statistical Classification of Diseases and Related Health Problems 10th Revision).

With family cooperation, she started medication, and her hallucinations and delusions improved following the use of risperidone (2 mg/day) and duloxetine (20 mg/day). Six months later, she began to smile and her appetite and sleep normalized. One year later she was able to eat meals by herself. At present, she can excrete by herself. Although her hallucinations, delusions, and psychomotor sluggishness have improved with medication, she continues to talk to herself and exhibits obsessive-compulsive behavior. She often throws tantrums when the order of daily activities set by her gets disturbed.

Cytogenetic examination

Conventional chromosomal G-banding revealed an interstitial deletion in 7q, described as 46,XX,del(7)(q31.3q31.3) (data not shown). Permission to analyze parental karyotyping could not be obtained. To confirm the precise deletion region, chromosomal microarray testing was performed with an Agilent 60 K Human Genome Comparative Genomic Hybridization Microarray platform (Agilent Technologies, Santa Clara, CA, USA) in accordance with previously reported methods.^{2,3} For this purpose, a peripheral blood sample was collected after obtaining written informed consent. Genomic DNA was extracted with the QIA quick DNA extraction kit (Qiagen, Hilden, Germany). The results are described as arr[hg19] 7q31.1q31.31(111,105,151-119,120,843)X1.

Discussion

In this study, an interstitial deletion in the 7q31 region was identified in a female patient with intellectual disability. In the deletion region, 25 registered RefSeq genes were

identified. The identified genes included FOXP2, DOCK4, MET, and WNT2. These genes play an important role in the function of central nervous system while affecting each other via the Wnt/β-catenin signaling pathway. Several studies have investigated the association between Wnt/β-catenin signaling pathway and autism spectrum disorders (ASD), and 7q31 has been considered a "hotspot" for ASD.⁴ The canonical Wnt pathway has been indicated to play a role in normal brain development and synaptic function and to affect dendrite growth and activity.5,6 Reportedly, Wnt and its receptors are involved in the embryonic development of central nervous system and are also highly expressed in adult brains, suggesting that their expression may affect higher brain functions and that Wnt signaling may be involved in higher brain functions. Therefore, genes affected by the Wnt pathway may be involved in the development of psychiatric disorders, such as schizophrenia or bipolar disorder.^{7,8}

Meanwhile, several previous studies have reported deletion cases similar to that reported here (Figure 2). Among them, cases with *FOXP2* deletion have mild-to-moderate

developmental delays (Table 1).^{2,9-16} FOXP2 encodes a transcription factor comprising a polyglutamine tract and a forkhead DNA-binding domain.¹⁷ FOXP2 protein may regulate the expression of other genes in the cortico-basal ganglia and cerebellar circuits of the brain.¹⁸ Disruptions in FOXP2 are associated with language, autistic, and cognitive disorders.^{19,20} Therefore, developmental delay in such patients is likely to be derived from FOXP2 haploinsufficiency. Interestingly, there is a report that FOXP2 may also regulate DISC1, which has attracted the most attention as a candidate gene for schizophrenia.²¹ However, there are both positive and negative reports on the association between FOXP 2 and schizophrenia.²²⁻²⁵ Further, in several previous studies as stated above including FOXP2 (Figure 2), schizophrenia has not been reported. DOCK4, a guanine nucleotide exchange factor (GEF) for Rac, is a component of the β -catenin destruction complex. Dock4 is expressed in the rat hippocampal neurons and plays an important role in supine formation.26 It has also been reported as candidate gene for autism, dyslexia, and schizophrenia.27-29 *MET* is the receptor for hepatocyte growth factor (HGF).

