Attenuated transcriptional response to pro-inflammatory cytokines in schizophrenia hiPSC-derived neural progenitor cells

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Abstract

Maternal immune activation (MIA) during prenatal development is an environmental risk factor for psychiatric disorders including schizophrenia (SZ). This link is particularly well established in the case of schizophrenia. Converging lines of evidence from human and animal model studies have suggested that elevated cytokine levels in the maternal and fetal compartments are an important indication of the mechanisms driving this association. However, there is variability in susceptibility to the psychiatric risk conferred by MIA, likely influenced by genetic factors. How MIA interacts with a genetic profile susceptible to SZ is challenging to test in animal models. To address this gap, we examined whether differential gene expression responses occur in forebrain-lineage neural progenitor cells (NPCs) derived from human induced pluripotent stem cells (hiPSC) generated from three individuals with a diagnosis of schizophrenia and three healthy controls. Following acute (24 hour) treatment with either interferon-gamma (IFNy; 25 ng/µl) or interleukin (IL)-1 β (10 ng/µl), we identified, by RNA sequencing, 3380 differentially expressed genes (DEGs) in the IFNy-treated control lines (compared to untreated controls), and 1980 DEGs in IFNy-treated SZ lines (compared to untreated SZ lines). Of these, 359 genes responded differently to IFNy treatment in SZ relative to control lines, which were enriched for genes differentially expressed in adult SZ cases and in pathways related to the regulation of antigen processing, neuronal postsynaptic density, and the presynapse. There were no differentially expressed genes in the IL-1βtreatment conditions after Benjamini-Hochberg correction. Gene set enrichment analysis however showed that IL-1β impacts immune function and neuronal differentiation. Collectively, our data suggest that a) SZ NPCs show an attenuated transcriptional response to IFNy treatment compared to control NPCs; b) IL-1ß may be less relevant to NPC cultures than IFNy and c) the genes differentially regulated in schizophrenia lines are primarily mitochondrial, loss-of-function, pre- and post-synaptic genes. Our findings particularly highlight early synaptic development as a key target for future research into the mechanisms of association between maternal immune activation and schizophrenia risk.

Key words: Maternal immune activation, inflammation, cytokine, IL-1β, IFNγ, differential gene expression, neurotransmission, prenatal development, neurodevelopment

Highlights

- There is a significant transcriptional response to IFNγ treatment in human induced cortical neural progenitor cells (NPCs)
- Differential expression implicates mitochondrial complex genes in this IFNγdependent response, which are underexpressed in response to treatment in schizophrenia (SZ) NPCs.
- SZ NPCs do not upregulate pre- and post-synaptic genes in response to IFNγ as much as control NPCs
- IL-1β may be less relevant to NPC cultures than IFNγ due to low IL1 receptor expression in NPCs.

Introduction

Activation of the maternal immune response during pregnancy is a known risk factor for neurodevelopmental disorders - especially autism and schizophrenia - in the offspring [1-5]. Although the precise molecular mechanisms driving this association remain unclear, exposure of the developing fetal brain to pro-inflammatory cytokines remains a promising candidate for study [2, 6-8]. Cytokines are cell signalling proteins that help immune cells to form coordinated responses to infection. Whilst their function in the peripheral immune system is well documented, there is growing evidence that cytokines also play an important role in brain development and that maternally-derived cytokines could affect the developing foetal brain [2, 6-8]. Emerging evidence from human studies suggest that elevated levels of canonical pro-inflammatory cytokines such as interferon gamma (IFNy) [2, 9] and interleukin-1 beta (IL-1 β) [7, 10] have been detected in the plasma of individuals with a diagnosis of schizophrenia [9, 11]. Furthermore, serum levels of IL-1β are elevated in the mothers of offspring who later develop psychosis [12]. Animal models of maternal immune activation (MIA) also provide evidence of elevated IFNy and IL-1ß levels in maternal serum, as well as the serum and brains of fetuses [6, 13]. Moreover, in mice exposed to MIA, offspring who are susceptible (those that show, as adults, deficits in social and cognitive functioning as seen in schizophrenia) have elevated plasma levels of IL-1ß (among other cytokines), compared to control mice and mice that are resilient to MIA [14].

However, the impact and outcome following prenatal immune activation is heterogeneous between individuals [5, 14, 15]. For example, recent work in an MIA

mouse model highlights the existence of subgroups of MIA-exposed offspring that show dissociable behavioural, transcriptional, neuroimaging, and immunological profiles [14, 16]. This is consistent with epidemiological studies in human cohorts: not all foetuses exposed to MIA will go on to develop schizophrenia [1, 17, 18]. It is therefore likely that MIA interacts with other factors, such as genetic background, to modulate the risk of developing schizophrenia. Indeed, the two-hit model of schizophrenia suggests that an amalgam of genetic risk and environmental insult is necessary to alter neurodevelopment enough to ultimately precipitate the symptoms of the disorder [19-21].

Importantly, if there are individual differences in human responses to immune activation due to genetic variability, there will certainly be differences in such responses between human and animal systems, given that they are even more genetically divergent. Although animal studies have provided important mechanistic insights, interactions between schizophrenia genetic burden and MIA cannot be fully recapitulated by rodent models: species differences in gene expression cannot be discounted [22-25]. It is therefore important to test the impact of gene-environment interactions in human model systems, such as human induced pluripotent stem cells (hiPSCs). These can be generated by reprogramming somatic cells, including hair keratinocytes and skin fibroblast biopsies, from patient cohorts or healthy controls [26-28]. The resulting hiPSCs can then be differentiated into multiple relevant cell types that retain the genetic make-up of the donor [29, 30] – including neural progenitor cells (NPCs), early precursors to neurons which are highly prevalent in the fetal brain [31]. There is evidence that hiPSCderived NPCs closely resemble fetal brain tissue, recapitulating the neurodevelopmental hallmarks of the late first trimester/early second trimester stage [2, 29, 30, 32-34]. These

facets make hiPSC-NPCs uniquely placed to model human fetal neurodevelopmental mechanisms and gene-environment interactions *in vitro* [33, 35].

Previous work from our group showed that transient IFNy treatment of hiPSC-NPCs from healthy controls increases neurite outgrowth (a cellular phenotype associated with neurodevelopmental disorders) and disproportionately alters the expression of genes associated with schizophrenia and autism [2]. The aim of the current study is therefore to understand how specific cytokines (IFNy and IL-1 β) implicated in the association between MIA and schizophrenia risk influence transcriptional responses in early cortical neural progenitors. We use induced cortical neural progenitors with forebrain identity, as there is extensive evidence of prefrontal cortical abnormalities in patients with schizophrenia [7, 36, 37], and experiments in animal models have shown that exposure to inflammatory cytokines alters proliferation and differentiation of neural progenitors [10, 38]. We hypothesise that NPCs derived from patients with schizophrenia will respond differently to IFNy and IL-1 β compared to cells from healthy donors. If so, this may shed light on the mechanisms by which maternal immune activation increases the risk of developing schizophrenia.

