

A polygenic burden of rare disruptive mutations in schizophrenia

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Schizophrenia is a common disease with a complex aetiology, probably involving multiple and heterogeneous genetic factors. Here, by analysing the exome sequences of 2,536 schizophrenia cases and 2,543 controls, we demonstrate a polygenic burden primarily arising from rare (less than 1 in 10,000), disruptive mutations distributed across many genes. Particularly enriched gene sets include the voltage–gated calcium ion channel and the signalling complex formed by the activity–regulated cytoskeleton–associated scaffold protein (ARC) of the postsynaptic density, sets previously implicated by genome–wide association and copy–number variation studies. Similar to reports in autism, targets of the fragile X mental retardation protein (FMRP, product of *FMR1*) are enriched for case mutations. No individual gene–based test achieves significance after correction for multiple testing and we do not detect any alleles of moderately low frequency (approximately 0.5 to 1 per cent) and moderately large effect. Taken together, these data suggest that population–based exome sequencing can discover risk alleles and complements established gene–mapping paradigms in neuropsychiatric disease.

Genetic studies of schizophrenia (MIM 181500) have demonstrated a substantial heritability^{1,2} that reflects common and rare alleles at many loci. Genome-wide association studies (GWAS) continue to uncover common single nucleotide polymorphisms (SNPs) at novel loci³. Rare or *de novo* genic deletions and duplications (copy-number variants (CNVs)) have been firmly established, including risk variants at 22q11.2, 15q13.3 and 1q21.1 (refs 4, 5). One notable outcome of these large-scale, genome-wide investigations is the degree of polygenicity, consistent with thousands of genes and non-coding loci harbouring risk alleles^{3,6-9}.

Nonetheless, progress has been made in implicating biological systems and quantifying shared genetics among related psychiatric disorders (for example, refs 10, 11), such as identifying common variants in calcium ion channel genes affecting schizophrenia and bipolar disorder¹² and *de novo* CNVs affecting genes encoding members of the postsynaptic density (PSD) proteome¹³, in particular members of the neuronal ARC protein and *N*-methyl-D-aspartate receptor (NMDAR) postsynaptic signalling complexes.

Here we apply massively parallel short-read sequencing to assay a substantial portion of variation that previously was essentially invisible: rare coding point mutations (single nucleotide variants (SNVs)) and small insertions and deletions (indels). Although previous schizophrenia studies have applied sequencing, the results have been inconclusive, reflecting limited sample sizes or a focus on small numbers of candidate genes^{14–17}. Exome-sequencing studies of *de novo* mutations published to date have neither demonstrated an increased rate in schizophrenia, nor conclusively implicated individual genes^{18,19}, although some data suggest a link with particular classes of gene, such as those with

higher brain expression in early fetal life¹⁹. *De novo* studies in intellectual disability^{20,21} and autism^{22–25} have, however, made considerable progress in identifying large-effect alleles and the underlying gene networks.

In this study, we sought to identify the alleles, genes or gene networks that harbour rare coding variants of moderate or large effect on risk for schizophrenia by exome sequencing 5,079 individuals, selected from a Swedish sample of more than 11,000 individuals. Previous analyses of the full sample (Supplementary Information section 1) have demonstrated an enriched burden of rare CNVs and a polygenic common variant component³. We generated high-coverage exome sequence to ensure sufficient sensitivity to detect and genotype alleles observed in only one heterozygous individual (singletons, implying an allele frequency of \sim 1 in 10,000, although the true population frequency will typically be rarer).

The high baseline rate of rare, neutral mutations makes it difficult to detect rare alleles that increase risk for common diseases²⁶. Although power can be increased by jointly testing groups of variants in a gene²⁷, association testing across all genes is likely to be under-powered at current sample sizes. Indeed, a recent application of population-based exome sequencing in autism did not identify genes²⁸, despite moderately large sample size and the success of the *de novo* paradigm. Furthermore, many confirmed results from candidate-gene sequencing studies of nonpsychiatric disease still fall short of exome-wide significance²⁹.

We therefore adopted a top-down strategy in which we studied a large set of genes with a higher likelihood of having a role in schizophrenia, on the basis of existing genetic evidence (Supplementary Information section 7). We focused on \sim 2,500 genes implicated by unbiased,

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large-scale genome-wide screens, including GWAS, CNV and *de novo* SNV studies, testing for enrichment of rare alleles in cases. To prioritize individual genes, we characterized emerging signals with respect to the genes and frequency and type of mutations. We coordinated analysis with an independent trio exome-sequencing study (Fromer *et al.*³⁰, this issue) and note key points of convergence below.

After alignment and variant calling of all samples jointly, we removed 11 subjects with low-quality data along with likely spurious sites and genotypes (Supplementary Information sections 2 and 3). Per individual, 93% of targeted bases were covered at \geq 10-fold (81% at \geq 30-fold). The final data set comprised 2,536 cases and 2,543 controls (Extended Data Table 1a and Extended Data Fig. 1a). Cases and controls had similar technical sequencing metrics, including total coverage, proportion of deeply covered targets, and overall proportion of non-reference alleles (Extended Data Table 1b). We observed 635,944 coding and splice-site passing variants of which 56% were singletons. Using Sanger sequencing and Exome Chip data on these samples, we determined high specificity and sensitivity for singletons (Supplementary Information section 3).

We annotated variants with respect to RefSeq and combined five in silico algorithms to predict missense deleteriousness (Extended Data Table 1c and Supplementary Information section 4). As expected, allelic types more likely to affect protein function showed greater constraint: 69% of nonsense variants were singletons, compared to 58% of missense and 51% of silent variants. Primary analyses tested (1) disruptive variants (nonsense, essential splice site and frameshifts, n = 15,972 alleles with minor allele frequency (MAF) < 0.1%); (2) disruptive plus missense variants predicted to be damaging by all five algorithms (n = 50,369); and (3) disruptive plus missense variants predicted to be damaging by at least one algorithm (n = 233,575). These groups are labelled disruptive, NS_{strict} and NS_{broad} , in which NS indicates nonsynonymous. We also stratified most analyses by allele frequency: (1) singletons; (2) up to 0.1% (ten or fewer minor alleles); and (3) up to 0.5% (50 or fewer minor alleles). In the main gene set analyses, we empirically corrected for multiple testing over the nine combinations of these factors (Supplementary Information section 7).

The most significant SNV or indel association ($P = 5 \times 10^{-8}$) was for a common missense allele in CCHCR1, in the major histocompatibility complex (MHC), a known risk locus; this top SNP was in linkage disequilibrium with many other schizophrenia-associated SNPs in the MHC. All $P < 10^{-5}$ variants were for either common alleles or a

few instances of likely aberrant variants that had escaped earlier filtering (Supplementary Information section 5). We performed two series of gene-based tests: a one-sided burden test of an increased rare allele rate in cases, and the SNP-set (sequence) kernel association test (SKAT 27), which allows for risk and protective effects. For both tests, the distribution of gene-based statistics broadly followed a global null (Extended Data Fig. 1b).

Considering only disruptive variants, the genic test yielding the lowest nominal P value was for KYNU (kynureninase), showing ten variants in cases and zero in controls (Extended Data Table 2 and Supplementary Table 1); one novel nonsense mutation at chr2:143713804 (g.468T>A; p.Y156*) was observed in seven cases and not present in either the Exome Variant Server (http://evs.gs.washington.edu/EVS/) or 1000 Genomes Project (http://www.1000genomes.org/). Although previous studies have suggested links between the kynurenine pathway and schizophrenia³¹, our P value of 1.7×10^{-3} does not withstand correction for multiple testing, even if considering only the 246 genes with \geq 10 rare disruptive mutations capable of achieving a nominally significant result.

