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Genome-wide Association Analysis of Schizophrenia and Vitamin D Levels Shows Shared Genetic Architecture and Identifies Novel Risk Loci

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Low vitamin D (vitD) levels have been consistently reported in schizophrenia (SCZ) suggesting a role in the etiopathology. However, little is known about the role of underlying shared genetic mechanisms. We applied a conditional/conjunctive false discovery rate approach (FDR) on large, nonoverlapping genome-wide association studies for SCZ (N cases = 53 386, N controls = 77 258) and vitD serum concentration (N = 417 580) to evaluate shared common genetic variants. The identified genomic loci were characterized using functional analyses and biological repositories. We observed cross-trait SNP enrichment in SCZ conditioned on vitD and vice versa, demonstrating shared genetic architecture. Applying the conjunctive FDR approach, we identified 72 loci jointly associated with SCZ and vitD at conjunctive FDR < 0.05. Among the 72 shared loci, 40 loci have not previously been reported for vitD, and 9 were novel for SCZ. Further, 64% had discordant effects on SCZ-risk and vitD levels. A mixture of shared variants with concordant and discordant effects with a predominance of discordant effects was in line with weak negative genetic correlation ($r_g = -0.085$). Our results displayed shared genetic architecture between SCZ and vitD with mixed effect directions, suggesting overlapping biological pathways. Shared genetic variants with complex overlapping mechanisms may contribute to the coexistence of SCZ and vitD deficiency and influence the clinical picture.

Key words: GWAS/genetic overlap/conditional/conjunctive false discovery rate/polygenic architecture

Introduction

Schizophrenia (SCZ) is a severe mental disorder characterized by psychotic symptoms with onset in early adulthood and large global disease burden.¹ SCZ has a high heritability. However, the underlying disease mechanisms are not yet fully understood.¹ A role of vitamin D (vitD) in SCZ pathophysiology has been supported by 2 decades of research.² The primary hypothesis, proposing low vitD level as a risk factor for SCZ, was based on epidemiological evidence linking SCZ with winter births (low maternal vitD), urban upbringing, and migration to high latitudes causing reduced sun exposure.²⁻⁴ Moreover, neonatal vitD deficiency was associated with higher risk for SCZ^{5,6} and low vitD level has been linked to poor prognostic markers in individuals with a psychotic disorder, including more severe negative symptoms^{7,8} and poorer cognitive functioning.⁹

The potential role of vitD in the pathogenesis of SCZ is further supported by its involvement in brain development and function.^{2,10} The main source (~90%) of vitD is production of its D3 form (cholecalciferol) from 7-dehydrocholesterol in the skin by exposure to ultraviolet

light.^{11,12} Additionally, vitD can be derived from some dietary sources such as fish liver oil (eg, cod), fatty fish (eg, mackerel), egg yolks, beef (eg, kidney), pork (eg, liver), mushrooms (eg, shiitake) as well as fortified orange juice and dairy products.¹³ VitD binds to its cytoplasmic receptor (VDR) which subsequently forms a heterodimer with the retinoic acid receptor and, after translocation to the nucleus, regulates the expression of target genes.² Regarding CNS-specific effects, vitD modulates prefrontal cortex development via non-genomic mechanisms, namely through modulation of L-type voltage-gated calcium channels.^{14,15} Animal models have indicated that vitD may also exert anti-proliferative and pro-apoptotic effects in the developing brain² which is associated with altered expression of cyclin-dependent kinases and cyclin D1.¹⁶ Developmental vitD deficiency results in abnormal architecture and/or size of rodent brain structures including the hippocampus and lateral ventricles.² It has been reported that developmental vitD deficiency downregulates some neurotrophic factors (eg, nerve growth factor and glial-derived neurotrophic factor) which play a critical role in neurite outgrowth and dendritic arborization.^{2,17} Moreover, vitD modulates the inflammatory response and the glucocorticoid system, as evidenced by elevated maternal inflammatory cytokines and glucocorticoid levels in rodent models of vitD deficiency.^{2,18} On a behavioral level, developmental vitD deficiency in rats is associated with impaired mother-offspring interaction, disturbed ultrasonic vocalization, stereotyped behavior, sensitivity to psychomimetics, and impaired learning and memory.²

Several lines of evidence implicate vitD in a plethora of developmental processes linked with the pathogenesis of SCZ.² Genetic variants may impact these neurobiological processes directly, and indirectly via systemic vitD levels (eg, through the gut vitD absorption feedback loop or the tan response), sensitivity to environmental factors (eg, a low sun exposure), as well as the resistance of the developing brain to vitD deficiency. Thus, it is important to scrutinize potential shared genetic architecture. SCZ is highly heritable, with broad-sense heritability estimated to be 0.64–0.81.^{19,20} Most of the genetic risk is attributed to thousands of common genetic variants with small effect sizes.²¹ While the most recent GWAS of SCZ identified 270 loci, the majority of genetic risk variants have not yet been discovered.²² The genetic loci associated with SCZ implicate processes related to brain development, signal transduction, and inflammatory response in the etiology of SCZ.²⁰ The heritability of vitD levels has been estimated in twin and familial studies to be within a broad range of 0%–85%.²³ A recent GWAS of vitD identified 143 loci associated with vitD concentration implicating among others genes associated with lipid metabolism, and estimated the SNP heritability to be 0.32.²⁴ The potential genetic link between vitD and SCZ has been investigated using genetic correlation, which was estimated to

