Article

The Role of the Major Histocompatibility Complex Region in Cognition and Brain Structure: A Schizophrenia GWAS Follow-Up

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Objective: The authors investigated the effects of recently identified genome-wide significant schizophrenia genetic risk variants on cognition and brain structure.

Method: A panel of six single-nucleotide polymorphisms (SNPs) was selected to represent genome-wide significant loci from three recent genome-wide association studies (GWAS) for schizophrenia and was tested for association with cognitive measures in 346 patients with schizophrenia and 2,342 healthy comparison subjects. Nominally significant results were evaluated for replication in an independent case-control sample. For SNPs showing evidence of association with cognition, associations with brain structural volumes were investigated in a large independent healthy comparison sample. Results: Five of the six SNPs showed no significant association with any cognitive measure. One marker in the major histocompatibility complex (MHC) region, rs6904071, showed independent, replicated evidence of association with delayed episodic memory and was significant when both samples were combined. In the combined sample of up to 3,100 individuals, this SNP was associated with widespread effects across cognitive domains, although these additional associations were no longer significant after adjusting for delayed episodic memory. In the large independent structural imaging sample, the same SNP was also associated with decreased hippocampal volume.

Conclusions: The authors identified a SNP in the MHC region that was associated with cognitive performance in patients with schizophrenia and healthy comparison subjects. This SNP, rs6904071, showed a replicated association with episodic memory and hippocampal volume. These findings implicate the MHC region in hippocampal structure and functioning, consistent with the role of MHC proteins in synaptic development and function. Follow-up of these results has the potential to provide insights into the pathophysiology of schizophrenia and cognition.

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In recent years, the advent of genome-wide association studies (GWAS) has advanced the field of complex disease genetics, including that of major mental illnesses. The first truly large-scale GWAS in schizophrenia were published in 2009 by the International Schizophrenia Consortium (1), the Molecular Genetics of Schizophrenia study (2), and SGENE-plus (3), which also obtained additional data resulting in a sample of up to 50,000 individuals (case subjects and healthy comparison subjects). Taken together, three regions of association were identified at the genomewide significant threshold: 1) a region on chromosome 6p22.1 that includes the major histocompatibility complex (MHC); 2) the transcription factor 4 (*TCF4*) locus on chromosome 18; and 3) the neurogranin (*NRGN*) locus on

chromosome 11. Although the association signals formally implicate genomic regions rather than specific genes, the roles of *NRGN* and *TCF4* in brain function and development (3–5) are compatible with the hypothesis that increased risk for schizophrenia is related to altered function of these genes. However, the association at 6p22.1 is within a region of exceptionally high linkage disequilibrium (LD) spanning hundreds of genes, any of which could contain the functional susceptibility variant or variants. The identification of causative variants is made challenging by such high LD, although investigators using conditional analysis have observed at least two statistically independent associations in the MHC region (3).

This article is discussed in an Editorial by McGuffin and Power (p. 821)

The identification of associated SNPs is only a first step toward elucidating the neurobiological pathways involved in schizophrenia. Attaining this goal requires the functional consequences of polymorphisms to be identified at the level of the protein, cell, neural circuit, or alternative human phenotypes. Studies of neurocognition and structural brain imaging offer a way of exploring the action of polymorphisms on behavioral neural circuits at the whole organism level (6). Widespread cognitive deficits are characteristic features of schizophrenia, and their amelioration is increasingly seen as central to improving the functional outcome of patients (7). In this study, we set out 1) to investigate the effects of GWAS-supported variants on cognitive phenotypes in schizophrenia patients and healthy comparison subjects and replicate any associations, and 2) to investigate the relationships between variants with cognitive effects and relevant brain structure volumes using a sample that was independent from the cognitive data collection sample. We previously reported results for rs12807809 (NRGN locus) in a subsample of the individuals included in this study (8). It should be noted that the design of this study predates the publication by the Psychiatric Genomewide Association Study Consortium of additional genome-wide significant findings in schizophrenia (9).

Method

Cognition Sample Characteristics

German sample. The German sample consisted of 346 patients with a DSM-IV diagnosis of schizophrenia and 2,342 healthy comparison subjects. Two hundred sixteen of the patients (62%) and 545 of the comparison subjects (23%) formed part of the SGENE-plus study (3). The patients were ascertained from mental health services in the Munich area. Exclusion criteria were a history of head injury or neurological disease. Patients, ages 18–80 years, were interviewed with the Structured Clinical Interview for DSM-IV (SCID) (10), and the interviews were rated by psychiatrists or psychologists.

