A human neurodevelopmental model for Williams syndrome

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Williams syndrome is a genetic neurodevelopmental disorder characterized by an uncommon hypersociability and a mosaic of retained and compromised linguistic and cognitive abilities. Nearly all clinically diagnosed individuals with Williams syndrome lack precisely the same set of genes, with breakpoints in chromosome band 7q11.23 (refs 1-5). The contribution of specific genes to the neuroanatomical and functional alterations, leading to behavioural pathologies in humans, remains largely unexplored. Here we investigate neural progenitor cells and cortical neurons derived from Williams syndrome and typically developing induced pluripotent stem cells. Neural progenitor cells in Williams syndrome have an increased doubling time and apoptosis compared with typically developing neural progenitor cells. Using an individual with atypical Williams syndrome6,7, we narrowed this cellular phenotype to a single gene candidate, frizzled 9 (FZD9). At the neuronal stage, layer V/VI cortical neurons derived from Williams syndrome were characterized by longer total dendrites, increased numbers of spines and synapses, aberrant calcium oscillation and altered network connectivity. Morphometric alterations observed in neurons from Williams syndrome were validated after Golgi staining of post-mortal layer V/VI cortical neurons. This model of human induced pluripotent stem cells8 fills the current knowledge gap in the cellular biology of Williams syndrome and could lead to further insights into the molecular mechanism underlying the disorder and the human social brain.

This study included participants with a clinical diagnosis of Williams syndrome (WS): individuals harbouring typical gene deletions in the Williams–Beuren syndrome critical region1,9 (WS17, 25, 77 and 79) and an individual with atypical WS with a partial deletion (pWS88) as well as typically developing (TD) participants (TD55, 59, 63 and 70) (Fig. 1a and Extended Data Fig. 1a). After a series of cognitive and social pathologies in humans, remains largely unexplored. Here we investigate neural progenitor cells and cortical neurons derived from Williams syndrome and typically developing induced pluripotent stem cells. Neural progenitor cells in Williams syndrome have an increased doubling time and apoptosis compared with typically developing neural progenitor cells. Using an individual with atypical Williams syndrome6,7, we narrowed this cellular phenotype to a single gene candidate, frizzled 9 (FZD9). At the neuronal stage, layer V/VI cortical neurons derived from Williams syndrome were characterized by longer total dendrites, increased numbers of spines and synapses, aberrant calcium oscillation and altered network connectivity. Morphometric alterations observed in neurons from Williams syndrome were validated after Golgi staining of post-mortal layer V/VI cortical neurons. This model of human induced pluripotent stem cells8 fills the current knowledge gap in the cellular biology of Williams syndrome and could lead to further insights into the molecular mechanism underlying the disorder and the human social brain.

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The impact of the genome-wide Williams–Beuren syndrome chromosome region deletion was determined by unbiased RNA sequencing (RNA-seq) (Extended Data Fig. 3a–d). Differential expression analyses revealed misregulated genes among the three genotypes (Extended Data Fig. 3e–h, Extended Data Table 1 and Supplementary Tables 3–9). Gene ontology (GO) analyses of NPCs and neurons revealed biological processes relevant to the condition (Extended Data Fig. 3i, i, Extended Data Table 2 and 3 and Supplementary Table 10). Remarkably, ’cell adhesion,’ ’axon guidance’ and ’cell maturation’ were also among the top-ranking categories detected in an independent publication using WS NPC gene expression analysis13.

As suggested by the NPC global gene expression analyses, during the culture maintenance, typical WS NPCs became confluent more slowly than TD NPCs (Fig. 2a). After plating the same number of NPCs, we verified that the number of typical WS NPCs on day 4 was less than the TD NPCs (Fig. 2b and Extended Data Fig. 4a). To rule out the possibility that the difference in heterogeneity of iPSC-derived NPCs could result in this observation, the NPC population was fully characterized and no difference between WS and TDs were found (Fig. 1e–g and Extended Data Fig. 4b). We also used single-cell gene expression profiling to access the homogeneity of the NPCs (Fig. 2c–e and Extended Data Fig. 4c–g). We further investigated the proliferation of WS NPCs by performing BrdU labelling, immunostaining and fluorescence-activated cell sorting (FACS) (Extended Data Fig. 4h, i). Since no difference was found, we assessed apoptosis in WS NPCs using DNA fragmentation (propidium iodide) and caspase assay (Extended Data Fig. 4j). We found a significant increase in subG1

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(Fig. 2f, g) and caspase-positive populations (Fig. 2h, i) in WS NPCs, indicating increased apoptosis. **FZD9** is expressed in NPCs14 (Fig. 2j) and has been shown to regulate cell division and programmed cell death in different cell types15,16. In our study, **FZD9** was hemizygously deleted in the participants with typical WS, but retained in atypical pWS88 (Fig. 1a and Extended Data Fig. 1b). Thus, we hypothesized that **FZD9** regulates human NPC apoptosis. We transduced TD NPCs with a lentivirus carrying either short hairpin RNA (shRNA) against **FZD9** (shFZD9) or non-specific shRNA (shControl) and WS NPCs with lentiviruses carrying a **FZD9** cDNA construct (Fig. 2k, l). TD NPCs transduced with shFZD9 showed a reduction in the number of cells on day 4 (Fig. 2m) and an increase in the subG1 population (Fig. 2n) and caspase activity (Fig. 2o) compared with TD NPCs expressing the shControl. Similar results were observed in atypical pWS88 (Extended Data Fig. 4k–n). Restoring **FZD9** expression in typical WS NPCs brought the number of NPCs on day 4/day 0 to a similar level to TD NPCs. It also significantly reduced the apoptotic population to the TD level.

Since several Wnt genes were downregulated in WS NPCs (Extended Data Tables 2 and 3) and **FZD9** can be activated by Wnt ligands, we tested if we could rescue the NPC viability by treating cells with the GSK3 inhibitor CHIR98014 (ref. 17). First, we confirmed that the canonical Wnt pathway was affected in WS by measuring the Axin2 and SP5 expression levels, two universal Wnt target genes18,19. Both genes were significantly downregulated in WS cells compared with TDs (Fig. 2p, q). By treating WS NPCs with CHIR98014, we were able to rescue cell viability (Fig. 2r). Together, our results indicate a role for **FZD9** in NPC viability.

Our protocol generated a consistent population of forebrain neurons, confirmed by the pan-neuronal and subtype-specific cortical markers.
such as CTIP2 (layers V/VI) and SATB2 (layer III) (Fig. 1h, i). The neuronal population was also characterized by single-cell gene expression profiling (Fig. 3a–c and Extended Data Fig. 5a–f), revealing mostly glutamatergic neurons, with a small population of GABAergic (γ-aminobutyric-acid-releasing) neurons and glia (Fig. 3a). We did not detect significant variability in these subtypes of neurons expressing target genes or in the expression levels of several markers for cortical layers and neurotransmitters among the genotypes. However, we did detect differences in the expression of specific genes in these populations (Fig. 3b, c) that could lead to specific alterations in mature neurons. We focused specifically on markers for cortical layers V/VI, since pathologies affecting these layers have been reported in disorders with compromised social functioning, such as autism.

We found that typical WS iPSC-derived CTIP2-positive neurons had significantly higher total dendritic length, dendrite number and number of dendritic spines than TDs (Fig. 3d–g and Extended Data Fig. 6a–m). Interestingly, atypical pWS88 neurons were morphologically similar to TD neurons except for dendrite number (Fig. 3f). To determine whether the WS neuronal phenotype was cell autonomous or dependent on other cells or culture conditions, we recorded the dendritic growth over time. The result showed a faster dendritic growth rate in WS neurons compared with TD or pWS88 (Extended Data Fig. 6n–r).

### Figure 2 | Defect in apoptosis of WS-derived NPCs owing to haploinsufficiency of FZD9

- **a**: Representative images showing the difference in confluency between TD, typical WS and pWS88 iPSC-derived NPCs on day 4. Scale bar, 100 μm.
- **b**: Ratio of NPC number on day 4 over day 0 relative to TD. Violin plots of representative genes expressed in NPCs from single-cell analyses. Principal component analysis (PCA) was used to compare the expression levels in individual cells on the basis of the first two principal components.
- **c**: Percentage of cells expressing NPC, neuronal (RBFOX3) and neural crest (PAX7, contaminant population) related genes. WS and TD iPSC-derived NPCs show similar percentages of cells expressing target genes over defined cycle threshold (C) control values.
- **d**: Representative propidium iodide histogram showing an increase in subG1 population in typical WS NPCs.
- **e**: Percentage of population with high caspase activity.
- **f**: FZD9 protein expression in TD iPSC-derived NPCs. Schematic of FZD9 gain/loss of function experiments in NPCs. Expression level of FZD9 protein after treatment with shFZD9, shControl and FZD9 overexpression vectors, assessed by western blot analysis. 
- **g**: Ratio of NPC number on day 4 over day 0 relative to TD (m), percentage of subG1 population (n) and percentage of population with high caspase activity (o) when TD NPCs were treated with shFZD9 and shControl, and WS NPCs were overexpressed with FZD9.
- **h**: Rescue of WS NPC viability after CHIR98014 treatment. All data are shown as mean ± s.e.m. © 2016 Macmillan Publishers Limited, part of Springer Nature. All rights reserved.
Also, no differences were observed in the total dendritic length, segment number or spine density using NPCs plated at different cellular densities (Extended Data Fig. 6 s–u).