be -0.086 .²⁴ However, genetic correlation fails to quantify mixtures of effect directions across variants since it is unable to differentiate scenarios of no genetic overlap from overlap with a balance of shared variants with concordant (a given variant has same direction of effect in both traits) and discordant effects (a given variant increases the risk of 1 trait while decreasing the risk of the second).²⁵ This limitation is overcome by the conjunctive false discovery rate (conjFDR) approach which build on an empirical Bayesian statistical framework.^{26–28}

In the current study, we applied the conjFDR method to large nonoverlapping GWASs of SCZ²² and vitD levels²⁴ to investigate their shared polygenic architecture. An improved understanding of the overlapping genetic architecture and jointly associated loci between SCZ and vitD regulation may implicate pathophysiological processes of SCZ and may form the basis of preventive measures and identify new targets for drug development.

Methods and Materials

GWAS Samples

GWAS summary statistics for SCZ were obtained from the Psychiatric Genomics Consortium (53 386 cases and 77 258 controls),²² and for vitD serum concentration (417 580 individuals) from <https://cnsgenomics.com/content/data>. See [Supplementary Materials](#) for more details. GWASs investigated in the study were approved by the ethics committees, and informed consent was obtained from all participants.

Statistical Analyses

To estimate the polygenicity (number of causal variants) and discoverability (expected squared effect size of a given causal variant) for SCZ and vitD we applied univariate MiXeR.²⁹ Univariate MiXeR is based on an analysis of the linkage disequilibrium (LD) structure from a detailed reference panel to estimate polygenicity and discoverability for a given trait from GWAS summary statistics.^{25,29} and has been successfully applied for human phenotypes, including psychiatric disease and brain architecture.^{30,31} Consistent with previous studies, polygenicity is expressed as the number of causal variants with strongest effects required to explain 90% SNP heritability.

We visualized cross-trait SNP enrichment by generating conditional QQ plots which display the distribution of *P*-values for the primary trait conditioned on significance levels in a secondary trait (ie, SCZ trait conditioned on significance levels in vitD serum concentration phenotype and vice versa). The QQ plots were constructed after random pruning averaged over 500 iterations to control for spurious enrichment. In every iteration, 1 SNP was randomly selected for each LD block ($r^2 > 0.1$) and corresponding *P*-values were used to construct the empirical cumulative distribution functions. To avoid potential

bias associated with complex LD structure, we excluded SNPs within 2 regions (major histocompatibility complex region 6:25119106-33854733; 8:7200000-12500000). An empirical cumulative distribution of nominal P -values for a given primary trait was computed for all SNPs and for SNPs with significance levels below the indicated cutoffs for the secondary trait (corresponding to $P < .1$, $P < .01$, $P < .001$). Statistical enrichment of the given primary trait (cross-trait enrichment) is indicated by successive leftward deflection in the conditional QQ plot as levels of association with the secondary trait increase.^{28,32,33}

To assess genetic overlap between phenotypes and to boost SNPs discovery rate, we used conditional FDR (condFDR) and conjunction FDR (conjFDR) approaches. The conjFDR method identifies loci with statistical pleiotropy.²⁸ The method builds on an empirical Bayes mixture model,³⁴ with 2 distributions (null and non-null effect loci) and has a higher replication rate compared to methods based on the infinitesimal model (all loci have an effect³⁴). This approach has been applied previously to identify novel genetic loci for a series of psychiatric and somatic phenotypes, as well as to detect shared genetic loci between diverse phenotypes.³⁵⁻³⁹ CondFDR is defined as the posterior probability that a SNP of interest is null for the primary trait given that the P -values for both traits are as small or smaller than the observed P -values. ConjFDR identifies shared genetic loci and is defined as the posterior probability that a SNP is null for either 1 trait or both traits simultaneously, given the P -value for traits are equal to or smaller than the observed P -values. A conservative estimate of the conjFDR is defined by the maximum of 2 condFDR values (trait 1 conditioned on trait 2 and *vice versa*).⁴⁰ We used FDR significance cutoffs for condFDR and conjFDR in line with previous studies, at 0.01 and 0.05, respectively.^{27,31} Due to extensive LD within these regions, we excluded SNPs located within the MHC and 8p23.1 inversion before fitting the cond/conjFDR model to avoid inflation of the estimated parameters.³⁵ Although previous studies demonstrated that conjFDR findings for human phenotypes are well replicated in independent samples,^{41,42} in-depth experimental analyses are required to determine the mechanisms involved.^{43,44}

We applied the FUMA protocol to identify independent genomic loci.⁴⁵ We defined independent significant SNPs as those which reached a conjFDR < 0.05 and were independent from each other at $r^2 < 0.60$. To create a subset of lead SNPs, we selected independent significant SNPs with LD with each other at $r^2 < 0.10$. The range of each genomic locus was limited to SNPs in LD ($r^2 \geq 0.60$) with any of the independent significant SNPs within the locus. Subsequently, we merged all loci that were less than 250 kb apart to define distinct genomic loci and marked SNPs with the lowest P -value as a lead SNP of the locus. We used the 1000 Genomes Project European ancestry haplotype reference panel to annotate LD information.