Healthy comparison subjects of German descent, ages 18–80 years, were randomly selected from population registers from the Munich area. Participants underwent an extensive screening process, described previously (11), to exclude those with neurological or psychotic disorders and those who had first-degree relatives with psychotic disorders. In the case of participants older than 60 years, the Mini-Mental State Examination (12) was employed to exclude individuals with possible cognitive impairment.

Irish replication sample. The Irish sample consisted of 377 patients with schizophrenia and 145 comparison subjects. One hundred three patients (but no comparison subjects) formed part of the International Schizophrenia Consortium study (13). Patients with schizophrenia, ages 18–65 years, were recruited from across Ireland, and the diagnosis was confirmed by trained psychiatrists using the SCID (10). Comparison subjects, ages 18–65 years, were recruited by local media advertisements. Exclusion criteria were the same as used in the German sample.

All patients with schizophrenia and comparison subjects were unrelated white Caucasians of German or Irish ancestry. All participants provided informed consent in accordance with the relevant ethics approvals.

Cognitive Assessment

The study was designed by selecting tests to represent domains of cognition that are known to be compromised in schizophrenia: IQ, episodic memory, working memory, and attention/vigilance. Given that we intended to seek replication of significant results, we selected identical or near identical tests within cognitive domains for the German and Irish samples. The number of tests and measures within each cognitive domain was limited to minimize multiple testing, mirroring our previous research (11). The German sample was used as a discovery sample, and markerphenotype combinations that were nominally significant were taken forward for replication in the Irish sample.

For the German sample, IQ was measured by the full German Wechsler Adult Intelligence Scale–Revised (WAIS-R) (14). Episodic memory was assessed using the immediate and delayed logical memory tests from the German Wechsler Memory Scale–Revised (WMS-R) (15). Verbal and spatial working memory were assessed using the digit span test from the WAIS-R and the spatial span test from the WMS-R. Attention/vigilance was assessed using the Continuous Performance Test, 3–7 Version (16). Test results were available in up to 342 patients and 2,342 comparison subjects.

For the Irish sample, IQ was measured using selected subtests (vocabulary, similarities, block design, and matrix reasoning) from the Wechsler Adult Intelligence Scale, 3rd edition (WAIS-III) (17). We measured episodic memory using the immediate and delayed logical memory tests from the Wechsler Memory Scale, 3rd edition (WMS-III) (18). Working memory was measured using the letter-number sequencing test from the WMS-III and the spatial working memory test from the Cambridge Neuropsychological Test Automated Battery (19). Attention/vigilance was measured using the three-letter condition of the Continuous Performance Test, Identical Pairs Version (20), but an insufficient number of comparison subjects completed this task, making it unavailable for this group. Test results were available for up to 377 patients and 148 comparison subjects. These procedures and the selected tests reproduce our previous research (11).

Selection of SNPs. We identified a nonredundant panel of SNPs representing all genome-wide significant signals from the combined studies of the International Schizophrenia Consortium (13), Molecular Genetics of Schizophrenia (2), and SGENE-plus (3). Among the multiple correlated signals at chromosome 6, we selected the most significantly associated variant from the International Schizophrenia Consortium and Molecular Genetics of Schizophrenia studies, rs13194053. In the German sample, we typed rs6904071, a perfect proxy for rs13194053 according to HapMap. In the SGENE-plus study, rs6932590 was the most significantly associated variant and was included in our panel. Also from that study, analysis conditioned on rs6932590 further revealed a strong, statistically independent signal at rs3131296 and, independent of both of these markers, a weak association signal at rs13219354; both of these SNPs were included in our analysis. We also selected the genome-wide significant SNPs at NRGN (rs12807809) and TCF4 (rs9960767) as reported by SGENE-plus.

We therefore analyzed the associations with cognition for a panel of six SNPs in total. The four individual MHC SNPs were not in high LD (1,000 genomes, $r^2 < 0.8$) and covered the statistically independent schizophrenia association signals that were identified in this region.