A polygenic burden of rare coding variants

We evaluated a polygenic burden of rare coding variants in cases, first selecting 2,546 genes (~10% of the exome) on the basis of previous genetic studies that we proposed to be enriched for schizophreniaassociated mutations (Supplementary Information section 6). Sources included genome-wide CNV studies^{5,13}, GWAS^{3,12,32} and exome sequencing of de novo mutations 18,19,30. In our sample, these genes had a significantly higher rate of rare (MAF < 0.1%), disruptive mutations in cases compared to controls ($P = 10^{-4}$ for 1,547 versus 1,383 mutations). The enrichment was unlikely to represent technical or ancestry-related artefact because the P values controlled for potential differences in exomewide burden in cases and controls, and because we observed no differences exome wide (P = 0.24). Furthermore, enrichment P values were empirically derived by permuting phenotypes within subgroups of cases and controls, matched on exome-wide identity-by-state, experimental batch and sex; the above result withstood correction for multiple testing (Table 1). We observed similar results for rarer (singletons, $P = 8 \times 10^{-4}$) and more frequent (MAF < 0.5%, $P = 2 \times 10^{-4}$) alleles. We also observed case enrichment for the strictly defined set of damaging mutations (NS_{strict}, $P = 1.5 \times 10^{-3}$) but not the broader set (NS_{broad})

Table 1 | Gene set analysis of primary schizophrenia candidate gene sets

Variant type	Gene set/subset	n genes	Singletons	MAF < 0.1%	MAF < 0.5%
Disruptive NS _{strict} NS _{broad}	Primary	2,546	0.0008 0.0059 0.0986	0.0001 0.0015 0.1295	0.0002 0.0110 0.1126
Disruptive	SCZ de novo genes				
	Exome sequencing (disruptive) ^{18,19,30}	87	0.0319	0.0007	0.0003
	Exome sequencing (nonsyn.) ^{18,19,30}	611	0.0053	0.0011	0.0055
	Copy number variants				
	de novo CNV genes ¹³	234	0.0234	0.0039	0.0124
	SCZ-associated CNV genes ⁵	345	0.3308	0.4596	0.4376
	GWAS				
	Voltage-gated calcium channel genes ¹²	26	0.0019	0.0214	0.0212
	Common SNPs ($P < 10^{-4}$ intervals) ³	479	0.1794	0.0368	0.0037
	miR-137 targets ³²	446	0.6573	0.5609	0.4747
	Synaptic genes				
	PSD (human core) ¹³	685	0.0808	0.1154	0.1256
	ARC ¹³	28	0.0016	0.0014	0.0014
	NMDAR network ¹³	61	0.0158	0.0251	0.0252
	PSD-95 ¹³	65	0.0017	0.0009	0.0010
	mGluR5 ¹³	39	0.1327	0.0900	0.0902

Enrichment test empirical *P* values for rare (singleton; MAF < 0.1%; MAF < 0.5%) variants from disruptive, NS_{strict} and NS_{broad} sets. *P* values represent the relative case enrichment compared to average exomewide case/control difference. Bolded values are significant at *P*^{corrected} < 0.05. Initial comparison corrects (based on the empirical distribution of minimum *P* values) for the nine correlated tests (top panel). The bottom panel focuses on the 12 subsets of the primary gene set, for disruptive variants only as they showed the greatest enrichment for the entire primary set. Again, bold values are significant after correcting for the 36 tests performed.

SCZ, schizophrenia.

This enrichment suggests a polygenic burden of rare variants. Although not so marked as to be detectable at the exome-wide level given the sample size, it is relatively concentrated in genes that were found to be associated with schizophrenia by other methods. The mean allelic effect was not large: in the primary comparison, the odds ratio was 1.12 (1.04-1.20, 95% confidence interval) for each MAF < 0.1% disruptive mutation; 46% of cases carried one or more allele in this primary set (0.62 per case) compared to 41% of controls (0.55 per control). At two extremes, the modest mean effect could represent either that a subset of mutations are fully penetrant or that every allele is associated but increases risk by only 12%, similar to common alleles from GWAS. To extract subsets of potentially stronger-effect alleles, we individually tested the constituent gene sources (Table 1 and Extended Data Fig. 1c), focusing on disruptive variants as they showed the strongest omnibus enrichment. For disruptive mutations, eight out of 12 sets were nominally significant (P < 0.05), indicating that the initial observation was not driven by a single category.

ARC, PSD-95 and calcium ion channel genes

Three of the smaller significantly enriched sets (the ARC and PSD-95 (encoded by DLG4) complexes and calcium ion channel genes) had odds ratios > 5. We observed enrichment ($P = 1.6 \times 10^{-3}$) of disruptive mutations among the 28 ARC complex genes: nine mutations in nine genes (all singletons) in cases and zero in controls, yielding an odds ratio of 19.2 (2.4-2,471, 95% confidence interval; Extended Data Table 2). Along with the NMDAR gene set (also significantly enriched), ARC genes largely accounted for the overall PSD enrichment $(P = 4 \times 10^{-8})$ in ref. 13, in which four ARC genes had one or more de novo CNVs. Of note, in an independent exome-sequencing study in trios, Fromer et al. 30 found that the ARC gene set was enriched $(P = 5 \times 10^{-4})$ for nonsynonymous de novo SNVs and indels, with four genes harbouring six mutations (Extended Data Table 7). The other PSD gene set with strong enrichment $(P = 9 \times 10^{-4})$; odds ratio = 5.1, 1.8–19.2, 95% confidence interval) was the PSD-95 complex, which contains 65 genes and overlaps with ARC. PSD genes are very highly conserved and have critical roles in excitatory neural signalling components, as well as dendrite and spine plasticity. Further categorization of neuronal genes on the basis of subcellular localization¹³ (Extended Data Table 3a) or associated mouse and human phenotypes³³ did not vield further enrichment.

The other subset yielding a large odds ratio of 8.4 (2.03-77, 95% confidence interval) was the 26 voltage-gated calcium ion channel genes (12 cases, one control; disruptive singletons, $P = 2 \times 10^{-3}$, although the effect is attenuated when including recurrent alleles: 15/8 cases/ controls, P = 0.021, see Extended Data Table 2). The singleton enrichment was predominantly driven by the pore-forming α_1 and auxiliary $\alpha_2\delta$ subunits; of the α_1 subunits, the Ca_V1/L-type genes carried the most case mutations, including two in CACNA1C, a gene implicated by GWAS of bipolar disorder and schizophrenia^{3,10}. Calcium signalling is involved in many cell functions including the regulation of gene expression³⁴ and is critical for modulating synaptic plasticity³⁵. In a secondary analysis of proteins found in the nano-environment of the calcium channel³⁶, we observed independent enrichment for other ion channel transporters (Supplementary Table 1), odds ratio 9.1 (2.2–83) for singletons $(P = 1 \times 10^{-3}; 13/1)$ disruptive alleles).

Convergence with de novo studies

A line of convergence across studies was that genes carrying nonsynonymous de novo mutations 18,19,30 were enriched for rare disruptive mutations in cases ($P = 1 \times 10^{-3}$; Table 1 and Extended Data Table 6a, b). We observed a similar result for the smaller class of genes carrying disruptive *de novo* mutations ($P = 7 \times 10^{-4}$, from 47 genes in our study); these genes included *UFL1* (5/0 disruptive mutations, P = 0.03; 7/0 NS_{strict} , P = 0.008), SYNGAP1 (4/0 NS_{strict} , P = 0.04) and SZT2 (18/9 $NS_{\text{strict}}P = 0.049$). SYNGAP1 (synaptic Ras GTPase activating protein 1) is a component of the NMDAR PSD complex³⁷ and mutations in this gene are known to cause intellectual disability and autism³⁸.

Genes under previously associated CNV regions did not show significant enrichment of rare disruptive mutations, although there was an enrichment of NS_{strict} mutations (P = 0.0044; Extended Data Table 4). Of the 11 CNV regions, only the 3q29 locus, which contains multiple genes including *DLG1* (ref. 4), was significant (P = 0.0006) and withstood correction for multiple testing.

Autism/intellectual disability genes and FMRP targets

We next tested, as a single set, the 2,507 genes representing autism and intellectual disability candidates (Supplementary Information section 6), which yielded only nominal significance (P < 0.05) for disruptive and NS_{strict} variants and no test survived correction for multiple testing (Table 2). Considering the 12 constituent sets, genes from autism de novo

Variant type	Gene set/subset	n genes	Singletons	MAF < 0.1%	MAF < 0.5%
Disruptive	Autism/ID	2,507	0.029	0.043	0.049
VS _{strict}			0.052	0.008	0.013
NS _{broad}			0.532	0.619	0.287
		n genes	Min. P°	orrected (for $9 \times 12 =$	108 tests)
Disruptive, NS _{strict} and NS _{broad}	De novo genes (exome sequencing)				
	Autism (disruptive) 22-25	128		1.000	
	Autism (nonsyn.) 22-25	743		1.000	
	ID (disruptive) ^{20,21}	30		0.070	
	ID (nonsyn.) 20,21	132		0.995	
	Neurodevelopmental candidates				
	ASD candidates ³⁹	112		1.000	
	ID candidates ³⁹	196		1.000	
	Autism PPI networks				
	CHD8 network ²⁴	6		1.000	
	49-gene network ²⁴	49		1.000	
	74-gene network ²⁴	74		1.000	
	Fragile X mental retardation protein targets				
	Darnell targets ⁴⁰	788		0.010	
	Ascano targets ⁴²	939		0.997	
	Ascano FMRP/autism overlap ⁴²	93		0.993	

Enrichment test empirical Pvalues for the secondary (autism/intellectual disability) gene set. As in Table 1, the top panel shows uncorrected Pvalues; tests significant after multiple test correction are in bold (that ted > 0.05). Because no class of variant is significant after multiple test correction for the omnibus test (top panel), we applied and corrected for all 108 tests (nine conditions by 12 subsets) in the bottom panel. The single-category FMRP targets (Darnell et al. 40) mainly reflect disruptive and NS_{strict} singleton enrichment. ASD, autism spectrum disorder; ID, intellectual disability; PPI, protein-protein interaction.

studies showed no enrichment (Extended Data Fig. 1c), despite greater sample size and number of disruptive *de novo* mutations. There was no evidence for autism or intellectual disability genes curated from the literature³⁹ or for genes in the protein–protein-interaction-derived subnetworks built around autism *de novo* mutations²⁴.