Materials and methods

Participants

This study included hiPSC lines derived from six participants: three individuals with a diagnosis of schizophrenia (cell lines 138_SZM_09, 044_SCZ_04 and 115_SCZ_01) (Supplementary Figure 1) and three healthy donors with no history of psychiatric illness (cell lines M1_CTR_04, M2_CTR_42, M3_CTR_36S – previously described in [32, 33]). Participants were recruited as part of the Patient iPSCs for Neurodevelopmental Disorders (PiNDs) study (REC No 13/LO/1218). Participants with a diagnosis of schizophrenia were recruited at the Maudsley Hospital, London. The collection of data used for this research was approved by the NHS Research Ethics Committee at the South London and Maudsley (SLaM) NHS Research and Development Office. All participants gave written informed consent before contributing to the study. A diagnosis of schizophrenia was established based on International Classification of Diseases (10th revision) [39] with the diagnosis (code F20), assessed using the Operational Criteria checklist [40] by a psychiatrist on the basis of information recorded by the clinical team following psychiatric interview. Healthy, unaffected individuals were selected as controls on the basis of having no history of psychiatric disorders [33].

Reprogramming of keratinocytes

Hair root samples were collected by plucking occipital scalp hair (~10+ roots per participant) and submerging these in Mouse Embryonic Fibroblast medium containing 50 µg/mL Gentamycin and 15 mM HEPES buffer (Gibco). The roots were then transferred to Geltrex[™]-coated 4-well plates (ThermoFisher), and outgrowth promoted, by supplementing with hair medium (Dulbecco's Modified Eagle's Medium (DMEM)

Advanced (Sigma Aldrich), GlutaMAX[™] (ThermoFisher), 10% FBS (Clonetech), HEPES buffer and Gentamycin), to establish primary keratinocytes. The keratinocytes were subsequently reprogrammed into human induced pluripotent stem cell (hiPSC) lines. This transformation was induced by introducing Sendai viruses encoding Yamanaka Factors (human OCT4, SOX2, KLF4 and C-MYC), using a CytoTune-iPS 2.0 Sendai expressing Reprogramming Kit (ThermoFisher, A16517). The treated keratinocytes were plated onto an irradiated MEF feeder layer (Millipore) and supplemented Epilife medium. After ten days, Epilife medium was exchanged for hES medium, which was comprised of KO-DMEM/F12 supplemented with 20% knock-out serum, non-essential amino acids, Glutamax, β-mercaptoethanol (all from Life Technologies) and bFGF (10 ng/mL; Peprotech). After two more weeks, reprogrammed colonies were selected and plated on Nunc multi-plates (Thermo Scientific) coated with Geltrex (Life technologies) and supplemented with E8 media (Life Technologies).

Maintenance of hiPSCs

The successfully reprogrammed hiPSCs were incubated in hypoxic conditions (5% CO2, 5% O₂) at 37°C and maintained in StemFlex[™] media (Gibco) on 6-well NUNC[™] plates (ThermoFisher) coated with Geltrex[™] (ThermoFisher). Cells were passaged (at a ratio between 1:6 and 1:18) upon reaching 60-70% confluency. During passage, cells were washed with room temperature Hank's Balanced Salt Solution (HBSS) and incubated at 37°C with Versene (EDTA) solution (Lonza) for 3-5 minutes, then replated in new Geltrex[™]-coated NUNC[™] plates.

Directed differentiation of hiPSCs

The six hiPSC lines used in this study were then differentiated into forebrain cortical neural progenitor cells (NPCs) by dual SMAD inhibition [32, 33]. In preparation for neuralisation, hiPSCs were passaged onto 6-well NUNC[™] plates coated with Geltrex[™] at a 3:2 ratio and maintained under hypoxic conditions for ~24–48 hrs until they approached 100% confluence. Directed differentiation was then initiated by changing StemFlex[™] medium to neuralisation medium containing N2:B27 (N2 medium and B27 medium at a 1:1 ratio) supplemented with 100 nM LDN193189 (Sigma Aldrich) and 10 µM SB431542 (Sigma Aldrich) for dual SMAD inhibition. N2 medium consisted of DMEM/F12 (Dulbecco's Modified Eagle's Medium/Nutrient Mixture F12 Ham; Sigma Aldrich), supplemented with 1X GlutaMAX[™] and 1X N2 supplement (ThermoFisher). B27 medium consisted of Neurobasal® medium (ThermoFisher), 1X GlutaMAX[™] (ThermoFisher) and 1X B27 supplement without vitamin A (ThermoFisher).

The neuralised cells were then incubated under normoxic conditions (37°C, 5% CO₂, 20% O₂). Neuralisation medium was replenished every 24 hours from day 0 to day 7. At the end of this 7-day neuralisation period, neuralisation medium was replaced with N2:B27 (without inhibitors), which was replenished every 24 hours from day 8 onwards. The neuralised cells were passaged four times: on day 7, day 12, day 15/16 and day 20/21. The passage procedure was, briefly, as follows: cells were washed with room temperature HBSS (ThermoFisher) and treated with Accutase (ThermoFisher) and incubated for 3–4 minutes at 37°C. The cells were then collected with the Accutase and mixed with room temperature DMEM/F12 (at a 2:1 ratio) and centrifuged at 1250 RPM for two minutes to separate the cells and Accutase. Cells were plated on new 6-well

NUNC[™] plates coated with Geltrex[™]. Passaging ratios were 1:1 for neural passaging 1 and 2, and 2:3 for neural passaging 3. To enhance cell survival, 10 µM protein kinase (ROCK) inhibitor (Sigma Aldrich), was added for 24 hours with the plating medium at each neural passage. After neural passage 3, cells were frozen in 10% DMSO (dimethyl sulfoxide). Cryovials were stored at -80°C for 24–48 hours in Mr. Frosty containers (to control freezing rate) before being transferred to liquid nitrogen.

For the final stages of neural passaging, cryovials were thawed in a 37°C water bath for 1 minute. The cell suspension was transferred to a 15 ml tube containing DMEM/F12 and centrifuged at 1250 RPM for 2 minutes. The cell pellet was resuspended in 3 mL of N2:B27 supplemented with 10 μ M ROCK inhibitor and plated in GeltrexTMcoated 6-well NUNCTM plates. From this point on, the following inhibitors were added to the NPC media (to make N2:B27-FGF): 10 ng/mL bFGF (basic Fibroblast Growth Factor; Peprotech), 100 μ M β -mercaptoethanol (Life Technologies), 5 μ g/mL insulin (Life Technologies), 1X non-essential amino acids (Life Technologies), 200 μ M ascorbic acid (Sigma Aldrich). The cells were then expanded at a 1:3 ratio (in 2–5 neural passages) to prepare three wells of each line (one for each experimental condition).

Successful reprogramming of hiPSCs was validated as described in previous studies [32, 34, 41]. Pluripotency of all hiPSCs was confirmed by immunocytochemistry, differentiation of embryoid bodies into the three characteristic germ layers [42-45] (Supplemental Figure 1), and PluriTest analysis of Illumina HT12v4 transcriptome array data (<u>https://www.pluritest.org</u>) [46]. Alkaline phosphatase activity was further used to assess the pluripotency of hiPSCs using an alkaline phosphatase expression kit (Milipore). Genome integrity of hiPSC lines was assessed by an Illumina Human

CytoSNP-12v2.1 beadchip array and analysed using KaryoStudio software (Illumina, San Diego, CA).

