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ORIGINAL RESEARCH

Association of Two Variable Number of Tandem Repeats in the Monoamine Oxidase A Gene Promoter with Schizophrenia

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Correspondence: Akitoyo Hishimoto Department of Psychiatry, Yokohama City University Graduate School of Medicine, 3-9 Fukuura, Kanazawa, Yokohama, 236-0004, Japan Tel +81-45-783-2540 Fax +81-45-783-2540 Email hishipon@yokohama-cu.ac.jp **Background:** Monoamine oxidase-A (MAO-A) decomposes dopamine and serotonin, and decreased MAO-A expression increases monoamine levels and is related to the pathophysiology of schizophrenia. Previous studies have reported that variable number of tandem repeats (VNTR), namely, upstream (u)VNTR, and some single nucleotide polymorphisms (SNPs) in the *MAOA* gene are associated with schizophrenia.

Methods: We investigated the two VNTRs and their related SNPs (rs6323 and rs1137070) in the *MAOA* gene promoter in 859 patients with schizophrenia and 826 healthy controls. Distal (d)VNTR and uVNTR were genotyped with fluorescence-based fragment polymerase chain reaction assays, and rs6323 and rs1137070 with TaqMan SNP genotyping assays.

Results: Neither the genotype nor allelic frequency of the VNTRs or SNPs showed significant differences between the schizophrenia and control groups. On the other hand, analysis of the dVNTR-uVNTR-rs6323-rs1137070 haplotype showed significant association for nine repeats (9R)-3R-T-C in female patients (corrected p = 0.0006, odds ratio [confidence interval] = 2.17 [1.446–3.257]).

Conclusion: Our findings provide novel evidence that *MAOA* gene polymorphisms are associated with an increased risk of developing schizophrenia in females.

Keywords: haplotype, monoamine oxidase A, polymorphism, schizophrenia, variable number of tandem repeats

Introduction

Schizophrenia is a severe psychiatric disorder that affects approximately 1% of the global population.¹ Schizophrenia has high heritability and is associated with complex poly genic factors.² Pharmacological studies have indicated that a dysfunction of dopaminergic neurons could contribute to the development of schizophrenia.¹ Dopamine degradation is catalyzed by monoamine oxidase (MAO) and catechol-o-methyltransferase (COMT) in the brain.^{3,4} Many studies have investigated the association of MAO and COMT with schizophrenia.^{4–7}

There are two types of MAO, MAO-A and MAO-B, both of which contribute to the degradation of dopamine.³ MAO-A has primary and minor isoforms; however, the functional differences between these isoforms remain unknown.⁸ The *MAOA* and *MAOB* genes are located adjacently to each other on the X chromosome, in the opposite direction.³ MAO-A has been reported to play an important role in mental illnesses such as schizophrenia.^{9,10} There is an upstream (u) variable number of tandem repeats (VNTR) in the *MAOA* gene promoter. uVNTR is located 1.2 kb

© 2021 Tanifuji et al. This work is published and licensed by Dove Medical Press Limited. The full terms of this license are available at https://www.dovepress.com/terms work you hereby accept the Terms. Non-commercial uses of the work are permitted without any further permission from Dove Medical Press Limited, provided the work is properly attributed. For permission for commercial use of this work, please see paragraphs 42 and 5 of our Terms (https://www.dovepress.com/terms.php). upstream of the *MAOA* gene, and is present in 3 repeats (R), 3.5R, 4R, or 5R. The 3.5R and 4R are high-expression alleles, and the 2R, 3R, and 5R are low-expression alleles.¹¹ Several studies have reported that the uVNTR and its related single nucleotide polymorphisms (SNPs) of the *MAOA* gene are associated with schizophrenia;^{12–15} however, these results are inconsistent with each other.^{16–18}

Recently, a novel VNTR, namely distal (d)VNTR has been identified in the *MAOA* gene promoter region. dVNTR is located approximately 500 bp upstream of uVNTR and present in 8R, 9R, 10R, 11R or 12R. dVNTR from 8R to 11R were found to be associated with uVNTR, and the corresponding transcripts were evaluated. 9R and 10R are associated with the highest and lowest levels of transcription, respectively, whereas 8R and 11 R show a moderate level of transcription.¹⁹ In addition, it was demonstrated that dVNTR and uVNTR are involved in the expression of the two MAO-A isoforms, wherein dVNTR increases the expression of the primary isoform that had little connection to uVNTR, and both VNTRs reduce the expression of a minor isoform that comprised a fraction of the total.⁸

In neuropsychiatric disorders, the combination of dVNTR and uVNTR was reported to be associated with nicotine dependence.²⁰ However, there is no study that explored the association between the two VNTRs and other mental illnesses such as schizophrenia. In this study, we investigated the association of the two *MAOA* gene promoter VNTRs, and their related SNPs, with schizophrenia.