Functional Annotation

Novel loci were defined as loci which were not physically overlapping with findings from the original GWAS, GWAS Catalog by National Human Genome Research Institute-European Bioinformatics (NHGRI-EBI⁴⁶) and previous cond/conjFDR studies.³¹ We applied positional mapping by physical position (10-kb window) and expression quantitative trait locus association (eQTL) as gene-mapping strategies for candidate SNPs with cond/conjFDR ≤ 0.10 .⁴⁵ The direction of allelic effects in SNPs identified by conjFDR were investigated by comparing the z-scores in SCZ original GWAS against the z-scores in vitD levels original GWAS. Only genes identified by both strategies, after exclusion of those located within regions of complex LD, were provided for subsequent gene-set analysis. Gene-set enrichment analysis was performed using FUMA GENE2FUNC functionality.

Results

Univariate MiXeR

Univariate MiXeR analyses established a higher heritability (SCZ = 0.36; vitD = 0.076), higher polygenicity (SCZ = 8700 SNPs accounting for 90% of the heritability; vitD = 400), and lower discoverability (SCZ = 6.48e-05; vitD = 3.34e-04) for SCZ than for vitD levels. Moreover, it was estimated that the SNPs which were genome-wide significant in the original GWASs explained 4.5% (SCZ) and 52.9% (vitD) of the trait variance associated with SNPs.

QQ Plots of SCZ SNPs Conditioned on Association with vitD and Vice Versa

The conditional QQ plot for SCZ GWAS conditioned on vitD levels ([figure 1A](#)) demonstrated enrichment across different levels of significance for vitD. The earlier leftward deflection from the dashed line (no enrichment) indicated a greater proportion of true associations for a given nominal SCZ P -value. Successive leftward shifts for decreasing nominal vitD P -value thresholds demonstrated that the proportion of non-null effects in SCZ increases as the strength of the association with vitD increases. Similarly, the conditional QQ plot displayed strong polygenic enrichment for vitD given the strength of association with SCZ ([figure 1B](#)).

Applying the condFDR approach (condFDR < 0.01), a total of 286 distinct genomic loci were associated with SCZ conditional on their association with vitD ([Supplementary figure 1](#) and [Supplementary table 5](#)). Sixteen of these loci have not been identified in the original SCZ GWAS,²² the NHGRI-EBI catalog⁴⁶ or the previous cond/conjFDR studies ([Supplementary table 1](#)). At condFDR < 0.01 , we identified 240 loci associated with vitD levels conditional on SCZ ([Supplementary figure 2](#), [Supplementary table 6](#)). Of these, 97 loci have not been

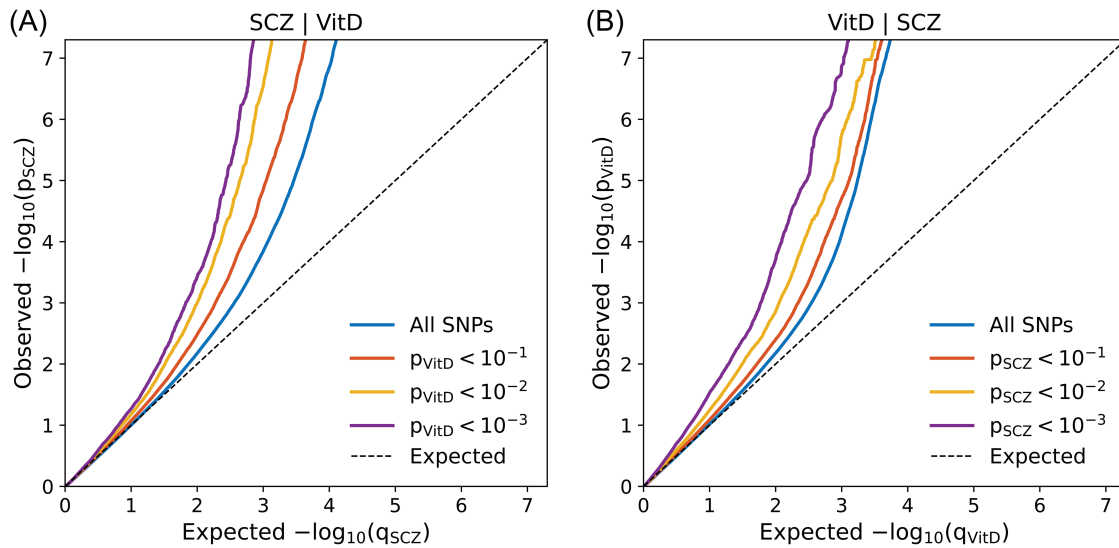


Fig. 1. Conditional QQ plots illustrating cross-phenotype polygenic enrichment between schizophrenia (SCZ) and vitamin D levels (vitD). (A) SCZ trait is conditioned on vitD (B) vitD is conditioned on SCZ.