The nominal omnibus signals arose largely from the Darnell et al. list of FMRP targets⁴⁰. FMRP is encoded by the gene FMR1 (the locus of the Mendelian fragile X syndrome repeat mutation) and is an RNAbinding protein that regulates translation and is needed at synapses for normal glutamate receptor signalling and neurogenesis⁴¹. Targets of FMRP are enriched for *de novo* mutations in autism^{22,40,42}; here we find significant enrichment of disruptive singletons ($P = 1.4 \times 10^{-3}$; 289/223 case/control count; odds ratio = 1.3). These FMRP targets overlap with PSD genes (Extended Data Table 3b), although were still enriched independently (Supplementary Information section 6). In addition, these genes were enriched in GWAS of this sample ($P < 10^{-3}$, Supplementary Information section 9). Whereas the Darnell list is derived from mouse brain, a second recently reported FMRP target list⁴² was generated from cultured human embryonic kidney cells, using a different experimental approach (Supplementary Information section 6). This list has relatively little overlap with Darnell targets and, in contrast to the Darnell list, does not show any enrichment for rare case mutations, for GWAS loci, or comparable overlap with PSD genes (Extended Data Table 3b).

Our results are perhaps surprising: unlike Fromer *et al.*³⁰, we did not observe direct evidence for overlap at the individual gene level with autism and intellectual disability, despite CNV studies showing pleiotropic effects of individual loci. Nonetheless, at the broader level of gene sets, all three disorders showed enrichment for FMRP targets; autism and intellectual disability *de novo* mutations also showed strong enrichment in several PSD complexes enriched in our study, including NMDAR, PSD-95 and (for intellectual disability) ARC. At the least, our results suggest that any overlap is far from complete, although more refined analyses in larger samples will be needed before a clearer picture can emerge of which genes and pathways are shared and which are specific to one disease.

Characterizing enrichment by variant type

To further characterize the observed enrichment with respect to mutational function and frequency, we created a single 'composite' set of 1,796 genes comprising all members of the most prominently enriched sets (Supplementary Table 2). Rare disruptive mutations in this set were present in 990 cases and 877 controls (for singletons, 645–530). Cases carrying rare disruptive mutations did not appear to be phenotypically or clinically unusual in terms of sex, ancestry, history of drug abuse, general medical conditions plausibly aetiologically related to psychosis, or epilepsy, although they did have a higher rate of admissions noting comorbid intellectual disability compared to other cases (P = 0.009; Extended Data Table 2b).

Figure 1 shows composite set enrichment across a range of conditions. As this set merges other sets showing enrichment, it necessarily shows enrichment; it was not, however, due to confounding effects of ancestry, sex or experimental wave (Supplementary Information section 8). It was primarily driven by singleton nonsense mutations across a large number of genes, as it was removed or greatly attenuated when either singleton or nonsense mutations were excluded. Considered alone, neither splice-site, frameshift, missense, silent or noncoding mutations showed enrichment at P < 0.01. Different ways of defining damaging missense mutations did not substantively affect results. Considering only nonsynonymous coding variants present on Exome Chip, we did not observe enrichment. Rather, enrichment mainly reflected novel variants (Extended Data Table 5b), which is expected as most rare variants in our study are novel. We also took an alternative approach, whereby instead of filtering variants on frequency, we excluded genes with any control disruptive variants before calculating the burden of case alleles; the composite set was still highly enriched ('case-unique' in

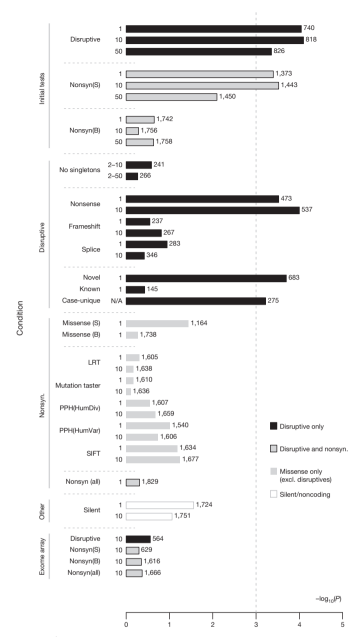


Figure 1 | Composite set gene set analysis, stratified by mutation type. Statistical significance (x axis) for the composite gene set stratified by type and frequency of mutation and other variables. Numbers to the right of each bar represent the number of genes with at least one mutation in that category for the composite set. (S) represents strictly defined damaging missenses; (B) represents the broadly defined group. Nonsyn (all) represents all nonsynonymous mutations. Numbers to the left of the bars (1, 10, 50) represent the minor allele count threshold (i.e. 1 indicates a singleton-only analysis); here the ranges 2-10 and 2-50 represent analyses that excluded singletons; N/A indicates that no allele-wise threshold was used. The source of deleteriousness prediction algorithms (LRT, MutationTaster, PPH and SIFT) is described in the Supplementary Information. For the exome array contrasts, Exome Chip sites were tested using the exome sequence calls.

Fig. 1; see Extended Data Table 5b and Supplementary Information section 7). Finally, the enrichment could not be attributed to only a small number of variants or genes (Extended Data Fig. 2a).

These findings do not preclude potentially important effects from other classes of rare variation in specific genes or other gene sets, although exploratory analyses of generic gene sets (for example, based on Gene Ontology terms) did not unambiguously identify novel signals after correction for multiple testing (Supplementary Information section 7).

We found preferential enrichment in genes with high brain expression, but not for genes with a prenatally biased developmental trajectory (Extended Data Fig. 3). In fact, greater enrichment came from postnatally biased genes. Finally, although greatly attenuated compared to disruptive mutations, other categories displayed nominal (0.01 < P < 0.05) enrichment in Fig. 1 and strictly defined damaging missense mutations alone showed enrichment for ARC and NMDAR gene sets (32/15 for ARC, P=0.007; Extended Data Tables 5a and 7). Although rare coding alleles other than ultra-rare nonsense mutations will undoubtedly contribute to risk, it will probably prove harder still to elucidate such effects.

Rare variants, CNVs and common GWAS variants

We quantified the relative impact of common SNPs (indexed by a genome-wide polygene score from independent GWAS samples³²), rare CNVs (the burden of genic deletions) and disruptive mutations in the composite set. Considering the same 5,079 individuals, all three classes of variation were uncorrelated and significantly, independently and additively enriched in cases compared to controls. From logistic regression, the relative effect sizes (reduction in model R^2) were 5.7%, 0.2% and 0.4% for GWAS, rare CNV and rare coding variants, respectively (Supplementary Information section 8). Although not a complete assessment, it indicates that for the current sets of identifiably enriched alleles, common GWAS variants account for an order-of-magnitude more heritability than this set of rare variants does. However, these estimates will be diluted to varying degrees, owing to associated variants being included. As a consequence of this, and also the fact that true risk variants outside of composite set genes were not considered here, this estimate represents a conservative lower bound on the contribution of rare coding variation.

Discussion

We have demonstrated a polygenic burden that increases risk for schizophrenia, primarily comprising many ultra-rare nonsense mutations distributed across many genes. Implicating individual genes remains challenging, as genes that contributed to the highest-ranked sets typically had unremarkable P values, often around 0.5 with the gene containing only one or two rare mutations. Nonetheless, we were able to detect several small and highly enriched sets, notably of genes related to calcium channels and the postsynaptic ARC complex. Across these ~ 50 genes, $\sim 1\%$ of cases carried a rare disruptive mutation likely to have a considerable impact on risk. However, reported effect sizes will have a tendency to over-estimate true population values (Supplementary Information section 5).

We add to previous work that has implicated disruption of synaptic processes in schizophrenia 13 . The PSD is comprised of supramolecular multiprotein complexes that detect and discriminate patterns of neuronal activity and regulate plasticity processes responsible for learning 43 . Members of the membrane-associated guanylate kinase (MAGUK) family of scaffold proteins, such as PSD-95, have a key role in assembling $\sim\!2$ MDa complexes comprising calcium channels, including the glutamate-gated NMDAR, voltage-gated calcium channels and ARC 36,44 . The genetic disruption of MAGUKs and their associated components result in specific cognitive impairments in mice and humans 45 . One possibility is that the genetic risk identified here reflects altered tuning in calcium-dependent signalling cascades, triggered by NMDAR 46 and L-type calcium channels 47 , mediated by postsynaptic MAGUK signalling complexes driving ARC synthesis.