Materials and Methods

Participants

We recruited 859 patients with schizophrenia and 826 healthy controls of Japanese descent from the city of Kobe in Japan. The demographic and clinical characteristics of the participants are given in Table 1. At least two psychiatrists diagnosed every patient based on the criteria listed in the Diagnostic and Statistical Manual of Mental Disorders, 4th Edition (DSM-IV) or DSM-5, and performed unstructured interviews and reviews of the patient's medical records at each hospital. The control participants were interviewed by a psychiatrist and screened for psychiatric disorders on the basis of unstructured interviews; the inclusion criteria were not having a present, past, and family history (first degree relatives)

of psychiatric disorders or substance abuse diagnosis (excluding nicotine dependence).

We implemented this study design and all related procedures in accordance with the Declaration of Helsinki. This study was approved by the Ethical Committee for Genetic Studies of Kobe University Graduate School of Medicine. Written informed consent was obtained from all the participants prior to the commencement of the experiments.

Genotyping of uVNTR and dVNTR in the MAOA Gene Promoter

Peripheral blood samples were drawn from the participants, and DNA was extracted using QIAamp DNA Blood Midi Kit (Qiagen Inc., Valencia, CA, USA). The quantity and purity of the DNA were assessed via NanoDrop (Thermo Fisher Scientific, Waltham, MA, USA) and the DNA samples were stored at -80 °C until analysis. For the MAOA uVNTR and dVNTR genotyping, we performed the fluorescence-based fragment polymerase chain reaction (PCR) assav according to previous studies.^{8,20} The MAOA uVNTR PCR assay volume (10 µL) contained 1 ng of genomic DNA, 5 µL of AmpliTag Gold Master Mix (Applied Biosystems, Foster City, CA, USA), and 15 pmol each of the following primers: 5'-GAA CGG ACG CTC CAT TCG GA-3' as a forward primer labeled with 6- Fluorescence (FAM) and 5'-ACA GCC TGA CCG TGG AGA AG-3' as a reverse primer (Invitrogen, Carlsbad, CA, USA). Thermal cycling comprised 10 min of initial denaturing at 95 °C followed by 40 cycles of 95 °C for 30 s, 55 °C for 30 s, and 72 °C for 30 s, and a final extension step of 7 min at 72 °C. The MAOA dVNTR PCR assay volume (20 µL) contained 10 ng of genomic DNA, 10 µL of AmpliTag Gold Master Mix with 10% GC enhancer (Applied Biosystems), 0.1 µM of 7 deazadGTP (Bio Labs, NewYork, NY, USA) and 25 pmol of each the primers: 5'-GGG TTA AGC GCC TCA GCT TC-3' as a forward primer labeled with 6-FAM and 5'-CAA GAG TGG ACT TAA GGA AGC AG-3' as a reverse primer (Invitrogen). Thermal cycling comprised 10 min of initial denaturing at 95 °C followed by 10 cycles of 95 °C for 20 s, the touchdown annealing steps from 65-56 °C for 20 s, and 72 °C for 30 s, 35 cycles of 95 °C for 20 s, 55 °C for 20 s, and 72 °C for 30 s, followed by a final extension step of 7 min at 72 °C. Both PCR the products were analyzed using SeqStudio Genetic Analyzer (Applied Biosystems) and GeneMapper Software version 6 (Applied Biosystems).