An increase in the number of dendritic spines per neuron could lead to an increase in synaptic contacts and, therefore, synaptic activity, which could result in functional alterations. WS neurons had significantly more glutamatergic excitatory synapses compared with TD and pWS88 (Fig. 3h, i and Extended Data Figs 6v and 7a) and an increased frequency of calcium transients with a higher percentage of signalling neurons in WS cultures (Fig. 3j–m and Extended Data Fig. 7b–f). Using multi-electrode array (MEA) electrophysiology, our data showed that WS neuronal cultures had a significant increase in spike frequency compared with TD-derived neurons (Fig. 3n, o and Extended Data Fig. 7g, h).

In an attempt to place our iPSC findings in the larger context of the cortical morphology of human participants at the gross anatomical and cellular levels, we conducted two sets of additional experiments to test predictions based on NPC and neuronal differences found in vitro. In addition to the total volume reduction in WS brains previously reported, multivariate analyses of variance (MANOVA) from...
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Figure 4 | Neuroanatomical and morphological alterations in WS human brains. **a**, Statistical parametric map of the vertex-wise group differences between TD and WS in cortical surface area (left hemisphere shown) assessed by structural MRI scans. Colour scales indicate the $P$ value for statistical test; blue, decrease; grey, no difference. Statistics are displayed on a template group-averaged cortical surface rendering of TD adult participants. **b**, Reduction in overall cerebral cortical surface area in WS. Data are shown as mean $\pm$ s.e.m.; $n$, number of brains analysed. ****$P < 0.01$, one-sided unpaired Student’s $t$ test. **c**, Representative images of post-mortem layer V/VI pyramidal neurons using Golgi staining (top) and their corresponding tracing (bottom) from TD and WS. **d–g**, Morphometric analysis showing significant increases in total dendritic length (d), dendritic spine numbers (e), dendritic segment number (f) and number of branching points (g) in WS compared with TD post-mortem cortical layer V/VI pyramidal neurons. Data are shown as mean $\pm$ s.e.m.; $n$, number of traced neurons. $*P < 0.05$, **$P < 0.01$, ***$P < 0.001$, two-sided unpaired Student’s $t$ test (d); two-sided unpaired Mann–Whitney test (e–g).

The morphometric data in combination with the increased glutamatergic gene expression and number of co-localized synaptic puncta observed in neurons from WS suggest that an increased number of synapses may result in the altered network activity, which could contribute to the characteristic behaviour of individuals with WS. Our study reveals that the WS phenotypes described here are the foundation for the understanding of the complex human social behaviour. This approach provides an additional strategy to study the cellular and molecular underpinnings of complex human attributes, such as language in a social environment.

Online Content | Methods, along with any additional 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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Supplementary Information is available in the online version of the paper.

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Author Contributions A.R.M., T.C. and C.A.T. designed the experiments and wrote the manuscript with input from K.S. and all authors. T.C. processed NPCs, generated and characterized iPSCs, NPCs and neurons, and performed cell number, proliferation, and apoptosis experiments as well as FZD9 knockdown and overexpression and statistical analysis. C.A.T. performed C1 single-cell analyses, synaptic quantification, calcium imaging, cell density experiments, live neuronal morphology analysis and statistical analysis. B.C.F. performed MEA recording, PCR for retrovirus silencing and Wnt pathway gene-expression analysis. B.C.F. and S.E.R. prepared astrocytes for co-culture experiments, NPC characterization by flow cytometry and CHIR 98014 experiments. K.S. designed all morphometry experiments with B.H.-M. and B.J., and co-wrote the manuscript to link the various levels of investigation from the whole-brain imaging findings to the cellular level. L.S. prepared Golgi staining for post-mortem neurons with help from K.L.H. and B.J. B.H.-M. obtained morphometric data on iPSC-derived neurons and post-mortem neurons. D.X.Y., M.C.M., C.A.T. and L.M. performed calcium transient experiments and statistical analysis. T.T.B. performed brain scan and statistical analysis with help from A.M.D. C.B. performed electrophysiological tests. M.D., W.W., P.L. and Y.M.S. performed neurocognitive and social tests. A.J., Y.M.S., and M.C.A. performed analyses and interpretation of social/neurocognitive tests. R.H.H. performed bioinformatics analysis. L.D. and J.R.K. confirmed deletion of all cells from participants with WS who donated them for reprogramming. E.H., U.B., F.H.G., K.S. and A.R.M. edited the manuscript for publication.

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 A.R.M. (muotri@ucsd.edu) or K.S. (ksemende@ucsd.edu).
METHODS

Participants for behavioural study and source of cells for reprogramming. The study protocols were approved by University of California San Diego and Salk Institute IRB/EscrO committees (protocols 141223ZF and 95-0001, respectively). Four TD individuals (ages 8–19 years) and five individuals with WS (ages 8–14 years; Extended Data Fig. 1a) were included in the analysis: four of the latter group had typical WS gene deletions and one (pWS88) had a partial deletion in the WS region. Informed consents were obtained from all participants or their parents as appropriate. Genetic diagnosis of WS was established using fluorescent in situ hybridization probes for elastin (ELN), a gene consistently associated with the deletion in the typical WS region. All of the participants with WS having confirmed genetic deletion exhibited the medical and clinical characteristics of the WS phenotype, including previously established cognitive, behavioural and physical features associated with the syndrome. A diagnosis of WS was confirmed on the basis of the Diagnostic Score Sheet (DSS) for WS (American Academy of Paediatrics Committee on Genetics, 2001), with a particular focus on the cardiovascular abnormalities and the characteristic facial features associated with the ELN deletion. The scores for the participants were at the mean for WS (9) or higher, with the individual with partial deletion in the WS chromosomal region (pWS88) scoring lower than the individuals with typical WS deletion. Similarly, pWS88 reported fewer symptoms with connective tissue and growth, his cognitive scores were slightly higher than the typical individuals with WS, and he did not demonstrate the disparity between verbal and visual–spatial abilities typical of WS. However, pWS88 did display behavioural and developmental features consistent with WS, including developmental delay, over-friendliness and anxiously.

Behavioural and neurocognitive tests. The participants were administered standard tests to quantify their non-verbal and verbal abilities, as well as versions of the WS cognitive and social profiles to capture the distinct pattern of strengths and weaknesses both within and across domains associated with the WS cognitive and social phenotype. Details of the tests and the measures tapping into the two profiles are presented in Extended Data Fig. 1. The WS cognitive profiles for the five participants with WS were constructed by calculating the log of predictive likelihood ratios under assumed normality for age-appropriate TD versus WS classifications on the basis of verbal and performance IQ (VIQ and PIQ), Beery Developmental Test of Visual–Motor Integration (VMI) and Peabody Picture Vocabulary Test (PPVT) standard scores, subject to availability. Predictive distributions were based on the published normative mean and s.d. for each of the tests employed, whereas for the WS classification the predictive distributions were determined using data from n = 81 (VIQ and PIQ), n = 56 (VMI) and n = 97 (PPVT) participants in a broader WS sample (described in Extended Data Fig. 1d). A tobit model was used to estimate parameters for individuals with WS on the VMI owing to the presence of floor effects. The WS social profiles for the five participants with WS were previously constructed using measures of social approach behaviour, emotionality/empathy and language use.

Confirmation of WS deletion. Quantitative PCR was used to define the breakpoints of deleted regions in DNA isolated from iPSCs, or lymphoblast cell lines for participants with WS, with probes spanning from CALN1 to WBSCR16 and template DNA. Taqman expression assays detecting the WS region genes were designed and synthesized with sequences shown in Supplementary Table 11. Quantitative PCR was performed on ABI PRISM 7900HT system and the results were analysed using SDS 3.2.