Although we cannot yet use rare mutations to partition patients into more homogeneous clinical subgroups, this will remain a central goal for future sequencing studies. The few population-based commondisease exome-sequencing studies published to date, in psychiatric (for example, ref. 28) and non-psychiatric (for example, ref. 48) diseases, have not been successful in finding individual genes showing significant enrichment. Our study yields similar findings for individual genes, but yields positive results when considering gene sets. These current findings are likely to foreshadow the definitive identification of individual

genes in larger cohorts, following the trajectory of GWAS and other genetic studies of complex disease.

METHODS SUMMARY

Sample ascertainment. Cases with schizophrenia were identified through the Swedish Hospital Discharge Register³. Case inclusion criteria: ≥ 2 hospitalizations with a discharge diagnosis of schizophrenia, both parents born in Scandinavia, age ≥ 18 years. Case exclusion criteria: hospital register diagnosis of any disorder mitigating a confident diagnosis of schizophrenia. Controls were randomly selected from Swedish population registers. Control inclusion criteria: never hospitalized for schizophrenia or bipolar disorder, both parents born in Scandinavia, age ≥ 18 years. All subjects provided informed consent; institutional human subject committees approved the research.

Sequencing. The samples (2,536 cases, 2,543 controls) were sequenced using either the Agilent Sure-Select Human All Exon Kit (29 Mb, n=132) or the Agilent Sure-Select Human All Exon v.2 Kit (33 Mb). Sequencing was performed by IlluminaGAII or Illumina HiSeq2000. Sequence data were aligned and variants called by the Picard (http://picard.sourceforge.net)/BWA⁴⁹/GATK⁵⁰ pipeline. Validation of selected variants used Sanger sequencing. On the basis of validation and Exome Chip data, we estimated high sensitivity and specificity of singleton calls. BAM and VCF files are available in the dbGaP study phs000473.v1 (http://research.mssm.edu/statgen/sweden).

Analysis. We used PLINK/SEQ (http://www.ncbi.nlm.nih.gov/projects/gap/cgibin/study.cgi?study_id=phs000473.v1.p1) to annotate variants according to RefSeq gene transcripts (UCSC Genome Browser, http://genome.ucsc.edu). Single-site association used Fisher's exact test; primary gene-based association used a burden test and the sequence kernel association test²⁷. Analyses controlled for ancestry and quality control metrics. Gene sets were evaluated on the empirical distribution of the sum of individual gene burden statistics, and incorporated an empirical correction for multiple testing. Odds ratios with 95% confidence intervals used penalized maximum likelihood (Firth's method) for low cell counts. See Supplementary Information for further details. Summary results are posted at http://research.mssm. edu/statgen/sweden/.

Online Content Any additional Methods, Extended Data display items and Source Data are available in the online version of the paper; references unique to these sections appear only in the online paper.

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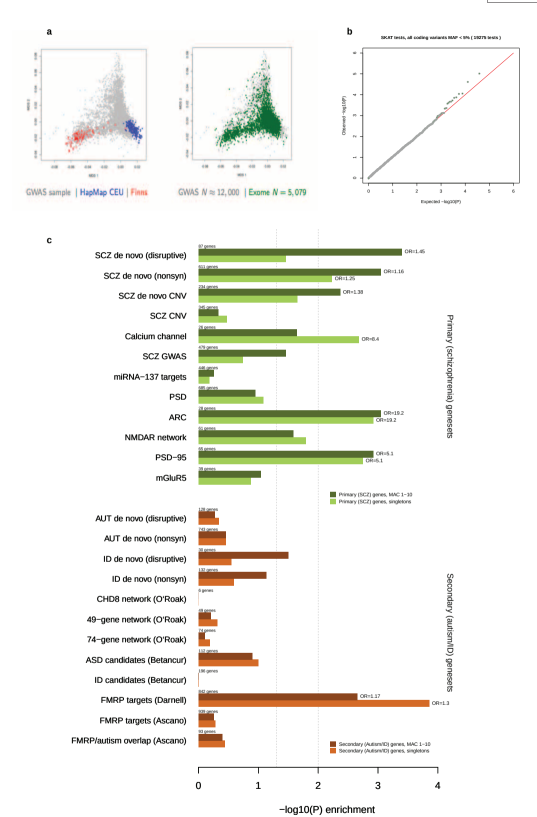
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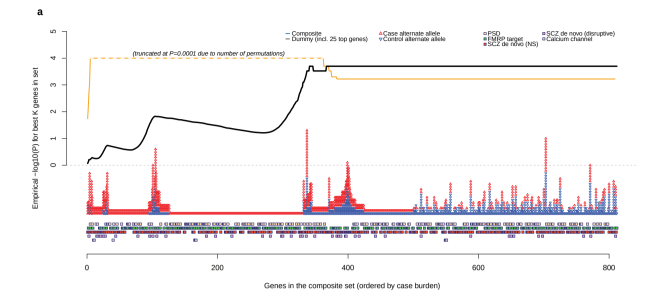
Author Information Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to S.M.P. (shaun@broadinstitute.org)



Extended Data Figure 1 | Ancestry and association summaries.

a, Multidimensional scaling plot of ancestry in the full Swedish GWAS sample, in which each point represents one individual; the left panel superimposes HapMap CEU and Finnish samples and the right panel highlights (in green) the subset of the full Swedish sample for whom we have exome sequence data. **b**, Quantile–quantile plot for gene-based SKAT results (MAF < 5% coding variants). Similar, or more conservative, profiles were obtained for other subsets of variants. **c**, Case enrichment of rare (MAF < 0.1%) and singleton disruptive mutations for the constituent sets of the primary/schizophrenia gene

set (top panel in green) and the secondary (autism/intellectual disability) gene set (bottom panel in orange). The primary set is enriched in cases (MAF < 0.1%; disruptive mutations, $P=10^{-4}$; singletons, $P=8\times10^{-4}$; significant after correction for multiple testing) whereas the autism/intellectual disability set shows only a modest trend (P=0.04 for MAF < 0.1% and P=0.03 for singletons) and is not significant after correction. x axis represents $-\log_{10}(P)$; OR, odds ratio. Number of genes is for total in the set (whether or not they had a rare variant).



		# admissions >0 admissions			nissions	>1 adm	nissions	>5 admissions	
Characteristic of composite set carriers (cases only		OR	Р	OR	Р	OR	Р	OR	Р
Hospital ad	dmissions								
	Duration of hospitalization for SCZ	1.02	0.610	1.02	0.512	1.02	0.625	1.02	0.66
	Total number of admissions	1.00	0.957	1.00	0.968	1.02	0.789	1.02	0.69
	Year of first admission	1.00	0.861	1.00	0.884	1.00	0.895	1.00	0.924
	Year of most recent admission	1.01	0.230	1.01	0.246	1.01	0.235	1.01	0.25
	Drug abuse	1.00	0.401	0.96	0.675	0.90	0.400	0.77	0.12
	General medical condition plausibly								
	etiologically related to psychosis	0.99	0.596	1.12	0.421	0.96	0.831	0.85	0.664
	Epilepsy	1.03	0.494	0.76	0.348	0.81	0.582	1.01	0.99
	Intellectual disability	1.05	0.009	1.41	0.044	1.59	0.019	2.03	0.01
emograp	hics								
	Male	0.89	0.187	0.90	0.199	0.90	0.222	0.90	0.23
	In homogeneous subset	1.11	0.287	1.10	0.323	1.10	0.316	1.11	0.28
	Finnish ancestry	0.95	0.736	0.94	0.684	0.95	0.719	0.96	0.78