	CTL (n = 826)	SCZ (n = 859)	p-value
Sex (male/female)	394/432	446/413	0.083 ^a
Age/all (years), median (IQR)	53.0 (36.0, 67.0)	55.0 (43.0, 65.0)	0.049 ^b
Age/male, median (IQR)	52.0 (34.8, 67.0)	55.0 (43.0, 64.0)	0.078 ^b
Age/female, median (IQR)	55.0 (38.0, 67.0)	56.0 (44.0, 66.0)	0.248 ^b
Age of onset/all (years), median (IQR)	-	24.0 (20.0, 30.0)	
Age of onset/male, median (IQR)	-	23.0 (19.0, 30.0)	
Age of onset/female, median (IQR)	-	25.0 (20.0, 31.5)	

Notes: We collected precise information about age from the clinical records of 810 (98%) out of 826 healthy controls and 843 (98%) out of 859 patients with schizophrenia, as well as about the age of onset from the clinical records of 683 (80%) out of 859 patients with schizophrenia. ^aWe evaluated *p*-value with the χ^2 -test between the schizophrenia and control groups. ^bWe evaluated *p*-value with the Mann-Whitney *U*-test between the schizophrenia and control groups. **b**We evaluated *p*-value with the Mann-Whitney *U*-test between the schizophrenia and control groups. **b**We evaluated *p*-value with the Mann-Whitney *U*-test between the schizophrenia and control groups.

Genotyping of rs6323 and rs1137070

We used TaqMan SNP genotyping assays, rs6323 (Assy ID: ANEP6VZ) and rs1137070 (Assy ID:C___8878813_20), obtained from Thermo Fisher Scientific database (<u>http://www.thermofisher.com</u>) as described previously.²¹ Genotyping was performed on a 7500 Real-Time PCR System (Applied Biosystems) according to the manufacturer's protocol.

Statistical Analysis

The data were analyzed using R version 4.0.0 (R development core team, Vienna, Austria) and EZR version 1.42 (Saitama Medical Center, Jichi Medical University, Saitama, Japan). We used Haploview version 4.2 (Dlay Lab, Broad Institute Cambridge, MA, USA)²² to analyze allele/haplotype frequencies and genetic association in females. Differences between the groups were analyzed using χ^2 and Mann-Whitney *U*-tests. We examined genotype–based associations, and alleles and haplotypes using the Cochran–Armitage trend test and χ^2 _test, respectively, and permutation tests based on 10,000 replications were performed for the correction, as necessary. The threshold for statistical significance was defined as a two tailed p < 0.05.

Results

Since the X chromosome contains the *MAOA* gene, we analyzed each sex separately. We tested VNTR polymorphisms via Hardy–Weinberg equilibrium (HWE) in females with Fisher's exact test. (dVNTR, control p = 0.783 and schizophrenia p = 0.984; uVNTR, control p = 0.780 and schizophrenia p = 0.108; rs6323, control p = 0.397 and schizophrenia p = 0.606; rs1137070, control p = 0.233 and schizophrenia p = 0.584). The genotype and allelic frequency

for dVNTR, uVNTR, rs6323, and rs1137070 are shown in Tables 2 and 3. Neither the genotype nor allelic frequency of the VNTRs or SNPs was significantly different between the schizophrenia and control groups.

The association analysis of dVNTR-uVNTR-rs6323rs1137070 haplotype is given in Table 4. The distribution of the haplotype comprising two VNTRs and the two SNPs showed a significant association for 9R-3R-T-C (p = 0.0001) and 10R-4R-G-T (p = 0.0139) in the female patients. Permutation tests based on 10,000 replications were performed, and significant differences were seen in the 9R-3R-T-C (p = 0.0006) haplotype, but not in the 10R-4R-G-T (p = 0.0799). Participants with the 9R-3R-T-C haplotype had 2.17 times increased odds of developing schizophrenia (odds ratio [confidence interval] = 2.17 [1.446–3.257], p = 0.0006). The haplotype containing dVNTR(9R) showed clearly significantly differences, when analyzed using the sliding window method (Table 5).

Discussion

In this study, we investigated whether the two VNTRs (dVNTR and uVNTR) and two SNPs (rs6323 and rs1137070) in the *MAOA* gene promoter are associated with schizophrenia. We found that the distribution of a haplotype consisting of the two VNTRSs and two SNPs was significantly associated with schizophrenia in females.

Although multiple studies have investigated the association of uVNTR and SNPs in the *MAOA* gene with schizophrenia, their results are not consistent.^{12–18} Furthermore, no studies have investigated the association between dVNTR and schizophrenia. To the best of our knowledge, this is the first study to investigate the association of the two *MAOA* VNTRs with schizophrenia.