Cell collection, reprogramming and characterization. We avoided invasive sample collection methods such as skin biopsy or blood withdrawal by taking advantage of the natural loss of deciduous teeth as a source of somatic cells. We chose to reprogram dental pulp cells (DPCs) because these cells develop from the same set of early progenitors that generate neurons. Furthermore, the neurons derived from DPCs would be a distinct cell type compared with those generated from skin fibroblast-derived iPSCs, serving the purpose of this study. Deciduous teeth were collected when they fell out and were shipped to our laboratory in DMEM 1× (Mediatech) with 4% Pen/Strep (Mediatech). Dental pulp was pulled out, washed in PBS with 4% Pen/Strep and incubated in 5% TrypsLe (Gibco) for 15 min. Pulp was partly dissociated using needles and plated in culture medium (DMEM/F12 50:50, 15% FBS, 1%NEAA, 1% fungizone and 2% Pen/Strep). In 1–4 weeks, DPCs migrated out of the pulp and could be passaged and frozen as stock. DPCs in early passage (two and three) were reprogrammed using pMXs retroviruses expressing Yamanaka transcription factors (obtained from Addgene, Cambridge, Massachusetts)112. After 4 days, transduced DPCs were tetraysinized, plated on mouse embryonic fibroblasts and cultured using human embryonic stem cell (hESC) medium.

Karyotyping. All G-band karyotyping analyses were performed by Molecular Diagnostics Service (San Diego, California) and Children’s Hospital Los Angeles (Los Angeles, California).

Genotyping. Two hundred nanograms of DNA were processed and hybridized to the Illumina Human Core Human Exome BeadChip following manufacturer’s instructions. Illumina GenomeStudio V2011.1 with the Genotyping Module version 1.9.4 was used to normalize data and call genotypes using reference data provided by Illumina. Illumina’s cnv Partition and gada R packages were used instead of the Infinium copy number module. Following detection, the B Allele Frequency (RAF) and Log R Ratio (LRR) distributions were manually checked to determine additional CNVs not detected by the software. Sample identification/relatedness was assessed by comparing called genotypes for each sample. The absolute number of different genotypes was counted and the Euclidean distances were calculated to identify relatedness of the samples.

Teratoma assay. Dissociated iPS colony were centrifuged and resuspended in 1:1 matrigel and phosphate buffer saline solution. The cells were injected subcutaneously in nude mice. After 1–2 months, teratomas were dissected, fixed and sliced. Sections were stained with haematoxylin and eosin for further analysis. Protocols were previously approved by the University of California San Diego Institutional Animal Care and Use Committee.

Neural induction and neuronal differentiation. iPS were cultured on matrigel-coated dishes and fed daily with mTeSR for 7 days. On the next day, mTeSR was substituted by N2 medium (DMEM/F12 supplemented with 0.5% N2-Supplement (Life Technologies), 1 mM dorsomorphin (Tocris) and 1 mM SB431542 (Stemgent)) for 1–2 days. iPS colonies were lifted off, cultured in suspension on the shaker (95 r.p.m. at 37°C) for 8 days to form embryoid bodies and fed with N2 media. Embryoid bodies then were mechanically dissociated, plated on a matrigel-coated dish and fed with N2B27 medium (DMEM/F12 supplemented with 0.5% N2-Supplement, 0.5% B27-Supplement (Life Technologies), 1% penicillin/streptomycin and 20 ng/ml G4F-2). The emerging rosettes were picked manually, dissociated completely using accutase and plated on a poly-ornithine/laminin-coated plate. NPCs were expanded in N2B27 medium and fed every other day. To differentiate NPCs into neurons, G4F-2 was withdrawn from the N2B27 medium. NPCs and neurons were characterized for stage-specific markers by immunostaining and flow cytometry (NPCs only), expression profile by single-cell RT–PCR and RNA sequencing and electrophysiological property (neurons).

Total RNA extraction. Total RNA of DPGs, iPSCs, NPCs and neurons was extracted using TRIzol reagent (Life Technologies) according to the manufacturer’s protocols. Contaminating DNA in RNA samples was removed using TURBO DNase (Life Technologies) according to the manufacturer’s protocols. Quality and quantity of RNA-treated RNA were assessed using NanoDrop 1000 (Thermo Scientific).

PCR for exogenous retrovirus DNA silencing. RNA was extracted from iPScs as previously described using Trizol reagent (Life Technologies). CDNA was generated from the RNA using SuperScript III protocol according to the manufacturer’s instructions. PCR was performed using primers listed below at the following cycles: 94°C for 10 min; 35 repeats of 94°C for 30 s, 62°C for 30 s and 72°C for 1 min; and finally, 72°C for 7 min. As a positive control, the IOMX plasmid of the four vectors used on the reprogramming of the cells was placed along the samples as well as water as a negative template control for amplification. As an additional positive control for the endogenous genes, two HSIC lines were used along with our iPScs: H1 and HUES6 cells. Primers used were as follows. Endo-cMyc: forward, TTG AGG GCC ATC GTC GCG GGA; reverse, GCC TCG TCT GAA GGA AGG AIA TCC. Endo-Klf4: forward, GAA ATT CGG CTC CGG CTA CGA; reverse, CTG TGT GTT GTG AGT AGT GCC. Endo-OCt4/3: forward, GCT TCT CAC GAG CGC CCC GC; reverse, TGC GGG GCG ACA TGG GGA GAT C. Endo-SOx2: forward, GCC GAC TGG AAA CTT TTG TCG; reverse, GCC AGC GTG TAC GGG AGG ACG GAG TGA. Endo-Oct4 transgenes pMXs-TgUS: forward, GAT GCT GGA GCC AAT GAC; reverse, GCG TCC TGC CCT GGA CCA TTA. Endo-Klf4: forward, GAA TCG CCC TCG AGC TCC. Endo-Oct4 pMXs-Oct4/3-TgDS: reverse, TAC GGA CCT TGG AGA ATC CA. Endo-SOx2 pMXs-SOx2-TgDS: reverse, GAC GTC TCT GGA CCA TTT ATT AGA. Endo-Oct4 pMXs-Oct4/3-TgDS: forward, GTG GTC GTA CCG GGA AAT AC. Endo-Oct4 pMXs-Oct4/3-TgDS: reverse, TAC GGA CCT TGG AGA ATC CA. Endo-SOx2 pMXs-SOx2-TgDS: reverse, GAC GTC TCT GGA CCA TTT ATT AGA. Endo-Oct4 pMXs-Oct4/3-TgDS: forward, GAT GCT GGA GCC AAT GAC; reverse, GCG TCC TGC CCT GGA CCA TTA. Endo-Klf4: forward, GAA ATT CGG CTC CGG CTA CGA; reverse, CTG TGT GTT GTG AGT AGT GCC. Endo-OCt4/3: forward, GCT TCT CAC GAG CGC CCC GC; reverse, TGC GGG GCG ACA TGG GGA GAT C. Endo-SOx2: forward, GCC GAC TGG AAA CTT TTG TCG; reverse, GCC AGC GTG TAC GGG AGG ACG GAG TGA. Endo-Oct4 transgenes pMXs-TgUS: forward, GAT GCT GGA GCC AAT GAC; reverse, GCG TCC TGC CCT GGA CCA TTA. Endo-Klf4: forward, GAA ATT CGG CTC CGG CTA CGA; reverse, CTG TGT GTT GTG AGT AGT GCC. Endo-OCt4/3: forward, GCT TCT CAC GAG CGC CCC GC; reverse, TGC GGG GCG ACA TGG GGA GAT C.
Microarray. Three hundred nanograms of total extracted RNA from each sample were subjected to microarray by using the Affymetrix GeneChip one-cycle target labelling kit (Affymatix, Santa Clara, California) according to the manufacturer’s recommended protocols. The resultant biotinylated cRNA was fragmented and then hybridized to the GeneChip Human 1.0 ST Array (764,885 probes, 28,869 genes, 19,734 gene-level probe sets with putative full-length transcript support (GenBank and RefSeq)) on the basis of human genome, Hg18. Arrays were prepared at the University of California DNA Core Facility. Arrays were also arrayed by the Affyl (Affymetrix pre-processing)²⁹ Bioconductor software package for microarray data. Data were then normalized by the RMA (robust multichip averaging) method to background-corrected and normalized probe levels to obtain a summary expression of normalized values for each probe set. Normalized microarray samples were then clustered by a hierarchical approach based on a matrix of distances. Normalized expression data were used to create a distance matrix that was calculated on the basis of Euclidean distance between the transcripts over a pair of samples representing a variation between two samples. Having the distances for all pairs of samples, a linkage method is used to cluster samples in a dendrogram by using calculated distances (sample expression similarities). This method also creates a heat map to graphically show the expression correlation between the samples.