Genotyping

The SNPs were genotyped in the German sample and in a proportion of the Irish sample using the Sequenom iPLEX Gold system. The call rate for this genotyping was >95% in both samples. The remainder of the Irish sample was genotyped using a Taqman SNP Genotyping Assay on a 7900HT Sequence Detection System (Applied Biosystems). The call rate for the Taqman genotyping was >95%. For all genotyping, both patient and comparison samples were in Hardy-Weinberg equilibrium for all SNPs (p>0.05).

Statistical Analysis

The association between markers and cognitive phenotypes was tested using linear regression, assuming an additive model, using SPSS, version 14 (21). Genotypes and patient/comparison status were entered as independent variables, and age and gender were entered as covariates when appropriate (i.e., unless scaled test scores were used). Linear regression was also performed in patient and comparison samples separately in order to examine differential effects between case and comparison subjects. Markers with nominally significant associations ($p \le 0.05$) in the German sample were taken forward for replication in the Irish samples. For significant markers, we also performed a casecontrol logistic regression analysis within the German discovery and Irish replication cohorts.

In addition to seeking replication, we combined the German and Irish samples to examine the association between SNPs and combined cognitive domain test scores in as large a sample as possible. We derived standardized z scores for each of the cognitive tests separately in each sample. These scores were then combined for equivalent tests across the two samples to give study-wide cognitive domain z scores. Linear regression was used in the combined sample using z scores as the dependent variable and genotype as the independent variable, adjusting for case status, age, and gender as appropriate.

The power of this design varies depending on the selected polymorphism and cognitive test. However, as an example, the German comparison sample has 80% power to detect a risk allele accounting for 0.34% of variance in IQ at the 5% significance threshold.

Structural Imaging Analysis

For those SNPs demonstrating a replicated association with cognition, we examined associations with cortical structure volumes that were derived from structural MRI. The imaging sample was part of the Brain Imaging Genetics project; a full description of the sample and procedures for genotyping and neuroimaging is provided in detail elsewhere (22). Briefly, 892 white Caucasian healthy young adults (369 men and 523 women; ages 18-35 years) underwent structural MRI and provided saliva samples that were genotyped using the Affymetrix GeneChip SNP, 6.0 array; genotypes were imputed using the MaCH software package (23). MRI acquisition was on 1.5-T (N=416) and 3-T (N=476) scanners, and segmenting into gray and white matter was performed with the VBM 5.1 toolbox, version 1.19, with the SPM software package (22). Informed by the neurocognitive tests and available imaging structural phenotypes, we chose to analyze total gray matter volume and total hippocampal volume. Gray matter volume was chosen because several of the cognitive tests involve generalized cognitive functioning or are reliant on multiple, rather than specific, brain regions. Hippocampal volume was chosen as a brain structure that has specific relevance to aspects of cognition tested, particularly episodic and spatial memory. Prefrontal structural volumes would have been of interest given our cognitive battery but were unavailable for this analysis. Using linear regression, SNPs that were significantly associated with cognition were assessed for association with these brain structural volumes. The analyses were adjusted for field strength, age, and gender; the analysis of hippocampal

volume was corrected for total brain volume, and the analysis of gray matter volume was corrected for total white matter volume.

Results

Association With Cognition

German sample. The results from the German data set are summarized in Table 1. As expected, patients with schizophrenia performed significantly below comparison subjects on all administered cognitive tests. Of the six SNPs examined, only one (rs6904071; equivalent by $r^2=1$ to rs13194053, the top hit in the International Schizophrenia Consortium and Molecular Genetics of Schizophrenia studies) showed a nominally significant association with any of the cognitive tests. For the nonsignificant SNPs, genotype mean scores for the cognitive tests of IQ, working memory, episodic memory, and attention/vigilance are presented in Tables S1–S5 in the data supplement that accompanies the online edition of this article.

The detailed results for the significantly associated marker, rs6904071, in the German sample are presented in Table 2 (and as part of Figure 1). Nominally significant associations were found with IQ (B=1.060, p=0.046), spatial working memory (B=0.448, p=0.049), delayed episodic memory (B=1.488, p=0.011), and attention (B=0.15, p=0.001). Notably, for all cognitive tests, individuals homozygous for the schizophrenia risk G allele performed more poorly than heterozygotes, who in turn performed more poorly than nonrisk A homozygotes.

