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Identification of de novo mutations in prenatal neurodevelopment-associated genes in schizophrenia in two Han Chinese patient-sibling family-based cohorts

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Abstract
Schizophrenia (SCZ) is a severe psychiatric disorder with a strong genetic component. High heritability of SCZ suggests a major role for transmitted genetic variants. Furthermore, SCZ is also associated with a marked reduction in fecundity, leading to the hypothesis that alleles with large effects on risk might often occur de novo. In this study, we conducted whole-genome sequencing for 23 families from two cohorts with unaffected siblings and parents. Two nonsense de novo mutations (DNMs) in GJC1 and HIST1H2AD were identified in SCZ patients. Ten genes (DPYSL2, NBPF1, SDK1, ZNF595, ZNF718, GCNT2, SNX9, AACS, KCNQ1, and MSI2) were found to carry more DNMs in SCZ patients than their unaffected siblings by burden test. Expression analyses indicated that these DNM implicated genes showed significantly higher expression in prefrontal cortex in prenatal stage. The DNM in the GJC1 gene is highly likely a loss function mutation (pLI = 0.94), leading to the dysregulation of ion channel in the glutamatergic excitatory neurons. Analysis of rare variants in independent exome sequencing dataset indicates that GJC1 has significantly more rare variants in SCZ patients than in unaffected controls. Data from genome-wide association studies suggested that common variants in the GJC1 gene may be associated with SCZ and SCZ-related traits. Genes co-expressed with GJC1 are involved in SCZ, SCZ-associated pathways, and drug targets. These evidences suggest that GJC1 may be a risk gene for SCZ and its function may be involved in prenatal and early neurodevelopment, a vulnerable period for developmental disorders such as SCZ.

Introduction
Schizophrenia (SCZ) is a severe psychiatric disorder that profoundly affects cognitive, behavior, and emotional processes, yet its etiology and pathophysiology are still largely unknown. The high heritability of SCZ suggests that genetic risk factors contribute to a significant proportion of the etiology. However, the marked reduction in fecundity in SCZ patients suggests the removal of risk variants with the largest effects from the population by natural selection. Thus these variants often occur de novo. Indeed, the strongest genetic risk factors for SCZ
identified so far are de novo large copy number variants (CNV)³.

The availability of next-generation sequencing permits the detection of de novo mutation (DNM) events at the genome level. Through large-scale sequencing in parent–offspring trios, DNM have been increasingly discovered from an array of severe neurodevelopmental disorders, including Autism Spectrum Disorders (ASD)⁴, Attention-Deficit Hyperactivity Disorder (ADHD)⁵, and epileptic encephalopathy⁶. For SCZ, Xu et al. showed a large excess of de novo non-synonymous changes and DNM presented with greater potential to affect protein structures and functions in SCZ patients;⁷ Girard et al. reported increased exonic DNM rate in SCZ patients;⁸ and Fromer et al. found that DNM in SCZ implicated synaptic networks and DNM-affected genes in SCZ overlapped with those mutated in other neurodevelopmental disorders⁹. Whole exome sequencing (WES) was applied predominantly in DNM identification thus far, however, few DNM studies have been based on whole-genome sequencing (WGS) as WES remains a cost-effective strategy.

However, only 1.22% of DNMNs are within exonic regions¹⁰ and meaningful mutations may occur outside of exons, such as in regulatory elements (i.e., transcriptional promoters, enhancers, and suppressors) thereby altering expression level of governed genes. Similarly, mutations within exon-intron junction regions may influence splice sites and thus lead to inappropriate expression of particular isoforms¹¹. Takata et al. reported that cis-acting splicing quantitative trait loci from prefrontal cortices of human were in linkage disequilibrium with SCZ genome-wide association study (GWAS) loci and DNMNs near splicing sites were enriched in SCZ patients as compared to controls¹²,¹³. Emerging roles of non-coding RNAs such as microRNA, circRNA, and lncRNA further call for attention to explore DNMNs in non-coding regions.

We therefore, conducted WGS on two cohorts of Chinese families with matched patient-sibling to capture all classes of DNMNs and to more fully describe the genetic architecture of SCZ. This study was conducted to identify potential DNMNs in SCZ in the Asian population, adding to the growing body of information regarding race-specific DNMNs.