Gene expression analyses by qPCR. RNA samples were reverse transcribed into cDNA using the Super Script III First Strand Synthesis System (Invitrogen, California) according to the manufacturer’s instructions. Reactions were run on the Bio-Rad detection system using Sybr-green master mix (Bio-Rad). Primers were selected from Primerbank; validated database (http://pga.mgh.harvard.edu/primerbank/) and specificities were confirmed by melting curve analysis through a Bio-Rad detection system. Sequences of the primers are described in Supplementary Table 12. Quantitative analysis used the comparative threshold cycle method.³⁰ GAPDH was used as housekeeping gene. Each sample was run in triplicate.

RNA-seq and global gene expression analyses. The RNA-seq analyses were previously described by our group.³¹ Briefly, RNAs were isolated using the RNeasy Mini kit (Qiagen). A total of 1,000 ng of RNA was used for library preparation using the Illumina TruSeq RNA Sample Preparation Kit. The RNAs were sequenced on Illumina HiSeq2000 with 50 bp paired-end reads, generating 50 million high-quality sequencing fragments per sample on average. For validation purposes of biological samples subjected to RNA-seq, hESC and iPSC samples available from the manufacturer were subjected to microarray by using the Affymatrix GeneChip one-cycle target labelling kit (Affymatrix, Santa Clara, California) according to the manufacturer’s recommended protocols. The resultant biotinylated cRNA was fragmented and then hybridized to the GeneChip Human 1.0 ST Array (764,885 probes, 28,869 genes, 19,734 gene-level probe sets with putative full-length transcript support (GenBank and RefSeq)) on the basis of human genome, Hg18. Arrays were prepared at the University of California DNA Core Facility. Arrays were also arrayed by the Affyl (Affymetrix pre-processing)²⁹ Bioconductor software package for microarray data. Data were then normalized by the RMA (robust multichip averaging) method to background-corrected and normalized probe levels to obtain a summary expression of normalized values for each probe set. Normalized microarray samples were then clustered by a hierarchical approach based on a matrix of distances. Normalized expression data were used to create a distance matrix that was calculated on the basis of Euclidean distance between the transcripts over a pair of samples representing a variation between two samples. Having the distances for all pairs of samples, a linkage method is used to cluster samples in a dendrogram by using calculated distances (sample expression similarities). This method also creates a heat map to graphically show the expression correlation between the samples.

DNA fragmentation analysis. One million NPCs were harvested to single-cell suspension in 1 mL PBS, then fixed by addition of 3 mL of 100% ethanol and stored at 4°C for at least 2 h. NPC pellets were washed once with 5 mL PBS. After removal of PBS, cells were resuspended in 1 mL of propidium iodide (PI) staining solution (0.1% (v/v) Triton X-100, 10 μg/mL PI and 100 μg/mL RNase A in 1 × PBS). WS and TD NPC samples were analysed by FACS on a Becton Dickinson LSRI, and using FlowJo v7.6.5 to cluster populations (cells with fragmented DNA) was examined using FlowJo flow cytometry analysis software.

Caspase assay. Caspase activity was measured using a Green FLICA Caspases 3 & 7 Assay Kit (ImmuNoChemistry Technologies). Briefly, NPCs were harvested, washed and stained with 1 × carboxyfluorescein Fluorochrome Inhibitor of Caspase Assay (FAM-FLICA) reagent, 10 μg/mL Hoechst and 10 μg/mL propidium iodide (PI). Samples were analysed on the NC-3000 using the pre-optimized Caspase Assay. The population with caspase activity was used to analyse for apoptosis.

Proliferation assay. NPC proliferation was assessed using BDF Pharmingen BrdU Flow Kits (BD Biosciences) according to the manufacturer’s protocol. Briefly, NPCs were incubated with 1 μM BrdU for 45 min at 37°C and harvested to single-cell suspension. NPCs were then fixed and permeabilized using BD Cytofix/Cytoperm Buffer and stained using FITC-conjugated anti-BrdU antibody and 7-aminoactinomycin D (7-AAD), a fluorescent dye for labelling DNA. Fluorescence-activated cell sorting (FACS) was done on LSReFortessa (BD Biosciences) and, to obtain the percentage of the BrdU-positive population, the cell-cycle profiles were analysed using FlowJo flow cytometry analysis software.

Construction and characterization of lentiviruses. Commercially available lentiviral vectors (pLKO.1) expressing short hairpin RNAs (shRNAs) against FZD9 were used as the background for GO annotation and enrichment analysis, taking statistical significance (P < 0.05) into account. The following primer pair: 5′-CCG AGA TCT TCG AGG TGT GTG GGG TTC TCC -3′; 5′-ATC TTT CGG ATG TGG AAG AGG-3′. For rescue experiments, FZD9 CDNA was amplified from TD NPC cDNA as template by the following primer pair: 5′-CCG AGA TCT TCG AGG TGT GTG GGG TTC TCC AAA G-3′; 5′-TCT AGA GCA ACC ATG CTC GAG CCG CCT GTC-3′. The reaction was performed using Phusion High-Fidelity DNA polymerase (New England Biolabs) according to the manufacturer’s protocol. The FZD9 cDNA was cloned into a lentiviral vector driven by the ubiquitin promoter followed by a self-cleave peptide and GFP sequence. The specificity and efficiency of shRNA-control, shRNA-FZD9, and the FZD9-WT constructs were verified by co-transfection into HEK-293 cells. Cell lysates were collected and analysed by western blot analysis with anti-FZD9 antibodies (Aviva OAEC02415, 1:1,000).

CHIR-98014 treatment. CHIR-98014 (Selleckchem) was resuspended according to manufacturer’s instructions into 10 mM stock using DMSO and then diluted to 10 μM. Final concentration used in cell culture was 100 nM of CHIR-98014, whereas the vehicle cells received only DMSO. For qPCR experiments, NPCs were propagated in six-well plates until 70% confluence and then treated with CHIR-98014 for 6 h to have their RNA collected using Trizol as previously described. For the NPC counting experiment, cells were seeded in six-well plates as described in the presence of CHIR-98014 or DMSO, in triplicates (TD and WS). After 48 h, the culture medium was changed and treatment was repeated. Cells were collected and counted after 96 h of incubation.

Astrocyte differentiation. The TD NPCs were lifted into suspension and maintained on a shaker (95 r.p.m.) to form neurospheres for 3 weeks. For the first week, the TD NPCs were lifted into suspension and main-
z-stack random images for all individuals and from two different experiments. Slides were analysed under a fluorescence microscope (Z1 Axio Observer Apotome, Zeiss). Only puncta in proximity of MAP2-positive processes were scored.

**Single-cell qRT-PCR and analysis.** Specific target amplification was performed in individual dissociated NPCs or 6-week-old neurons using C1 Single-Cell and BioMark HD Systems (Fluidigm), according to the manufacturer’s protocol and as described previously[37,44]. Briefly, single cells were captured on a C1 chip (10- to 17-μm cell size) using a LIVE/DEAD C1 viability kit (Invitrogen). Cytotoxicity test (Life Technologies). After lysis, RNA was reverse transcribed into cDNA with validated amplicon-specific DELTAGene Assays (Supplementary Table 13) using SuperScript III RT Platinum Taq Mix. Specific target amplification was performed by 17 cycles of 95°C denaturation for 15s and 60°C annealing and amplification for 4 min. Each preamplified cDNA was mixed with 2× SoFast EvaGreen Supermix with Low ROX (Bio-Rad) and then pipetted into an individual sample inlet in a 96.96 Dynamic Array IFC chip (Fluidigm). DELTAGene primer pairs (Supplementary Table 13) were diluted and pipetted into individual assay inlets in the same 96.96 Dynamic Array IFC chip. Quantitative PCR results were analysed using Fluidigm's Real-Time PCR Analysis software using the linear (derivative) baseline correction method and the automatic (gene) Ct threshold method with 0.65 curve quality threshold. Hierarchical clustering heat map, PCA analyses, violin plots of log(expression of C values) (limit of detection = 24) and ANOVA statistical analysis were performed using Singular Analysis Toolset 3.0 (Fluidigm).