Table 1. Novel Schizophrenia (SCZ) Genomic Loci also Associated with Vitamin D Levels (vitD) at Conjunctural FDR < 0.05

Locus	Chr	Lead SNP	A1/ A2	Nearest gene	Functional category	P value SCZ	Odds Ratio SCZ	P value vitD	Beta vitD	conjFDR
12	2	rs56393175*	T/G	<i>TM4SF20</i>	intergenic	3.04E-06	1.05	3.18E-05	-1.12E-02	1.09E-02
20	3	rs112936817*	T/G	<i>DGKG</i>	intronic	1.16E-04	0.95	3.13E-05	1.33E-02	4.90E-02
23	4	rs10516786	G/A	<i>AFF1</i>	intronic	5.66E-05	0.96	2.52E-05	-1.12E-02	3.37E-02
34	8	rs12546328	T/C	<i>UBXN2B</i>	intronic	1.03E-04	1.04	7.97E-05	8.40E-03	4.60E-02
35	8	rs4734176*	G/A	<i>EMC2</i>	intronic	6.63E-05	1.04	1.98E-04	-8.13E-03	3.66E-02
39	10	rs11001489*	G/A	<i>C10orf11:RP11-367B6.2</i>	ncRNA_intronic	3.41E-05	0.94	2.24E-04	1.20E-02	3.73E-02
40	11	rs1950039	T/C	<i>SPON1</i>	ncRNA_intronic	9.67E-05	0.97	6.68E-21	1.91E-02	4.47E-02
54	14	rs79621605	C/T	<i>CDKLI</i>	intronic	1.17E-04	0.91	2.13E-06	-2.62E-02	4.92E-02
68	19	rs72999390	A/G	<i>UPFI</i>	intronic	3.79E-06	0.90	8.80E-05	-1.81E-02	2.09E-02
70	20	rs6020924*	T/G	<i>KCNGB1</i>	intergenic	5.97E-05	0.95	5.90E-06	1.30E-02	3.46E-02

Note: The most strongly associated SNPs in independent genomic loci shared between SCZ and vitD at conjFDR < 0.05. Chromosomal position (Chr), nearest gene, and functional category, as well as P values and effect sizes (Odds ratio and Beta for SCZ and Vitamin D, respectively) from original summary statistics. The effect allele is A1. Novel for vitD levels marked with *.

identified earlier, neither in the original vitD GWAS,²⁴ nor in the NHGRI-EBI catalog⁴⁶ (Supplementary table 2).

Applying the conjFDR approach, a total of 72 loci were jointly associated with SCZ and vitD at conjFDR < 0.05 (Figure 2, Supplementary table 4). Evaluation of the allelic effect direction for the 72 lead SNPs showed that 46 (64%) had discordant effect directions on SCZ-risk and vitD levels, ie, increased the risk for SCZ while reducing vitD levels or vice versa.

Our conjFDR analysis displayed 9 novel loci for SCZ (table 1), not previously identified.^{22,46} Among the novel SCZ loci, 1 locus (lead SNP rs112936817, 3:186075556-186079990) was included in the GWAS catalog due to reported associations with chronotype⁴⁷ and morningness.⁴⁸ The other loci (lead SNP rs12546328, 8:59321609-59384675) were associated with narcolepsy

with cataplexy.⁴⁹ Moreover, we identified 40 novel loci for vitD levels (table 2), many of which were reported in the GWAS catalog for traits associated with vitD (eg, lead SNP rs6779146, 3:135798658-136507818,⁵⁰⁻⁵² lead SNP rs78004870, 16:89736352-89886392⁵³; table 3).

Functional Analyses

Functional annotation of all shared lead SNPs identified in conjFDR analysis displayed that the majority (81%) were intergenic or intronic, while 3 were exonic and 6 were associated with noncoding RNA (Supplementary table 1). Seven lead SNPs (rs2464196, rs2267791, rs72999390, rs61182735, rs10514040, rs13024749, rs112936817) had a CADD score above the threshold score of 12.37, indicative of deleteriousness.⁶¹ Among those 7 SNPs with

Table 2. Novel Vitamin D Levels Genomic Loci also Associated with Schizophrenia (SCZ) at Conjunctive FDR < 0.05