Acute treatment with pro-inflammatory cytokines

NPCs were treated for ~24 hours in three treatment conditions: IFNy, IL-1 β or vehicle. Media was fully removed and replaced with 3 mL per well of treatment media (N2:B27-FGF, supplemented as follows). IFNy wells were treated with 25 ng/µL IFNy (Abcam); the IL-1 β wells with 10 ng/µL IL-1 β (Abcam), as in [2]; and the control wells with vehicle (unsupplemented N2:B27-FGF media). After 24 hours, cells were lysed and collected in TRIzol® reagent (Thermo Fisher) and rapidly frozen on dry ice. The frozen samples were stored at -80°C until RNA extraction.

RNA extraction and sequencing

RNA was extracted from the eighteen samples in two batches (to ensure durations of exposure of each sample to extraction reagents were well controlled). Both batches of extractions were conducted on the same day, by the same experimenter. The batches were randomised for experimental group (batch 1: lines M1_CTR, M2_CTR, 138_SZM; batch 2: lines M3_CTR, 044_SCZ, 115_SCZ), using the RNeasy Plus Mini Kit (QIAGEN), according to the manufacturer's instructions. Extracted RNA was sent for sequencing at GENEWIZ® Ltd. Strand-specific, paired-end RNA sequencing with Poly(A) selection was performed using the Illumina® NovaSeq platform, at a read length of ~30 million reads per sample.

Quality control of RNA sequence and gene expression data

Initial quality control checks of raw RNA sequence data were conducted using the FastQC software Babraham from Bioinformatics (www.bioinformatics.babraham.ac.uk/projects/fastqc). Sequence reads were then aligned to the latest version of the human reference genome (hg38) using the STAR (Spliced Transcripts Alignment to a Reference) alignment tool [47]. The number of reads mapped onto each gene in Ensembl's gene annotations for hg38 (version 99) was counted using FeatureCounts [48]. We plotted the distribution of log₁₀-transformed counts-per-million (CPMs) and, by visual inspection, set a threshold of $log_{10}CPM = 0.6$ for filtering out lowly expressed genes in order to minimise technical noise and reduce the multiple-testing burden (Supplementary Figure 7). After applying that threshold, 15060 out of 60642 genes were left for downstream analysis. TMM (trimmed mean of M-values)normalization (Robinson & Oshlack, Genome Biology, 2010) was then applied on the gene counts, gene-expression values were log₂-transformed, and observational-level theoretical variances were calculated using 'voom' for precision-weighting [49].

Differential gene expression

In order to evaluate potential sources of overall gene expression variation, we performed Principal Component Analysis (PCA) on the *voom*-transformed gene expression [49], plotting the samples along the first three principal components (Supplementary Figure 2). We then used the 'variancePartition' R package [50] to estimate the contribution to the variance in expression of each gene for the following variables: study subject, clinical group (schizophrenia (SZ)/control), treatment condition

(IFN- γ /IL-1 β /vehicle), donor age, and the fraction of all RNA sequencing 'reads' that were mapped to genes (i.e., 'assigned percent').

To prepare the expression data for linear mixed effects modelling, we applied *voomWithDreamWeights* ('variancePartition' R package) to the expression-level-filtered TMM-normalized counts. We then conducted linear mixed model regressions using *dream* ('variancePartition'), which allows modelling of interindividual variability by adding individual identifiers as a random effect in the regression model, as shown below [50, 51]. We also included the 'assigned percent' for each sample as a covariate, as we observed this variable to have a non-negligible effect on the overall variance of gene expression (Supplementary Figure 3). Participant age was not included as a covariate on the assumption that reprogramming samples to stem cells negates age-related effects [52, 53]. Age did seem to contribute to variation in gene expression in the variancePartition analysis (Supplementary Figure 3), but we attributed this to noise due to the small size of our sample. All the participants were male, so gender was not included as a covariate. The final model with an interaction term between clinical group and treatment was as follows:

Y_i = Group*Treatment + Assigned percent + Individual ID

(Where "Group", "Treatment" and "Assigned percent" were fixed effects and "Individual ID" was a random intercept effect).

Using contrasts, the following differential gene expression signatures were generated:

A. Vehicle-treated SZ NPCs vs vehicle-treated control NPCs (i.e., between SZ and control NPCs treated with vehicle).

- B. **IFNγ-treated control NPCs vs vehicle-treated control NPCs** (i.e., the effect of IFNγ stimulation on gene expression in the control NPCs).
- C. **IFNγ-treated SZ NPCs vs vehicle-treated SZ NPCs** (i.e., the effect of IFNγ stimulation on gene expression in the SZ NPCs).
- D. Interaction effect of IFNγ treatment in schizophrenia NPCs vs in control NPCs (i.e., how the transcriptional response to IFNγ stimulation differs in SZ NPCs compared to control NPCs).
- E. IL-1β-treated control NPCs vs vehicle-treated control NPCs (i.e., the effect of IL-1β stimulation on gene expression in the control NPCs).
- F. IL-1β-treated SZ NPCs vs vehicle-treated schizophrenia NPCs (i.e., the effect of IL-1β stimulation on gene expression in the SZ NPCs).
- G. Interaction effect of IL-1 β treatment in SZ NPCs vs in control NPCs (i.e., how the transcriptional response to IL-1 β stimulation differs in SZ NPCs compared to control NPCs).

Approximation of residual degrees of freedom and subsequent calculation of moderated eBayes *t*-statistics was done using the Satterthwaite method in 'dream' [51].

Gene set enrichment analysis

Our gene set enrichment analyses (GSEA) included 935 unique gene sets: 519 immune-related and 421 nervous-system/neural function related (5 overlapping). Of these, 135 were obtained from previous literature [54-57] and the remaining from either the Molecular Signature Database (HALLMARK and Gene Ontology biological process

gene sets) or the pathway databases KEGG, PANTHER, Pathway Commons and Reactome (see Supplementary Tables 4A-G for a full list of gene sets). GSEA assesses whether genes belonging to specific pathways or predefined sets of genes are over-represented in the significant or peri-significant results of a differential expression analysis. We used a linear mixed effects regression-based competitive gene set enrichment approach using the GSEA tool (Subramanian et al, 2005). GSEA was run on the seven signatures generated by the DGE analysis. We applied the *fgsea* function of the R package 'fgsea' [58], using the standardized Z-score obtained in the differential expression analysis to rank the genes and running 100,000 permutations. All gene sets containing fewer than five genes were excluded. Multiple testing correction was performed within fgsea using the false discovery rate (FDR) method, and gene sets with an FDR < 0.05 were considered significant.

The resulting gene sets showed substantial constituent similarity (Supplementary Figure 4), so we clustered them based on the overlap of the genes that belong to each gene set. This was done by calculating the Jaccard Similarity Index (which quantifies the intersection of two lists) between all pairs of significantly enriched gene sets and then applying a hierarchical clustering of gene sets based on the resulting dissimilarity matrix (1-Jaccard similarity). We then applied a cut-off of h=0.5 to the dendrograms to obtain clusters of significantly enriched gene sets (Supplementary Figure 5).

Enrichment of schizophrenia genes

To test whether differentially expressed genes in our experimental conditions were enriched for genes differentially expressed in *post-mortem* brain samples originating from SZ cases, we split genes by direction of effect (up- or downregulation) and assessed their overlap with genes differentially expressed in schizophrenia, according to Gandal et al [59]. Significance of this overlap was estimated using the Fisher's exact test in R, through the *GeneOverlap* package, assuming a genome size of 20000 protein-coding genes.