Irish replication sample. The SNP rs6904071, which showed significant associations with cognition in the German sample, was taken forward for replication (Table 3). Assuming a conservative one-tailed alpha of 0.0125 (0.05/4 [4=number of tests that were significant in the German sample]), the delayed episodic memory test displayed significant replication (WMS delayed logical memory; see Tables 2 and 3 and Figure 1). This association was in the same allelic direction as in the German sample, that is, the schizophrenia risk G allele was associated with worse performance in delayed episodic memory.

The detailed results of the separate patient and comparison group analyses in both the German and Irish samples are provided in Table S6 in the online data supplement.

We also examined whether this SNP, rs6904071, was associated with schizophrenia in the two samples. The SNP was nominally associated with schizophrenia in the Irish sample (odds ratio=1.49, p=0.023) but was not significantly associated with schizophrenia in the German sample (odds ratio=1.11, p=0.33). Notably, the allelic direction of association was the same in both samples and mirrored that in the original GWAS studies (although there was sample overlap as outlined above).

Combined sample. Table 4 summarizes the results for the combined analysis. This analysis confirms an association

	General C Abil	ognitive ity	Verbal W Mem	/orking ory	Spatial Working I Memory		Immediate Mem	e Episodic Iory	Delayed E Mem	Episodic ory	Attent Vigila	Attention/ Vigilance	
SNP	В	р	В	р	В	р	В	р	В	р	В	р	
rs6904071	1.060	0.046	0.242	0.077	0.448	0.049	0.553	0.284	1.488	0.011	0.150	0.001	
rs13219354	0.968	0.138	0.033	0.847	0.200	0.474	-0.275	0.663	0.525	0.465	0.102	0.063	
rs3131296	-0.107	0.866	-0.044	0.790	-0.369	0.219	0.563	0.403	0.615	0.423	0.107	0.067	
rs6932590	0.909	0.056	0.131	0.284	0.362	0.081	-0.335	0.473	0.002	0.997	0.077	0.053	
rs9960767 ^b	0.537	0.555	0.405	0.085	-0.100	0.794	-0.201	0.814	0.177	0.856	-0.127	0.073	
rs12807809	-1.03	0.060	-0.104	0.460	-0.064	0.791	0.309	0.570	-0.276	0.656	0.003	0.950	

TABLE 1. Results of Linear Regression of Genome-Wide Significant Single-Nucleotide Polymorphism (SNP) Associations With Cognitive Domains^a

^a Positive regression coefficient indicates schizophrenia risk allele associated with worse performance on cognitive test. Nominally significant p values were taken forward for replication.

^b For loci with minor allele frequencies <0.1, the analysis was redone by grouping together minor allele and heterozygote genotype groups. The results remained unchanged and none were significant.

TABLE 2. Detailed Results for rs6904071 in the German Sample in a Study of Single-Nucleotide Polymorphism Associations With Cognitive Domains

Cognitive				AA		AG		GG		Regression		
Function	Test or Subscale ^a	Sample	Ν	Mean	SD	Mean	SD	Mean	SD	Coefficient ^b	95% CI	р ^с
General cognitive ability	Full-scale IQ	Patients Comparison	342 2,244	106.5 114.5	17.8 13.9	100.8 113.8	17.7 14.5	99.3 113.0	18.5 14.5	1.060	0.020–2.101	0.046
Verbal working memory	WAIS digit span	Patients Comparison	342 2,244	16.4 14.7	4.6 4.0	14.0 14.3	3.5 3.9	13.3 14.2	3.7 3.8	0.242	-0.026 to 0.511	0.077
Spatial working memory	WMS-R spatial span	Patients Comparison	239 399	18.5 17.8	3.73 3.79	15.3 17.3	2.98 3.11	14.91 17.11	3.40 3.43	0.448	0.002–0.895	0.049
Immediate episodic memory	WMS-R immediate logical memory	Patients Comparison	240 400	29.2 31.6	8.6 7.4	24.8 30.6	8.2 6.5	23.9 30.7	9.0 6.6	0.553	-0.461 to 1.566	0.284
Delayed episodic memory	WMS-R delayed logical memory	Patients Comparison	239 400	30.0 36.7	11.5 9.1	28.8 34.7	9.4 6.6	25.8 34.1	10.5 7.4	1.488	0.341–2.635	0.011
Attention/ vigilance	CPT-IP (3–7 version)	Patients Comparison	346 517	4.59 4.90	0.67 0.49	4.28 4.85	0.83 0.56	4.10 4.71	0.95 0.58	0.150	0.063–0.236	0.001

^a WAIS=Wechsler Adult Intelligence Scale; CPT-IP=Continuous Performance Test, Identical Pairs Version; WMS-R=German Wechsler Memory Scale–Revised. Raw scores are provided for WMS-R tests; all other scores are scaled.

