Functional impact of a single nucleotide polymorphism in the 
OPRD1 promoter region

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Abstract

The δ-opioid receptor mediates rewarding effects of many substances of abuse. We reported an increased frequency of the minor G-allele of single nucleotide polymorphism (SNP) rs569356 [the only variant identified so far in the promoter region of the δ-opioid receptor gene (OPRD1)] in subjects with opioid dependence. In this study, we examined the functional significance of this variant. 

OPRD1 promoter region harboring SNP rs569356 was amplified by polymerase chain reaction (PCR) and inserted into a firefly luciferase reporter vector. HEK293 cells were co-transfected with these constructs and a renilla luciferase vector to control for transfection efficiency. Expression of firefly luciferase (driven by the OPRD1 promoter) was measured by a dual luciferase reporter assay and normalized by renilla luciferase expression. Moreover, alleles altering expression were further assessed for binding of human brain nuclear proteins by electrophoretic mobility shift assay (EMSA). The minor G-allele was associated with significantly greater expression levels of firefly luciferase than the major A-allele of SNP rs569356 (P=0.003). EMSA also showed specific gel shift bands, suggesting that SNP rs569356 is situated in the
binding site of potential transcription factors. These results suggest that the minor G-allele of SNP rs569356 may enhance transcription factor binding and increase OPRD1 expression.

**Keywords**
dual luciferase reporter gene assay; electrophoretic mobility shift assay; δ-opioid receptor gene; polymerase chain reaction; promoter polymorphism

**Introduction**

Opioid receptors (mainly μ-, δ- and κ-receptors) in the endogenous opioid system regulate neuronal activity via neurotransmission or neuromodulation.1 There is mounting evidence that opioid receptors directly mediate reward, tolerance, and dependence associated with opioids such as morphine and heroin.2,3 Opioid receptors are also indirectly involved in the reinforcing properties of non-opioid drugs (such as cocaine and alcohol) due to the intimate relationship between the opioid system and the mesolimbic dopamine system.4,5 The basal release of dopamine in the nucleus accumbens is modulated by the stimulatory μ- and δ-opioid systems and the inhibitory κ-opioid system.4

Variation in μ-, δ- and κ-opioid receptor genes may change the expression level or activity of opioid receptors, leading to increased risk for drug or alcohol dependence. Take the δ-opioid receptor gene as an example. Animal studies have demonstrated that the δ-opioid receptor is involved in addictive processes. When the mouse δ-opioid receptor gene is knocked out (by deletion of exon 2), morphine tolerance in mice is lost as well.6,7 The human δ-opioid receptor gene (OPRD1, located at 1p36.1-p34.3) is the first of the human opioid receptor genes to be cloned.8 It has a relatively conserved coding sequence. Only two coding sequence polymorphisms, a non-synonymous single nucleotide polymorphism (SNP) C80T (Cys27Phe or rs1042114) in exon 1 and a silent SNP T921C (Gly307Gly or rs2234918) in exon 3, have been identified.9,10 A study by Mayer et al.9 demonstrated a significant association of SNP T921C and heroin dependence in a German population. However, follow-up studies in another German population11 and two Han Chinese Populations12,13 did not support the finding. Our recent study14 also did not show evidence of association between SNP T921C and heroin dependence in European Americans (EAs). Nevertheless, a highly significant association of the non-synonymous SNP C80T and opioid dependence was observed. Additionally, the promoter SNP rs569356 (A/G) also showed a statistically significant association with opioid dependence by comparing both allele and genotype frequencies of the marker in cases and controls. The minor G-allele of SNP rs569356 was enriched in opioid dependent EA cases compared with EA controls.

The promoter SNP rs569356 is 1,968 bp upstream to the OPRD1 transcription start site and is the only documented variant identified in the promoter region of OPRD1 [refer to the NCBI SNP database http://www.ncbi.nlm.nih.gov/projects/SNP or the ABI SNPbrowser15]. Polymorphisms located in promoter regions are presumed to affect gene transcription by modifying the coordinated action of multiple regulatory proteins through complex protein-DNA and protein-protein interactions.16 Functional genetic variation in promoter region may alter the affinity of transcription factors and other modulatory proteins to bind to the
DNA sequence and thus influence the specificity and kinetics of the transcription process. Although the promoter SNP rs569356 is 2,289 bp apart from the non-synonymous SNP C80T, these two markers are in tight linkage disequilibrium (LD) ($r^2=0.833, D'=0.95$). Therefore, it is necessary to clarify whether SNP rs569356 is truly an independent disease locus for opioid dependence or its positive association results were due to co-inheritance of the non-synonymous SNP C80T.