To test whether the differentially expressed genes in our experiments were enriched for GWAS-supported genes, we performed gene-set enrichment analysis using MAGMA. The summary statistics file from the schizophrenia GWAS performed by Pardiñas et al [60] was downloaded and pre-processed using standard quality control procedures, where variants with minor allele frequency < 0.01, or those in the extended MHC region on chromosome 6, from 25-34 Mb, were removed. The GWAS variants were annotated to a list of protein-coding genes provided by the authors, which included genes located on chromosomes 1 to 22 and X, allowing a window of 35 kb upstream and 10 kb downstream of each gene, as described previously [61]. Gene-level enrichment analysis was performed to identify genes more likely to be associated with schizophrenia according to the GWAS results, using the European subset of the 1000 Genomes Phase 3 as reference panel. Subsequently, we tested whether there was an enrichment of genes differentially expressed in our experimental models (excluding those that were noncoding) within these results. All tests were corrected for multiple testing using the false discovery rate method, according to the number of gene sets analysed per condition (i.e.,

two gene lists [up- and downregulated genes] from two groups each [control vs. SZ cell lines] = 4 comparisons per analysis).

Results

Demographic and sample details

The demographic and clinical characteristics of the six participants are described in Table 1. Subjects were male and of White British, or 'Other White' background. Ages ranged from 33 to 55 years old. The patients were diagnosed with paranoid schizophrenia, and controls were selected on the basis of having no history of neuropsychiatric disorders.

Validation of hiPSCs and NPCs

All hiPSC lines differentiated into embryoid bodies with three characteristic germ layers and expression of pluripotency markers NANOG, OCT4, SSEA4 and TRA-1-81 (Supplementary Figure 1). For each participant, one clone was used for the NPC induction. All hiPSC lines successfully differentiated into NPCs as determined by immunostaining for known NPC markers βIII-tubulin and Nestin (Figure 1).

Sources of variation in gene expression

We observed that the greatest source of variability across all samples was individual differences between the patients, as seen with principal component (PC) 1 (Supplementary Figure 2 A-C, left panels), as well as in the VariancePartition (Supplementary Figure 3) comparison. The VariancePartition analysis also revealed that the other variables included in the model (while having a low impact on overall gene expression variation individually) together explained at least 50% of the expression variance (Supplementary Figure 3). The importance of the clinical group was further supported by the clustering of SZ and control samples along the 2nd and 3rd principal components of PCA analysis (Supplementary Figure 2C, right panel).

Differential expression of genes and gene set enrichment analysis (GSEA)

Of the seven comparisons we made, four showed significant differential gene expression at FDR < 0.05: the effect of diagnosis (Signature A), and the three IFN γ treatment conditions (Signatures B-D). There were no statistically significant (FDR 5%) differentially expressed genes (DEGs) as a result of acute IL1- β treatment. A full table of DEGs can be found in Supplementary Tables 3A-G. Immune-related gene sets were among the top ten most significantly enriched for all seven signatures, and synaptic transmission-related gene sets were among the top ten most significantly enriched for the seven signatures. A full table of DEGs and gene set enrichment analysis (GSEA) results for each signature are presented below.

Effect of schizophrenia diagnosis (Signature A)

We first investigated the gene expression differences observed in SZ relative to control lines, in vehicle-treated NPCs. We found only one statistically significant DEG (Figure 2A), *AL132709.7*, a human-specific lncRNA gene which was overexpressed in patient lines (*FDR* = 0.0395; logFC = 3.111). Our sample was underpowered to detect other DGEs, but the GSEA revealed 26 significantly enriched gene-sets. The top five

gene sets were enriched among nominally significant upregulated genes in this comparison (Figure 2B), and the gene set with the lowest *p*-value was 'Lek2015 loss-of-function (90)' (*FDR* = 0.00098; normalized enrichment score (NES) = 1.36; genes in gene set = 3007), which contains 3007 genes that are intolerant to loss-of-function variants. This is consistent with previous schizophrenia genetic association studies that find associations with the same loss-of-function gene sets [55, 56].

Effect of IFNy treatment in control NPCs (Signature B)

We observed 1847 upregulated and 1533 downregulated genes in control NPCs (total = 3380 genes, out of 15061 tested) in response to the IFNY treatment, relative to the vehicle-treated lines (Figure 3A and Supplementary Table 1). We observed significant upregulation of *STAT1* (*FDR* = 5.572 x 10⁻⁶; logFC = 5.680), *STAT2* (*FDR* = 5.045 x 10⁻⁶; logFC = 2.247) and *JAK2* (*FDR* = 0.001; logFC = 2.111) – consistent with activation of the IFNY signal transduction pathways) – as well as *IRF1* (*FDR* = 3.23 x 10⁻⁶; logFC = 7.022), a key downstream signalling target of this cytokine [2, 62]. The genes whose expression was most significantly altered by IFNY treatment were *IFI27* (*FDR* = 2.97 x 10⁻⁶; logFC = 6.067) and *CD274* (*FDR* = 2.97 x 10⁻⁶; logFC = 6.386), both upregulated. Three genes encoding guanylate-binding proteins were among those that showed the highest fold change (logFC): *GBP1* (*FDR* = 3.11 x 10⁻⁵; logFC = 13.113). The upregulation of these genes is consistent with the role of guanylate-binding proteins (especially *GBP1*) in the inflammatory response associated with IFNY [63]. Our findings for this comparison were

also consistent with recent work which found MHC-I related genes among the most differentially expressed in IFNγ-treated control neural progenitors and neurons [2]. In our results (Supplementary Table 3A), key MHC-I related genes such as *HLA-A*, *HLA-B* and *HLA-C* are all consistently upregulated in response to IFNγ treatment.

We additionally tested whether the genes differentially expressed due to IFNy treatment overlapped with genes known to be differentially expressed in patients with SZ, based on gene lists provided by [64] (split by direction of effect: up- or down-regulation) and using the R package GeneOverlap [65]). We observed a significant overlap between the genes downregulated in the IFNy-treated control NPCs with genes downregulated in SZ (p = 7.00×10^{-4} , FDR = 0.0014, odds ratio (OR) = 1.3), but not with those that were upregulated (p < 0.05). Similarly, the genes upregulated in our model overlapped with those upregulated in SZ ($p = 1.20 \times 10^{-8}$, FDR = 4.80 x 10⁻⁸, OR = 1.5), but not with those that were downregulated (p < 0.05). We performed another gene-set enrichment analysis, using MAGMA [61], to test whether DEGs in our model were overrepresented in GWAS summary statistics from a large-scale schizophrenia GWAS [55]. This analysis did not identify evidence that IFNy-regulated genes in our model were associated with interindividual genetic variation contributing to schizophrenia susceptibility (p < 0.05). Collectively, these results suggest that IFNy signalling may impact neurodevelopment in a way that predisposes to schizophrenia, but that this may be independent from genetic effects. One caveat of the enrichment analysis using MAGMA is that it excludes genes within the MHC region due to the complex linkage disequilibrium structure at this locus, even though many of the genes in the MHC region are relevant for the IFNy response.

We further observed 168 pathways enriched when comparing IFN γ -treated cells and untreated cells in control NPCs. The gene set with the lowest *p*-value (Figure 3B) was 'immune system process' from Gene Ontology (*FDR* = 0.0002; *NES* = 2.41; genes in gene set = 1235), which consists of genes involved in the development or functioning of the immune system. All of the top ten gene pathways for this signature were overexpressed amongst genes upregulated in response to IFN γ treatment. While most of these were related to the immune response initiated by cytokine exposure, we observed two that were, notably related to synaptic function: 'post-synaptic density, human core' and 'presynapse'.