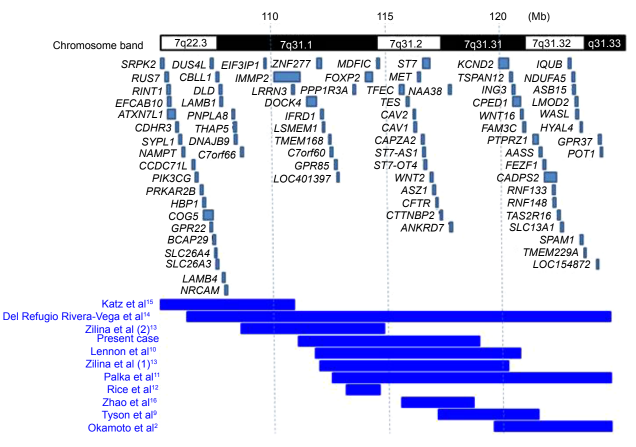


Figure 2 Genome map around 7q31 region depicting reported deletion regions.

Note: Deletion regions in previously reported patients and those in the present patient are depicted (blue bars). Genomic positions refer to build19.

	Katz	Del Refugio	Zilina		Present	Lennon et al ¹⁰	Zilina		Palka	Rice		Zhao	Tyson	Okamoto
	et al ⁱ⁵	Rivera-Vega et al ¹⁴	et al ^{!3} (2)		case		et al ¹³ (I)		et al''	et al ¹²		et al' ⁶	et al°	et al²
	2016	2015	2012			2007	2011		2012	2012		2016	2004	2011
Age	3y7m	ly9m	6y	28y	27y	77	Зу	28y	10y	4y10m	24y	4y	14y	3у
Gender	Σ	ч	Ъ	Ē	Ē		с	Ē	Ē	Σ	Ē	Ē	Σ	Σ
				(Mo)				(oM)			(Mo)			
Growth delay	+	+	I		I	I	+	I	I	I	I	I	+	+
Developmental delay	+	Mild	+	+	Mild	Mod	РоМ	I	Mild	+	I	+	I	I
ASD	AN	NA	AN		I	NA	AN	٩N	AN	AN	ΑN	AN	AN	+
Psychiatric symptoms	NA	NA	AN	Behavior	+	NA	NA	AN	NA	NA	ΝA	NA	ΝA	I
				abnormality										
Distinctive features	+	+	+		+	+	+	+	I	+	I	+	+	+
Deletion size (Mb)	12.0	23.0	6.0		8.0	9.0	8.0		14.0	2.0		3.0	5.0	5.0
									Mosaicism					
Chromosomal position														
Start	q22. I	q22.3	q31.1		q31.1	q31.1	q3 I. I		q31.1	q31.1		q31.2	q31.2	q31.31
End	q31.1	q32.I	q31.2		q31.31	q31.31	q31.31		q31.3	q31.2		q31.31	q31.32	q3 I.33
Deletion region (Mb)														
Start	66	107	601		112	112	112		113	113		116	117	120
End	Ξ	130	115		120	121	120		127	115		611	122	125
Inheritance	ΝT	DN	Inherited		NT	Inherited from	Inherited		AN	Inherited		ND	NT	DN
						parental-balanced								
						insertion								
FOXP2 involvement	I	+	+		+	+	+		+	+		I	I	I

HGF-MET signaling contributes to neuronal differentiation, cerebral cortex and cerebellum development, and axon growth. *MET* transcription is regulated by *FOXP2*,³⁰ and *DOCK4* and *MET* are related to the Wnt pathway through β -catenin.⁴ Finally, *WNT2* is a homologue of *WNT1*, and Wnt1 has been suggested to regulate genetic networks in midbrain dopaminergic neuron development.³¹ Wnt-1 protein has increased expression in the brains of individuals with schizophrenia.³² Similarly, *WNT2* may be associated with the development of schizophrenia.

FOXP2, *DOCK4*, *MET*, and *WNT2* are heterozygous in this case. We suspected that one possibility is that mutation or epigenetic change occurs somewhere in the contralateral allele and the disruptions of these plural genes affect each other and may be involved in the onset of schizophrenia in this patient.