The purpose of this study is to examine whether SNP rs569356 modulates OPRD1 transcription by changing the affinity of the promoter for transcription factors. We applied both the luciferase reporter gene assay and DNA-protein binding assay or electrophoretic mobility shift assay (EMSA) to examine the functional implication of OPRD1 promoter SNP rs569356.

**Materials and methods**

**DNA samples**

Genomic DNAs, extracted from peripheral blood cells of EA subjects, were included in our previous study regarding the association of OPRD1 variants and drug or alcohol dependence. Subjects were recruited at the University of Connecticut Health Center and the VA Connecticut Healthcare System-West Haven campus, where the study protocol was approved by the respective institutional review boards. All subjects gave written informed consent prior to participating in the study.

**Amplification of OPRD1 promoter region containing SNP rs569356**

A pair of primers was designed to amplify 2,250 bp of OPRD1 promoter region containing SNP rs569356 by polymerase chain reaction (PCR). The forward primer (Primer F1: 5’-TGTGTGCCACCGTGCCCAGCCTTTT-3’) was homologous to OPRD1 upstream sequence (−2,134 bp to −2,109 bp) and the reverse primer (Primer R1: 5’-GCCCGGCTGTCTCTGGCGCTGCT-3’) was complementary to part of the sequence (+97 bp to +120 bp) of OPRD1 exon 1. The PCR mixture contained 50 ng of genomic DNA, 2.5 mM of each dNTPs, 200 nM of each primer, 2.5 µl of dimethyl sulfoxide (DMSO) (Invitrogen, Carlsbad, CA, USA), 1x Pfu Ultra II reaction buffer (Stratagene, La Jolla, CA, USA), and 2.5 units of Pfu Ultra II fusion HS DNA polymerase (Stratagene, La Jolla, CA, USA) in a final volume of 50 µl. PCRs were run on a PTC-200 thermocycler (MJ Research, Waltham, MA, USA) and consisted of an initial denaturation step of 95°C for 2 min, followed by 39 cycles of a two-step reaction (95°C for 30 sec and 72°C for 2 min 30 sec).

**Construction of luciferase expression vectors**

PCR products were subcloned into pSMART HC Amp plasmid vector (1,833 bp) (Lucigen, Middleton, WI, USA). The pSMART plasmid is a transcription-free vector optimized for large or otherwise difficult to clone DNA fragments. Briefly, the gel purified PCR products were 5’ end phosphorylated by polynucleotide kinase (PNK) (New England Biolabs, Ipswich, MA, USA) and ligated into pSMART vector using T4 DNA ligase (New England Biolabs, Ipswich, MA, USA) according to the protocol of the PCR-SMART Cloning Kit (Lucigen, Middleton, WI, USA). The ligation mixture was transformed into DH5α.
competent *E. coli* cells (Invitrogen, Carlsbad, CA, USA) by heat-shock transformation. Clones were screened by restriction enzyme digestion using EcoRV (a pSMART polycloning site flanking the inserted DNA sequence) and Sca I [which can be used to differentiate the A-allele (uncut) and the G-allele (cut) of SNP rs569356]. Moreover, cloned *OPRD1* promoter fragments were sequenced from both ends using two forward primers and two reverse primers. The forward primer SL1 (5′-CAG TCC AGT TAC GCT GGA GTC-3′) and the reverse primer SR2 (5′-GGT CAG GTA TGA TTT AAA TGG TCA GT-3′) were included in the PCR-SMART Cloning Kit. An additional forward sequencing primer (Primer F2: 5′-TGGAATGGTTAGGCTTGTGGA-3′, homologous to *OPRD1* upstream sequence from −935 bp to −915 bp) and an additional reverse sequencing primer (Primer R2: 5′-GGAAGAGGATTCCGGATAAA-3′, complementary to *OPRD1* upstream sequence from −751 bp to −731 bp) were synthesized. The location of the four primers and *OPRD1* SNP rs569356 is depicted in Figure 1. Clones with the A-allele or the G-allele with otherwise identical sequences, and identical to the GenBank reference sequence (NT_004610.18) for *OPRD1*, were selected for construction of luciferase expression vectors.