Effect of IFNy treatment in schizophrenia NPCs (Signature C)

We observed 1061 upregulated and 919 downregulated genes in SZ cell lines (*FDR* < 0.05, total = 1980 genes, out of 15061 tested) in response to the IFNγ treatment, relative to the vehicle-treated SZ lines (Figure 4A and Supplementary Table 2). The genes whose expression was most significantly altered by IFNγ treatment in the SZ neural progenitors were *STAT2* (*FDR* = 1.46 x 10⁻⁵; logFC = 2.6103), *IFl27* (*FDR* = 1.74 x 10⁻⁵; logFC = 6.331) and *STAT1* (*FDR* = 1.74 x 10⁻⁵; logFC = 5.453). Once again, *IRF1* (*FDR* = 3.543 x 10⁻⁵; logFC = 7.278) and *JAK2* (*FDR* = 0.0033; logFC = 1.846) were also significantly upregulated. Here too, the highest logFC was shown by *GBP1* (*FDR* = 3.91 x 10⁻⁵; logFC = 12.308), followed by the pseudogene *GBP1P1* (*FDR* = 0.0001; logFC = 11.116).

We saw that genes upregulated in SZ NPCs treated with IFN γ (*FDR* < 0.05) also significantly overlapped with those known to be upregulated in SZ patients (p = 1.10 x 10⁻

¹⁴, FDR = 4.40 x 10⁻¹⁴, odds ratio (OR) = 1.9) (but not those that were downregulated). However, the downregulated genes in the model did not overlap with those downregulated (or upregulated) in cases, suggesting that the SZ NPCs may respond differently to IFNγ when compared to NPCs from unaffected individuals. Ultimately, we did not observe an enrichment of GWAS-supported variants within the genes differentially expressed in SZ NPCs after treatment, as observed in control NPCs (i.e., Signature B).

Nevertheless, there were 132 pathways enriched in the comparison of IFN γ -treated SZ cells (versus untreated SZ lines); the gene set with the lowest *p*-value (Figure 4B) was, again, 'immune system process' (*FDR* = 0.0002; *NES* = 2.366; genes in gene set = 1235). The results in this comparison show activation of similar pathways in response to IFN γ in SZ lines as seen in control lines in the previous comparison (Figure 4B). However, the transcriptional response appears attenuated, as we observed fewer DEGs overall.

Interaction effect of IFNy treatment and schizophrenia diagnosis (Signature D)

To get a general picture of whether the patient NPCs respond differently to IFNγ treatment compared to how control NPCs do, we first examined the overlap of DEGs between signatures B and C (presented as a Venn diagram in Supplementary Figure 6). It was evident that of the 4137 genes that responded to IFNγ in any of the two groups, only 1223 genes were in common to both, meaning that there are 2914 genes that appeared to respond differentially to IFNγ treatment between SZ and control NPCs. The following signature (Signature D) effectively assesses the same overlap; but subjects this

comparison to an additional test of statistical significance (essentially omitting any of the 2914 genes observed in the initial comparison that may have differentially responded to treatment by chance). For this interaction term, we performed multiple-testing correction on the *p*-value obtained for the 4137 genes that are differentially expressed in response to IFNy in any condition (controls and/or SZ cells). At *FDR* < 0.05 there were 359 genes that respond significantly differently to IFNy between control and SZ cells (Figure 5A; Table 2); most significantly the mitochondrial complex genes *NDUFA2* (*FDR* = 0.0003; logFC = -0.591) and *NDUFS3* (*FDR* = 0.0006; logFC= -0.330) and the lncRNA gene *AC092279.2* (*FDR* = 0.0006; logFC = 0.645). Indeed, the previous comparisons show that the mitochondrial genes are overexpressed in response to IFNy in the control NPCs (logFC = 0.405, *NDUFA2*; logFC = 0.145, *NDUFS3*) but are underexpressed in response to IFNy in SZ NPCs (logFC = -0.187, *NDUFA2*; logFC = -0.185, *NDUFS3*). *AC092279.2* shows the opposite profile, responding with underexpression in control cells (logFC = -0.517) and overexpression in SZ cells (logFC = 0.123).

There were 20 gene sets that were differentially expressed in this comparison. These gene sets were comprised of genes that showed different transcriptional responses to IFN γ in SZ NPCs compared to control NPCs (Figure 5B). The most significantly different of these gene sets were 'post-synaptic density (PSD), human core' (*FDR* =0.001; *NES* = -1.72; genes in gene set = 654), which includes several notable genes including the Alzheimer's risk gene *APOE*, autism and schizophrenia risk genes *NRXN1, CYFIP1* and *SHANK1-3*, NMDA receptor gene *GRIN1,* and *DLG4,* which encodes the postsynaptic density protein PSD-95; as well as a 'presynapse' gene set, which includes genes that regulate the pre-synaptic 'active zone' and synaptic vesicle

formation [54, 55, 66] – notable genes in this gene set include *SV2A, MAOA* and several Na+/K+ transport ATPase genes. In other words, genes influencing synaptic transmission showed a particularly attenuated response to IFNγ treatment in SZ NPCs.

Effect of IL-1 β treatment on gene expression (Signature E, F & G)

We observed no differentially expressed genes associated with the effect of IL-1 β treatment in either control (Signature E, Supplementary Figure 8A) or SZ (Signature F, Supplementary Figure 9A) NPCs (FDR > 0.05). We hypothesized that this could have been caused by the reduced expression of the main IL-1 β receptors in cells at the neural progenitor stage. We assessed the expression of the genes encoding the IL-1 β receptors *IL1R1, IL1R2,* and *IL1RAP,* and compared these to the expression of the IFN γ receptor genes *IFNGR1* and *IFNGR2*. We observed that the IL-1 β receptor genes exhibited visibly lower expression relative to the IFN γ receptors (Figure 6), which may explain the lower responsiveness of the NPCs to IL-1 β stimulation. In addition, these results suggest that IFN γ signalling may be more relevant than IL-1 β at this developmental stage and/or for this cell type (NPCs).

Nevertheless, we observed 123 and 112 gene sets enriched for signatures E and F, respectively (Supplementary Figure 8B and 9B). For signature E, 'regulation of immune system process' was the most significant term (FDR = 0.0005; NES = 1.55; genes in gene set = 888); and 'Lek2015 loss-of-function (90)' [54, 55, 66] was the most significant term for signature F (FDR = 0.0008; NES = -1.97; genes in gene set = 3007). The fact that there are significant gene set enrichment terms for this comparison despite there being

no DEGs suggests that there are indeed effects of IL-1 β on transcription, but that our sample is underpowered to detect these individually. This method incorporates the expression signal from peri-significant genes and restricts the number of tests performed, reducing the multiple testing burden (instead of analysing 15061 genes for differential expression, as in the DGE analysis, our enrichment analysis tests 895 gene sets).