**Calcium imaging.** Neuronal networks derived from human iPS cells were transplanted with lentiviruses carrying the Syn::RFP reporter construct. Cell cultures were washed with Krebs HEPES buffer (KHB) (10 mM HEPES, 4.2 mM NaHCO3, 10 mM dextrose, 1.2 mM MgSO4, 1.18 mM KH2PO4, 4.69 mM KC1, 118 mM NaCl, 1.29 mM NaCl, pH 7.3) and incubated with 2×5 mM Flu-o-4AM (Molecular Probes/Invitrogen, Carlsbad, California) in KHB for 40 min. Five thousand frames were acquired at 28Hz with a region of 256 pixels × 256 pixels (×100 magnification), using a Hamamatsu ORCA-ER digital camera (Hamamatsu Photonics K.K., Japan) with a 488 nm (FITC) filter on an Olympus IX81 inverted fluorescence confocal microscope (Olympus Optical, Japan). Images were acquired with MetaMorph 7.7 (MDS Analytical Technologies, Sunnyvale, California), processed and analysed using individual circular regions of interest (ROI) on ImageJ and Matlab 7.2 (Mathworks, Natick, Massachusetts). Syn::RFP neurons were selected after confirmation that calcium transients were blocked with 1 mM of tetrodotoxin (TTX). The amplitude of signals was presented as relative fluorescence changes (ΔF/ΔF) after background subtraction. The threshold for calcium spikes was set at the 95th percentile of the amplitude of all detected events.

**Electrophysiology.** For whole-cell patch-clamp recordings, individual cover slips containing live 1-month-old neurons were transferred into a heated recording chamber and continuously perfused (1 mL/min) with artificial cerebrospinal fluid bubbled with a mixture of CO2 (5%) and O2 (95%) and maintained at 25°C. Artificial cerebrospinal fluid contained (in mM) 121 NaCl, 4.2 KCl, 1.1 CaCl2, 1 MgSO4, 29 NaHCO3, 0.45 NaH2PO4·H2O, 0.5 NaHPO4, and 20 glucose (all chemicals from Sigma). Whole-cell recordings were performed using a digidata 1440A/Multiclamp 700B and Clampex 10.3 (Molecular devices). Patch electrodes were filled with internal solutions containing 130 mM K-glutamate, 4 mM KCl, 4 mM NaCl, 10 mM Na-HEPES, 0.2 mM K-EGTA, 0.3 mM GTP, 2 mM Mg-ATP, 0.2 mM CAMP, 10 mM d-glucose, 0.15% bicytin and 0.06% rhodamine. The pH and osmolarity were adjusted for physiological conditions. Data were all corrected for liquid junction potentials, electrode capacitances were compensated on-line in cell-attached mode and a low-pass filter at 2kHz was used. The access resistance of the cells in our sample was around 37 MΩ with resistance of the patch pipettes 3–5 MΩ. Spontaneous synaptic AMPA events were recorded at the reversal potential of Glutamate and could be reversibly blocked by AMPA receptor antagonist (10 μM NBQX, Sigma). Spontaneous synaptic GABA events were recorded at the reversal potential of Na+ and could be reversibly blocked by GABA_A receptor antagonist (10μM SR95531, Sigma).

**Multi-electrode array (MEA).** Using 12-well MEA plates from Axion Biosystems, we plated the same density of NPCs from TD and WS individuals in triplicate. Data were all corrected for liquid junction potentials, electrode capacitances were compensated on-line in cell-attached mode and a low-pass filter at 2kHz was used. The access resistance of the cells in our sample was around 37 MΩ with resistance of the patch pipettes 3–5 MΩ. Spontaneous synaptic AMPA events were recorded at the reversal potential of Glutamate and could be reversibly blocked by AMPA receptor antagonist (10 μM NBQX, Sigma). Spontaneous synaptic GABA events were recorded at the reversal potential of Na+ and could be reversibly blocked by GABA_A receptor antagonist (10μM SR95531, Sigma).

**Morphometric analysis of Golgi-impregnated neurons.** Cortical neurons from all six post-mortem brains were used in the study. Neurons included in the morphological analysis did not display degenerative changes[46]. Only neurons with fully impregnated soma, apical dendrites with present oblique branches and at least two basal dendrites with third-order segments were chosen for the analysis[47]. To minimize the effects of cutting on dendritic measurements, we included the neurons with cell bodies located near the centre of 120-μm-thick histological sections, with natural terminations of higher-order dendritic branches present where possible. More inclusion of the neurons completely within 120-μm sections biases the sample towards smaller neurons, leading to the underestimation of dendritic length[48], therefore, we applied the same criteria blinded across all WS and TD specimens, and we thus included the neurons with incomplete endings if they were judged to otherwise fulfill the criteria for successful Golgi impregnation. All neurons were oriented with apical dendrite perpendicular to the pial surface; impregnated pyramidal cells as well as magnopyramidal neurons were excluded from the analysis. Neuronal morphology was quantified along x-, y-, and z-coordinates using NeuroLucida version 10 software (MBF Bioscience, Williston, Vermont) connected to a Nikon Eclipse 80i microscope with a 40×0.75 numerical aperture objective. Tracings were conducted on both apical and basal dendrites, and the results reflect summed values for both types of dendrite per neuron. Following the recommendation that the applications of Sholl’s concentric spheres or Eayrs’ concentric circles for the analysis of neuronal morphology are not adequate when neuronal morphology is analysed in three dimensions[48], we conducted dendritic tree analysis with the following measurements[57,67]: (1) soma area—cross sectional surface area of the cell body; (2) dendritic length—summed total length of all dendrites per neuron; (3) dendrite number—number of dendritic trees emerging directly from the soma per neuron; (4) dendritic segment number—total number of dendritic spines/protrusion number—total number of dendritic spines per neuron; (6) dendritic spine/protrusion density—average number of spines per 20 μm of dendritic length; and (7) branching point number—number of nodes (points at the dendrite where a dendrite branches into two or more) per neuron. Dendritic segments were defined as parts of the dendrites between two branching points—between the soma and the first branching point
in the case of first-order dendritic segments, and between the last branching point and the termination of the dendrite in the case of terminal dendritic segments. Since the long formalin-fixation time could have resulted in degradation of dendritic spines, spine values might be underestimated and are thus reported here with caution. All of the tracings were accomplished blind to brain region and diagnostic status.

**Morphometric analysis of iPSC-derived neurons.** The iPSC-derived sample consisted of EGFP-positive 8-week-old neurons with pyramidal- or ovoid-shaped soma and at least two branched neurites (dendrites) with visible spines/protrusions. Protrusions from dendritic shaft, which morphologically resembled dendritic spines in post-mortem specimens, were considered and quantified as dendritic spines in iPSC-derived neurons. The neurites were considered dendrites on the basis of the criteria applied in post-mortem studies: (1) thickness that decreased with the distance from the cell body; (2) branches emerging under acute angle; and (3) presence of dendritic spines. In addition, only enhanced-GFP-positive neurites with nucleo-co-stained with CTP2, indicative of layer V/V1 neurones, and with the dendrites displaying evenly distributed fluorescent stain along their entire length, were included in the analysis. The morphology of the neurones was quantified along x-, y-, and z-coordinates using Neuro lucida version 9 software (MBF Bioscience, Williston, VT) connected to a Nikon Eclipse E600 microscope with a ×40 oil objective. No distinction was made between apical and basal dendrites, and the results reflect summed length values of all neurites/dendrites per neuron, consistent with what was done for the post-mortem neurones. The same set of measurements used in the analysis of Golgi-impregnated neurones was applied to the analysis of iPSC-derived neurones, and all of the tracings were accomplished blind to the diagnostic status and were conducted by the same rater (B.H.-M.). Intra-rater reliability was assessed by having the rater trace the same neuron after a period of time. The average coefficient of variation between the results of retracted neurons was 2% for soma area (SA), total dendritic length (TDL), dendritic segment number (DSN) and branching point number (BPN), and 3% for dendritic spine/protrusion number (DPN); there was no variation in tree/dendrite number (TN) in different tracings of the same neuron. The accuracy was further checked by having three individuals (B.H.-M., B.J., and L.S.) trace the same neuron.

**Brain imaging data acquisition and quality control.** MRI scanning was completed in 19 participants with WS (aged 19–43 years; mean 29.0, s.d. 8.8; 11 males, 8 females) and 19 TD comparison participants (aged 16–43 years; mean 26.2, s.d. 7.3; 8 males, 11 females). There was no significant difference between the groups in age (t=1.0, P<0.30) or in gender ratio (Pearson’s χ²=0.95, P<0.33). A standardized multiple modality high-resolution structural MRI protocol was implemented, involving three-dimensional T₁- and T₂-weighted volumes and a set of diffusion-weighted scans. Imaging data were obtained at the University of California San Diego Radiology Imaging Laboratory on a 1.5 T GE Sigma HDX 14.0M5 TwinSpeed system (GE Healthcare, Waukesha, Wisconsin) using an eight-channel phased array head coil. A three-dimensional inversion recovery spoiled gradient echo (IR-SPGR) T₁-weighted volume was acquired with pulse sequence parameters optimized for maximum grey/white matter contrast (echo time = 8.7 ms, repetition time = 720 ms, flip angle = 9°, difference in echo times = 750 ms, bandwidth = ±15.63 kHz, field of view = 225 × 225 mm, matrix = 192 × 192, voxel size = 1.25 mm × 1.25 mm × 1.25 mm). All MRI data were collected using prospective motion (PROMO) correction for non-distortion imaging. This method has been shown to improve image quality, reduce motion-related artefacts, increase the reliability of quantitative measures and improve the clinical diagnostic utility of MRI data obtained in children and clinical groups. Standardized quality control procedures were followed for both raw and processed data, including visual inspection ratings by a trained imaging technician and computer algorithms testing general image characteristics as well as aspects specific to each imaging modality, such as contrast properties, registrations and artefacts from motion and other sources. Participants included in the current analyses were only those who passed all raw and processed quality control measures.