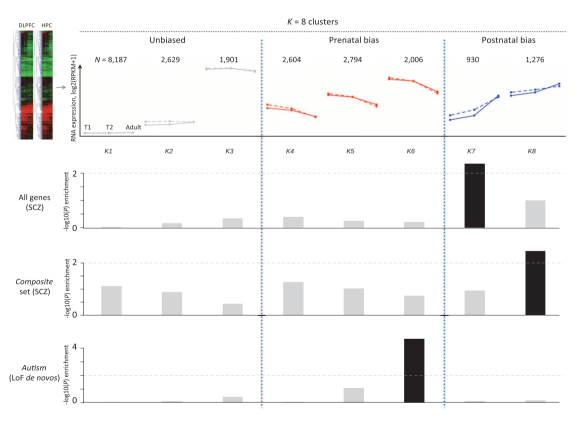
Extended Data Figure 2 | Genic and phenotypic subset analyses for the composite set. a, Individual gene-ranking of composite set genes. Genes are ranked by their case burden of rare disruptive mutations, from left to right, for the composite set. The squares along the bottom indicate to which sets each gene belongs. The red and blue triangles represent case and control counts for each gene. The lines above represent the statistical significance of the best test for this set: that is, the significance of the top K genes, evaluated by permutation. The black line represents results for the real data (disruptive MAF < 0.1%composite set analysis). The orange line represents the dummy condition, in which we artificially constructed a set in which the number of genes, statistical enrichment, odds ratio and case/control counts were similar to the real composite set. However, this set included the 25 top-ranked genes from individual gene-based tests (disruptive MAF < 0.1% variants), with the remainder selected at random. The profile of the best test line is markedly different between the real and dummy gene sets (note: truncated at P = 0.0001reflecting the number of permutations performed). Whereas the dummy P value climbs quickly and then drops to the final aggregate result, the true composite set line continues to climb after 200 genes, indicating that many genes with a single disruptive mutation contribute to the observed set

enrichment (rather than a relatively small proportion of the 1,796 genes accounting for the majority of the signal, as in the dummy set). b, Phenotypic characteristics of cases carrying mutations. Relationship between clinical and demographic measures in schizophrenia cases in relation to carrying one or more composite set disruptive risk alleles (MAF < 0.1%). Hospital Discharge Registry (HDR) data (ICD9 codes) were available on 979 of the 990 case carriers. All P values (uncorrected) are two-sided from a case-only joint logistic regression of carrier status (one or more risk alleles) on all admission and demographic variables including year of first and last admissions. The four pairs of columns represent analyses in which we varied the way in which the HDR admission data were represented (for drug abuse, general medication condition, epilepsy and intellectual disability). # admissions, independent variables are the untransformed number of admissions; > X admissions, independent variable is binary 0/1 variable representing whether individuals had more than X admissions. Of all clinical/demographic measures considered, we observed a nominally significant increased likelihood that cases carrying a disruptive allele in the composite set have increased rates of secondary diagnoses of intellectual disability compared to other cases (based on HDR ICD9 codes).

a

		Bias:	(higher in)			Prenata	al	Postr	natal
BrainSpan : 8-class	Geneset	Average level:	None K1	Low/average K2	High K3	Low K4	Higher K5	Highest K6	Low K7	High K8
BrainSpan . 6-class	n genes		8,187	2,629	1,901	2.604	2.794	2.006	930	1,276
	All genes		0.8653	0.7511	0.8719	0.4672	, -	0.3518	0.1639	0.0005
	Composite		0.1295	0.0447	0.3896		0.1395	0.3310	0.1862	0.0014
	PSD genes		0.1293	0.2103	0.1043		0.4265	0.8340	0.1002	0.001-
	FMRP targets		0.1245	0.2581	0.4230		0.5382	0.1616	0.3797	0.0008
	SCZ de novos (NS)		0.1138	0.1593	0.5044		0.1855	0.2325	0.2541	0.1367
BrainSpan : 4-class	002 de 110103 (110)		None/low		High		natal	0.2020	Postnata	
	n genes		2,748		3,688		540		1,058	
	All genes		0.8504		0.0020		6946		0.0255	
	Composite set		0.0322		0.0017		1535		0.0006	
	PSD		0.3982		0.1043	0.9	9692		0.0008	
	FMRP targets		0.0257		0.0100	0.	1918		0.0131	
	SCZ de novo (NS)		0.0500		0.0145	0.0	0359		0.0504	
Human Brain Transcripto	me :									
Xu et al. (ref. 19) 3-way cl	assifications, HPC			Unbiased		Pre	enatal		Postnata	ı
	n genes			6,578		6,	,841		8289	
	All genes			0.6467		0.7	7281		0.0517	
	Composite set			0.1009		0.2	2876		0.0001	
	PSD			0.3693		0.9	9691		0.1043	
	FMRP targets			0.5629		0.	1264		0.0022	
	SCZ de novo (NS)			0.0397		0.	1395		0.0108	
Human Brain Transcripto										
Xu et al. (ref. 19) 3-way cl	assifications, DLPFC			Unbiased			enatal		Postnata	ı
	n genes			6,255		- /	,669		8,784	
	All genes			0.5076			7538		0.0894	
	Composite set			0.2143			2054		0.0001	
	PSD			0.5949			8329		0.1043	
	FMRP targets			0.3205			4257		0.0005	
	SCZ de novo (NS)			0.1383		0.	1189		0.0043	





Extended Data Figure 3 | Stratified enrichment analysis P values by developmental trajectory of expression in brain (BrainSpan and Human Brain Transcriptome (HBT) data sets). a, Uncorrected Pvalues for a set of exploratory analyses in which we stratified genes in the enrichment analyses by their developmental profile of brain expression. We used four schemes to classify genes as 'brain expressed' and/or 'biased' with respect to prenatal or postnatal expression (see Supplementary Information section 6 for details). We merged data on the hippocampus and dorsolateral prefrontal cortex for the BrainSpan classifications; to mirror the classification of Xu et al.19 we kept separate these two groupings for the HBT data set. Results presented for MAF < 0.1% disruptive variants; similar results were obtained for singletons with the exception that the 'K4' prenatal enrichment signals were no longer significant. In general, the most consistent enrichment across variant classes, classification schemes and brain regions emerges for postnatally biased genes with high brain expression. b, Analysis of exome variants by developmental expression trajectory in human brain. Genes are grouped by

cluster analysis of human postmortem brain expression into eight developmental trajectories, using RNA-sequencing data from the BrainSpan project. The top row gives the number of genes per cluster and the cluster centres in log₂-scaled RPKM (reads per kilobase per million) values; solid and dotted solid lines indicate dorsolateral prefrontal cortex (DLPFC) and hippocampus (HPC), respectively. The bottom two rows show enrichment in the current study, relative to the exome-wide average, for singleton disruptive mutations in cases compared to controls, either subsetting all genes by expression profile (first row), or considering only genes in the composite set (second row). In both cases, we only observed nominally (P < 0.01) significant enrichment for genes that are postnatally biased. By contrast, a list of genes with loss-of-function (LoF) de novo mutations (compiled and reported in Fromer et al.30) shows strong enrichment for prenatal bias (see Fromer et al.30 for details on how de novo enrichment was calculated). Alternative approaches to classifying genes as prenatally or postnatally biased led to similar conclusions (Supplementary Information section 6).



Extended Data Table 1 | Sample and detected variant properties

а

Case/control status by sex (P = 5e-11)	Status	Male	Female	Female (%)
	Control	1291	1252	49%
	Case	1520	1016	40%
Case/control status by ancestry (P = 2e-12)	Ancestry	Control	Case	Case (%)
	Swedish	2356	2197	48%
	Finnish	139	274	66%

b

Sample and sequencing metrics	Cases	Controls	P (case vs.
oumpic and sequenomy metrics	04363	CONTROLS	oonii oij
n	2536	2543	
n (pre-QC)	2546	2545	
Total number of reads	100,532,755	100,079,333	0.62
Filtered, unique reads aligned	68,940,753	68,339,964	0.26
Filtered, unique bases aligned	5,106,614,996	5,070,497,844	0.34
Mean target coverage	89.98	89.55	0.53
Percentage of target bases covered > 10x	92.83	92.85	0.55
Percentage of target bases covered > 20x	87.30	87.30	0.93
Percentage of target bases covered > 30x	81.13	81.07	0.63
Percentage of targets w/out any bases covered at 2x	1.72	1.72	0.60
Mean number of non-reference genotypes per individual (unfiltered)	18772.9	18786.6	0.13
Mean number of on-target singletons per individual (unfiltered)	49.6	49.0	0.38
Mean dbSNP % per individual	98.3970%	98.3969%	1.00

C

Property	Variant type	n	Mean MAC	% singleton
All alternate alleles		635,944	103.37	56%
All alternate alleles		000,044	100.01	3070
Functional class				
Noncoding		61,416	142.03	53%
Silent		185,336	152.85	51%
Missense		342,561	69.52	58%
Non-essential splice site		25,450	127.04	54%
Nonsense		9,022	20.68	69%
Essential splice-site		4,394	16.18	70%
Frameshifting indel		3,461	9.46	79%
In silico annotation of missenses				
LRT		168,437	34.55	62%
MutationTaster		167,316	19.90	63%
PolyPhen2 (HumDiv)		130,719	28.84	62%
PolyPhen2 (HumVar)		91,156	24.74	64%
SIFT		140,345	43.85	61%
Primary variant groupings for analys	sis			
Singletons	Gene disruptive	12,047	1.00	100%
	NSstrict	36,542	1.00	100%
	NSbroad	160,229	1.00	100%
<0.1% MAF (1-10 alleles)	Gene disruptive	15,972	1.56	75.4%
	NSstrict	50,369	1.65	72.5%
	NSbroad	233,575	1.78	68.6%
<0.5% MAF (1-50 alleles)	Gene disruptive	16,523	2.24	72.9%
	NSstrict	52,545	2.51	69.5%
	NSbroad	248,217	3.04	64.6%

a, Numbers of individuals in the final data set, after individual-level QC. Finnish ancestry was inferred by multidimensional scaling. P values from Fisher's exact test. b, Technical metrics for the cases and controls (after individual-level QC); P values for two-sided test of case/control differences (t-test). c, Properties of variants detected by exome sequencing. Counts (N) and minor allele counts (MAC) for various classes of variant in the main exome data set, following all QC. Missense deleteriousness prediction algorithms and how they were combined are described in Supplementary Information section 4.