Locus	Chr	Lead SNP	A1/ A2	Nearest gene	Functional category	P value Vitamin D	Beta Vi- tamin D	P value SCZ	Odds Ratio SCZ	conjFDR
3	1	rs2842198	A/G	<i>HYI-AS1</i>	intergenic	2.13E-05	-8.68E-03	1.51E-07	0.95	8.40E-03
4	1	rs12567744	G/A	<i>CDKN2C</i>	intergenic	3.31E-04	-7.68E-03	1.03E-04	1.04	4.71E-02
7	1	rs12743378	A/G	<i>SDCCAG8</i>	intronic	6.55E-06	1.28E-02	5.10E-07	0.94	3.91E-03
10	2	rs999494	T/C	<i>EMX1</i>	intronic	8.85E-05	-9.90E-03	2.61E-07	0.95	2.10E-02
11	2	rs13024749	T/G	<i>LY75:LY75-CD302</i>	intronic	3.38E-04	-7.24E-03	3.97E-05	1.04	4.77E-02
12	2	rs56393175*	T/G	<i>TM4SF20</i>	intergenic	3.18E-05	-1.12E-02	3.04E-06	1.05	1.09E-02
13	3	rs61182735	A/G	<i>AC091493.2</i>	intergenic	4.85E-05	-8.11E-03	1.16E-07	1.05	1.43E-02
14	3	rs9831428	A/C	<i>RBMS3</i>	intronic	2.02E-04	7.52E-03	6.77E-05	0.97	3.70E-02
17	3	rs2241823	A/C	<i>ATXN7</i>	intronic	1.38E-04	8.30E-03	1.26E-07	1.05	2.76E-02
18	3	rs2088176	G/A	<i>ROBO2</i>	intronic	1.66E-04	7.56E-03	1.57E-05	0.96	3.11E-02
19	3	rs6779146	C/T	<i>MSL2</i>	intronic	1.04E-04	-9.21E-03	1.87E-06	1.05	2.32E-02
20	3	rs112936817*	T/G	<i>DGKG</i>	intronic	3.13E-05	1.33E-02	1.16E-04	0.95	4.90E-02
21	4	rs80279461	T/G	<i>PCDH7</i>	intergenic	1.15E-04	-1.09E-02	1.55E-05	0.95	2.47E-02
25	5	rs904612	A/G	<i>CTB-35F21.1</i>	ncRNA_ intronic	2.54E-06	-1.05E-02	4.05E-06	1.05	7.06E-03
27	6	rs9273493	T/C	<i>HLA-DQB1:HLA-DQB1-AS1:XXbac-BPG254F23.6</i>	ncRNA_ exonic	8.21E-06	-9.10E-03	2.35E-06	1.05	4.94E-03
30	7	rs274632	A/C	<i>GRM3</i>	intergenic	2.24E-04	-7.43E-03	7.98E-05	1.04	4.03E-02
32	7	rs568471	T/C	<i>RNU4-74P</i>	upstream	1.82E-04	1.15E-02	6.57E-05	0.95	3.64E-02
35	8	rs4734176*	G/A	<i>EMC2</i>	intronic	1.98E-04	-8.13E-03	6.63E-05	1.04	3.66E-02
37	8	rs10875482	C/T	<i>TSNARE1</i>	intronic	3.05E-04	-7.40E-03	9.96E-12	0.94	4.49E-02
39	10	rs11001489*	G/A	<i>C10orf11:RP11-367B6.2</i>	ncRNA_ intronic	2.24E-04	1.20E-02	3.41E-05	0.94	3.73E-02
41	11	rs2582894	G/T	<i>RP11-22P4.1</i>	intergenic	1.18E-04	-1.03E-02	1.78E-05	0.95	2.51E-02
42	11	rs6591434	G/A	<i>RP11-734C14.2</i>	intergenic	2.76E-04	-7.78E-03	8.61E-08	1.05	4.24E-02
43	11	rs484201	T/C	<i>MACROD1</i>	intronic	2.53E-04	-7.41E-03	4.44E-06	1.04	4.02E-02
45	11	rs7948931	C/T	<i>DLG2</i>	intronic	6.60E-05	8.08E-03	1.44E-05	1.04	1.74E-02
46	11	rs61123830	A/G	<i>GRAMD1B</i>	intergenic	4.25E-05	8.90E-03	7.09E-05	0.96	3.80E-02
48	12	rs10860964	C/T	<i>RP11-328J6.1</i>	intergenic	2.82E-05	8.59E-03	6.39E-09	0.95	1.01E-02
49	12	rs7313797	C/T	<i>KCTD10</i>	UTR5	1.55E-04	-7.58E-03	9.48E-05	0.97	4.42E-02
50	12	rs2464196	A/G	<i>HNF1A</i>	exonic	2.40E-04	8.05E-03	7.43E-05	1.04	3.89E-02
51	12	rs79478560	A/G	<i>DNAH10</i>	intronic	4.63E-06	-9.53E-03	3.89E-07	1.05	3.11E-03
53	14	rs17522122	T/G	<i>AKAP6</i>	intergenic	2.59E-04	-7.34E-03	1.08E-10	1.06	4.08E-02
56	15	rs8035957	C/T	<i>RASGRP1</i>	intronic	1.63E-05	-9.85E-03	8.35E-06	1.04	1.12E-02
57	15	rs12905223	C/T	<i>UBE2Q2P1:LINC00933</i>	ncRNA_ exonic	1.01E-04	-8.68E-03	1.69E-09	0.94	2.28E-02
60	16	rs116001925	T/C	<i>RP11-439II4.3</i>	intergenic	1.05E-05	1.88E-02	5.17E-05	0.92	3.21E-02
61	16	rs61733486	T/C	<i>PRMT7</i>	exonic	2.40E-04	-1.58E-02	2.87E-06	1.09	3.89E-02
62	16	rs78004870	T/G	<i>FANCA</i>	exonic	1.40E-06	1.05E-02	3.27E-06	0.96	6.13E-03
63	17	rs6502219	C/T	<i>ARHGAP44</i>	intronic	1.44E-05	9.38E-03	6.60E-06	1.04	9.66E-03
64	17	rs11263770	A/G	<i>MYO19</i>	intergenic	5.99E-05	-8.03E-03	4.83E-06	0.96	1.64E-02
65	17	rs6504584	T/G	<i>CALCOCO2</i>	UTR3	6.16E-05	8.42E-03	3.31E-05	0.96	2.51E-02
67	19	rs8103241	G/A	<i>NFIX</i>	intronic	2.36E-05	8.43E-03	5.07E-05	0.97	3.17E-02
70	20	rs6020924*	T/G	<i>KCNGB1</i>	intergenic	5.90E-06	1.30E-02	5.97E-05	0.95	3.46E-02