Polymorphism	CTL (n = 394)	= 394)			SCZ (n = 446)	= 446)			Chi Square	Allele p-value ^b	Odds Ratio (95% CI)	Power
	Genoty	Genotype Distribution ^a	ution ^a	Allele Freq	Genotyp	Genotype Distribution ^a	ution ^a	Allele Freq				
	x/x	-/×	-/-		x/x	-/×	-/-					
dVNTR _{8R-12R}												
8R		_	393	0.003		0	446	0.0	1.13	0.287	NA	0.225
9R		163	231	0.414		661	247	0.446	0.90	0.343	1.141 (0.868–1.502)	0.152
IOR		224	170	0.569		244	202	0.547	0.39	0.532	0.917 (0.698–1.204)	0.093
IIR		4	390	0.010		З	443	0.007	0.297	0.586	0.660 (0.147–2.968)	0.071
I 2R		2	392	0.005		0	446	0.0	2.27	0.132	NA	0.331
uVNTR _{2R-4R}												
2R		e	391	0.008		8	438	0.018	1.72	0.189	2.381 (0.627–9.036)	0.237
3R		234	160	0.594		245	201	0.549	1.70	0.193	0.833 (0.634–1.096)	0.259
4R		157	237	0.398		193	253	0.433	10.1	0.315	1.151 (0.874–1.516)	0.175
SNP rs6323 (G/T) G		219	175	0.556		238	208	0.534	0.42	0.519	0.914 (0.639–1.308)	0.093
SNP rs1137070 (T/C)												
F		224	170	0.569		240	206	0.538	0.78	0.376	0.884 (0.673–1.161)	0.145
Notes : ^a This column shows the reference allele homozygotes, heterozygotes, and others as x/x, x/-, and -/-, respectively. There is no x/x in male samples because the p -values with the χ^2 -test. If the nominal <i>p</i> -value significantly showed difference ($p < 0.05$), the precise <i>p</i> -value for multiple testing (10,000 permutations) is calculated. Abbreviations : MAOA, monoamine oxidase A; CTL, healthy controls; SCZ, schizophrenia; CI, confidence interval; NA, not applicable.	the reference he nominal か noamine oxid	allele homoz value signific lase A; CTL,	zygotes, hete antly showec healthy cont	rozygotes, and other 1 difference (p < 0.0 rols; SCZ, schizoph	s as x/x, x/-, 5), the precis enia; CI, con	and -/-, resp ie p-value foi fidence inter	ectively. Thei r multiple te: ~val; NA, not	re is no x/x in male sting (10,000 permu : applicable.	samples because the itations) is calculater	: MAOA gene is located d.	Notes: ^a This column shows the reference allele homozygotes, heterozygotes, and others as x/x, x/-, and -/-, respectively. There is no x/x in male samples because the MAOA gene is located in the X chromosome. ^b We evaluated allelic p -values with the χ^2 -test. If the nominal <i>p</i> -value significantly showed difference ($p < 0.05$), the precise <i>p</i> -value for multiple testing (10,000 permutations) is calculated. Abbreviations: MAOA, monoamine oxidase A; CTL, healthy controls; SCZ, schizophrenia; CI, confidence interval; NA, not applicable.	luated allelic

Table 2 Allelic and Genotypic Distribution of the Polymorphisms in the MAOA Promoter in Healthy Controls and Male Patients with Schizophrenia

tion of the Po	lymor	phisms	s in the	e MAOA Prom	oter in He	tion of the Polymorphisms in the MAOA Promoter in Healthy Controls and Female Patients with Schizophrei	Female Patient	s with Schizophre	
	scz	SCZ (n = 413)	413)		Z value	Genotype	Chi Square	Allele p-value ^c	
Allele Freq	Gen Disti	Genotype Distribution ^a	on ^a	Allele Freq		p-value ^b			
	x/x	-/x	-/-						
0.0	0	2	411	0.002	AN	NA	2.09	0.148	
0.398	76	187	150	0.410	0.49	0.623	0.26	0.608	
0.594	144	187	82	0.575	0.75	0.455	0.61	0.436	
0.006	0	6	404	0.011	AN	NA	1.34	0.247	
0.002	0	_	412	0.001	NA	AA	0.29	0.590	
0.006	2	4	407	0.010	0.80	0.422	0.84	0.359	
0.622	183	159	71	0.636	0.56	0.574	0.36	0.550	
0.372	67	158	188	0.354	0.73	0.468	0.59	0.441	