**Methods**

**Subjects**

Subjects were drawn from two distinct cohorts, Taiwan and Shanghai, of Han Chinese origin. Recruitment of subjects from the Taiwan cohort was described in previous publications¹⁴,¹⁵. The rationale for combining the two cohorts was to increase the power to detect DNMNs because each of the two cohorts has small sample size but they are ethnically homogeneous. Briefly, families with at least three siblings, two of whom were diagnosed with SCZ, were recruited in the Taiwan Schizophrenia Linkage Study (TSLS) from 1998 to 2002. All recruited subjects were interviewed using the Diagnostic Interview for Genetic Studies (DIGS)¹⁶, accompanied by the Family Diagnostic Interview for Genetic Studies (FIGS) (https://www.nimhgenetics.org/resources/clinical-instruments/figs/list-of-figs). Final diagnostic assessment was based on the criteria of the fourth edition of the Diagnostic and Statistical Manual (DSM-IV), joined with the record of DIGS, FIGS, interviewer notes, and hospital anamnesis. For the Shanghai cohort, families from the Bio-x SCZ Biobank with at least three siblings, two of whom were diagnosed with SCZ, were selected. All families from the Bio-x SCZ Biobank were recruited from the city of Shanghai and the provinces of Hebei, Liaoning and Guangxi from 2001 to 2003. All individuals with SCZ were interviewed by two independent psychiatrists and diagnosed according to DSM-IV criteria. A total of 23 families (10 from Taiwan and 13 from Shanghai), with SCZ patients, unaffected siblings, and parents, were used in this study (Fig. 1). For detailed demographic...
characteristics of all individuals in the 23 families, please refer to Supplementary Table S1. All subjects gave written informed consent with the approval of the local research ethics committees.

Whole genome sequencing

For the subjects of the Taiwan cohort, whole blood samples were collected with anticoagulant (EDTA) tubes and sent to the National Institute of Mental Health (NIMH) Repository and Genomics Resource (RGR). Lymphocytes from the whole blood samples were transformed into immortalized lymphoblastoid cell lines and stored. The DNA samples extracted from the cell lines were used for WGS. WGS was carried out on the Illumina HiSeq 2000 platform using paired-end chemistry with 75 base-pair read length through NovoGene, Inc. (Beijing, China). For detailed description please refer to previous publication. WGS data for the subjects of Taiwan cohort can be accessed in BioProject of NCBI: https://www.ncbi.nlm.nih.gov/bioproject/PRJNA551447. For the subjects from Shanghai cohort, whole blood samples were also collected with anticoagulant tubes. DNA was extracted from blood lymphocytes by standard procedures using FlexiGene DNA kits (Fuji, Tokyo, Japan). DNA libraries were prepared using protocols recommended by Illumina (Illumina, San Diego, CA). WGS was performed on Illumina HiSeq X Ten platform with 150 base-pair read length through Cloud Health Genomics Ltd. (Shanghai, China).

Quality control and variant calling

FastQC (v0.11.8) was used to perform quality checks on all samples across Taiwan and Shanghai cohorts (Supplementary Figure S1).

The GATK best practices of variant calling were applied to process all raw reads from both Taiwan and Shanghai cohorts. Raw sequencing reads in FASTQ format were aligned to the GRCh37 build of the human reference genome with BWA-mem. Then the aligned reads in BAM format were sorted, indexed, and marked with duplicate reads with Picard Tools. Reads containing indels were realigned with GATK's IndelRealigner tool. Next, GATK was used to perform Base Quality Score Recalibration (BQSR). Quality control after alignment was performed using Picard (v2.20.4) CollectAlignmentSummaryMetrics (Supplementary Table S2). Variant calling was performed across all samples with GATK (v4.1.1.0) HaplotypeCaller. For detailed variant calling pipeline, please refer to Supplementary Fig. S2. To further systematically remove potentially false positively called variants, we applied the following quality filters for each variant: (1) the quality score normalized by allele depth is <2; (2) the root mean square of the mapping quality is <40; (3) the strand bias is more than 60; (4) three consecutive variants were clustered within 10 bases; and (5) the FILTER tag is not PASS.

Kinship analysis, DNM calling, and annotation

We used PLINK to perform kinship analysis. Briefly, for a given family, if the father was shown to be within the third degree relative to the mother, the family would be excluded; if the child was not shown to be the first degree relative to the parents, the child would be excluded.

To ensure the DNM calls with high confidence, three tools, GATK PhyseByTransmission (PBT), TrioDeNovo (v0.6.6) and DeNovoGear (v.develop), were used to evaluate the calls. Only DNMs called by all three tools consistently were considered as candidate DNMs.

PBT was run with the following command:

```java
java -jar GenomeAnalysisTK.jar -T PhaseByTransmission -R human_gIk_v37_decoy.fasta -V parent_child_trio.vcf -prior 1.0e-8 -out PBT_result.vcf
```

TrioDeNovo was run with the following command:

```bash
triodenovo --ped parent_child_pedigree.ped --in vcf parent_child_trio.vcf --mu 1.0e-8 --out vcf TrioDeNovo_result.vcf
```

DeNovoGear was run with the following command:

```bash
dngen dnm --ped parent_child_pedigree.ped --in vcf parent_child_trio.vcf --mu 1.0e-8 --write DeNovoGear_result.vcf
```

After obtaining consistently called DNMs, the following five criteria were applied to retain high quality calls tagged with PASS in the FILTER annotation: (1) quality score is greater than or equal to 30; (2) genotypes of the parents are homozygotes; (3) genotype of the child is heterozygote; (4) phred-scaled maximum likelihood of heterozygote for the parents is less than 50; and (5) phred-scaled maximum likelihood of heterozygote of the child is 0. DNMs with MAF greater than 0.01 in East Asian populations were further excluded by querying gnomAD (https://gnomad.broadinstitute.org/) and Exac (http://exac.broadinstitute.org/). Please refer to Supplementary Fig. S2 for detailed pipeline of DNM calling and Supplementary Fig. S3 for the distributions of DNM quality scores. Remaining DNMs were annotated using ANNOVAR.

