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ORIGINAL ARTICLE System-based proteomic and metabonomic analysis of the $Df(16)A^{+/-}$ mouse identifies potential miR-185 targets and molecular pathway alterations

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Deletions on chromosome 22g11.2 are a strong genetic risk factor for development of schizophrenia and cognitive dysfunction. We employed shotgun liquid chromatography-mass spectrometry (LC-MS) proteomic and metabonomic profiling approaches on prefrontal cortex (PFC) and hippocampal (HPC) tissue from $Df(16)A^{+/-}$ mice, a model of the 22q11.2 deletion syndrome. Proteomic results were compared with previous transcriptomic profiling studies of the same brain regions. The aim was to investigate how the combined effect of the 22q11.2 deletion and the corresponding miRNA dysregulation affects the cell biology at the systems level. The proteomic brain profiling analysis revealed PFC and HPC changes in various molecular pathways associated with chromatin remodelling and RNA transcription, indicative of an epigenetic component of the 22g11.2DS. Further, alterations in glycolysis/ gluconeogenesis, mitochondrial function and lipid biosynthesis were identified. Metabonomic profiling substantiated the proteomic findings by identifying changes in 22q11.2 deletion syndrome (22q11.2DS)-related pathways, such as changes in ceramide phosphoethanolamines, sphingomyelin, carnitines, tyrosine derivates and panthothenic acid. The proteomic findings were confirmed using selected reaction monitoring mass spectrometry, validating decreased levels of several proteins encoded on 22g11.2, increased levels of the computationally predicted putative miR-185 targets UDP-N-acetylglucosamine-peptide Nacetylglucosaminyltransferase 110 kDa subunit (OGT1) and kinesin heavy chain isoform 5A and alterations in the non-miR-185 targets serine/threonine-protein phosphatase 2B catalytic subunit gamma isoform, neurofilament light chain and vesicular glutamate transporter 1. Furthermore, alterations in the proteins associated with mammalian target of rapamycin signalling were detected in the PFC and with glutamatergic signalling in the hippocampus. Based on the proteomic and metabonomic findings, we were able to develop a schematic model summarizing the most prominent molecular network findings in the $Df(16)A^{+/-}$ mouse. Interestingly, the implicated pathways can be linked to one of the most consistent and strongest proteomic candidates, (OGT1), which is a predicted miR-185 target. Our results provide novel insights into system-biological mechanisms associated with the 22q11DS, which may be linked to cognitive dysfunction and an increased risk to develop schizophrenia. Further investigation of these pathways could help to identify novel drug targets for the treatment of schizophrenia.

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INTRODUCTION

Hemizygous deletions in chromosome 22q11.2 occur predominately *de novo* and cause a deletion syndrome (22q11.2DS), characterized by a broad spectrum of physical and mental manifestations with a variable phenotype. The syndrome has an incidence of 1 in 2000–4000 live births. The physical phenotype is characterized by variably penetrant craniofacial and cardiovascular anomalies, immunodeficiency, short stature and hypocalcaemia. Furthermore, deletion carriers typically present with a range of behavioural and cognitive deficits and 25–30% develop schizophrenia during adolescence or early adulthood.¹ 22q11.2 deletions account for 1–2% of sporadic schizophrenia cases, exemplifying the role of rare mutations in disease susceptibility. Importantly, there are no major clinical differences in the core schizophrenia phenotype between individuals with schizophrenia who are 22g11.2 microdeletion carriers and those with an idiopathic disease onset.² A 1.5-Mb critical region, which confers haplo-insufficiency of 27 genes,^{3,4} including the candidate schizophrenia susceptibility genes (proline dehydrogenase (PRODH), catechol-O-methyltransferase (COMT) and DHHC palmitoyltransferase (ZDHCC8)), is largely conserved on mouse chromosome 16. This has facilitated generation of mouse models of the human 22q11DS, carrying hemizygous deletions in the syntenic region of chromosome 16. One of the best characterized models is the $Df(16)A^{+/-}$ mouse, which carries a hemizygous 1.3-Mb chromosomal deficiency on chromosome 16 (Df(16)A), encompassing 27 genes, which is syntenic to the minimal 1.5-Mb human 22q11.2 deletion.⁵ These mice develop deficits in spatial working memory, sensorimotor gating and fear conditioning as well as presenting with hyperactive behaviour.^{5,6} In addition, they show abnormalities in synaptic connectivity of hippocampal and cortical