The *OPRD1* promoter fragment was excised from the pSMART clones using restriction enzyme EcoRV. Then it was phosphorylated and inserted into the SmaI cloning site in the promoterless and enhancerless pGL3-basic luciferase reporter vector (Promega, Madison, WI, USA) by blunt-end ligation. The *OPRD1* promoter fragment was placed upstream of the firefly luciferase reporter gene. DNA was transformed into DH5α competent *E. coli* cells (Invitrogen, Carlsbad, CA, USA). The *OPRD1* promoter sequence and cloning orientation in pGL3-basic vector were determined via restriction analysis and direct sequencing using primers F1, F2, R1 and R2 described above.

**Transient transfection and dual-luciferase reporter assay**

HEK293 cells (purchased from the American Type Culture Collection or ATCC) were grown in 48-well cell culture plates (Becton Dickinson, New Jersey, USA; 3.5 × 10⁵ cells/well) for 24 hours. They were then transiently transfected with a promoterless luciferase vector (pGL3-basic) (Promega, Madison, WI, USA) or with a pGL3-basic vector with the *OPRD1* promoter sequence harboring the rs569356 A-allele or G-allele (construct rs569356_A-allele or construct rs569356_G-allele) using linear polyethylenimine (MW 25,000) (Polysciences, Warrington, PA, USA) based on the method of Reed et al.21 An internal control vector pTK-RL, which contained the renilla luciferase gene downstream of a minimal HSV-TK promoter (Promega, Madison, WI, USA), was included in the transfection mix (375 ng DNA in total, firefly construct : renilla construct = 10 : 1). Cells were harvested 48 hours after transfection and luciferase assays were performed with the Dual Luciferase Assay Kit (Promega, Madison, WI, USA) according to the manufacturer’s instructions. Cell lysate (10 µl) were evaluated first for firefly luciferase activity and then for renilla luciferase activity in a white opaque 96-well flat-bottom microplate (Becton Dickinson, New Jersey, USA) using a DTX 880 Series Multimode Detector (Beckman Coulter, Fullerton, California, USA). Firefly luciferase luminescence values were divided by renilla luciferase luminescence values from the same transfection to control for differences in transfection.
efficiency. For each transfection mix (pGL3+pTK-RL), the reporter assay was conducted four times in parallel.

**Electrophoretic mobility shift assay (EMSA) study**

EMSA studies were conducted to examine whether SNP rs569356 (A/G) influences the binding of human brain nuclear extract proteins (potentially containing OPRD1 transcription factors) to OPRD1 promoter sequence. Two double-stranded DNA oligonucleotides (21-bp) spanning OPRD1 promoter SNP rs569356 were synthesized. One contained rs569356 A-allele (5'-GTTTTACACAAATACTTTTTTG-3') and the other contained rs569356 G-allele (5'-GTTTTACACAGTACTTTTTTG-3'). EMSA was performed based on the instructions accompanying the DIG Gel Shift Kit, 2nd Generation (Roche, Mannheim, Germany). In brief, probes were first labeled with Digoxigenin-11-ddUTP (DIG-ddUTP) at the 3' end. 2 µl of labeled probes (4 ng/ul) was then incubated with 6 µl of human brain nuclear protein (5 µg/µl) (Active Motif, Carlsbad, CA, USA). To prevent nonspecific binding of brain nuclear proteins to the probe, nonspecific competitors poly[d(I-C)] (1 µg) and poly-lysin (0.1 µg) were added to the binding reaction (40 µl of volume in total). To establish specificity, 4 µl of unlabeled probes (100 ng/µl) was added in the probe (labeled)-protein mixture as competitors. Following electrophoresis and gel-blotting, the DIG labeled probes were visualized by an enzyme immunoassay using an anti-Digoxigenin-alkaline phosphatase conjugate and the chemiluminescent substrate CSPD. An X-ray film was exposed to fluorescence for 15–25 min at room temperature and then developed. The image on the X-ray film was scanned at a resolution of 400 dpi. The density of the gel shift bands was quantified using software NIH ImageJ.22 The EMSA experiment was replicated once.

**Bioinformatics analysis**

We queried the OPRD1 promoter sequence harboring SNP rs569356 against the transcription factor database TRANSFAC23 using the software MatInspector24 to determine if the promoter SNP rs569356 is located in the binding site of potential transcription factors.

**Statistical analysis**

Mean luciferase activity corresponding to each construct was compared using the one-way analysis of variance (ANOVA) in the SPSS 16.0 software package (SPSS Inc., Chicago, Illinois), and a one-step multiple comparison was performed using the Tukey Honestly Significant Difference (HSD) post-hoc test.