Polymerase chain reaction (PCR)-based Sanger sequencing validation

For the families where candidate DNMs were found, DNA from all members of the family was subjected to PCR-based Sanger sequencing by capillary electrophoresis according to standard molecular biology practices (ABI 3130 genetic analyzer, ThermoFisher Scientific). Primer3Plus was used to design the PCR primers. For the G/C1 mutation, the forward primer sequence was 5′- TTAGGTGGGGTTGGCCTG-3′ and the reverse primer sequence was 5′- CACCGTGAGACGACAA GAA -3′. For the HIST1H2AD insertion, the forward
primer sequence was 5′-CTCGTTTACTTGCCC TTGG -3′ and the reverse primer sequence was 5′-ACAACAAGAAGACCCGCATC -3′.

Reactions were performed on an Eppendorf MasterCycler (Eppendorf North America, New York, USA) under the following cycling conditions: denaturation at 95 °C for 3 min, 35 cycles of 95 °C for 15 s, 55 °C for 20 s, 72 °C for 30 s, and a final extension at 72 °C for 5 min. Sanger sequencing data was then analyzed using Chromas software (https://technelysium.com.au/wp/).

Identification of genes implicated by coding DNMs in SCZ

As the spontaneous background mutation rates vary greatly between genes, those carrying relatively more protein-coding DNMs in cases are not necessarily implicated by DNMs. To identify those genes implicated by protein-coding DNMs or of which the DNMs occurred higher than the background mutation rates, the R package denovolyzeR was used to analyze protein-coding DNMs based on a mutation model developed previously. Briefly, denovolyzeR estimates underlying mutation rate based on trinucleotide context and incorporates exome depth and divergence adjustments based on macaque-human comparisons over a ± 1-Mb window and accommodates known mutational biases, such as CpG hotspots. By applying the underlying mutation rate estimates, denovolyzeR generates prior probabilities for observing a specific number and class of mutations (synonymous, missense, nonsense, splice-site and frameshift) for a given gene.

To validate whether the genes implicated by coding DNMs were also implicated by rare variants, the population-based SCZ Swedish case-control cohort with whole exome sequences and variants called were used to perform the analysis. Variants with MAF >0.01 were excluded and only rare variants were retained subsequently. The association between the set of rare variants from Swedish case-control cohort located in the exons of a given gene implicated by coding DNM and phenotype was tested using SKAT. For a given gene, SKAT takes the rare variants across the gene as input and compute if the rare variants were more likely to be burdened in cases than in controls.

DNM burden test

To determine whether some genes carry more DNMs in SCZ patients than expected by chance, we performed the DNM burden test for each gene potentially implicated by DNM. Human brain-specific gene enhancer information was included from PsychENCODE (http://resource.psychencode.org/) and human-specific gene promoter information from Eukaryotic Promoter Database (EPD, https://epd.epfl.ch/index.php) for a given gene. DNMs occurring in its gene body, which includes both exons and introns, brain-specific gene enhancer, and gene promoter regions were considered as burdens. For a given gene, we compared the number of DNMs mapped to these regions in SCZ patients with the number of DNMs mapped to these regions in unaffected siblings, and then assessed the significance of the comparison using 10,000 within-sibship case–control label-swapping permutations. P value was calculated as the proportion of permutations with relative risk (RR) as or more extreme than in the observed data.

Developmental expression of DNM-implicated genes

Multiple lines of evidence have shown that prenatal maternal infection, malnutrition, and stress are risk factors for SCZ. The neurodevelopmental model of SCZ posits that a perturbation in early brain development leads to an altered brain developmental trajectory that is sensitive to molecular changes associated with development and environmental experience, consequently converging on the emergence of SCZ in early adulthood. It was hypothesized that DNMs drove dysfunction of genes in early brain development and that this dysfunction confers risks for subsequent SCZ. To determine whether DNM-implicated genes (loss-of-function DNM genes and DNM-burdened genes) were involved in early brain development, human brain developmental expression data from BrainCloud and BrainSpan was evaluated. The expression data of BrainCloud consisted of data derived from the prefrontal cortices of 269 individuals, whereas the expression data of BrainSpan consisted of 42 brain specimens across 13 developmental states in 8–16 brain structures. BrainCloud expression data were examined by comparing the expression of a DNM gene in the prenatal stage to the expression in postnatal stage and BrainSpan expression data was evaluated by leveraging the developmental effect scores curated in a previous publication. A developmental effect score measures the effect of age on expression per gene per brain structure, with a higher developmental effect score of a gene in a given brain structure indicating a stronger involvement of brain development in that structure. By 10,000 gene label-swapping permutation, P value of a DNM gene for a given brain structure was calculated as the proportion of permutations with developmental effect scores as or more extreme than the observed value. DNMs were then further evaluated in brain developmental expression data from other species (macaque and mouse).