Acknowledgments

We would like to thank the patient's family for their written informed consent to publish this case report and photographs. We also acknowledge the DECIPHER database for checking overlapping deletions. We are grateful to the technicians from our laboratories, including Ms Yumiko Ondo for her skillful help. This research was supported by the Practical Research Project for Rare/Intractable Diseases from the Japan Agency for Medical Research and Development (AMED), and the Japan Society for the Promotion of Science (JSPS) KAKENHI grant number 15K09631 (TY).

Disclosure

The authors report no conflicts of interest in this work.

References

- 1. Freedman R. Schizophrenia. N Engl J Med. 2003;349(18):1738-1749.
- Okamoto N, Hatsukawa Y, Shimojima K, Yamamoto T. Submicroscopic deletion in 7q31 encompassing CADPS2 and TSPAN12 in a child with autism spectrum disorder and PHPV. *Am J Med Genet A*. 2011; 155A(7):1568–1573.
- Sangu N, Shimojima K, Takahashi Y, Ohashi T, Tohyama J, Yamamoto T. A 7q31.33q32.1 microdeletion including LRRC4 and GRM8 is associated with severe intellectual disability and characteristics of autism. *Hum Genome Var.* 2017;4:17001.
- 4. Kalkman HO. A review of the evidence for the canonical Wnt pathway in autism spectrum disorders. *Molecular Autism*. 2012;3(1):10.
- Lie DC, Colamarino SA, Song HJ, et al. Wnt signalling regulates adult hippocampal neurogenesis. *Nature*. 2005;437(7063):1370–1375.
- Hiester BG, Galati DF, Salinas PC, Jones KR. Neurotrophin and Wnt signaling cooperatively regulate dendritic spine formation. *Mol Cell Neurosci*. 2013;56:115–127.
- 7. Wilson SW, Houart C. Early steps in the development of the forebrain. *Dev Cell*. 2004;6(2):167–181.
- Ciani L, Salinas PC. WNTs in the vertebrate nervous system: from patterning to neuronal connectivity. *Nat Rev Neurosci*. 2005;6(5):351–362.