**Results**

**SNP rs569356 A-allele and G-allele differentially regulate OPRD1 promoter activity**

Because the OPRD1 promoter region was subcloned into the pGL3 expression vector upstream to the firefly luciferase reporter gene using a blunt-end ligation, four possible expression vectors were obtained. As shown in Figure 2, two constructs (one with the A-allele and another with the G-allele) had the forward cloning orientation and another two constructs (one with the A-allele and another with the G-allele) had the reverse cloning orientation.
Before HEK293 cells were used for luciferase reporter gene assay, we verified that OPRD1 was expressed in these cells at a moderate level using the Affymetrix expression array study (Simen and Mane, 2009, unpublished observations). Thus, HEK293 cells express the necessary proteins to maintain OPRD1 activity. For each of the five expression vectors (Constructs 1–4 and pGL3-basic, used as a negative control, see Fig. 2), four independent reporter assays were conducted in replication. The raw data were presented in Supplementary material Table S1. To adjust for variation in transfection efficiency and cell density, the firefly fluorescence reading was normalized by the renilla fluorescence reading (as a ratio of firefly to renilla) (Fig. 3). A one-way ANOVA revealed a main effect of constructs (pGL3-basic vector vs. A-allele_forward vector vs. G-allele_forward vector: $F_{(2,11)}=179.37, P<0.001$). Both the A-allele_forward and the G-allele_forward constructs (with OPRD1 promoter sequence in the correct orientation) showed considerably higher luciferase activity (A-allele_forward vector vs. pGL3-basic vector: $P<0.001$; G-allele forward vector vs. pGL3-basic vector: $P<0.001$) compared to the promoterless pGL3-basic vector (by the Tukey HSD test). These results support the conclusion that the 2,250 bp region upstream of OPRD1 that we cloned possesses strong promoter activity. When the OPRD1 promoter sequence was placed in the opposite orientation, firefly luciferase activity was suppressed (Fig. 3). In the forward orientation, the G-allele construct was associated with significantly higher luciferase activity than the A-allele construct ($P=0.003$) (by the Tukey HSD test), suggesting that the A to G polymorphism at position −1,968 bp of the OPRD1 regulatory region increases the activity of the OPRD1 promoter.

**Binding of brain nuclear proteins to OPRD1 promoter sequence harboring rs569356**

Because the G-allele construct was associated with significantly higher luciferase activity, we hypothesized that the G-allele might bind to certain activating transcription factors more effectively than the A-allele. Double-stranded DNA probes encoding the OPRD1 rs569356 A-allele or G-allele were labeled with digoxigenin and mixed with a human brain nuclear extract in binding buffer. As shown in Figure 4A, human brain nuclear proteins retarded the migration of probes for both the G-allele (lane 5) and the A-allele (lane 8). The interaction could be inhibited by competition with unlabeled probe for the G-allele (lane 6) or the A-allele (lane 9). Moreover, both the G-allele probe and the A-allele probe showed two gel shift bands of similar size. Since the two gel shift bands for the G-allele probe or the A-allele probe were very close, their density was measured together. As shown in Figure 4B, the density of the gel-shift band for the G-allele probe is 1.61 times higher than that of the A-allele probe. Additionally, a probe for Oct2a and purified Oct2a protein (Fig. 4A, Lanes 1–3) were included as a positive control to test the quality of the reagents. The EMSA replication results were shown in Supplementary material Figure S1 and Table S2. The mean ratio ($±$S.D.) of the G-allele gel shift band density to the A-allele gel shift band density was 1.64 ($±$0.04).

**Bioinformatics analysis results**

The A-allele of SNP 569356 was in the binding site of two transcription factors [Fork head homologous X binds DNA with a dual sequence specificity (FHXA and FHXB): tttacACAAtacttt; SWI/SNF related, matrix associated, actin dependent regulator of chromatin, subfamily a, member 3 (Smarca3): cagACTTttt)]. The G-allele of SNP rs569356
was in the binding site of three transcription factors [NK3 homeobox 1 (NKX3-1): ttatacaGTACttttg; zinc finger proliferation 1 (Zipro 1): cAGTAcat; Smarca3: cagtACTTttt] (The underlining shows the allele (A or G) and capital letters represent the core binding site).