Genetic susceptibilities of DNM-implicated genes in SCZ and SCZ-related traits

To determine whether DNM-implicated genes are among the loci found by the GWAS of SCZ or SCZ-related traits, we used GWAS summary statistics from SCZ, ASD, ADHD, bipolar disorder (BD), major
depressive disorder (MDD)\textsuperscript{43}, intelligence\textsuperscript{44}, educational attainment (EA)\textsuperscript{45}, cognitive performance (CP)\textsuperscript{45}, and smoking and drinking\textsuperscript{46}. The criterion for annotating a single nucleotide polymorphism (SNP) to a given gene was that the GWAS SNP is located within 10 kb of the gene boundaries.

**Co-expression and enrichment analysis**

To explore the potential functions or pathological pathways affected by DNM-implicated genes, expression data from BrainCloud was leveraged by retrieving genes highly co-expressed (Pearson correlation coefficient $\beta > 0.8$ or $< -0.8$) with the DNM-implicated gene to construct the co-expression network. To identify enrichments in gene ontologic features, biological pathways, diseases, and drug targets, WebGestalt was used (http://www.webgestalt.org/)\textsuperscript{47}. In WebGestalt, genes co-expressed with the candidate DNM gene were input as the target gene set and all genes in the BrainCloud expression data were input as the reference gene set. Enrichments with B-H FDR-corrected $P$ values $< 0.05$ were considered significantly enriched. To use B-H procedure to control for a desired type I error level $\alpha$, we sorted the $P$ values to find the largest $k$ such that $P_{(k)} \leq k/ma$, where $m$ is the number of hypotheses to be tested. Hypotheses with $P > P_{(k)}$ would be rejected\textsuperscript{48}.

**Cell-type specific expression analysis**

To examine cell-type specific expression of DNM-implicated genes, brain tissue single nucleus RNA-seq (snRNA-seq) data of middle temporal gyrus (MTG), of which the cells have been sub-typed from Allen Brain Atlas (https://celltypes.brain-map.org/rnaseq) and single cell expression data from PsychENCODE were utilized. Single cell expression data from PsychENCODE were merged from multiple brain regions, including frontal cortex, visual cortex and cerebellar hemisphere\textsuperscript{49}. Raw read count data were normalized by log-transformation using R package Seurat\textsuperscript{50}. For a given DNM gene, the dominant cell type(s) with high expression was determined by pair-wise Wilcoxon test.

**Results**

The goal of our study was to identify DNM disturbed genes, of which the dysfunctions can contribute to the pathogenesis of SCZ. We identified coding DNM-implicated genes and genes carrying more DNM burdens in SCZ patients than their unaffected siblings. These DNM-implicated genes were potentially detrimental and subsequent analyses were performed to explore their pathological effects (Fig. 1).

**Identification of DNMs and genes implicated by DNMs**

To ensure parental unrelatedness and that the children are indeed biological offspring of their parents, kinship analysis was performed for individual family. For the Taiwan cohort, one family (Family ID: 35-04560) was excluded as the father is within the third degree relative of the mother, one family (Family ID: 35-93405) was excluded as the father was not found to be related to any of the children and one child in one family was excluded as he was found to be unrelated to his parents (Individual ID: 35-02497-01) (Supplementary Table S1). Three children from Shanghai cohort were excluded as they were unrelated to their respective parents (Individual IDs: CHG000225, CHG000236, and CHG000246) (Supplementary Table S1). A total of 21 families were retained for analyzes. For detailed information of the families recruited and the individuals in each family in the Taiwan and Shanghai cohorts and the families or individuals excluded due to unrelatedness, please refer to Supplementary Table S1.