**MRI data post-processing.** Image post-processing and analysis were performed using FreeSurfer software suite (http://surfer.nmr.mgh.harvard.edu/). Surface-based cortical reconstruction and subcortical volumetric segmentation procedures have been shown elsewhere. Briefly, a three-dimensional model of the cortical surface was generated using MRI scans with four attributes: white matter segmentation; tessellation of the grey/white matter boundary; inflation of the folded, tessellated surface; and computation of topological defects. Cortical thickness was measured using the distances from each point on the white matter surface to the pial surface. Cortical surface area was measured at the pial surface for the entire cerebrum and for each parcel of the Desikan and Destrieux atlases.

**Statistical analysis.** Means ± s.e.m. for each parameter were obtained from samples described in Supplementary Table 1. There were no statistical methods used to pre-determine sample size. The experiments were not randomized. All of the tracings were accomplished blind to brain region and diagnostic status. All statistical analyses were done using Prism (Graphpad). Before statistical analysis comparing means between three to five unmatched groups of data, normal distribution was tested using D’Agostino and Pearson omnibus normality test and variance similarity was tested using Bartlett’s test for equal variances. Means of three to five unmatched groups, where normal distribution and equal variances between groups were confirmed, were statistically compared using one-way ANOVA and Tukey’s post hoc test. Otherwise, a Kruskal–Wallis test and Dunn’s multiple comparison test were used. Before statistical analysis comparing means between two unmatched groups of data, normal distribution was tested using D’Agostino and Pearson omnibus normality test and variance similarity was tested using an F test to compare variances. To compare the means of two groups where normal distribution and similar variance between groups were confirmed, Student’s t test was used. Otherwise, a Mann–Whitney test was used. Significance was defined as *P* < 0.05, **P** < 0.01, ***P*** < 0.001 and ****P*** < 0.0001.
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Extended Data Figure 1 | Participants with WS in iPSC study and their neurocognitive and social profiles. a, Summary of scores on the Diagnostic Score Sheet (DSS) for individuals with WS. b, Table showing allele number of genes in WS-deleted region in each participant obtained from qPCR. c, Summary of all neurocognitive and social behavioural tests used on this study. d, e, WS neurocognitive profiles. Log of predictive likelihood ratio for iPSC participants (identified by participant number) calculated as the log of the ratio of the likelihoods for each individual test score based on the predictive distributions for TD individuals and those with WS (d). Values less than 0 indicate depressed scores consistent with expectations for WS. Predictive distributions for TD participants used published norms (means and standard deviations with assumed normality). Predictive distributions for individuals with WS were calculated using available WS data (VIQ/PIQ n = 81, VMI n = 56, PPVT n = 97) (e), assuming normality and least squares estimation, and according to the procedures described elsewhere. WS parameter estimates for the VMI were calculated using censored regression owing to several individuals with WS scoring at the instrument floor. f, Description of population included in Benton Face Recognition and Judgment of Line Orientation in Fig. 1b (TD n = 22 versus WS n = 65). g, Boxplots for WS (red) and TD (blue) participants on complex syntax (WS n = 45; TD n = 47) and social evaluation (WS n = 44; TD n = 49). Red and blue circles depict scores more than 1.5 times the interquartile range away from the median.
Extended Data Figure 2 | Generation and characterization of iPSCs.

a, Summary of reprogramming protocol using retrovirus carrying Yamanaka transcription factors (see Supplementary Information for details). Scale bar, 200 μm. 
b, Representative images of iPSCs expressing pluripotent markers including Nanog, Lin28, Oct4 and SSEA4 assessed by immunofluorescence staining. Scale bar, 200 μm. 
c, Expression of three germ-layer markers in iPSC-derived embryoid bodies (EBs); PAX6 (ectoderm), MSX1 (mesoderm) and AFP (endoderm) assessed by semiquantitative RT–PCR. TBP, housekeeping control. 
d, Cluster analysis showing correlation coefficients of microarray profiles of three WS DPCs, three TD DPCs, three WS iPSCs, three TD iPSCs and one ESC. 
e, Representative PCR showing silencing of the four transgenes (exogenous) in iPSCs. 
f, Representative images of teratoma from iPSCs showing tissues of three germ layers; neural rosettes (ectoderm), cartilage (mesoderm), muscle cells (mesoderm) and goblet cells (endoderm). 
g, Representative image of iPSC chromosomes showing its genetic stability assessed by G-banding karyotype analysis. 
h, i, Spontaneous synaptic GABA events (h) and spontaneous synaptic AMPA events (i) in 1-month-old iPSC-derived neurons.
Extended Data Figure 3 | See next page for caption.
Extended Data Figure 3 | Global gene expression analysis during neuronal differentiation. a, PCA plot of embryonic stem cells (ES), induced pluripotent stem cells (iPS), neuronal progenitor cells (NPC) and neurons (NE) for TD, WS and pWS88. c, Euclidian matrix distance-based heat map and hierarchical clustering-based dendrogram of ES, NPC and NE cells for WD, WS and pWS88 samples. Expression variability between samples is indicated by Z-score, varying from green (negative variation) to red (positive variation). c, Euclidian matrix distance-based heat map and hierarchical clustering-based dendrogram of pluripotency gene markers for ES, NPC and NE cells for TD, WS and pWS88 samples. d, Euclidian matrix distance-based heat map and hierarchical clustering-based dendrogram of neuronal gene markers for iPS, NPC and NE cells for TD, WS and pWS88 samples. Expression variability between samples is indicated by Z-score, varying from green (negative variation) to red (positive variation). e, Specific cell type-based clustering analysis of biological replicates subjected to RNA-seq for the WS-related genes in three stages during differentiation (iPS, NPC and NE). f, Fold change variation of WS-related genes in different cell lines. Ideogram of chromosome 7 (band 7q11.23) corresponding to the commonly deleted region with the WS-related genes. Fold change variation of normalized WS-related gene expression in NPCs and neurons (NE) compared with TDs. Non-represented fold change corresponds to those genes having high expression variability between biological replicates, or having very low expression values. g, Expression of FZD9 gene in iPSC, NPCs and neurons from TD and WS. Error bars, s.e.m. h, Venn diagram showing correlation of significant differentially expressed genes between TD, pWS88 and WS during neuronal differentiation. Significantly enriched GO terms found for downregulated (red histogram) and upregulated (blue histogram) differentially expressed genes between TD and WS in NPC. Significantly enriched GO terms found for downregulated (red histogram) and upregulated (blue histogram) differentially expressed genes between TD and WS in neurons (NE). Vertical line (black) corresponds to a significant P value (<0.05). i, Enriched GO metabolic process terms found in NPC of WS samples correlated with the GO found by a similar comparison performed in ref. 13.
Extended Data Figure 4 | See next page for caption.
Extended Data Figure 4 | Defect in WS NPC apoptosis and role of FZD9.