432 168 82 427

0 184 187

0 80 0 0

8R 9R 10R 11R

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×'

×/×

dVNTR_{8R-12R}

Distribution^a Genotype

430

5 2

uVNTR_{2R-4R}

with Schizophrenia Table 3 Allelic and Genotypic Distribution

CTL (n = 432)

Polymorphism

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Power

Odds Ratio (95% CI)

0.157 0.054

0.123 0.055

0.081

0.926 (0.763-1.124) 1.893 (0.632–5.671)

0.522 (0.047–5.772)

1.052 (0.866–1.278)

٩Z

2R	0	ß	427	0.006	7	4	407	0.010	0.80	0.422	0.84	0.359	1.680 (0.547–5.157)	0.097
3R	177	183	2	0.622	183	159	71	0.636	0.56	0.574	0.36	0.550	1.062 (0.872–1.294)	0.062
4R	69	183	180	0.372	67	158	188	0.354	0.73	0.468	0.59	0.441	0.925 (0.758–1.128)	0.078
5R	0	_	431	0.001	0	_	412	0.001	AN	AN	0.001	0.975	1.046 (0.653–16.751)	AN
SNP rs6323 (G/T) G	156	161	191 85	0.582	142	188	83	0.571	0.43	0.667	0.20	0.655	0.957 (0.789–1.161)	0.051
SNP rs1137070 (T/C) T	161	161 185 86	86	0.589	144	187	82	0.575	0.47	0.640	0.24	0.625	0.953 (0.785–1.156)	0.061
Notes: "This column shows the reference allele homozygotes, heterozygotes, genotypic p -values with the Cochran-Armitage trend test. "We evaluated allelic calculated.	ws the r e Cochra	eferenc∈ ın-Armit	e allele h tage trer	nomozygotes, heter nd test. ^c We evalua	ozygote ted allel	ss, and c ic p-valu	others as es with	x/x , $x/-$, and $-/-$, the χ^2 -test. If the	respectively. nominal p-val	There is no x/x in male submitted to the significantly showed di	amples because th fference ($p < 0.05$)	e MAOA gene is loca the precise <i>p</i> -value fo	and others as x/x, x/-, and -/-, respectively. There is no x/x in male samples because the MAOA gene is located in the X chromosome. ^b We evaluated p -values with the χ^{2-} test. If the nominal <i>p</i> -value significantly showed difference ($p < 0.05$), the precise <i>p</i> -value for multiple testing (10,000 permutations) is	e evaluated (

Abbreviations: MAOA, monoamine oxidase A; CTL, healthy controls; SCZ, schizophrenia; Cl, confidence interval; NA, not applicable.