In this study, 70.71 ± 6.83 de novo point mutations and 6.31 ± 3.64 de novo indel mutations with high confidence per individual were identified. The observed de novo point mutation rate of $1.145 \times 10^{-8}$ was consistent with the neutral expectation of $1.140 \times 10^{-8}$ ($P = 0.95$, two-sided exact binomial test)$^{23}$. The observed de novo indel mutation rate of $1.022 \times 10^{-9}$ was consistent with the neutral expectation of $1.420 \times 10^{-9}$ ($P = 0.500$, two-sided exact binomial test)$^{23}$. All DNMs were checked visually by Integrative Genomics Viewer (IGV). Overall, the DNMs occurred in non-coding regions predominantly in intergenic regions (Supplementary Fig. S4). No obvious difference was observed between unaffected siblings and SCZ patients with regards to the distributions of DNM locations relative to genes (Supplementary Fig. S4). No obvious differences were observed between unaffected siblings and SCZ patients with regards to the percentages of DNMs occurring in exons (unaffected siblings: 29.85% versus SCZ patients: 30.29%) and introns (unaffected siblings: 1.53% versus SCZ patients: 2.03%). The nonsynonymous-to-synonymous ratio in SCZ patients was not found to be different from the ratio in unaffected siblings$^{51}$, which might be attributed to the limited sample size. Two nonsense loss-of-function DNMs implicating $GJC1$ and HIST1H2AD respectively were identified in SCZ patients (Table 1 and Supplementary Table S3). The two loss-of-function DNMs were visually verified by IGV (Supplementary Figs. S5, S6) and then were confirmed by Sanger sequencing (Supplementary Figs. S7, S8). To examine whether the occurrence rates of nonsense loss-of-function DNMs in $GJC1$ and HIST1H2AD in SCZ patients were higher than expected by chance, we compared the nonsense mutation probabilities of $GJC1$ and HIST1H2AD to those of all genes calculated in Samocha et al.$^{52}$. Nonsense DNMs were not prone to occur in DNA regions of $GJC1$ and HIST1H2AD (Supplementary Fig. S9). No loss-of-function DNMs were identified in unaffected siblings on the whole genome scale (Supplementary Table S4).
Except for the two nonsense loss-of-function DNMs, we also investigated other DNMs in coding regions to find damaging DNMs because DNMs occurring in coding regions may change protein structures physically (Fig. 1 and Supplementary Table S4). A detailed list of these DNMs is shown in Supplementary Table S3. To identify genes implicated by these protein-coding DNMs, denovoR was used. Due to the relatively small sample size of this study, Bonferroni correction was used to reduce the falsely identified DNM-implicated genes. Only GJC1 and HIST1H2AD carrying nonsense DNMs were significantly identified to be implicated by DNMs (Supplementary Table S5). We applied another background mutation rate-based model, referred to as the chimpanzee-human divergence model, which also showed GJC1 and HIST1H2AD were significantly implicated by the nonsense DNMs ($P$ value $= 1.56 \times 10^{-5}$ for GJC1 and $P$ value $= 1.82 \times 10^{-5}$ for HIST1H2AD)\(^5\). Of note, the GJC1 nonsense DNM was predicted to be extremely close to the most severe 0.1% of mutations by combined annotation dependent depletion (CADD) (Supplementary Table S3; mutations with CADD phred-like scores $\geq 30$ is the most severe 0.1% of mutations)\(^5\). Moreover, GJC1 falls into the haploinsufficient category with a high probability of loss-of-function intolerance (pLI) of 0.94, therefore is extremely intolerant of loss-of-function variation\(^5\).

To investigate whether GJC1 and HIST1H2AD were also implicated by rare variants as DNMs in SCZ, we analyzed SCZ Swedish case–control cohort comprised of 4969 SCZ patients and 6245 controls. Rare variants in the exons of GJC1 were significantly associated to SCZ (Table 2).

**Table 1** Loss-of-function DNMs identified in schizophrenia patients.

| Individual ID  | Chr | Position (hg19) | Gene  | Reference allele | Mutant allele | Mutation type | Amino acid substitution |
|---------------|-----|----------------|-------|-----------------|---------------|---------------|------------------------|
| 35-50505-02   | 17  | 42,882,819     | GJC1  | G               | A             | Nonsense      | p.Q123X                |
| 35-06277-01   | 6   | 26,199,170     | HIST1H2AD | A            | ACTTTACCCAG  | Nonsense      | p.V101AfsX2             |

Chr chromosome.

**Table 2** Rare variant association tests for the loss-of-function DNMs identified in schizophrenia patients.

| Gene     | Rare variant frequency in SCZ | Rare variant frequency in control | SKAT $P$ value |
|----------|------------------------------|----------------------------------|----------------|
| GJC1     | 0.0115                       | 0.0091                           | $5.60 \times 10^{-3}$ |
| HIST1H2AD| 0.0029                       | 0.0043                           | $6.28 \times 10^{-1}$ |

Identification of genes burdened with DNMs in SCZ patients

Previous studies have shown that 99% of the DNMs occurred in noncoding regions\(^10\). DNMs occurring in regulatory regions can potentially disturb the bindings of transcriptional factors and thus influence gene expression. If a gene or regulatory regions of the gene carry more DNMs in SCZ patients than their unaffected siblings, this gene may have a role in disease predisposition. Based on this assumption, for a given gene, we compared the number of DNMs that occurred in brain-specific enhancer, promoter, and gene body in SCZ patients to the number of DNMs in unaffected siblings by within-sibship case-control label-swapping permutation. Before calculation, to reduce the falsely identified genes by chance, we excluded genes with low total DNM counts in both SCZ patients and unaffected siblings ($\leq 3$). Due to the relatively small sample size, Bonferroni correction was used to reduce the false-positive genes. Ten genes, DPYSL2, NBPF1, SDK1, ZNF595, ZNF718, GCNT2, SNX9, AACS, KCNQ1, and MS12, were identified to carry more DNMs in SCZ patients than their unaffected siblings. Multiple DNMs implicated enhancers for DPYSL2, NBPF1, SNX9, and MS12 (Supplementary Table S6), suggesting the dysregulation of these genes may predispose an individual to SCZ. One single enhancer can regulate the expression of multiple genes. Therefore, single DNM implicating an individual enhancer can disturb the expression of multiple genes. We leveraged transcription factor-enhancer-target gene linkage data from PsychENCODE to establish the gene regulatory network disturbed by the DNMs implicating the enhancers identified by DNM burden test (Fig. 2). For detailed list of the DNMs found in the ten genes reported here, please refer to Supplementary Table S6. Interestingly, no DNM occurred in promoters of the ten genes.