^b B coefficient with respect to schizophrenia risk allele (G) adjusted for age and sex as appropriate.

^c Nominally significant association results taken forward for replication in Irish sample.

between rs6904071 and five of the six cognitive tests at a nominally significant level. Notably, the combined analysis results indicate association with delayed episodic memory and attention/vigilance at a level that would survive adjustment for multiple testing of all six SNPs and six cognitive tests (p<0.0014). Table 4 also outlines the results for schizophrenia patients and healthy comparison subjects considered separately. rs6904071 was associated with performance on cognitive measures in both patients and comparison subjects.

Because the replicated finding and the most significant association in the combined analysis was for delayed episodic memory, we repeated the combined analysis for the other cognitive domains while adjusting for delayed episodic memory to investigate whether the other cognitive associations were independent of the association with this cognitive domain. The sample for the adjusted analysis was smaller (maximum N=1,161), but rs6904071 remained significantly associated with the same cognitive domains in this sample in unadjusted analyses. However, after adjusting for delayed episodic memory performance, none of the other cognitive domains remained significantly associated (see Table S7 in the online data supplement for results of this analysis).

The association of rs6904071 genotype with demographic and clinical variables in both samples is summarized in Table S8 in the online data supplement. No significant differences in age, gender distribution, and education by genotype were found in patients or comparison subjects in either sample. Furthermore, in patients, no significant differences were observed between genotype groups in antipsychotic medication dosage in either sample.

Structural Imaging

As the only significant SNP to emerge from the analysis with cognition, rs6904071 alone was taken forward to the structural imaging analysis. This SNP was significantly

880

associated with total hippocampal volume (B=0.080; 95% confidence interval [CI]=0.006–0.154, p=0.035) and with total gray matter volume (B=-5.93; 95% CI=-0.37 to -11.49, p=0.036). The schizophrenia risk allele, which was associated with worse cognition, was associated with smaller hippocampal volume, which was the predicted direction of association. The finding for gray matter was in the counterintuitive direction, in that the risk allele was associated with increased gray matter volume.

Discussion

At the time this study was conducted, the polymorphisms that we examined represented all of the genomewide significant common variants thus far reported for schizophrenia, with the exception of rs1344706 at ZNF804A, on which we have previously reported (11). In taking the GWAS results forward to examine their effects on cognition, we sought to overcome the limitations of previous cognitive and genetic studies by testing SNP associations with a priori selected cognitive tests in large samples of healthy subjects and schizophrenia patients and by seeking replication of results in a sample with equivalent tests. Despite this design, we failed to show any association with cognition for five of the six schizophrenia risk SNPs tested. TCF4 and NRGN are good candidate genes for cognition given their putative roles in neurodevelopment and the processes involved in cognition (4, 5). It is therefore of particular interest that neither of these risk variants was associated with cognitive performance in this study. Given that our study had the power to detect an allelic effect accounting for 0.3% of variance in IQ in the German comparison sample alone, it is unlikely that these alleles have effects of this magnitude on human cognition, although we cannot exclude smaller effects.

This lack of effect at the behavioral and cognitive level for the majority of the tested SNPs is perhaps to be expected, given recent evidence that the overlap between the genetics of schizophrenia and cognition is less than previously proposed and that the majority of the genetic variance for schizophrenia does not overlap with that of cognition (24, 25). This is in keeping with other research suggesting a limited role in cognition for common candidate variants for schizophrenia (26).