Polymorphism	CTL	(n = 826)	SCZ	(n = 859)	Chi Square	p-value ^a	Odds Ratio (95% CI)	Power
	n	Frequency	n	Frequency				
Male (CTL, n = 39	4; SCZ, 1	n = 446)		1			•	1
8R-3R-G-T	I	0.003	0	0.00	1.13	0.287	NA	0.225
9R-3R-T-C	11	0.028	23	0.052	3.01	0.083	1.893 (0.911–3.935)	0.418
9R-3R-G-T	5	0.013	7	0.016	0.13	0.714	1.241 (0.391-3.940)	0.054
9R-4R-T-C	142	0.360	165	0.370	0.08	0.774	1.042 (0.786–1.381)	0.048
9R-4R-G-T	I.	0.003	2	0.004	0.22	0.637	1.770 (0.160–19.598)	0.118
9R-4R-T-T	4	0.010	2	0.004	0.95	0.330	0.439 (0.080-2.411)	0.189
I0R-2R-G-T	3	0.008	8	0.018	1.72	0.189	2.381 (0.627-9.036)	0.237
IOR-3R-T-C	16	0.041	12	0.027	1.22	0.270	0.653 (0.305-1.398)	0.204
IOR-3R-G-T	195	0.495	201	0.451	1.64	0.200	0.837 (0.638-1.099)	0.246
I0R-4R-T-C	I	0.003	6	0.013	3.02	0.083	5.359 (0.642-44.708)	0.351
I0R-4R-G-T	8	0.002	17	0.038	2.30	0.130	1.912 (0.816-4.480)	0.962
IOR-4R-T-T	1	0.003	0	0.000	1.13	0.287	NA	0.225
IIR-3R-G-T	4	0.010	2	0.004	0.95	0.33	0.439 (0.080-2.412)	0.189
IIR-4R-G-T	0	0.000	I.	0.002	0.88	0.347	NA	0.127
I2R-3R-G-T	2	0.005	0	0.000	2.27	0.132	NA	0.331
Female (CTL, n = 4	432; SCZ	Z, n = 413)						
9R-2R-T-C	I	0.001	2	0.003	0.59	0.442	2.100 (0.189–23.091)	0.153
9R-3R-T-C	37	0.043	74	0.090	14.59	0.0001 (0.0006)	2.170 (1.446–3.257)	0.973
9R-3R-G-T	П	0.012	11	0.014	0.05	0.822	1.101 (0.475–2.554)	0.055
9R-4R-T-C	289	0.336	244	0.295	3.19	0.074	0.829 (0.674–1.018)	1.000
9R-4R-T-T	3	0.003	5	0.006	0.59	0.444	1.740 (0.414–7.303)	0.151
9R-4R-G-T	3	0.003	2	0.003	0.05	0.822	0.693 (0.116-4.160)	NA
9R-5R-T-C	I	0.001	1	0.001	0.001	0.976	1.044 (0.065–16.713)	NA
I0R-2R-G-T	4	0.005	6	0.007	0.36	0.549	1.558 (0.438–5.541)	0.078
IOR-3R-G-T	457	0.531	402	0.486	3.33	0.068	0.837 (0.691–1.013)	0.455
IOR-3R-T-C	23	0.027	25	0.030	0.15	0.696	1.121 (0.632–1.987)	0.056
IOR-3R G-C	I	0.001	2	0.002	0.37	0.541	2.085 (0.189-23.037)	0.078
10R-4R-G-T	19	0.022	36	0.044	6.05	0.0139 (0.0799)	2.001 (1.14–3.511)	0.717
I0R-4R-T-C	5	0.005	4	0.005	0.03	0.865	0.832 (0.223-3.110)	NA
IIR-3R-G-T	5	0.006	8	0.010	0.84	0.359	1.680 (0.547–5.158)	0.151
I2R-3R-G-T	2	0.002	1	0.001	0.29	0.589	0.521 (0.047-5.759)	0.075

Table 4 Haplotypic Distribution of Polymorphisms in the MAOA Promoter in Controls and Patients with Schizophrenia

Notes: ^aWe evaluated haplotypic *p*-values with the χ^2 -test. If the nominal *p*-value significantly showed difference (p < 0.05), the precise *p*-value for multiple testing (10,000 permutations) is calculated. The boldface indicates a significant difference.

Abbreviations: MAOA, monoamine oxidase A; CTL, healthy controls; SCZ, schizophrenia; CI, confidence interval; NA, not applicable.

We found that the dVNTR(9R)-uVNTR(3R)-rs6323(T)rs1137070(C) haplotype was associated with schizophrenia in females. Previous studies have reported dVNTR(9R) as a high-expression allele,¹⁹ uVNTR(3R) as a low-expression allele,¹¹ rs1137070(C) as a relatively high-expression allele,²³ and that rs6323(T)-rs1137070(C) haplotype can lead to decreased MAO-A expression, leading to an increased risk of developing schizophrenia.^{14,15,24} These results including those of the current study also appear to be inconsistent. Only considering the function of each allele, and that low-expression alleles increase dopamine, which increases the risk of schizophrenia, is insufficient to clarify the molecular mechanism of schizophrenia.^{7,9,11,13,19} Schizophrenia is a multifactorial genetic disorder, and thus we need to investigate haplotypes with multiple variants and have a different viewpoint.¹⁵ The current study shows that several variants in the *MAOA* locus had a combined effects on the risk of developing schizophrenia. In the similar catabolic enzyme of dopamine, the variants allele affects dopamine levels in a specific region of the brain, which is related

Markers	Global p-value ^a /Odds Ratio (95% Cl)		
	Two Markers	Three Markers	Four Markers
dVNTR (9R)			
	0.0004 (0.0015)/1.924 (1.336-2.770)		
uVNTR (3R)		0.0002 (0.0010)/2.119 (1.417-3.167)	
	0.0006 (0.0034)/1.778 (1.273-2.483)		0.0001 (0.0006)/2.170 (1.446-3.257)
rs6323 (T)		0.0005 (0.0017)/1.808 (1.293-2.530)	
	0.6620/1.044 (0.860–1.267)		
rs1137070 (C)			