a, Ratio of NPC number on day 4 over day 0 relative to TD. Data are shown as mean ± s.e.m.; n, number of clones.
b, High percentage (>95%) of Sox1/Sox2-positive and Pax6/Nestin-positive cell population was comparably observed in TD, typical WS and pWS88 NPCs assessed by FACS. Data are shown as mean ± s.e.m.; n, number of clones.
c, Microfluidics of C1 chip used to capture live single cells (calcein+ cell).
d, Outlier exclusion based on the recommended/default limit of detection value of 24, analysed by Fluidigm Singular 3.0. Outliers were removed manually on the basis of the sample median log2(expression) values.
e, Representative example of non-normalized Ct plot, indicated with the rectangle in the heat map. Cells are shown in rows and genes in columns. The range of cycle threshold (Ct) values is colour coded from low (blue) to high (red) and absent (black).
f, Violin plots of all 96 genes showing the comparison between TD and WS NPCs from the single-cell analyses (log2(expression) values). The majority of genes show unimodal expression distribution.
g, Volcano plot of single-cell expression data. Plot illustrates differences in expression patterns of target genes of iPSC-derived NPCs. The dotted lines represent more than or equal to 3.0-fold differentially expressed genes between the groups at P < 0.05 (unpaired two-sample t-test).
h, Schematic diagram summarizing NPC preparation for proliferation assay and representative scatter plot showing cells in each cycle phase (G1, S and G2/M).
i, No significant differences in percentage of the BrdU-positive population between TD, typical WS and pWS88 NPCs.
j, Schematic diagram summarizing NPC preparation for apoptosis analysis and representative analysed data for DNA fragmentation (left) and caspase assay (right).
k–m, Changes in ratio of NPC number on day 4 over day 0 relative to TD (k), percentage of subG1 population (l) and percentage of population with high caspase activity (m) of pWS88 NPCs when treated with shFZD9 and shControl.
n, Increase in cell number day 4/day 0 upon overexpression of FZD9 in WS iPSC-derived NPCs. Data are shown as mean ± s.e.m. for each individual; n, technical replicates. For i and k–m, data are shown as mean ± s.e.m.; n, number of clones, *P < 0.05, **P < 0.01, ***P < 0.001, one-way ANOVA and Tukey’s post hoc test (i), Kruskal–Wallis test and Dunn’s multiple comparison test (k–m).
Extended Data Figure 5 | Single-cell analysis of WS and TD iPSC-derived neurons. a, b, Outlier exclusion based on limit of detection = 24, analysed by Fluidigm Singular 3.0. Outliers were removed manually on the basis of the sample median log2(expression) values. c, Heat map of number of genes with ANOVA P-value < 0.05 (82 genes in total). d, Unsupervised hierarchical clustering of 672 single-cell of WS and TD iPSC-derived neurons identified cell sub-populations not linked with the genotype. e, PCA projections of the 96 genes, showing the contribution of each gene to the first two PCs. f, Violin plots of all 96 genes showing the comparison between TD, WS and pWS88 neurons from the single-cell analyses (log2(expression) values). Cells are shown in rows and genes in columns. Log2(gene expression levels) were converted to a global Z-score (blue is the lowest value, red is highest). Genes were clustered using the Pearson correlation method and cells were clustered using the Euclidean method.
Extended Data Figure 6  |  See next page for caption.
Extended Data Figure 6 | Morphometric analysis of WS-derived CTIP2-positive cortical neurons. a, Summary of preparation of neurons for evaluation by morphometric analysis. b, Representative images of EGFP- and CTIP2-positive neuron (arrowhead) and tracing. Scale bar, 200 μm. c–f, No significant differences in dendritic segment numbers (c), number of branching points (d), dendritic spine density (e) and soma area (f) between TD, typical WS and pWS88 were observed. g–m, Morphometric analysis shown as individual participant for total dendritic length (g), dendritic tree number (h), dendritic spine number (i), dendritic segment number (j), number of branching points (k), dendritic spine density (l) and soma area (m). n, Four-week-old neurons were dissociated and plated to trace total neurite length every hour, for a total of 6 h. Representative images of traced neurons plated after 0 and 6 h from TD, typical WS and atypical pWS88 iPSC-derived neurons. o–r, Morphometric analysis showing significant differences among TD, typical WS and pWS88 in the initial neurite growth velocity (6 h period). r, Morphometric analysis shown for individual participants for neurite growth velocity for 6 h interval. n, Number of traced neurons. s–u, No significant changes were observed in the total dendritic length (s), dendritic segment number (t) and dendritic spine number (u) of TD neurons plated in different densities (300–1,200 cells per square millimetre). v, Individual channels of puncta quantification of post- and presynaptic markers (Homer1/Vglut1). Scale bar, 2 μm. For c–m and o–u, data are shown as mean ± s.e.m.; n, number of traced neurons, *P < 0.05, **P < 0.01, Kruskal–Wallis test (c–f), one-way ANOVA and Tukey’s post hoc test (o–q, r–u).
Extended Data Figure 7 | Alteration in calcium transient in WS iPSC-derived neurons and morphometric analysis of cortical layer V/VI pyramidal neurons in post-mortem tissue. 

a. Puncta quantification of post- and presynaptic markers. The synaptic proteins Vglut (presynaptic) and Homer1 (postsynaptic) were used as markers and only co-localized puncta on MAP2+ cells were quantified and graphed. Data are shown as the mean ± s.e.m.; n, number of neurons.

b. Summary of preparation of neurons for calcium transient analysis. Representative images of live neuronal culture expressing RFP driven by synapsin promoter and the uptake of Fluo-4AM calcium dye.

c. Blockade of calcium transient by TTX inhibition of synaptic activity. Representative images of calcium transient in single neurons (RFP-positive, arrowhead) from TD (top), typical WS (middle) and pWS88 (bottom). Number in the lower right of each figure represents each time point (seconds) when change in Fluo-4AM occurs.

d. Calcium transient analysis shown as individual for frequency (e) and percentage of signalling neurons (f). Data are shown as mean ± s.e.m.; n, number of fields analysed.

g. MEA analyses revealed an increase in spontaneous neuronal spikes. Data show individual clones.

h. Raster plot of TD and WS iPSC-derived neurons analysed by multi-electrode array.

i. Details of individuals used for the analysis.

j–l. No significant differences in dendrite number (j), dendritic spine density (k) and soma area (l) between TD and typical WS were observed. Data are shown as mean ± s.e.m.; n, number of traced neurons, two-sided unpaired Student’s t test.

m–s. Morphometric analysis shown for each individual for total dendritic length (m), dendritic spine number (n), segment number (o), branching point number (p), dendrite number (q), dendritic spine density (r) and soma area (s). Data are shown as mean ± s.e.m.; n, number of traced neurons.

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Extended Data Table 1 | List of top ten most significant differentially expressed genes in WS compared with TD for NPC and neurons

### NPC: TD x WS

| Gene name | Description | Fold-change | p-value |
|-----------|-------------|-------------|---------|
| SCN4A     | sodium channel, voltage gated, type IV alpha subunit | -11.92      | 9.72E-10 |
| SLC7A14   | solute carrier family 7, member 14 | -15.85      | 1.85E-12 |
| SLC38A5   | solute carrier family 38, member 5 | -8.51       | 5.67E-09 |
| ADGRA2    | adhesion G protein-coupled receptor A2 | -30.50      | 3.26E-08 |
| SLC1A6    | solute carrier family 1 (high affinity aspartate/glutamate transporter), member 6 | 8.03        | 1.56E-08 |
| CXCL12    | chemokine (C-X-C motif) ligand 12 | -7.51       | 3.33E-08 |
| SLC30A3   | solute carrier family 30 (zinc transporter), member 3 | 10.28       | 8.32E-09 |
| SLC8A2    | solute carrier family 8 (sodium/calcium exchanger), member 2 | -16.84      | 5.36E-13 |
| HTR1B     | 5-hydroxytryptamine (serotonin) receptor 1B, G protein-coupled | -10.96      | 8.20E-09 |
| SLC24A2   | solute carrier family 24 (sodium/potassium/calcium exchanger), member 2 | -13.25      | 4.43E-10 |

### NPC: TD x pWS88

| Gene name | Description | Fold-change | p-value |
|-----------|-------------|-------------|---------|
| GABRA3    | gamma-aminobutyric acid (GABA) A receptor, alpha 3 | 14.71       | 6.30E-10 |
| SYT13     | synaptotagmin XIII | 6.55        | 1.87E-08 |
| PPP2R2C   | protein phosphatase 2, regulatory subunit B, gamma | 15.74       | 7.69E-09 |
| CELF4     | CUGBP, Elav-like family member 4 | 7.27        | 1.22E-07 |
| TRIM67    | tripartite motif containing 67 | 15.08       | 6.52E-10 |
| ADRA2A    | adrenoceptor alpha 2A | 8.44        | 1.42E-07 |
| JAKMP1    | janus kinase and microtubule interacting protein 1 | 19.56       | 7.21E-09 |
| CA10      | carbonic anhydrase X | 74.54       | 8.50E-10 |
| LHFP4L4   | lipoma HMGIC fusion partner-like 4 | 8.31        | 5.71E-08 |
| ACSL6     | acyl-CoA synthetase long-chain family member 6 | 7.07        | 7.63E-08 |