Note: The most strongly associated SNPs in independent genomic loci shared between SCZ and vitD at conjFDR < 0.05. Chromosomal position (Chr), nearest gene, and functional category, as well as P values and effect sizes (Odds ratio and Beta for SCZ and Vitamin D, respectively) from original summary statistics. The effect allele is A1. Novel for SCZ marked with *.

CADD scores above 12.37, 5 had opposite effects on SCZ-risk and vitD levels. Moreover, 4 SNPs among those 72 lead SNPs shared between SCZ and vitD (conjFDR) displayed low RDB score (1b and 2b), suggestive of regulatory functionality.⁶² Two shared loci were located within genomic regions with complex LD, namely the extended MHC region (lead SNP rs9273493, 6:32628247-32501395) and chromosomal region 8p23.1 (lead SNP rs6999466, 8:9390529-10283748). Given the effect of complex LD,

these loci are evidence of the involvement of the MHC and 8p23.1 regions in the studied traits rather than of any specific gene.

Functional annotation of all 10556 SNPs in the loci shared between SCZ and vitD (conjFDR < 0.10) displayed that 84.7% of them were intronic or intergenic, while 1.3% were exonic. Having excluded regions with complex LD (MHC and 8p23.1 loci) we applied FUMA to link the 10196 candidate SNPs in the 70 shared loci

Table 3. Novel for Vitamin D Levels Loci—the GWAS Catalog Traits Associated with Vitamin D Metabolism

Locus	Lead SNP	Genomic position	Previously reported to GWAS catalog trait(s)
17	rs2241823	3: 63862842-64003983	Glomerular filtration rate ⁵⁴
19	rs6779146	3: 135798658-136507818	Body mass index, ⁵⁰ Liver enzyme levels, ⁵¹ Waist-hip ratio ⁵²
25	rs904612	5: 139053438-139095473	Fat-free mass ⁵⁵
45	rs7948931	11: 82987208-83335764	Oily fish consumption ⁵⁶
49	rs7313797	12: 109849297-110042348	Oily fish consumption, ⁵⁶ Total cholesterol levels ⁵⁷
50	rs2464196	12: 121384495-121471337	Gamma glutamyl transferase levels, ⁵⁸ Total cholesterol levels ⁵⁸
51	rs79478560	12: 124388511-124496316	Waist-hip index, ⁵⁹ Body mass index ⁶⁰
53	rs17522122	14: 33292743-33309495	Body mass index ⁵²
57	rs12905223	15: 84672181-85280792	Waist circumference adjusted for body mass index ⁵⁰
62	rs78004870	16: 89736352-89886392	Low tan response ⁵³
64	rs11263770	17: 34825861-34952964	Body mass index ⁵²

Note: Genomic position: Chromosome: lead SNP pair base lower threshold position—upper threshold position.

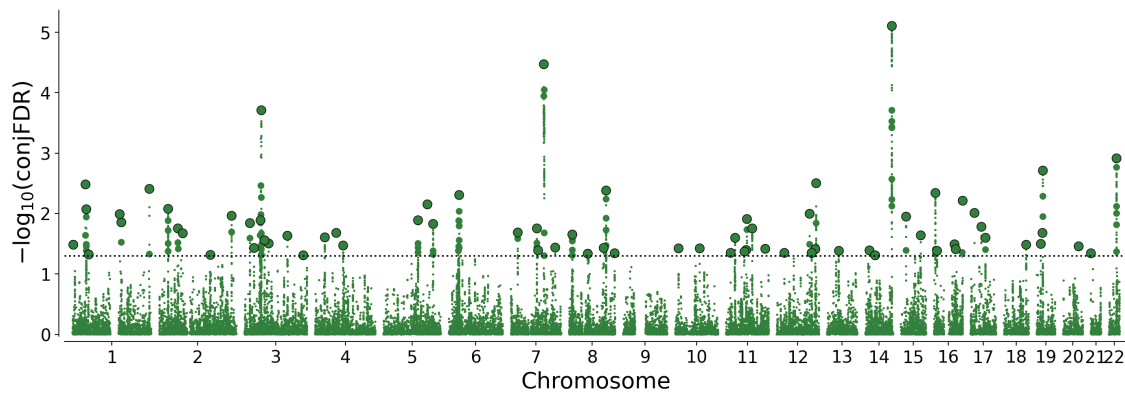


Fig. 2. “ConjFDR Manhattan plot” showing common genetic variants jointly associated with schizophrenia and vitamin D levels at conjunctional false discovery rate (conjFDR) < 0.05. The y-axis shows the $-\log_{10}$ transformed conjFDR. Chromosomal position is presented along the x-axis. The threshold for significant shared associations (conjFDR < 0.05) is represented by the horizontal dotted line. Independent SNPs are indicated by a black perimeter.