Extended Data Table 2 | Genes prioritized as more likely to harbour large-effect alleles

Class	Gene	Singletons	MAF < 0.1%	Notes
ARC/PSI) complex			
AINO/I OL	CYFIP1	1/0	1/0	SCZ de novo (CNV)
	BAIAP2	1/0	1/0	SCZ de novo (NS)
	DLG1	1/0	1/0	SCZ de novo (NS), SCZ de novo (CNV)
	SLC25A3	1/0	1/0	
	GLUD1	1/0	1/0	
	CAMK2A	1/0	1/0	FMRP target
	ATP1B1	1/0	1/0	AUT de novo (disruptive); FMRP target
	IQSEC2	1/0	1/0	ID de novo (disruptive); FMRP target
	MBP	1/0	1/0	FMRP target
	Total	9/0	9/0	
		P = 0.0016	P = 0.0014	
		OR = 19.2	OR = 19.2	
		(2.4 - 2471)	(2.4 - 2471)	
PSD-95 g	neneset			
8	ABLIM1	1/0	1/0	
	ACO2	1/0	1/0	FMRP target
	ANKS1B	3/1	3/1	· ···· · · · · · · · · · · · · · · · ·
	ATP1B1	1/0	1/0	AUT de novo (disruptive); FMRP target
	ATP5A1	1/0	1/0	FMRP target
	BAIAP2	1/0	1/0	SCZ de novo (NS)
	CAMK2A	1/0	1/0	` ,
				FMRP target
	CAMK2B	2/0	2/0	FMRP target
	DLG1	1/0	1/0	SCZ de novo (NS), SCZ de novo (CNV)
	GAPDH	1/0	1/0	
	IQSEC2	1/0	1/0	ID de novo (disruptive); FMRP target
	NRXN1	1/0	1/0	SCZ de novo (NS); AUT de novo (disruptive); FMRP target
	PRDX1	0/1	0/1	
	PRDX2	0/1	0/1	
	SUCLA2	1/0	1/0	AUT de novo (disruptive)
	SYNGAP1	1/0	1/0	SCZ de novo (disruptive); ID de novo (disruptive); FMRP targe
	Total	17/3	17/3	
		P = 0.0017	P = 0.0009	
		OR = 5.1 (1.8 - 19.2)	OR = 5.1 (1.8 - 19.2)	
Voltage-g	ated calcium ion	channel genes		
	CACNA1B	1/0	1/0	FMRP target
	CACNA1C	2/0	2/0	SCZ & BP GWAS hit
	CACNA1H	1/0	3/0	
	CACNA1S	2/0	2/3	SCZ & AUT de novos (NS)
	CACNA2D1	1/0	1/0	PSD
	CACNA2D1	3/0	3/0	1 30
	CACNA2D2	0/0	3/0	ALIT de nove (digruptive)
				AUT de novo (disruptive)
	CACNA2D4	1/0	2/4	
	CACNB2	0/1	0/1	202
	CACNB4	1/0	1/0	PSD
	Total	12/1	15/8	
		P = 0.0021 OR = 8.4 (2.03 - 77)	P = 0.021 OR = 2.1 (0.97 - 4.9)	
		,		
Top disru	ptive gene-based			
	KYNU	3/0	10/0	
		P = 0.13	P = 0.0017	
			OR = 21.2 (2.7 - 2725)	

Individual gene case/control counts, odds ratios and *P* values for genes from primary gene sets with odds ratios >5, and *KYNU* (top-ranked individual gene). Odds ratios are calculated using Firth's method (penalized maximum likelihood logistic regression) and shown with 95% confidence intervals. *P* values are empirical, uncorrected one-sided burden tests. FMRP target annotations are based on the Darnell *et al.*⁴⁰ list only. Supplementary Table 1 lists singleton variant and genotype information for the genes listed here.

Extended Data Table 3 | Extended results for all PSD gene sets

а

			Disruptive		_	NS strict			NS broad	
Set	n genes	Singletons	MAF < 0.1%	MAF < 0.5%	Singletons	MAF < 0.1%	MAF < 0.5%	Singletons	MAF < 0.1%	MAF < 0.5%
	_	_			_					
PSD (human core)	685	0.0729	0.1019	0.1083	0.0058	0.1045	0.1285	0.0866	0.5827	0.3743
ARC	28	0.0016	0.0013	0.0014	0.0004	0.0018	0.0047	0.2830	0.4607	0.3542
NMDAR network	61	0.0154	0.0229	0.0225	0.0001	0.0007	0.0005	0.0012	0.1426	0.0420
mGluR5	39	0.1299	0.0861	0.0862	0.0628	0.0837	0.0900	0.0302	0.2192	0.1329
PSD-95	65	0.0015	0.0008	0.0008	0.0027	0.0204	0.0393	0.2992	0.3722	0.1147
Pre-synapse	431	0.0187	0.0983	0.1458	0.2327	0.1811	0.3600	0.1306	0.7597	0.6515
Pre-synaptic active zone	173	0.0518	0.0487	0.0482	0.6162	0.6641	0.7082	0.8439	0.9918	0.9554
Synaptic vesicle	344	0.1030	0.3133	0.4151	0.1439	0.1093	0.2466	0.0375	0.3423	0.2718
Cytoplasm	271	0.5851	0.1793	0.1034	0.8983	0.5351	0.6007	0.1861	0.1084	0.1192
Early Endosomes	17	0.8917	0.7860	0.7826	0.2891	0.2420	0.2139	0.1472	0.4019	0.4123
Endoplasmic Reticulum	97	0.3005	0.1882	0.2612	0.6615	0.2805	0.5036	0.6194	0.5044	0.7878
ER/Golgi-derived vesicles	94	0.4258	0.2678	0.3644	0.3001	0.4977	0.6239	0.3063	0.6944	0.7649
Golgi	31	0.5130	0.5493	0.5481	0.1998	0.0921	0.1338	0.0074	0.1628	0.4178
Mitochondrion	197	0.0141	0.0259	0.0178	0.4351	0.0860	0.0671	0.6999	0.2112	0.6079
Nucleus	167	0.1790	0.3029	0.2900	0.0626	0.1512	0.2728	0.2006	0.3340	0.3196
Plasma membrane	50	0.7940	0.5659	0.5635	0.9416	0.8059	0.8091	0.8944	0.5531	0.3028
Recycling Endosomes/										
trans-Golgi network	68	0.1502	0.0944	0.1556	0.5349	0.4514	0.5359	0.0902	0.0862	0.1862

b

		PSD (human core) (ref. 13)		FMRP targets (Darnell, ref. 40)		FMRP targets (Ascano, ref. 42)	
	n	n	%	n	%	n	%
PSD (human core)	685			170	25%	80	12%
ARC	28	-		16	57%	2	7%
NMDAR network	61			32	52%	5	8%
mGluR5	39			25	64%	7	18%
PSD-95	65			30	46%	4	6%
Pre-synapse	431	213	49%	87	20%	31	7%
Pre-synaptic active zone	173	121	70%	50	29%	10	6%
Synaptic vesicle	344	162	47%	72	21%	27	8%
Cytoplasm	271	77	28%	16	6%	23	8%
Early Endosomes	17	6	35%	2	12%	1	6%
Endoplasmic Reticulum	97	13	13%	5	5%	4	4%
ER/Golgi-derived vesicles	94	24	26%	7	7%	4	4%
Golgi	31	2	6%	2	6%	4	13%
Mitochondrion	197	57	29%	6	3%	12	6%
Nucleus	167	19	11%	7	4%	19	11%
Plasma membrane	50	16	32%	6	12%	5	10%
Recycling Endosomes/trans-Golgi network	68	19	28%	3	4%	7	10%
Total	1509	685	45%	170	11%	80	5%

a, Full PSD gene set association results. For all nine (three annotation levels by three frequency levels), P values for enrichment of all gene sets described and tested in Kirov et al. 13. In addition to the PSD genes (top five rows), enrichment statistics for presynaptic genes, and neuronal genes clustered on the basis of subcellular location are given. Although the P values presented are uncorrected, we performed this analysis correcting for all 9 × 17 = 153 tests (by considering the distribution of the minimum empirical P value across tests and sets, as described in the Supplementary Information). The values in bold are significant (P^{corrected} < 0.05) after correction for multiple testing. Both ARC and NMDAR network are significant after multiple test correction, for the singleton NS_{atrict} category. (Note: for ARC the disruptive singleton category is, as reported in the primary test, highly significant and withstands correction for multiple testing in that context; in this broader, less focused analysis it yields P^{corrected} = 0.17; the majority of P^{corrected} values (not shown) are 1.00.) b, PSD and FMRP-target gene sets: descriptive statistics and overlap. Overlap between Darnell et al. 40 and Ascano et al. 42 FMRP targets and PSD genes: for example, 57% (16 out of 28) of ARC genes are in the Darnell FMRP list. By contrast, only 7% (2 out of 28) are in the Ascano list. There is a similar trend across the three other major PSD subsets considered here: NMDAR network, PSD-95 and MGluR5 genes. Conversely, 22% of Darnell targets are in the PSD (human core) compared to only 9% of Ascano targets.