As there were gene sets significantly enriched for these signatures, we also tested whether genes that were differentially expressed at a more lenient threshold of *FDR* < 0.1 were enriched for genes differentially expressed in SZ. We observed that genes upregulated in SZ NPCs treated with IL-1 β showed a significant overlap with those upregulated in schizophrenia cases (p = 0.0013, FDR = 0.0052, odds ratio (OR) = 1.6). Further, using MAGMA, we observed that genes downregulated in the SZ NPCs were enriched with GWAS-supported variants associated with SZ (β = 0.19, SE = 0.06, P = 8.56 x 10⁻⁴, FDR = 3.42 x 10⁻³). While these results corroborate a role for IL-1 β signalling in schizophrenia, it is likely that the effects of this cytokine on NPCs are limited due to the lack of other cell types (such as microglia) in the culture system used or occur at another this involves another developmental stage.

The interaction effect between IL-1 β and SZ (Signature G) also did not yield any significantly differentially expressed genes (Supplementary Figure 10A) but did yield 15 gene sets that were significantly enriched. Of these, 'regulation of ligase activity' had the lowest *p*-value for signature G (*FDR* =0.0015; *NES* = -2.15; genes in gene set = 121; Supplementary Figure 10B). These enrichment results also show a suppression of transcriptional response to IL-1 β exposure in SZ NPCs, with pathways regulating the

post-synaptic density and presynapse once again amongst the top ten (Supplementary Figure 10B).

Discussion

In this study, we sought to assess how the cytokines interferon-gamma (IFN γ) and interleukin-1 beta (IL-1 β) interact with genetic profiles associated with schizophrenia, to better understand the increased susceptibility to schizophrenia seen in offspring of mothers exposed to infection during pregnancy. We hypothesised that cortical neural progenitor cells (NPCs) derived from patients with schizophrenia would respond differently to IFN γ or IL-1 β exposure compared to those of healthy controls.

We performed a preliminary analysis to evaluate the transcriptomic differences between our schizophrenia and controls NPCs without treatment stimulation and found only one differentially expressed gene. This result contrasts with the at least 80 DEGs found in schizophrenia patient-derived NPCs by Evgrafov et al [66]. However, the fact that our sample size is small and that we use a more stringent significance threshold (*FDR* < 0.05 as opposed to *FDR* < 0.1) may account for that difference. The identification of 26 gene-sets significantly enriched for this comparison further supports the idea that, while there are noticeable differences between schizophrenia and control NPCs at the transcriptomic level, our study may be underpowered to detect those for each gene individually.

We then assessed whether IFN γ treatment alters transcriptional responses in control neural progenitors and found that there were 3380 significant differentially expressed genes in response to IFN γ treatment. This is interesting itself, as the cell cultures used in this study do not contain glial cells – supporting the notion presented by

our group in a recent study [2] that human NPCs can launch an immune response independent of microglia. Immune responses in the brain are thought to be predominantly mediated by glia [67], but the fact that NPCs are themselves responsive to a proliferation of IFNy indicates that immunity in the brain extends beyond glial cells. This is also consistent with [68], who show that neuronal co-culture with activated microglia is sufficient to induce deficits in the neurons – suggesting that cytokines might come from glia, but neurons can respond to them independently. Moreover, in both control and schizophrenia NPCs, IFNy treatment activated the canonical JAK-STAT signalling pathway, as would typically be seen in response to viral infection. Our findings for this comparison were also consistent with recent work which found MHC-I related genes among the most differentially expressed in IFNy-treated control neural progenitors and neurons [2]. In our results (Supplementary Table 3A), key MHC-I related genes such as HLA-A, HLA-B and HLA-C are all consistently upregulated in response to IFNy treatment. However, the genes most significantly upregulated on IFNy exposure were IFI27 and CD274. IFI27 encodes Interferon Alpha Inducible Protein 27, which is involved in interferon-induced apoptosis and is considered to be a biomarker that differentiates between viral and bacterial infection [69]. CD274 encodes a receptor ligand that binds to PD-1 receptors on T-cell surfaces, inhibiting T-cell activation and antibody production an essential process for preventing autoimmunity [70]. Interestingly, there is some evidence that IFI27 is differentially expressed in transgenic mice that exhibit schizophrenia-like behaviours [71]. Similarly, CD274 is a member of several gene-sets found to be enriched in a study of de novo copy number variant associated to

schizophrenia risk [72]. In light of this evidence, these genes may be promising candidates for future studies exploring the link between MIA and schizophrenia risk.

In schizophrenia NPCs, there were fewer differentially expressed genes in response to IFNγ treatment: only 1980. This may be because these cells are not able to activate in response to IFNγ to the same extent as controls. Physiologically, this could mean that cells from healthy donors exhibit a compensatory transcriptional response to infection, which is inhibited in cells from donors with schizophrenia. The two genes showing the most divergent response to IFNγ in SCZ and control cells (*NDUFA2* and *NDUFS3*) being mitochondrial complex I genes further suggests that schizophrenia donor cells are relatively driven to conserve energy in response to an infection, while healthy donor cells are able to expend more energy to restore health [68]. Indeed, there is some recent evidence for differences in expression of mitochondrial genes in rodents susceptible to MIA [14]. This may also be related to pre-existing deficits in mitochondrial function in schizophrenia lines, as it is well established that mitochondrial dysfunction contributes to the pathophysiology of schizophrenia [73].

The gene sets significantly enriched for IFNy treatment in both schizophrenia and control NPCs largely converged in function, as expected, upon immune regulation. The gene sets that responded most differently to IFNy in schizophrenia lines were those regulating the postsynaptic density, presynapse, and presynaptic active zone. This result shows that the genes involved in the aberrant response to immune activation by schizophrenia NPCs are involved in synaptic transmission, which fits with previous work by Shatz [82], as well as models of schizophrenia from other fields of neuroscience, including the concept of schizophrenia as a disorder of synaptic 'dysconnection' in

computational neuroscience – a promising bridge between two very different but equally rich views of the same disorder. The dysconnection hypothesis suggests a dysregulation of neuromodulation (particularly across glutamatergic synapses) lies at the core of the various factors contributing to schizophrenia susceptibility [74, 75].

Our results did not reveal any significant DEGs in response to IL-1 β treatment in SCZ or control NPCs. This is likely due to the low expression of the IL1 receptor 1 (IL1R1) gene that we observed, as blocking IL1R1 has previously been shown to significantly reduce the influence of IL-1 β on NPCs [10]. However, this low IL-1 receptor expression was surprising in light of previous work showing high *IL1R1* (but not *IL1R2*) expression in rat ventral mesencephalon neural progenitors [10]. It is possible that there is lower *IL1R1* expression is upregulated at a later stage of prenatal neurodevelopment in humans than in rats, demonstrating the significance of leveraging human systems in the study of human neurodevelopment. It is also possible that that, by the 24-hour time point, any effect induced by IL-1 β may have been lost – i.e., the effects of IL-1 β may have been lost – i.e., the effect h

Nevertheless, pathway analyses did reveal significant enrichment of gene sets in response to IL-1 β treatment, suggesting that there were transcriptional effects in both schizophrenia and control NPCs in response to IL-1 β , but our sample size only allowed their identification at the gene set level (at which the multiple-testing burden is smaller). The GSEA analyses revealed substantially different gene set enrichment profiles for the IL-1 β signatures in schizophrenia and control NPCs. Among the significantly enriched gene sets in control lines treated with IL-1 β were (as with IFN γ) genes regulating the immune response, the presynapse and the post-synaptic density – all upregulated.