DNMs occurred in genes involving in early brain development

In the following analyzes, coding DNM-implicated genes (GJC1 and HIST1H2AD, Supplementary Table S5) and DNM-burdened genes (Table 3) were as assumed to
be detrimental DNM genes to explore their potential functions that might be related to SCZ.

Early neurodevelopmental events have been implicated in the pathogenesis of SCZ. Genes implicated in SCZ function in processes important to fetal brain development. To determine whether the detrimental DNM genes, including GJC1 and HIST1H2AD implicated by nonsense mutations (Supplementary Table S5) and genes carrying more DNMs in SCZ patients (Table 3), were involved in brain development, we leveraged the developmental expression data of prefrontal cortices (PFC), one of the most highly implicated brain regions in SCZ, from BrainCloud. Nine of the 13 detrimental DNM genes, including the two loss-of-function DNM-implicated genes GJC1 and HIST1H2AD, and seven DNM-burdened genes DPYS2L, NBPF1, SDK1, ZNF595, ZNF718, and SNX9, showed biased higher expression in prenatal stage compared to that of the expression in postnatal stage (Fig. 3). The large proportion of 9/12 potentially detrimental DNM genes with higher expression in prefrontal cortices was more than expected by chance (Supplementary Fig. S10, P = 0.05, hypergeometric test). We next sought to examine whether the involvements of detrimental DNM genes in PFC development were preserved in other brain regions. The genes with available developmental data in BrainSpan were analyzed. It was found that GJC1, SDK1, and GCNT2 were still involved in the developments of other brain regions (Supplementary Fig. S11). To examine whether the involvements of detrimental DNM genes in human brain development were conserved across species, we first queried the brain developmental expression data of macaque, a primate species evolutionarily close to Homo sapiens. Only DNM genes with expression data of macaque were analyzed. GJC1, HIST1H2AD, DPYS2L, SDK1, and MSI2 showed higher expression in prenatal stage than the expression in postnatal stage (Supplementary Fig. S12). We then queried the brain developmental expression data of mouse. Since the developmental data of mouse were limited in postnatal stage, we applied linear regression to examine the change

![Fig. 2 Gene regulatory network disturbed by the DNMs implicating the enhancers identified by DNM burden test. Light green arrow nodes are the transcription factors. The yellow round nodes are the enhancers perturbed by the DNMs. The Orange round nodes are the genes. The edge between transcription factor and enhancer represent the transcription factor can bind to the enhancer without the perturbation of DNM. The edge between enhancer and gene represents the enhancer can enhance the expression of the gene without the perturbation of DNM. Red labeled genes were those identified by DNM burden test.](image)
of DNM gene expression level with time. The expression levels of \textit{GJC1}, \textit{DPYSL2}, \textit{SDK1}, and \textit{AACS} significantly decreased with time (Supplementary Fig. 13), which implied their involvements in prenatal neurodevelopment of mouse. Specifically for \textit{GJC1}, Leung et al. showed that \textit{GJC1} displayed high expression in embryonic stage followed by a massive postnatal decrease in the rat midbrain-floor, where dopaminergic neurons were mostly populated\textsuperscript{57}.

Common variants in detrimental DNM genes may influence risks of SCZ-associated traits

To determine whether the detrimental DNM genes are involved more broadly in SCZ, we asked whether common variants present in the DNM genes confer risk for SCZ and SCZ-associated traits. We extracted all SNPs that occur within 10 kb of the DNM genes. For a given trait, the SNP with the minimum \( P \) value mapped to the gene was used to represent the gene risk on the trait. Genes with minimum SNP \( P \) values less than the suggestive threshold of \( 1 \times 10^{-5} \) were considered as risk

Table 3  Genes significantly enriched with higher DNM burden in schizophrenia patients than unaffected siblings.

| Gene symbol | DNM count in unaffected siblings | DNM count in schizophrenia patients | \( P \) value | Bonferroni \( P \) value |
|-------------|---------------------------------|-------------------------------------|-------------|------------------------|
| DPYSL2      | 0                               | 6                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| NBPF1       | 0                               | 6                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| SDK1        | 1                               | 5                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| ZNF595      | 1                               | 5                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| ZNF718      | 1                               | 5                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| GCNT2       | 0                               | 5                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| SNX9        | 0                               | 5                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| KCNQ1       | 1                               | 4                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| AACS        | 1                               | 3                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |
| MS2         | 1                               | 3                                   | \( 1 \times 10^{-4} \) | \( 3.4 \times 10^{-3} \) |

Fig. 3  Detrimental DNM genes implicated in early brain development in PFC. \( a \) \textit{GJC1}; \( b \) \textit{HIST1H2AD}; \( c \) \textit{DPYSL2}; \( d \) \textit{NBPF1}; \( e \) \textit{SDK1}; \( f \) \textit{ZNF595}; \( g \) \textit{ZNF718}; \( h \) \textit{KCNQ1}; \( i \) \textit{SNX9}. Black dots represent samples from prenatal stage. Blue dots represent samples from postnatal stage. \( P \) values were derived from Wilcoxon rank sum test by comparing gene expression from prenatal stage to postnatal stage.
GJC1 co-expressed with multiple potassium channel genes and is a potential target for SCZ