Extended Data Table 4 | Association results for individual CNV regions

		Di	sruptive	NS	strict	
Group	Genes	Singletons	MAF < 0.1%	Singletons	MAF < 0.1%	
CNV loci	All	0.3279	0.4557	0.0843	0.0044	
	1g21.1	0.4533	0.6966	0.3205	0.1832	
	2p16.3	0.4775	0.3580	0.4703	0.2750	
	3q29	0.1054	0.0068	0.0123	0.0006	
	7q36.3	0.8642	0.5750	0.6411	0.2688	
	7q11.23	0.7199	0.6800	0.3329	0.2207	
	15q11.2	0.3208	0.1616	0.5138	0.1362	
	15q13.3	0.0883	0.3976	0.3672	0.2746	
	16p13.11	0.4194	0.3775	0.8346	0.4124	
	16p11.2	0.1613	0.1240	0.0655	0.0974	
	17q12	0.7377	0.5205	0.4313	0.1385	
	22q11.21	0.9386	0.9977	0.5456	0.8754	
	Gene	A/U	Gene name			
3q29 genes	DLG1	5/0	discs, large homolog 1	(Drosophila)		
	RNF168	5/1	ing finger protein 168,	E3 ubiquitin protein	ligase	
	CEP19	2/0	centrosomal protein 19	kDa		
	LRRC33	2/0	eucine rich repeat con	taining 33		
	PAK2	2/0	o21 protein (Cdc42/Ra	c)-activated kinase 2	2	
	PCYT1A	5/2	ohosphate cytidylyltran	sferase 1, choline, a	ılpha	
	PIGX	6/3	phosphatidylinositol gly	can anchor biosynth	nesis, class X	
	FBXO45	1/0	-box protein 45	•		
	NCBP2		nuclear cap binding pro	otein subunit 2, 20k)a	
	PIGZ	1/0	phosphatidylinositol gly	can anchor biosynth	nesis, class Z	
	TFRC	1/0	ransferrin receptor (p9	0, CD71)		
	ZDHHC19	1/0	zinc finger, DHHC-type	containing 19		
	C3orf43	4/2	chromosome 3 open re	ading frame 43		
	MFI2	15/12	antigen p97 (melanoma	a associated) identif	ed by monoclonal	antibodies 133.2 and 96.5
	SLC51A		solute carrier family 51	·	-	
	BDH1		3-hydroxybutyrate dehy			
	TCTEX1D2	0/1	Tctex1 domain containi	ng 2		
	WDR53	0/1	ND repeat domain 53	-		

Focused enrichment analysis of genes under schizophrenia-associated CNV regions. The top panel presents omnibus P values testing all genes/regions (bold indicates significance after correction for the four tests, $P^{\text{corrected}} = 0.016$ for NS_{strict} MAF < 0.1% variants). This enrichment arises solely from the 3q29 locus (middle panel; bold indicates significance after correction of the 44 tests performed, $P^{\text{corrected}} = 0.024$ for 3q29). Genes and NS_{strict} case/control counts for the 3q29 region (bottom panel).

Extended Data Table 5 | Further stratification of enrichment analyses by class of variant

		Disruptiv	e singletons		Damaging missense (strict) singleton				
Primary geneset	P	n	A/U	OR	P	n	A/U	OR	
All primary genes	0.0008	905	852/716	1.20	0.0393	1357	2080/2001	1.04	
SCZ de novo genes (refs. 18,19,30)									
Exome sequencing (disruptive)	0.0349	40	56/38	1.48	0.5613	53	121/120	1.01	
Exome sequencing (nonsyn)	0.0059	332	384/309	1.25	0.2776	393	750/736	1.02	
Copy number variants (refs. 5,13)									
de novo CNV genes	0.0224	64	61/40	1.53	0.0593	90	125/112	1.12	
SCZ-associated CNV genes	0.3378	72	65/55	1.19	0.0310	111	148/119	1.25	
GWAS (refs. 3,5,12)									
Voltage-gated calcium channel genes	0.0021	9	12/1	8.40	0.4629	18	37/35	1.06	
Common SNPs (P < 1e-4 intervals)	0.1832	185	165/146	1.14	0.9246	268	359/395	0.91	
miR-137 targets	0.6643	140	98/100	0.99	0.1415	263	376/361	1.05	
Synaptic genes (ref. 13)									
PSD (human core)	0.0824	219	172/145	1.19	0.0070	394	646/581	1.12	
ARC	0.0012	9	9/0	19.20	0.0069	19	32/15	2.14	
NMDAR network	0.0162	17	17/5	3.42	0.0003	34	76/45	1.70	
PSD-95	0.0018	16	17/3	5.10	0.0218	34	44/30	1.47	
mGluR5	0.1335	10	9/3	3.02	0.1715	22	52/36	1.45	

b

	Case-ur	nique burden an	alysis	Known variants					Novel v		
Geneset	P	n genes(A/U)	A(U)	Singlet <i>P</i>	ons n	MAF <	0.1% <i>n</i>	Singlet <i>P</i>	ons n	MAF <	0.1% : n
Geneset		II genes(A/O)	A(U)								
Composite	0.0006	829(275/214)	378(297)	0.3733	145	0.2202	226	0.0002	683	0.0005	74
Primary	0.0022	1026(325/265)	440(367)	0.1003	191	0.0417	299	0.0074	831	0.0058	91
SCZ de novo genes (refs. 18,19,30)											
Exome sequencing (disruptive)	0.0018	47(16/6)	29(12)	0.5514	13	0.2196	24	0.0362	35	0.0010	40
Exome sequencing (nonsyn)	0.0037	371(108/80)	159(116)	0.3647	94	0.2064	144	0.0142	302	0.0071	32
Copy number variants (refs. 5,13)											
de novo CNV genes	0.1267	79(25/17)	32(24)	0.0156	13	0.0116	24	0.0819	59	0.0679	6
SCZ-associated CNV genes	0.7971	90(20/23)	24(32)	0.0355	23	0.0069	34	0.6081	63	0.9187	76
GWAS (refs 3,5,12)											
Voltage-gated calcium channel	0.0129	9(5/1)	10(1)	0.4922	2	0.7077	3	0.0006	7	0.0022	8
P < 1e-4 intervals	0.1079	211(65/55)	91(78)	0.0394	44	0.0409	69	0.4425	164	0.1882	18
miR-137 targets	0.4498	156(52/50)	67(60)	0.9939	14	0.9972	22	0.3846	133	0.2757	14
Synaptic genes (ref. 13)											
PSD (human core)	0.2234	244(92/79)	113(109)	0.5348	25	0.3072	40	0.1091	205	0.1629	
ARC	0.0008	9(9/0)	9(0)		0		0	0.0016	9	0.0013	ç
NMDAR network	0.0105	21(13/4)	18(4)	1.0000	1	0.6905	2	0.0075	16	0.0085	
PSD-95	0.0137	16(13/2)	14(2)	0.1218	1	0.1559	1	0.0034	15	0.0022	
mGluR5	0.1363	11(7/1)	8(1)		0	0.1458	1	0.1427	10	0.1826	11
Secondary (autism/ID)	0.0916	1249(348/314)	479(471)	0.1679	226	0.3543	352	0.1834	1041	0.0807	114
De novo genes (exome sequencing)											
Autism (disruptive) (refs. 22-25)	0.662	65(17/17)	20(26)	0.4463	18	0.0781	23	0.6161	50	0.7009	56
Autism (nonsyn) (refs. 22-25)	0.220	407(101/96)	143(154)	0.3198	89	0.4487	133	0.6656	336	0.4960	36
ID (disruptive) (refs. 20, 21)	0.262	8(4/1)	4(2)	1.0000	1	0.2747	3	0.3558	8	0.0578	8
ID (nonsyn) (refs. 20,21)	0.052	69(22/18)	35(28)	0.1303	14	0.0934	26	0.5331	62	0.3368	6
Neurodevelopmental candidates											
ASD candidates (ref. 39)	0.110	37(12/6)	16(7)	0.5824	9	0.7543	14	0.0484	24	0.0429	2
ID candidates (ref. 39)	0.994	88(14/28)	16(38)	0.6553	16	0.7056	24	0.9488	74	0.9556	82
Autism PPI networks											
CHD8 network (ref. 24)	1.000	1(0/1)	0(1)		0		0	1.0000	1	1.0000	1
49-gene network (ref. 24)	0.796	19(3/7)	4(16)	0.4755	5	0.7326	7	0.6081	30	0.7231	33
74-gene network (ref. 24)	0.654	33(6/13)	10(28)	0.6438	4	0.6667	4	0.7285	17	0.8411	19
Fragile X mental retardation protein targets		(-, -3)	-(/	2.2.30			•	50			
Darnell (ref. 40) targets	0.022	341(131/95)	169(133)	0.3048	39	0.3889	61	0.0007	288	0.0022	30
	0.440	E47(104/401)	107(200)	0.5571	83	0.7281	128	0.5261	439	0.4089	48
Ascano (ref. 42) targets	0.449	517(134/131)	107 (200)	0.5571	03	0.7201	120	0.5261	439	0.4003	40