However, none of these gene sets were among those most significantly enriched in IL-1 β -treated schizophrenia lines; instead, almost all of these were sets of genes involved in central nervous system development and neuronal morphogenesis – all downregulated. The genes that responded differently to IL-1 β in schizophrenia vs control NPCs were enriched in gene sets involved in cell division, antigen presentation and, once again, synaptic transmission. The most interesting finding to emerge from all these analyses was indeed that genes involved in synaptic transmission respond differentially to IFN γ and IL-1 β exposure in schizophrenia NPCs compared to control NPCs

The current study was primarily limited by the relatively small sample size, which would warrant future replication studies. It was also surprising to find that donor age had some influence over the variance in gene expression in the sample, as one would also expect a negation of age-related epigenetic effects [79]. It is likely that this is due to noise (again as a result of the small sample size), which emphasises the importance of replicating these findings. We also administered only a single, acute dose of cytokine treatment to our NPC cultures: in future studies, it would be interesting to examine the effect of chronic treatment. For IL-1 β effects, future studies looking at different developmental time points and cell types could elucidate this cytokine's role in neurodevelopmental processes associated with schizophrenia.

In summary, having conducted the first transient cytokine exposure study using hiPSC-derived neural progenitor cells from patients with schizophrenia, we have found that immune activation induced by IL-1 β and IFN γ elicits transcriptional changes that may alter the course of subsequent neurodevelopment. There were two particularly significant take-home messages from this study, as follows. First, there does appear to be a significant

transcriptional response to IFNy treatment in NPCs, with differential expression implicating mitochondrial complex genes, which are underexpressed in response to treatment in schizophrenia lines. Second, our findings highlight pre- and post-synaptic genes as differentially expressed in response to IFNy, and differentially regulated in response to treatment in schizophrenia NPCs. In other words, schizophrenia NPCs do not upregulate synaptic genes in response to a cytokine challenge as much as control NPCs do. This is also consistent with previous literature, including large-scale transcriptome-wide association studies [56] and the recent Psychiatric Genetics Consortium study showing synaptic genes to be the most enriched for schizophrenia risk [81]. This is particularly interesting as NPCs do not have synapses. It could be that these early changes impact synaptic development after these cells differentiate into neurons. Indeed, our previous research shows that IFNy induces molecular and cellular changes in NPCs that persist even when these cells differentiate into neurons [2]. Our findings exemplify differences in how the brains of people with schizophrenia may have responded to infection or inflammation during prenatal development and suggest immune insults early in life can alter neurotransmission. Finally, we identify new gene targets for future research on the influence of maternal immune activation on schizophrenia susceptibility and resilience.

Author contributions: A.B. wrote the manuscript, conducted the cell culture and experimental treatments, and worked with H.I. on quality control of the genetic data and statistical analyses. H.I. worked on the analyses as well as imputation and quality control of the genetic data and edited the manuscript. P.R., L.D.P and R.N. assisted with the cell culture and experimental treatments. R.R.D conducted the *GeneOverlap* analyses and

edited the manuscript. P.J.M.D., C.S., R.N., and D.P.S. set up the experimental pipelines, conducted cell culture and cellular analysis of iPSC and neural progenitor cells, G.M., C.I., R.N. and J.P. were involved in obtaining ethical approval, sample collection and reprogramming of keratinocyte samples. S.B. was responsible for recruitment and clinical evaluation of patients whose hair samples were used in this study. A.V., T.P, E.B., J.P. and D.P.S. edited and provided guidance on the manuscript and interpretation of the results. D.P.S is the principal investigator, corresponding author and PhD supervisor to A.B., oversaw the project and developed the study with A.B., provided essential guidance throughout, as well as writing and editing the manuscript and figures. All of the co-authors provided intellectual input to the study.

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Figure Legends

Figure 1. Validation of neural progenitor cells. Successful differentiation to neural progenitor state was confirmed by staining at Day 20 for NPC markers, Nestin and β -III-tubulin. DAPI was used for baseline nuclear staining. Scale bar = 50µm

Figure 2. Expression differences between NPCs from cases vs. controls (signature A) at the gene and pathway level. A. The y-axis here shows statistical significance (-log10 p-value) of differential expression of genes in untreated cells from patient donors compared to gene expression in cells from untreated control donors. The x-axis shows the log₂ fold change of expression of those genes in schizophrenia cell lines vs control cell lines. **B.** The top 10 significantly enriched gene set clusters (the gene set with the lowest p-value in each cluster is labelled on the x-axis). Data-points are sized according to significance (-log10 p-value) and coloured according to normalised enrichment score (NES), with blue indicating downregulation and red indicating upregulation.

Figure 3. Expression differences between NPCs from IFNγ treated versus untreated control NPCs (signature B) at the gene and pathway level. A. The volcano plot shows, on the y-axis, the statistical significance (-log10 p-value) of differential expression of genes in IFNγ-treated control NPCs compared to untreated control NPCs. The x-axis is the magnitude of change (log₂ fold change) in expression of those genes due to after IFNγ treatment. **B.** The top 10 significantly enriched gene set clusters (the gene set with the lowest *p*-value in each cluster is labelled on the y-axis). Please see Supplementary Spreadsheets 4A-G for full lists of enriched gene sets for each of the signatures. Datapoints are sized according to significance (-log₁₀ p-value) and coloured according to normalised enrichment score (NES), with blue indicating downregulation and red indicating upregulation. **Figure 4. Expression differences between NPCs from IFNγ treated versus untreated schizophrenia NPCs (signature C) at the gene and pathway level. A.** The volcano plot shows, on the y-axis, the statistical significance (-log₁₀ p-value) of differential expression of genes in IFNγ-treated schizophrenia (SCZ) NPCs compared to untreated SCZ NPCs. The x-axis is the magnitude of change (log₂ fold change) in expression of those genes due to after IFNγ treatment. **B.** The top 10 significantly enriched gene set clusters (the gene set with the lowest *p*-value in each cluster is labelled on the y-axis). Please see Supplementary Spreadsheets 4A-G for full lists of enriched gene sets for each of the signatures. Data-points are sized according to significance (-log₁₀ p-value) and coloured according to normalised enrichment score (NES), with blue indicating downregulation and red indicating upregulation.

Figure 5. Interaction effect between IFNy-treatment and diagnostic group on gene expression (Signature D). A. The scatterplot shows IFNy response results for Signature in the 4137 genes that responded differentially to IFNy in Signatures B and C. DEGs for control cells are on the *x*-axis and DEGs for schizophrenia cells are on the *y*-axis. The data are coloured by signed $-\log_{10}$ FDR obtained for the interaction term (with blue indicating downregulation and red indicating upregulation). The 359 significant genes that are significant in the interaction are labelled. **B.** The top ten significantly enriched gene set clusters (the gene set with the lowest *p*-value in each cluster is labelled on the *y*-axis). Data-points are sized according to significance ($-\log_{10} p$ -value) and coloured according to normalised enrichment score (NES), with darker blue indicating greater downregulation.

Figure 6. Expression of cytokine receptors in schizophrenia cell lines compared to

controls. A. Shows the differential gene expression in schizophrenia (SCZ) compared to controls (CON) the four cytokine receptor genes that passed low-expression filtering. Values that lie above minimum expression threshold ($\log_2 CPM = 0.6$) are shown in red. **B.** Shows the differential gene expression in schizophrenia (SCZ) compared to controls (CON) the four cytokine receptor genes that passed low-expression filtering. Values that lie above minimum expression threshold ($\log_2 CPM = 0.6$) are shown in red.