Next, we focused on the loss-of-function DNM gene GJC1, a member of the connexin gene family, which showed strong evidence of SCZ susceptibility (Supplementary Table S8). DNM s in other brain-associated connexin genes were also found in psychiatric patients (Supplementary Table S9). GJC1 showed higher expression in prenatal stage than the expression in postnatal stage in PFC and multiple other brain regions, and its involvement in early brain development was conserved across species (Fig. 3; Supplementary Figs. S11–S13). Moreover, common variants present in GJC1 confer risks for intelligence, educational attainment, cognitive performance and alcohol abuse, which were associated with SCZ (Supplementary Table S7).

To explore the pathogenic effects of GJC1, we first sought to identify the genes that might be impacted by the dysfunction of GJC1. We performed co-expression analysis with Pearson correlation coefficient $>|0.8|$ or $<-0.8$ in PFC to identify these genes (Fig. 4a and Supplementary Table S10). Then we leveraged the genes co-expressed with GJC1 to perform enrichment analysis to identify functions or pathways involved by GJC1. Interestingly, the genes co-expressed with GJC1 were enriched in SCZ, SCZ drug zuclopenthixol and SCZ-associated functions or pathways, including potassium ion transport (Fig. 4b and Supplementary Table S11). Of note, multiple potassium ion channel genes were negatively co-expressed with GJC1 (Fig. 4a), suggesting their dysfunctions may be subsequent to the dysfunction of GJC1.

To determine whether the expression of GJC1 was specific to a certain cell type, we performed cell type specific analysis of GJC1 expression. GJC1 was predominantly expressed in glutamatergic excitatory neurons (Fig. 4c, d, refer to Supplementary Tables S12, S13 for pair-wise $P$ values among different cell types derived from Wilcoxon rank sum test). This should be interesting as glutamatergic dysfunction has long been implicated in SCZ.

**Discussion**

In this study, we used two matched SCZ-sibling family cohorts of Han Chinese origin to investigate DNMs and DNM-implicated genes in SCZ. By integrating this information with publicly available data, including brain developmental expression profiles and summary GWAS statistics, we identified DNM-implicated genes that were involved in fetal neurodevelopment. These genes may confer risks for SCZ or SCZ-associated traits, and are potential drug therapy targets for SCZ.

Connexin 45 (Cx45) encoded by GJC1 is a component of the gap junction channel. It is one of the two connexin genes expressed predominantly in neurons. Our analyses showed that GJC1 was expressed highly in prenatal stage as compared to that in postnatal stage and was expressed predominantly in glutamatergic excitatory neurons in the human brain (Fig. 4c, d). For the nonsense mutation identified in our study, its occurrence might trigger the activation of surveillance pathway nonsense-mediated mRNA decay, which reduces aberrant proteins to be formed. The reduction of GJC1 may perturb the co-expressed genes and thus influence the potassium ion transport and axon/dendrite formations, especially in early neurodevelopment (Fig. 4a, b). Krüger et al. reported Cx45-deficient embryos exhibited striking abnormalities in vascular development and died between embryonic day 9.5 and 10.5. Kumai et al. reported that Cx45-deficient embryos displayed an endocardial cushion defect in early cardiogenesis and died of heart failure at around embryonic day 9.6. Nishii et al. showed that mice lacking Cx45 conditionally in cardiac myocytes displayed embryonic lethality and the requirement of Cx45 for developing cardiac myocytes. Recently, Homsy et al. reported shared genetic susceptibilities between congenital heart disease and neurodevelopmental disability. These studies indicate the critical role of Cx45 in embryonic development. Our results showed that GJC1 displayed biased high expression across multiple brain regions in prenatal stage, suggesting it is required for neurodevelopment. The high expression of GJC1 in prenatal/early stage in other species indicated its involvement in development is phylogenetically conserved. The dysfunction or loss-of-function can cause lethal effects, thereby such mutation would be eliminated by natural selection in evolution. Interestingly, all potassium ion channel genes were negatively co-expressed with GJC1. Therefore, the reduction of GJC1 indicates an increase or overactivity of potassium channels. Miyake et al. reported overexpression of Ether-a’-go-go potassium channel gene KCNH3 in the forebrain can impair the performances of working memory, reference memory and attention. Ghelardini et al. reported that administrations of potassium channel openers bring out amnesic effect which can be reversed by potassium channel blockers. This evidence suggested that the cognitive or memory deficits in this case were caused by the increased expression of potassium channel genes due to the reduction of GJC1 activity (Fig. 5). The various connexins of gap junctions may be capable of differentiating between the operation
qualities of the cognate synapses defined by the neurotransmitter types. Mitterauer et al. raised a hypothesis that if the function of glial gap junction proteins is lost, the brain is incapable of distinguishing between the same and different qualities of information processing and thus cause severe cognitive impairments in SCZ. Our findings support these hypotheses. Overall, the loss-of-function of Cx45 may be a driver of pathogenesis for some SCZ cases.