to 858 genes. Positional mapping aligned the SNPs to 627 genes while cis-eQTL mapping implicated 528 genes. Two hundred and 97 genes were implicated by both gene-mapping procedures. Seven of those genes are located within newly identified loci, specifically *TM4SF20* and *AGFG1* at chromosome 2, *C4orf36*, *RP11-397E7.4*, and *AFF1* at chromosome 4, *UBXN2B* and *EMC2* at chromosome 8.

Gene Ontology gene-set analysis for genes indicated by both positional mapping and eQTL indicated 2 significantly associated processes, namely, “hyaluronan metabolic process” and “synapse organization.” Further, the genes were significantly associated with “platelet dense granule lumen” cellular components and with 3 molecular functions gene sets, namely, “adenyl nucleotide binding,” “ribonucleotide binding,” and “drug binding” (Supplementary table 7).

Discussion

In the present study, we found that the genetic architecture of SCZ consists of approximately 25 times more

variants than vitD levels, probably reflecting a more complex genetic etiology of SCZ.³¹ The SNPs associated with SCZ and vitD levels are mutually enriched (QQ plots) demonstrating shared genetic architecture. We found a total of 72 loci jointly associated with SCZ and vitD level, 9 novel for SCZ, and 40 novel for vitD (conjFDR). Of the 72 loci shared between SCZ and vitD, 64% had discordant effects on SCZ and vitD levels. Further, functional analyses of the novel shared loci identified 2 genes (DGKG, EMC2), suggesting links between lipid metabolism, involved in 25OHD pathways and brain development. Together, the findings provide a better understanding of the overlapping genetic architecture and jointly associated loci between SCZ and vitD regulation beyond genetic correlation. These findings implicate pathophysiological processes of SCZ linked to vitD regulation which may form the basis of preventive measures and identify new targets for drug development.

Our finding highlights the complexity of the genetic relationship between SCZ and vitD levels, consistent with previously reported weak negative genetic correlation between the 2 traits.²⁴ Weak negative correlation is consistent

with genetic overlap if there is a mixture of concordant and discordant effect directions, since genetic correlation is a summary measure of the correlation of effect sizes. The sum of concordant and discordant effect directions therefore cancel each other out, resulting in a decrease in the absolute value of the genetic correlation estimate.⁶³ The observed distribution of directional effects among the shared loci supports epidemiological findings that suggest co-occurrence of low vitD levels with SCZ.^{2,64} Moreover, in connection with previous findings, this may be consistent with subpopulations of patients with SCZ in which vitD levels might frame clinical manifestation of psychosis and CVD comorbidity.^{65,66} Consequently, even though a Mendelian randomization analysis did not support a causal role for vitD in SCZ, it might be beneficial for future studies to assess specific subgroups of patients with SCZ with a high SNP concordance/discordance rate within genes implicated in both traits.²⁴

Further, our study supports the notion that genetic pleiotropy, ie, the association of a single gene or variant with more than 1 distinct phenotype plays a role in a range of human phenotypes. Extensive pleiotropy has been shown across brain-related traits and disorders, such as mental and cognitive traits, personality, and mental disorders.⁴⁴ Similarly, polygenic pleiotropy between SCZ and somatic traits including CVD risk factors has also been reported.^{26,31} Here, we provide evidence of pleiotropy between SCZ and vitD levels, suggesting novel molecular pathways related to SCZ. Of particular interest, the novel loci shared between SCZ and vitD levels were also found at or in proximity to the genes *DGKG*, *EMC2*, and *KCNGL*.

DGKG encodes diacylglycerol kinase γ which is a member of the type I subfamily of diacylglycerol kinases, known to phosphorylate diacylglycerol in the production of phosphatidic acid.⁶⁷ Phosphatidic acid, a product of the reaction catalyzed by diacylglycerol kinases, plays an important role in cell proliferation and survival, the processes that have been linked to both vitD deficiency in a developing brain and SCZ.^{2,68} Interestingly, both diacylglycerol and vitD may directly impact the activation of Ca^{2+} -permeable cation channels in neurons.^{14,69} Moreover, diacylglycerol kinase γ may influence the synthesis of phosphatidylinositol that has been linked to the pathogenesis of SCZ and other neurodevelopmental disorders, eg, autism spectrum disorders (please note that diacylglycerol kinases ϵ is the predominant enzyme of the phosphatidylinositol cycle).^{68,70,71} Diacylglycerol kinases provide precursors for the synthesis of cardiolipins, and an increased activity of anti-cardiolipin antibody accompanied by a higher activity of MMP9 has been detected in serum from patients with SCZ.^{72,73} *Dgkg* has been shown to be highly expressed in GABAergic interneurons (especially somatostatin expressing) and to regulate neurite outgrowth (maturation) of GABAergic interneurons in mice.⁷⁴ This is of particular interest in the context of the

hypothesized role of inhibitory local circuit neurons in the pathogenesis of SCZ.^{75,76} Diacylglycerol kinases may impact on vitD serum level through inhibiting lipid droplet formation by removing the substrate and lipid signal that triggers this process.⁶⁸ These processes may play an important role in vitD homeostasis since vitD receptors and key vitD enzymes (1α -hydroxylase and CYP 24) are accumulated within the membrane of lipid droplets.⁷⁷