a, Gene set analyses for damaging missense mutations only. For the primary gene set and the 12 constituent subsets, a comparison of disruptive versus (strictly defined) damaging missenses, that is, an independent set of variants. The ornnibus result for the primary test is modest (P = 0.04) and did not withstand correction for multiple testing: as illustrated in Fig. 1 and the main text, the bulk of the enrichment signal we observe comes from (singleton) disruptive mutations. Nonetheless, specific gene sets such as ARC and the NMDAR network are highly and independently enriched for missense variants. N represent take number of genes with at least one mutation of this class observed in the sample. A/U represent case/control counts of non-reference genotypes. OR represents the odds ratio (not corrected for exome-wide rates) estimated by Firth's method for sets with small cell counts. All tests are empirical and one-sided (higher values expected in cases) as described in the main text and Methods. b, Enrichment analyses of novel and case-unique disruptive mutations. For primary and secondary gene sets (and constituent subsets) as well as the composite set: results of alternative burden analyses. First, focusing only on genes without any control disruptive variants; no further frequency filter is imposed. Here 'N genes(A/U)' indicates the number of genes with at least one disruptive variants, followed by the number of genes with case-only genes: the test statistic is based on the empirical distribution of this count. The U in this field represents the similar quantity for controls (not explicitly used in the statistic). The second set of analyses represent standard burden/enrichment tests (that is, as Tables 1 and 2) but stratified for novel versus known disruptive variants, according to dbSNP and the Exome Sequencing Project/Exome Variant Server (ESP/EVS) database. Novel variants show greater enrichment, although most rare variants observed in our study (both in cases and in controls) are novel, so tests of no



Extended Data Table 6 | Gene set analysis of de novo genes from schizophrenia exome-sequencing studies

а

	•		Disruptive			NSstrict		NSbroad		
Set	n genes	Singletons	MAF < 0.1%	MAF < 0.5%	Singletons	MAF < 0.1%	MAF < 0.5%	Singletons	MAF < 0.1%	MAF < 0.5%
Fromer et al.										
(ref. 30) disruptive	63	0.1484	0.0075	0.0034	0.7401	0.7324	0.6264	0.3347	0.1660	0.2536
Fromer et al.										
(ref. 30) nonsyn	464	0.0004	0.0003	0.0016	0.0341	0.0057	0.0892	0.4688	0.4842	0.6547
Girard & Xu										
(refs. 18,19) disruptive	24	0.0342	0.0082	0.0082	0.0423	0.0412	0.0602	0.0774	0.1106	0.0891
Girard & Xu (refs. 18,19) nonsyn	151	0.6916	0.4124	0.4186	0.1510	0.1326	0.1285	0.2258	0.3420	0.1790
(1010: 10,10) 110110311	101	0.0310	0.4124	0.4100	0.1010	0.1020	0.1200	0.2200	0.0420	0.1750
Combined SCZ										
disruptive	87	0.0319	0.0007	0.0003	0.3355	0.3162	0.2692	0.1325	0.0757	0.1016
Combined SCZ										
nonsyn	611	0.0053	0.0011	0.0055	0.0192	0.0024	0.0379	0.3408	0.4064	0.4472

b

Gene	De novo study (type)	Test	n	A/U	P	Gene name
ALDH1L2	Fromer (ref. 30) nonsyn	disruptive	6	10/3	0.028	aldehyde dehydrogenase 1 family, member L2
CACNA1S	Fromer (ref. 30) nonsyn	NSstrict	23	28/15	0.031	calcium channel, voltage-dependent, L type, alpha 1S subunit
DLG1	Fromer (ref. 30) nonsyn	NSstrict	4	5/0	0.021	discs, large homolog 1 (Drosophila)
IGSF22	Fromer (ref. 30) nonsyn	disruptive	5	5/0	0.043	immunoglobulin superfamily, member 22
JARID2	Fromer (ref. 30) nonsyn	NSstrict	5	5/0	0.041	jumonji, AT rich interactive domain 2
LAMA4	Fromer (ref. 30) nonsyn	NSstrict	8	12/4	0.041	laminin, alpha 4
NBEA	Fromer (ref. 30) nonsyn	NSstrict	5	5/0	0.025	neurobeachin
POLL	Fromer (ref. 30) nonsyn	disruptive	4	4/0	0.042	polymerase (DNA directed), lambda
PTK2B	Fromer (ref. 30) nonsyn	NSstrict	4	4/0	0.044	PTK2B protein tyrosine kinase 2 beta
SHKBP1	Fromer (ref. 30) nonsyn	NSstrict	9	15/4	0.018	SH3KBP1 binding protein 1
SULF2	Fromer (ref. 30) nonsyn	NSstrict	5	7/0	0.007	sulfatase 2
SYNGAP1	Xu (ref. 19) LoF	NSstrict	4	4/0	0.043	synaptic Ras GTPase activating protein 1
SZT2	Xu (ref. 19) LoF	NSstrict	22	18/9	0.049	seizure threshold 2 homolog (mouse)
TANC1	Fromer (ref. 30) nonsyn	NSstrict	14	17/4	0.002	tetratricopeptide repeat, ankyrin repeat and coiled-coil containing 1
TEP1	Xu (ref. 19) nonsyn	NSstrict	26	25/14	0.048	telomerase-associated protein 1
UFL1	Fromer (ref. 30) LoF	NSstrict	4	7/0	0.008	UFM1-specific ligase 1
UFL1	Fromer (ref. 30) LoF	disruptive	2	5/0	0.029	UFM1-specific ligase 1

a, Test of case enrichment of rare variants in cases compared to controls, for genes with one or more de novos in Fromer etal. And/or Girard etal. The P values in bold are significant at P corrected < 0.05, correcting for all $3 \times 3 \times 6 = 54$ tests reported. b, Genes nominally significant (no correction) that had an observed de novo in one of the schizophrenia studies.



Extended Data Table 7 | Summary of observed likely N.A, deleterious variants in ARC genes across studies

		Curr	rent study			
ARC gene (n=28)		Disruptive	Damaging missense (strict)	de novo CNV (Kirov et al., ref. 13)	de novo SNV (Fromer et al., ref. 30)	de novo SNV in ID (refs. 20,21)
ACTN4			3/1			
ARF5						
ATP1A1			3/0			
ATP1A3			2/1			
ATP1B1		1/0	1/0			
BAIAP2		1/0			NS(x2)	
CAMK2A		1/0	1/0			
CRMP1			1/3			
CYFIP1		1/0	4/1	2 del; 2 dup		
DLG1		1/0	2/0	1 del	NS	
DLG2			2/3	2 del	LoF	
DLG4			1/2			NS
DLGAP1			2/0	1 del		
DLGAP2			1/0			
DPYSL2			0/1			
GLUD1		1/0	1/0			
GLUL			2/0			
GRIN1			2/0			
HSPA8					LoF & NS	
IQSEC1			4/1			
IQSEC2		1/0	0/1			LoF
MBP		1/0	0/1			
PKM2						
PLP1						
SLC25A3		1/0				
SLC25A4						
SLC25A5						
STXBP1						LoF, NS(x2)
	Counts:	9/0	32 / 15	8 CNVs	6 SNVs	5 SNVs
	P-value:	0.0016	0.0069	0.00025	0.0005	0.00002

For the 28 ARC genes, a summary of which genes had singleton disruptive, or damaging missense, variants in the current study, compiled alongside the genes with *de novo* CNVs or SNVs observed in Kirov *et al.* ¹³ or Fromer *et al.* ³⁰). The *P* values at the bottom indicate that in each comparison the ARC gene set was significantly enriched.