The strongest evidence we obtained for an association between schizophrenia risk alleles and cognitive measures was for SNP rs6904071, in the MHC region. The association was in the predicted direction—that is, the schizophrenia risk allele was associated with poorer cognition. The replicated association of this SNP with cognition was for the domain of delayed episodic memory. That phenotype was evaluated in the German and Irish samples using precisely the same test (delayed logical memory). The lack of replication for other cognitive domains may be due to a lack of formal equivalence between the domain tests in the two samples or as a result of the relatively smaller size of FIGURE 1. Delayed Episodic Memory Scores in German and Irish Patients With Schizophrenia and Healthy Comparison Subjects, by rs6904071 Genotype^a



^a WMS-R= German Wechsler Memory Scale–Revised; WMS-III=Wechsler Adult Intelligence Scale, 3rd edition.

the Irish replication sample. To counter the sample size limitation, we combined data across both samples using z scores for equivalent tests, and we demonstrated nominal association with five of the six cognitive domains in the combined sample. However, in follow-up adjusted analyses, the significant associations for the other cognitive domains could have been explained by the association with delayed episodic memory. This would suggest that the effect on cognition of whatever functional variant is tagged by rs6904071 is through biological pathways involved in episodic memory. It follows that the same pathways could be involved in the action of this polymorphism on elevated schizophrenia risk, although the possibility of pleiotropy cannot be ruled out by our analysis.

TABLE 3. Results	for rs6904071 in [•]	the Irish Replicatio	n Sample in a St	udy of Single-Nucle	otide Polymorphism	Associations
With Cognitive Do	omains					

Cognitive				AA		AG		GG		Regression		
Function	Test or Subscale ^a	Sample	Ν	Mean	SD	Mean	SD	Mean	SD	Coefficient ^b	95% CI	р ^с
General cognitive ability	Abbreviated full-scale IQ	Patients Comparison	300 148	92.5 131.0	19.3 11.5	91.5 121.5	18.0 14.9	90.5 121.8	18.1 14.6	1.389	-1.592 to 4.369	0.180
Spatial working memory	Cambridge Neuropsychological Test Automated Battery spatial working memory task	Patients Comparison	365 140	-1.08 0.26	0.92 0.83	-0.97 0.26	1.29 0.75	-0.96 0.17	1.34 0.84	0.005	-0.188 to 0.199	0.478
Delayed episodic memory	WMS-III logical memory delayed	Patients Comparison	377 145	6.5 16.2	3.3 1.6	7.7 13.4	3.2 2.5	7.0 13.0	3.3 2.6	0.575	0.083–1.067	0.011
Attention/ vigilance	CPT-IP (3 letters)	Patients Comparison	257 n/a	2.27	1.5	1.97	0.87	1.95	1.04	0.031	-0.209 to 0.271	0.399

^a Scaled test scores. WMS-III=Wechsler Memory Scale, 3rd edition; CPT-IP=Continuous Performance Test, Identical Pairs Version.

^b B coefficient with respect to schizophrenia risk allele (G) adjusted for age and sex as appropriate.

^c One-sided p value of regression.

TABLE 4. ASSOCIATION OF IS6904071 WITH Z SCORES OF COGNITIVE DOMAINS IN COMDINED GERMAN AND THIS SAMPL	TABLE 4.	Association	of rs6904071	With z Score	es of Cognitive	Domains in	Combined	German and	Irish Sa	ample
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		Comparison Subjects				Patients			Combined Sample ^a		
Cognitive Function	Participants	N	B (Regression Coefficient)	95% Cl	р	В	95% CI	р	В	95% CI	р
General cognitive ability ^b	Patients Comparison	642 2,392	0.072	0.006–0.139	0.034	0.113	-0.040 to 0.265	0.149	0.08	0.018–0.141	0.011
Verbal working memory ^c	Patients Comparison	711 2389	0.053	-0.020 to 0.126	0.158	0.145	0.018–0.271	0.025	0.072	0.009–0.136	0.026
Spatial working memory ^d	Patients Comparison	604 539	0.073	-0.057 to 0.203	0.269	0.025	-0.178 to 0.127	0.743	0.027	-0.073 to 0.127	0.596
Immediate episodic memory ^e	Patients Comparison	620 545	0.093	-0.027 to 0.213	0.128	0.104	-0.035 to 0.243	0.141	0.098	0.006–0.189	0.036
Delayed episodic memory ^f	Patients Comparison	616 545	0.156	0.042–0.270	0.007	0.180	0.042–0.317	0.01	0.166	0.077–0.255	2.66×10 ⁻⁴
Attention/ vigilance ^g	Patients Comparison	603 517	0.148	0.040-0.257	0.007	0.180	0.010-0.349	0.038	0.164	0.063–0.265	0.001

^a Adjusted for diagnostic status.

^b Full-scale IQ in German sample and abbreviated IQ in Irish sample.