EMC2 encodes one of the cytosolic subunits of the endoplasmic reticulum (ER) membrane protein complex, participating in the insertion and folding of protein into the ER,⁷⁸ and has been linked to SCZ^{79,80} (also DISC1^{81–84}). Misfolded ER proteins can induce ER stress—a cell death pathway.⁸⁵ A recent study emphasized a protective role of vitD against ER stress in hippocampal and cortical neurons.⁸⁶ Previous studies indicated that vitD might regulate lipid metabolism by inducing degradation of SREBP/SCAP complex located in ER.⁸⁷ ER stress mediates the effect of vitD on intestinal epithelial cells in the small intestine (the main site of vitD absorption) and may play an important role in the vitD level feedback loop.⁸⁸ *KCNGL* encodes potassium voltage-gated channel modifier subfamily G member 1. Expression of this ion channel has been associated with regions of the brain dedicated to cognitive functions, emotions, biological needs, and internally driven rhythms (eg, circadian rhythm).⁸⁹

Two of the candidate SNPs among the 9 novel SCZ loci jointly associated with vitD levels, rs4643531 (2:228243905) and rs79621605 (14:50655307) were nonsynonymous exonic variants within *TM4SF20* and *SOS2*, respectively. Nonsynonymous exonic variants are of particular interest as they potentially disrupt protein function by changing the amino acid sequence of the protein. *TM4SF20* encodes a 4-transmembrane protein, and single-exon deletion of *TM4SF20* has been associated with language delay and white matter hyperintensities.⁹⁰ *SOS2* encodes a protein involved in signal transduction pathways. Two SNPs have been associated with late-onset Alzheimer's disease.⁹¹ Moreover, variants in *SOS2* have been identified as a very rare cause of Noonan syndrome, in which intellectual/learning disabilities and skin abnormalities might be expressed among others symptoms.⁹² A functional rare variant in *SOS2* has also been associated with estimated glomerular filtration rate,⁹³ while a study in zebrafish *Sos2*-knockdowns have shown changes in renal development and function⁹³ that in consequence might modify vitD metabolisms.

The novel SCZ-risk loci positionally mapped to *AFF1*, *UBXN2B*, *CDKLI*, and *UPF1* were also found to be shared between SCZ and vitD levels. *AFF1* encodes AF4/FMR2 family, member 1 (AFF-1). AFF-1 protein has been demonstrated to play a role in dendrite reconnection following injury and to maintain regenerative potential during aging in *C. elegans* model.⁹⁴ Prior GWASs have associated a different SNP at *AFF-1* with risk of

Alzheimer's disease⁹⁵ and systemic lupus erythematosus.⁹⁶ *UBXN2B* encodes UBX domain protein 2B, an adaptor protein involved in mitotic spindle orientation, Golgi, and ER biogenesis. Further, a SNP at *UBXN2B* has been associated with the age of onset of excessive daytime sleepiness in narcolepsy.⁴⁹ *CDKL1* encodes cyclin-dependent kinase like 1 (CDKL1). In vivo studies in rats have shown that this protein is mainly expressed in glial cells, is upregulated in gliosis, and is probably involved in hyperphosphorylation of tau protein in Alzheimer's diseases.⁹⁷ *UPF1* encodes UPF1 RNA helicase and ATPase (UPF1) a part of multiprotein complex controlling mRNA nuclear export and surveillance. Which seems particularly interesting in view of a recent study assessing the role of RNA-binding protein (including UPF1) in the pathogenesis of SCZ.⁹⁸

There were limitations to the present study. The conjFDR analysis, applied in our study, identifies loci with a statistical pleiotropy, and further in-depth experimental analyses are required to investigate if detected signals directly influence studied phenotypes and which causal mechanisms are involved.^{43,44} Another limitation of our study is that both GWASs applied in the current study has been performed using European ancestry population.

In conclusion, the current study demonstrates extensive genetic overlap between SCZ and vitD level beyond genetic correlations and identifies several novel polygenic effects. The current findings provide insights into potential mechanistic relationships underlying the epidemiological and clinical associations between these 2 phenotypes. Interestingly, the novel loci shared between SCZ and vitD levels were found in the proximity of genes (*DGKG*, *EMC2*) which are connected to pathogenic neurodevelopmental processes, and to autism spectrum disorders.^{99,100} Moreover, our results suggest that vitD may influence the resilience of the developing brain to adverse factors. This is a potential direction for future animal studies and research in large birth cohorts.

Supplementary Material

Supplementary material is available at <https://academic.oup.com/schizophreniabulletin/>

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