Cell line	Diagnosis	Year diagnosed	Medication	Age	Gender	Ethnicity	Reprogrammed by
044	Schizophrenia	2011	Risperidone	33	Male	White British	Sendai virus
115	Schizophrenia	2010	Aripiprazol	43	Male	White British	Sendai virus
138	Schizophrenia	2008	Risperidone,	39	Male	Black British	Sendai virus
			Mirtazapine				
M1	Control	-	-	55	Male	White British	Lentivirus
M2	Control	-	-	35	Male	White British	Lentivirus
M3	Control	-	-	35	Male	White British	Sendai virus

Table 2. Top 20 genes significantly differentially expressed in IFN γ -treated compared to untreated cell lines in schizophrenia versus controls (Signature D) – all of these are downregulated. The right side of the table shows the effect of IFN γ treatment on the same genes in controls only, for comparison. Please see Supplementary Spreadsheet 3D for differential expression results for all genes in this comparison. A negative logFC indicates downregulation.

	IFNy effect in schizophrenia versus in controls				IFNy effect in controls					
Gene	Log Fold	Average ²	<i>p</i> -value	FDR ¹	Log Fold	Average	<i>p</i> -value ³	FDR ¹		
Symbol	Change	Expression	<i>p</i>		Change	Expression	<i>p</i>			
NDUFA2	-0.591	4.976	1.00E-06	3.27E-04	0.405	4.976	0.000	0.000		
NDUFS3	-0.330	6.229	5.00E-06	5.84E-04	0.145	6.229	0.000	0.005		
SS18L2	-0.587	4.809	6.00E-06	5.84E-04	0.153	6.628	0.001	0.009		
TMEM14C	-0.411	6.628	7.00E-06	5.84E-04	0.342	4.809	0.000	0.002		
AC092279.2	0.645	3.906	9.00E-06	6.19E-04	-0.517	3.906	0.000	0.001		
BEX2	-0.291	5.997	2.30E-05	1.38E-03	0.307	5.997	0.000	0.000		
RBX1	-0.414	6.291	3.20E-05	1.64E-03	0.445	6.291	0.000	0.000		
MPLKIP	-0.506	5.272	4.30E-05	1.93E-03	0.322	5.272	0.000	0.004		
COX6A1	-0.605	5.983	4.90E-05	1.95E-03	0.418	5.983	0.000	0.003		
UQCRQ	-0.669	5.772	5.50E-05	1.96E-03	0.392	5.772	0.000	0.006		
AL033519.3	-1.604	0.605	6.90E-05	2.26E-03	0.651	0.605	0.004	0.026		
ALG14	-0.597	3.048	1.29E-04	2.26E-03	0.176	3.048	0.024	0.082		
ATP5F1E	-0.714	7.683	2.50E-04	2.26E-03	0.430	7.683	0.001	0.012		
BPNT1	-0.661	4.865	8.50E-05	2.26E-03	0.307	4.865	0.001	0.012		
BTF3	-0.371	8.866	1.91E-04	2.26E-03	0.168	8.866	0.003	0.025		
CA3	-0.936	1.623	2.51E-04	2.26E-03	0.089	1.623	0.432	0.574		
CHCHD2	-0.430	6.979	1.60E-04	2.26E-03	0.378	6.979	0.000	0.003		
COA3	-0.507	5.485	2.22E-04	2.26E-03	0.490	5.485	0.000	0.002		
COPS9	-0.804	5.617	9.30E-05	2.26E-03	0.512	5.617	0.000	0.007		
1		1								

COX7C -0.552 7.975 1.50E-04 2.26E-03 0.409 7.975	0.000	0.005
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¹ False Discovery Rate

² Expression of the gene in (TMM-normalized) log₂ CPMs (counts-per-million) averaged across all samples

³ Uncorrected *p*-values

Table 3. Top 20 gene sets significantly overrepresented among DEGs in IFNγ-treated compared to untreated cell lines in schizophrenia versus controls (Signature D). The right side of the table shows the effect of IFNγ treatment on the same genes in controls only, for comparison. Please see Supplementary Spreadsheet 4D for differential expression results for all genes in this comparison.

	IFNy effect in schizophrenia versus in controls					IFNγ effect in controls		
Gene Set	Database	<i>p</i> -value	ES ¹	NES ²	Number of genes in set	<i>p</i> -value	ES ¹	NES ²
PSD human core	OP	1.07E-05	-0.32686	-1.719	654	1.19E- 05 1.25E-	0.297	1.764
Presynapse	OP	1.11E-05	-0.37736	-1.944	465	05 1.30E-	0.369	2.136
Synaptic vesicle	OP	1.15E-05	-0.35428	-1.787	353	05 1.31E-	0.360	2.028
Neutrophil degranulation	Reactome	1.16E-05	-0.33523	-1.681	330	05 1.43E-	0.444	2.483
Presynaptic active zone	OP	1.27E-05	-0.44027	-2.068	177	05 1.53E-	0.402	2.085
Regulation of ligase activity	GO	1.34E-05	-0.44658	-1.994	121	05 1.53E-	0.505	2.402
Positive regulation of ligase activity Cdc20:Phospho-APC/C mediated	GO	1.37E-05	-0.47269	-2.056	102	05 1.58E-	0.505	2.402
degradation of Cyclin A	Reactome	1.43E-05	-0.54536	-2.224	70	05	0.599	2.661

APC/C:Cdh1 mediated degradation of								
Cdc20 and other APC/C:Cdh1 targeted						1.58E-		
proteins in late mitosis/early G1	Reactome	1.43E-05	-0.54725	-2.232	70	05	0.601	2.668
APC/C:Cdc20 mediated degradation of						1.60E-		
Securin	Reactome	1.44E-05	-0.57565	-2.316	65	05	0.642	2.806
						1.60E-		
Activation of NF-kappa-B in B cells	Reactome	1.44E-05	-0.53276	-2.137	64	05	0.648	2.826
						1.60E-		
Autodegradation of Cdh1 by Cdh1:APC/C	Reactome	1.45E-05	-0.57029	-2.281	63	05	0.650	2.825
E3 ubiquitin ligases ubiquitinate target						1.64E-		
proteins	Reactome	1.48E-05	-0.57381	-2.195	50	05	0.549	2.273
						1.62E-		
Hedgehog ligand biogenesis	Reactome	2.93E-05	-0.55385	-2.166	56	05	0.684	2.900
Cross-presentation of soluble exogenous	Reactome					1.65E-		
antigens (endosomes)		3.01E-05	-0.58254	-2.170	44	05	0.748	3.007
Antigen processing: Ubiquitination &						1.34E-		
Proteasome degradation	Reactome	3.56E-05	-0.34313	-1.697	283	05	0.448	2.466
Antigen processing and presentation of						1.43E-		
exogenous peptide antigen via MHC1	GO	4.42E-05	-0.53871	-2.093	54	05	0.549	2.833
						1.57E-		
Downstream TCR signaling	Reactome	7.04E-05	-0.4687	-1.958	80	05	0.634	2.889
						2.80E-		
Ligand gated channel activity	GO	7.05E-05	0.40826	1.952	85	05	-0.482	-2.412
						1.65E-		
Reactive oxygen species pathway	Hallmark	7.51E-05	-0.5478	-2.050	45	05	0.579	2.338

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Α



–log10Pvalue

4.5

4.9

NES

4.9

В



Α



Log Fold Change

В







В

Α