^c Wechsler Adult Intelligence Scale digit span in German sample and letter-number sequencing in Irish sample.

^d Wechsler Memory Scale spatial span in German sample and Cambridge Neuropsychological Test Automated Battery spatial working memory in Irish sample.

^e Immediate logical memory in both samples.

^f Delayed logical memory in both samples.

^g 3–7 version of the Continuous Performance Test used in German sample and three-letter version of the Continuous Performance Test, Identical Pairs Version, used in Irish sample.

By an order of magnitude, delayed episodic memory was the most strongly associated cognitive domain, and rs6904071 was associated with performance in this domain in both patients and comparison subjects. Performance on the delayed logical memory test is known to be sensitive to hippocampal pathology and resection (27, 28) and is correlated with hippocampal volume in clinical samples (29) as well as in the relatives of patients with schizophrenia (30). Thus, our finding of association with decreased hippocampal volume links rs6904071 to both structure and function. Given the known prior relationship between the cognitive and structural phenotype, the fact that we observed an association with hippocampal structure in a sample independent of the sample from which the cognitive measures were obtained provides independent validation of the associations. This finding adds to the evidence suggesting an association between SNPs in the MHC region and structural brain volumes, including the hippocampus (31, 32). rs6904071 was also associated with increased gray matter volume. The interpretation of this result is not as straightforward as the hippocampal finding and may well be a false positive finding as a result of chance. While schizophrenia is typically associated with decreased gray matter volume, other neurodevelopmental disorders, such as autism, may be characterized by increased gray matter volume, particularly in individuals with low IQ (33). An inverse relationship between structural volume and performance has been posited for various cognitive abilities (34), although whether this proves relevant for the association between rs6904071 and gray matter volume will require further exploration.

The associated SNP (rs6904071) lies within the extended MHC region on chromosome 6. The prospect of this region harboring genetic risk variants for schizophrenia has long been recognized (35) and has some support in research (9, 36, 37). Despite this, the association between the MHC region and schizophrenia has been criticized as likely a result of population stratification (38). To counter this criticism, our study used a subset of participants in the German sample for whom genome-wide genotyping was available (N=1,171) and derived principal components using the Eigenstrat method (39). The association between rs6904071 and cognition was robust to the addition of one, two, or three principal components, none of which affected the estimated effect size for any of the cognitive tests (see Table S9 in the online data supplement). This indicates that population stratification is unlikely to explain these results, particularly when taken together with the finding of replication in an independent sample.

It is possible that the association between schizophreniaassociated markers at the MHC region and cognition and brain structure may implicate altered immune function in each of these phenotypes. Of over 160 genes in the MHC region, roughly 30% are immune related, and the region has been implicated in numerous immune-related diseases (40). The possible link between immune function and schizophrenia is reinforced by evidence suggesting co-segregation between a range of autoimmune conditions and schizophrenia (41).

However, in addition to immune-related explanations, there are alternative potential mechanisms underpinning the associations we have observed. Our signal lies within the MHC class I region. MHC class I proteins have an established role in adaptive immunity, but findings also point to an involvement of this complex in synaptic development and plasticity (42). Specifically, MHC class I influences hippocampal long-term potentiation (43), providing a potential link between rs6904071, altered hippocampal structure and function, and a major mechanism of learning and memory. Furthermore, MHC class I protein expression co-localizes postsynaptically with postsynaptic density protein 95 in dendrites of the hippocampus and has been found to play a role in synaptic morphology and function (44). In the absence of evidence that the associations we have observed reflect altered function at MHC class I, we mention this as a caution against assuming that the association implicates immunity. Given the LD structure at this region, identifying the direct pathophysiological implications of the genetic associations will prove challenging.

To summarize, in following up genome-wide significant schizophrenia loci, we have demonstrated that the most significant risk variant from the International Schizophrenia Consortium and Molecular Genetics of Schizophrenia studies (1, 2) is associated with cognition, specifically delayed episodic memory. We have also both replicated this finding and shown that the same allele is associated with hippocampal volume in a manner consistent with known relationships between cognition and hippocampal structure. The associated polymorphism is in the MHC region, but until the source of the association signal is more precisely identified, it is impossible to adjudicate between several theoretically plausible mechanisms linking the association to schizophrenia and cognition in the MHC. The ultimate resolution of this question has the potential to indicate novel mechanisms linking cognitive deficits to schizophrenia.

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