- Tyson C, McGillivray B, Chijiwa C, Rajcan-Separovic E. Elucidation of a cryptic interstitial 7q31.3 deletion in a patient with a language disorder and mild mental retardation by array-CGH. *Am J Med Genet A*. 2004;129A(3):254–260.
- Lennon PA, Cooper ML, Peiffer DA, et al. Deletion of 7q31.1 supports involvement of FOXP2 in language impairment: clinical report and review. *Am J Med Genet A*. 2007;143A(8):791–798.
- Palka C, Alfonsi M, Mohn A, et al. Mosaic 7q31 deletion involving FOXP2 gene associated with language impairment. *Pediatrics* 2012; 129(1):e183–e188.
- Rice GM, Raca G, Jakielski KJ, et al. Phenotype of FOXP2 haploinsufficiency in a mother and son. *Am J Med Genet A*. 2012;158A(1): 174–181.
- Zilina O, Reimand T, Zjablovskaja P, et al. Maternally and paternally inherited deletion of 7q31 involving the FOXP2 gene in two families. *Am J Med Genet A*. 2012;158A(1):254–256.
- 14. Del Refugio Rivera-Vega M, Gomez-Del Angel LA, Valdes-Miranda JM, et al. A Novel 23.1 Mb interstitial deletion involving 7q22.3q32.1 in a girl with short stature, motor delay, and craniofacial dysmorphism. *Cytogenet Genome Res.* 2015;145(1):1–145.
- Katz OL, Krantz ID, Noon SE. Interstitial deletion of 7q22.1q31.1 in a boy with structural brain abnormality, cardiac defect, developmental delay, and dysmorphic features. *Am J Med Genet C Semin Med Genet*. 2016;172(2):92–101.
- Zhao J, Noon SE, Krantz ID, Wu Y. A de novo interstitial deletion of 7q31.2q31.31 identified in a girl with developmental delay and hearing loss. *Am J Med Genet C Semin Med Genet*. 2016;172(2):102–108.
- Lai CS, Fisher SE, Hurst JA, Vargha-Khadem F, Monaco AP. A forkhead-domain gene is mutated in a severe speech and language disorder. *Nature*. 2001;413(6855):519–523.
- Enard W. FOXP2 and the role of cortico-basal ganglia circuits in speech and language evolution. *Curr Opin Neurobiol*. 2011;21(3):415–424.
- Reuter MS, Riess A, Moog U, et al. FOXP2 variants in 14 individuals with developmental speech and language disorders broaden the mutational and clinical spectrum. *J Med Genet*. 2017;54:64–72.
- 20. Vernes SC, Spiteri E, Nicod J, et al. High-throughput analysis of promoter occupancy reveals direct neural targets of FOXP2, a gene mutated in speech and language disorders. *Am J Hum Genet*. 2007;81(6):1232–1250.
- Walker RM, Hill AE, Newman AC, et al. The DISC1 promoter: characterization and regulation by FOXP2. *Hum Mol Genet*. 2012;21(13): 2862–2872.
- 22. Sanjuán J, Tolosa A, González JC, et al. Association between FOXP2 polymorphisms and schizophrenia with auditory hallucinations. *Psychiatr Genet.* 2006;16(2):67–72.
- Tolosa A, Sanjuán J, Dagnall AM, et al. FOXP2 gene and language impairment in schizophrenia: association and epigenetic studies. *BMC Med Genet*. 2010;22(11):114.
- Rao W, Du X, Zhang Y, et al. Association between forkhead-box P2 gene polymorphism and clinical symptoms in chronic schizophrenia in a Chinese population. *J Neural Transm (Vienna)*. 2017;124:891–897.
- Yin J, Jia N, Liu Y, et al. No association between FOXP2 rs10447760 and schizophrenia in a replication study of the Chinese Han population. *Psychiatr Genet.* 2018;28(2):19–23.
- Ueda S, Negishi M, Katoh H. Rac GEF Dock4 interacts with cortactin to regulate dendritic spine formation. *Mol Biol Cell*. 2013;24(10): 1602–1613.
- Pagnamenta AT, Bacchelli E, de Jonge MV, et al. Characterization of a family with rare deletions in CNTNAP5 and DOCK4 suggests novel risk loci for autism and dyslexia. *Biol Psychiatry*. 2010;68(4):320–328.
- Maestrini E, Pagnamenta AT, Lamb JA, et al. High-density SNP association study and copy number variation analysis of the AUTS1 and AUTS5 loci implicate the IMMP2L-DOCK4 gene region in autism susceptibility. *Mol Psychiatry*. 2010;15(9):954–968.
- Alkelai A, Lupoli S, Greenbaum L, et al. DOCK4 and CEACAM21 as novel schizophrenia candidate genes in the Jewish population. *Int J Neuropsychopharmacol.* 2012;15(4):459–469.

- Mukamel Z, Konopka G, Wexler EL, et al. Regulation of MET by FOXP2, genes implicated in higher cognitive dysfunction and autism risk. *J Neurosci*. 2011;31(32):11437–11442.
- Wurst W, Prakash N. Wnt1-regulated genetic networks in midbrain dopaminergic neuron development. J Mol Cell Biol. 2014;6(1):34–41.
- Miyaoka T, Seno H, Ishino H. Increased expression of Wnt-1 in schizophrenic brains. Schizophr Res. 1999;38(1):1–6.

Neuropsychiatric Disease and Treatment

Publish your work in this journal

Neuropsychiatric Disease and Treatment is an international, peerreviewed journal of clinical therapeutics and pharmacology focusing on concise rapid reporting of clinical or pre-clinical studies on a range of neuropsychiatric and neurological disorders. This journal is indexed on PubMed Central, the 'PsycINFO' database and CAS, and is the official journal of The International Neuropsychiatric Association (INA). The manuscript management system is completely online and includes a very quick and fair peer-review system, which is all easy to use. Visit http://www.dovepress.com/testimonials.php to read read quotes from published authors.

Submit your manuscript here: http://www.dovepress.com/neuropsychiatric-disease-and-treatment-journal and the second second

Dovepress