Table 5 Haplotype Analysis of the MAOA Promoter in Controls and Female Patients with Schizophrenia

Notes: ^aWe evaluated haplotypic *p*-values with the χ^2 -test. If the nominal *p*-value significantly showed difference (*p* < 0.05), the precise *p*-value for multiple testing (10,000 permutations) is calculated. The boldface indicates a significant difference. **Abbreviations:** MAOA, monoamine oxidase A; CI, confidence interval.

to the risk of developing schizophrenia.⁴ There is a possibility that low levels of MAO-A contribute to hyperfunction in the mesolimbic pathway resulting in positive symptoms of schizophrenia, whereas high levels of MAO-A contribute to hypofunction in the mesocortical pathway cortex leading to negative symptoms of schizophrenia with the combined effects of variant alleles.^{5,7} Therefore, the 9R-3R-T-C haplotype may have different functions for MAO-A expression dependent on brain regions (Figure 1). Further studies focusing on differences in the brain regions are required to determine the effects on MAO-A expression. Indeed, previous studies have reported that uVNTR is associated with the activity of different brain regions, contributing to the developments of psychiatric disorders.^{24,25}

It was previously reported that uVNTR (low-expression alleles)-rs6323(T)-rs1137070(C) haplotype is associated with schizophrenia,^{14,15} in line with our findings. We found that the odds ratio of the 9R-3R-T-C haplotype (corrected p = 0.0006, odds ratio = 2.170) was higher than that of the 3R-T-C-haplotype (corrected p = 0.0017, odds ratio = 1.808) (Table 5), indicating that the haplotypes containing dVNTR (9R) may increase the role of MAO-A in the pathophysiology

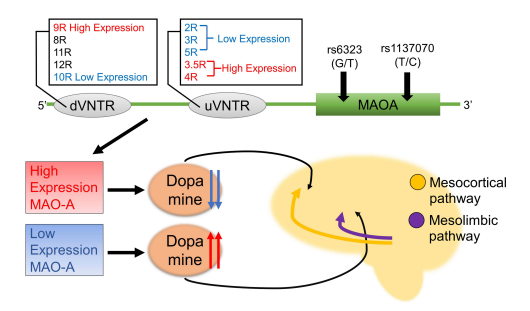


Figure 1 Two MAOA gene promoter VNTRs, and their related SNPs have combined effects on the pathogenesis of schizophrenia. For uVNTR, 3.5R and 4R are highexpression alleles, and the 2R, 3R, and 5R are low-expression alleles. For dVNTR, 9R and 10R are high-expression alleles, 8R and 11R are moderate expression alleles, and 12R is unknown. Low levels of MAO-A contribute to increased dopamine levels in the mesolimbic pathway resulting in positive symptoms, and high levels of MAO-A contribute to decreased dopamine levels in the mesocortical pathway cortex leading to negative symptoms with the combined effects of two VNTRs and two SNPs.

of schizophrenia. It is important to emphasize the combined effects that several variants had on the risk of schizophrenia and that finding new variants may help increase our current knowledge of the molecular mechanism underlying this disease.

Although recent large genome-wide association studies (GWASs) have shown that various SNPs are associated with schizophrenia, 2^{26-28} they did not include the *MAOA* gene polymorphisms that we considered in this study. Therefore, our findings warrant reconsideration of the previous studies while also bearing in mind the effects of VNTRs and haplotypes as well as sex differences.

Our study has several limitations. First, our sample size was relatively small, and our cohort comprised participants of only Japanese descent. Further studies with large samples sizes and different populations are required to validate our findings. Second, we did not consider longitudinal effects or detailed history of the symptoms, such as whether negative or positive symptoms were dominant.

Conclusion

To the best of our knowledge, this is the first study reporting the association of the two *MAOA* gene promoter VNTRs, and their related SNPs with schizophrenia. Our findings show that the dVNTR(9R)-uVNTR(3R)-rs6323(T)rs1137070(C) haplotype was associated with schizophrenia in females, which may increase the risk of developing and help reveal the molecular mechanism of this disease.

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Disclosure

The authors report no conflicts of interest in this work.

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