Discussion

We used both the luciferase reporter gene assay and EMSA to examine the function of OPRD1 promoter SNP rs569356. Our findings indicate that SNP rs569356 (A/G) affects the activity of the OPRD1 promoter. Luciferase reporter assays demonstrated that the 2,250 bp fragment upstream of OPRD1 possesses strong promoter activity. OPRD1 promoter region with either rs569356 A-allele or G-allele can drive reporter gene expression at a substantial level (Fig. 3). Moreover, significantly higher luciferase activity was found for the rs569356 G-allele in comparison to the A-allele. This finding suggests that SNP rs569356 is functional and the substitution of a G for an A can enhance OPRD1 promoter activity, possibly by enhancing transcription factor binding.

Although we are unable to determine whether OPRD1 promoter activity observed in HEK293 cells (derived from human embryonic kidney cells) reflects promoter activity in neural cells, microarray results demonstrated that OPRD1 is expressed in HEK293 cells at a moderate level. Thus, transcription factors required for OPRD1 transcription are expressed in HEK293 cells and the observed effects of rs569356 likely reflect OPRD1 promoter activity in neuronal cells. In this study, we only tested OPRD1 promoter activity under basal unstimulated conditions. It is possible that additional effects of rs569356 are evident when promoter activity is stimulated, for example, under conditions of elevated protein kinase A activity.

The results from our EMSA assays, which were conducted with human brain nuclear proteins, paralleled those observed from our promoter assays. EMSA studies revealed that human brain nuclear proteins interact with probes containing OPRD1 promoter sequence with either the A-allele or the G-allele of SNP rs569356. Both the A-allele and the G-allele probes showed two gel-shift bands, but the binding of the G-allele probe to nuclear proteins was 1.64±0.04 (mean±S.D.) times that of the A-allele probe (Fig. 4, Fig. S1 and Table S2). The enhanced DNA-protein binding may be responsible for the increased OPRD1 promoter activity that we observed in our promoter assays. Consistent with the observation that SNP rs569356 only slightly modifies the binding of transcription factors to OPRD1 promoter sequence, both the A-allele construct and the G-allele construct showed high expression levels of the reporter gene. However, differential expression levels associated with these two alleles were observed as well (Fig. 3).

Bioinformatics analysis revealed that SNP rs569356 may affect the binding of a number of transcription factors including FHXA and FHXB, Smarca3, NKX3-1, and Zipro 1. In this study, we did not use super-gel shift assays to verify these interactions because antibodies specific for them are unavailable. Future studies could utilize immunoprecipitation plus massively parallel DNA sequencing (ChIP-Seq) to test whether these transcription factors are involved in the formation of transcriptional complexes at the rs569356 site.
Functional characterization of gene variants is important for interpreting the genetic mechanism of diseases. SNPs have been widely used in candidate gene and genome-wide association studies to understand the genetic basis of complex disorders. However, the biological relevance from SNP analyses is not easily interpretable. Thus, it is critical to study the function of disease-associated genetic variants. Previous studies have demonstrated that opioid receptor availability is positively correlated with levels of craving for abused substances. These studies have also shown significant regional increases in opioid receptor availability in early abstinence from dependence on cocaine, alcohol, or opioids when compared with controls. Thus, elevated OPRD1 expression levels may increase the rewarding effect of drugs of abuse, leading to increased vulnerability to substance dependence (including opioid dependence). The finding from the present study that the minor G-allele of SNP rs569356 increases OPRD1 promoter activity supports our previous genetic association results that the minor G-allele is a genetic risk factor for opioid dependence.

Our findings may have important implications for the pharmacotherapy of substance dependence. Lowering the expression, activity, or occupancy of the δ-opioid receptor in patients with substance dependence may facilitate recovery from dependence. In fact, nonselective opioid receptor antagonists such as naltrexone (with a high affinity for the μ- and κ-opioid receptors but a low affinity for the δ-opioid receptor) and naloxone (with an extremely high affinity for the μ-opioid receptors but a low affinity for the δ- and κ-opioid receptors) have been used in the management of alcohol and/or opioid dependence. It is possible that δ-opioid receptor antagonists or drugs that down-regulate δ-opioid receptor expression will show efficacy in the treatment of substance dependence. Animal studies have provided some support for this hypothesis. Nielsen et al. found that a novel δ-opioid receptor antagonist SoRI-9409, a derivative of naltrexone and with a high affinity for the δ-opioid receptor, produced a selective and long-lasting decrease in ethanol consumption in heavy drinking rats. A recent study also demonstrated that the δ-opioid receptor antagonists naltrindole and naltriben prevented the enhanced response to the conditioned rewarding effects of morphine in rats. Given that SNP rs569356 influences OPRD1 promoter activity and possibly OPRD1 expression levels, future pharmacogenetic studies need to take into consideration its genetic effect on the response to treatment of substance dependence with δ-opioid receptor antagonists.