**HIST1H2AD**, the other nonsense DNM gene close to HLA locus, encodes the member D in the histone cluster 1 H2A family. Studies have demonstrated that several SCZ candidate genes are especially susceptible to changes in transcriptional activity as a result of histone modification. Therefore, a deficit in the histone itself can implicate multiple SCZ candidate genes. Moreover, epigenetic regulation effects of histone deacetylase inhibitors were potentially suggested to treat SCZ, which implies the fundamental role of histone in the pathology of SCZ.

Gene level DNM burden test identified a list of genes carrying more DNMs in SCZ patients than their unaffected siblings. In the literature, there were experimental or clinical evidences demonstrating their potential association with SCZ. Luan et al. reported SNPs in MSI2 are
strongly associated with SCZ in the Chinese population. In the GWAS of PGC, the SNPs in MSI2 were also associated with SCZ (Supplementary Table 7). Another DNM-burdened gene, DPYSL2, is a member of the collapsin response mediator protein (CRMP) family. CRMP forms homo- and hetero-tetramers and facilitates neuron guidance, growth, and polarity. It also plays a role in synaptic signaling through interactions with calcium channels. Lee et al. revealed that DPYSL2 was downregulated in the PFC and hippocampus of prenatally stressed adult rats that underperformed in behavioral tests. Bruce et al. conducted potassium channel-targeted SNP association analyses with SCZ and SCZ-associated phenotypes. rs8234 in KCNQ1, a DNM-burdened gene in our study, was associated to processing speed. Geschwind et al. found that patients with voltage-gated potassium channel complex antibody (VGKCC-Abs) had particular impairment in memory and executive functions when they evaluated cognitive function and imaging data in patients with VGKCC-Abs associated encephalopathy. Interestingly, the impaired brain functions coincide with the two most implicated brain regions in SCZ, dorsolateral prefrontal cortex and the hippocampus. This evidence demonstrated the efficacy of our DNM burden test by incorporating enhancer and promoter regions into consideration. It also indicated DNMs contributing to the genesis of disease are not limited in coding regions, underlying the value of WGS.

Except for HIST1H2AD, common variants in these identified DNM genes were not strongly associated to psychiatric diseases. However, a few of them, GJC1, HIST1H2AD, and SDK1, were strongly associated to SCZ-associated traits, including intelligence, educational attainment, smoking and drinking. It suggested the significance of these DNM genes in neurodevelopment and that the lethal mutations occurred in these genes were eliminated by marked reduction of fecundity in psychiatric patients.

The results should be interpreted with caution because of limited samples in the pilot study. A few parents with other mental illness (depression, mania, psychosis, or drug use) from the Taiwan cohort may complicate the interpretation of the DNMs if the mental illness shares certain genetic liability with SCZ. Nevertheless, the association of the nonsense DNM gene GJC1 with SCZ is deemed reliable: First, the nonsense DNM in GJC1 is from a family with healthy parents; second, the association of GJC1 with SCZ is supported by the analyzes of rare variants of an independent exome sequencing dataset where SCZ patients have more rare variants in the gene than controls. Second, with its intolerance of loss-of-function index of pLI = 0.94, the loss-of-function mutation is very likely to have detrimental consequences in the regulation of ion channels which have been implicated in the pathogenesis of SCZ. To validate our findings, future studies with more and independent Chinese family samples are necessary.

There are some differences between the samples from Shanghai and Taiwan cohorts. Although the DIGS and FIGS were assessed in the Taiwan cohort, the well-established system for the classification of mental disorders, DSM, was applied to make the final diagnosis for both cohorts. Therefore, the differences in clinical assessments had minimal impact on the diagnosis. With regards to the demographics, both cohorts were of Han Chinese group. For political reason, Taiwan was separated from mainland China and stopped social and economic exchange since 1949, the effect of separation was negligible from evolution perspective. Furthermore, there was evidence that genetic susceptibility of schizophrenia is broadly shared by different ethnic populations. For example, Lam et al. revealed recently that genetic risk for
SCZ is highly correlated between East Asian and European ancestries (genetic correlation = 0.98 ± 0.03)\(^8,9\). The platforms used to sequence the two cohorts were different, however, our highly strict DNM pipeline made DNM calling consistent between the Shanghai and Taiwan cohorts, as all DNMs were assessed by three different software in follow-up analyzes. Overall, these minor differences between the two cohorts should not impact our conclusions.

In summary, we identified a list of DNM-implicated genes which are involved in prenatal neurodevelopment. Common variants in these DNM-implicated genes had been previously reported to be associated with SCZ and related traits in previous GWASs, which is consistent with our analyzes. DNMs implicating the enhancers may contribute to pathogenesis of SCZ by dysregulating the expression of genes. GJC1 is one of these DNM-implicated genes that is primarily expressed in glutamatergic neurons and may be involved in the modulation of ion channel functions, which has also been implicated in SCZ in previous studies\(^8,9,10\). Overall, our study provided new evidence that DNMs have a significant role in SCZ. Further study of these DNM-implicated genes with functional analyzes could lead to better understanding of the pathology of SCZ.

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