### Neuron: TD x pWS88

| Gene name | Description | Fold-change | p-value |
|-----------|-------------|-------------|---------|
| CRYM      | crystallin, um | 7.68        | 3.21E-05 |
| RASL12    | RAS-like, family 12 | 13.41       | 6.04E-08 |
| PDZM1     | PDZ and LIM domain 1 | 5.44        | 1.00E-05 |
| ZSCAN10   | zinc finger and SCAN domain containing 10 | 58.80       | 6.22E-06 |
| ANO1      | anoctamin 1, calcium activated chloride channel | 9.61        | 1.78E-06 |
| DUSP23    | dual specificity phosphatase 23 | 5.09        | 0.00014507 |
| SLC33A6   | solute carrier family 16 (monocarboxylate transporter), member 5 | 33.27       | 8.90E-05 |
| KRT19     | keratin 19, type I | 9.94        | 0.000130985 |
| TMEM30B   | transmembrane protein 30B | 17.09       | 2.11E-07 |
| KIF18B    | kinesin family member 18B | 4.50        | 0.000217123 |

### Neuron: TD x WS

| Gene name | Description | Fold-change | p-value |
|-----------|-------------|-------------|---------|
| FAM19A5   | family with sequence similarity 19 (chemokine (C-C motif)-like), member A5 | 1076.39    | 2.23E-10 |
| TPM2      | tropomyosin 2 (beta) | -5.88       | 2.99E-08 |
| SCN4A     | sodium channel, voltage gated, type IV alpha subunit | -45.81     | 1.63E-07 |
| IGSF21    | immunoglobulin superfamily, member 21 | 8.20        | 2.33E-07 |
| TNN1      | tropinin T 2 (cardiac) | -29.87      | 3.58E-07 |
| PXMP4     | peroxisomal membrane protein 4, 24kDa | -7.39       | 3.83E-05 |
| ZSCAN10   | zinc finger and SCAN domain containing 10 | 35.64       | 9.74E-05 |
| MYOZ1     | filamin-, Actinin- And Telethonin-Binding Protein | -6.64       | 0.000119159 |
| LAD1      | ladinin 1 | -24.81      | 0.000213205 |
| PHOSPHO1  | phosphatase, Orphan 1 | 33.95       | 0.000266061 |
### Extended Data Table 2 | Most significant (P < 0.05) enriched GO terms in NPC of WS compared with tD samples

| GO TERM | GO Description | Log (p-value) | P-value |
|---------|----------------|--------------|---------|
| GO:0038416 | phenotypic feature | 18.342667 | 4.52E-10 |
| GO:0006956 | protein structure organization | 18.851644 | 1.97E-17 |
| GO:0007010 | cell adhesion | 17.708687 | 2.90E-17 |
| GO:0038188 | extracellular region organization | 13.458765 | 1.25E-14 |

**Down-regulated genes in WS compared to TD in NPC cells**

| GO TERM | GO Description | Log (p-value) | P-value |
|---------|----------------|--------------|---------|
| GO:0005396 | clathrin-mediated endocytosis | 3.383902 | 2.90E-05 |
| GO:0005901 | lipid metabolic process | 2.336182 | 0.005489 |
| GO:0005329 | fibril organization | 3.642526 | 0.006640 |
| GO:0005709 | protein organization | 1.672066 | 0.0134|
| GO:0005150 | interferon-gamma-mediated signaling pathway | 1.692204 | 0.015387 |
| GO:0005456 | extracellular region | 1.649850 | 0.020329 |
| GO:0005676 | transmembrane transport | 1.873092 | 1.34E-03 |

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### Extended Data Table 3 | Most significant (P < 0.05) enriched GO terms in neurons of WS compared with TD samples

#### Down-regulated genes in WS compared to TD in neurons

| GO TERM                          | GO Description                        | -log10(P-value) | P-value | Genes                                                                 |
|---------------------------------|---------------------------------------|-----------------|---------|----------------------------------------------------------------------|
| ion binding                     |                                       | 2.99584255      | 0.0010086 | BAX1B, FM07, V52Q, DISQ2, RFC2.C2, ME1, ITTF2A2, ADAMTS12, MYL3, TNC22, ENOS3, PDZ1, AC2B, TRIM38, PCDH18S, TN1C1, ACG22CAT, ACG12S, MYL2, ACG12W, ZNF395, ATRIP, MAL3, GSTM1, ACTA1, SHRXF2, ACTA1, ADY15, CAPR13, ZNF283, ZNF514, ACG12Q, GSTM4, SLNF21, RHOJ, JNK3, ZNF826, SERPINAS1, 1.00A4, ZNF514, ZNF72, ZNF72, ZNF73, ZNF578 |
| cytoskeleton protein binding    |                                       | 2.64132966      | 0.00144104 | USC1C1, MYBP22, TNC2X1, TNC11, TNC17, MYH2, ACTA1, ACTA1, MYH21, TPM2 |
| circulatory system process      |                                       | 2.42826763      | 0.00274472 | ELN, TNQ1C1, ACTA1, ACT1C1, C34                                       |
| response to stress              |                                       | 2.27051445      | 0.00563986 | TBNXSR, BAX1B, V52Q, RFC2.C2, ATPT2A2, IL4, LAT2, MYL3, TNC22, CAT, ACG12S, MYL2, ACG12W, MAL3, GSTM1, PPARA, ALOX15, ACG11, BATF2, CD43, BPHAX15, SHRXF2, IFITM2, SERPINAS1, L.HA-DRBS1, L.HA-DRBS2, POUSP1 |
| structural molecule activity     |                                       | 2.01566209      | 0.00964553 | MYG2, ELN, MYBP22, MYL3, MYH2, TNC1C1, ACTA1, LAD1, TPM2              |
| cytofilm                        |                                       | 1.67077895      | 0.0211311 | USC1C1, MYBP22, MYL3, TNC22, STX1A, ERF-1L, ENOS3, TRIM38, TN1C1, TNC17, MYH2, GSTM1, PPARA, ACTA1, ACT1C1, ALG1C5, GSTM1, RHOJ, ADY15, TNC17, TPM2, POUSP1 |
| peroxisome                      |                                       | 1.57493369      | 0.02661313 | PXR4, CAT, ACG22C          |
| Molecular function               |                                       | 1.54859974      | 0.03282788 | USC1C1, TBNXSR, SCNA4, BAX1B, FM07, ACG12W, V52Q, MYL2, DISQ2, PROQ2, ELN, RFC2.C2, ATRIP, MAL3, GSTM1, ACG12Q, GSTM4, ACG12W, PROQ2, MYL2, ACG12Q, TNC2X1, TNC11, MYH2, ACG12S, MYL2, ACG12W, GSTM1, PPARA, ALOX15, ACG11, BATF2, CD43, BPHAX15, SHRXF2, IFITM2, SERPINAS1, L.HA-DRBS1, L.HA-DRBS2, POUSP1 |
| Transferase activity, transferring shikl or any other than methyl groups | | 1.47536534 | 0.03346653 | GSTM1, GSTM4 |
| ATPase activity                 |                                       | 1.46244853      | 0.03473731 | RFC2, ATRIP2A2, TNC17, ACG22C, ATRIP2B3, ACTA1C1 |
| lipase activity                 |                                       | 1.40625589      | 0.03923878 | RFC2, ACG12S, TRIM38, SLC22A6, SHRXF2, ZNF21 |
| photosynthesis                  |                                       | 1.38734532      | 0.04598787 | RFC2 |
| extracellular space             |                                       | 1.32271586      | 0.0477842 | AOPP, ENOS3, ACG12W, ADY15, ACT1C1, ACT1C1, CYTL1, SERPINAS1, 1.00A4, 1.00A4 |

#### Up-regulated genes in WS compared to TD in neurons

| GO TERM                          | GO Description                        | -log10(P-value) | P-value | Genes                                                                 |
|---------------------------------|---------------------------------------|-----------------|---------|----------------------------------------------------------------------|
| nucleic acid binding transcription factor activity | | 2.49095017 | 0.00396898 | PTX1, T7FAP2, HAND1, NRS42, RPS6, ZSCAN15, KL.LALK1, DMBX1, FOXB2 |
| Peptide activity                |                                       | 1.53713232      | 0.02394539 | HPN, PRPS1B, RHBH2, ADAMTS15, LPSP41, TPRPSS2 |
| embryo development              |                                       | 1.35647681      | 0.04418999 | HPN, HAND1, NRS42, KL.F4 |
| generation of precursor metabolites and energy | | 1.33043536 | 0.0465726 | ALDOC, MT-ND2, MT-ND4, MT-ND1 |
| anatomical structure development | | 1.31245495 | 0.0487019 | PTX1, T7FAP2, HPN, ALDOC, HAND1, NRS42, RPS6, ZSCAN15, KL.LALK1, DMBX1, FOXB2 |