Promoter activity may be regulated by several independent or correlated variants. For example, Zhao et al. studied the association of variants in the promoter region of the Chitinase 3-like 1 gene (CHI3L1) and schizophrenia. They found that an at-risk haplotype showed low promoter activity but a protective haplotype showed high promoter activity, and these haplotypes consisted of alleles of three correlated loci in the promoter region. Although SNP rs569356 is the only common variant identified so far, we cannot exclude the possibility that rare variants in the OPRD1 promoter region may exert an even larger effect on OPRD1 transcription than SNP rs569356. Therefore, deep-sequencing of OPRD1 promoter for rare variants is necessary.

Taken together, this study demonstrated that SNP rs569356 is a functional variant which alters OPRD1 promoter activity. Although OPRD1 promoter SNP rs569356 and OPRD1
non-synonymous SNP C80T (in exon 1) are highly linked, they are likely to be two independent genetic variants which influence the vulnerability to opioid dependence through two different mechanisms. The former one may affect promoter activity and thus compromise OPRD1 transcription (as shown in this study). The latter one may directly change δ-opioid receptor activity by changing the amino acid sequence of the receptor protein. We are now in the process of investigating the function of the non-synonymous variant C80T by receptor binding assays. Additionally, future pharmacotherapy of opioid dependence must consider the effect of these two OPRD1 variants on the outcome of treatment.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1. Schematic representation of the δ-opioid receptor gene (*OPRD1*)
Primers F1, F2, R2, and R1 are complementary to *OPRD1* sequence at −2,134 bp to −2,109 bp, −935 bp to −915 bp, −751 bp to −731 bp, and +97 bp to +120 bp, respectively. SNP rs569356 (A/G) is located at −1,968 bp in the promoter region of *OPRD1*. 
Fig. 2. Construction of four pGL3 expression vectors

OPRD1 promoter fragment containing SNP rs569356 A-allele or G-allele was excised from a pSMART plasmid vector and subcloned into a pGL3 expression vector by blunt-end ligation. OPRD1 promoter sequence was placed upstream to the firefly luciferase reporter gene (luc+).
Fig. 3. Luciferase reporter gene assay of the effect of SNP rs569356 (A/G) on OPRD1 promoter activity
Four constructed vectors (A-allele_reverse, A-allele_forward, G-allele_reverse, and G-allele_forward) and the pGL3-basic vector were co-transfected respectively with pTK-RL (internal control vector harboring renilla luciferase gene) into HEK293 cells. Cell lysate was used to measure sequentially the activity of firefly luciferase and renilla luciferase. Firefly luminescence was normalized by renilla luminescence. Four independent replications of the experiment were performed. The difference in firefly luciferase activity between different vectors was analyzed by one-way analysis of variance (one-way ANOVA) and the Tukey HSD test. Y axis: ratios of firefly fluorescence to renilla fluorescence; X axis: five expression vectors.
Fig. 4. Electrophoretic mobility shift assay (EMSA)

A. The binding of human brain nuclear proteins (potentially containing transcription factors) with OPRD1 promoter sequence (harboring either the rs569356 A-allele or G-allele) was determined by the EMSA. Lanes 1–3: control probe and control factor Oct2A included in the DIG Gel Shift Kit (Roche, Mannheim, Germany). Lanes 4–6: OPRD1 rs569356 G-allele probe. Lanes 7–9: OPRD1 rs569356 A-allele probe. Lanes 1, 4, and 7: in the absence of brain nuclear extract, DIG-labeled probes showed no retardation; Lanes 2, 5 and 8: in the presence of brain nuclear extract, DIG-labeled probes showed retardation, and gel shift bands due to the slow migration of protein-probe complex were observed; Lane 3, 6, and 9: in the presence of brain nuclear extract and unlabeled probes (as competitors), DIG-labeled probes showed no retardation. B. The intensity of gel shift bands derived from the control
probe (lane 2), the G-allele probe (lane 5) and the A-allele probe (lane 8) was scanned at 400 dpi and quantified by software ImageJ22.