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Studies that have examined the effect of *Df(16)A* on transcriptional networks revealed downregulation of microRNA (miRNA) transcripts,^{5,8} resulting from the combined effect of two disrupted genes (*Dgcr8* and miR-185) in the critical region. *Dgcr8* hemizygosity leads to dysregulation in the production of up to 20% of all miRNAs, while miR-185 levels are reduced by 70–80% owing to a combined effect with the *Dgcr8* hemizygosity,⁹ which leads to an impaired maturation of the pri-miR-185 transcript. Based on the transcriptional profiling, the most robust miRNA target was a novel neuronal regulatory gene *2310044H10Rik/Mirta22*, which was confirmed to be specific for miR-185 and has been followed up in functional studies. It was shown that this gene encodes a protein, which is located in the Golgi apparatus and in vesicles and tubular-like extensions of dendrites, mediating at least some of the effects of the 22q11.2 deletions on dendrite and spine formation.¹⁰

To date, no other direct miRNA targets have been reported for the 22q11.2DS and it is still not known how the reported transcriptomic^{5,11} alterations affect cell biology at the proteomic and metabonomic level. In the disease pathology, proteomic and metabonomic networks are likely to be regulated through the combined interactions of the 22q11.2 deletion gene products, which involve the hemizygosity of four transcription factors and disruption of miRNA-regulated targets. This may involve dysregulation of other transcription factors, chromosome remodelling elements or changes in the expression of signalling proteins. Also, secondary compensation effects might have a role. Here we have carried out a combined shotgun proteomic and metabonomic profiling analysis of brain tissue from $Df(16)A^{+/-}$ mice to investigate the combined effects of miRNA dysregulation and disruption of genes residing in the 22q11.2 region. Our results provide novel insights towards a better understanding of the molecular pathophysiology of psychiatric disorders and cognitive dysfunction.

MATERIALS AND METHODS

For more detailed information, please refer to Supplementary Methods. A flowchart of the molecular profiling methods can be found in Supplementary Figure S1.

Animals

The $Df(16)A^{+/-}$ mice were produced and bred at Columbia University (New York, NY, USA) as described previously.⁵ Eight-week-old male mice were used for the analyses. Mutant mice carry a hemizygous 1.3-Mb chromosomal deficiency ($Df(16)A^{+/-}$), which ranges from Dgcr2 to the *Hira* gene and encompasses a segment syntenic to the 1.5-Mb human 22q11.2 deletion, including 27 protein-coding genes. All animal procedures were performed according to protocols approved by the Columbia University Institutional Animal Care and Use Committee under federal and state regulations.

Multiplex immunoassay profiling

Serum samples from 11 $Df(16)A^{+/-}$ and 10 *wild-type* mice (cohort 1) were analysed randomized and blinded using a multianalyte profiling platform, comprising multiplexed immunoassays for 58 analytes (Supplementary Table S1) in a Clinical Laboratory Improved Amendments (CLIA)-certified laboratory at Myriad-RBM (Austin, TX, USA) as described previously.^{12,13} Data quality was assessed via principal component analysis (PCA), data were checked for normality and significance analysis of mircroarray¹⁴ was performed.

Metabonomics

Metabonomic sample preparation and profiling of prefrontal cortical (PFC) and hippocampal (HPC) tissues from 11 $Df(16)A^{+/-}$ and 10 *wild-type* mice (cohort 1) was performed as described previously.¹⁵ In brief, tissue samples

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were prepared using a two-step process based on extraction into 'aqueous' and 'organic' phases for polar and nonpolar metabolites. Quality-control samples were prepared by combining an aliquot from each study sample to produce a representative sample, which was used for column conditioning and data quality assessment as described by Want et al.¹⁵ Aqueous extracts were analysed via nano-ultra-performance liquid chromatography-mass spectrometry (UPLC-MS) analysis using a Waters XEVO G2 O-TOF mass spectrometer coupled online to an Acquity UPLC-MS system (Waters, Milford, MA, USA). Metabolites were separated using a $2.1 \times 100 \text{ mm}^2$ (1.7 µm) HSS T3 Acquity column, and acquisition was performed in both positive ion mode and negative ion mode. Qualitycontrol samples were injected 10 times at the start of the analytical batch in order to condition the column, and then after every 10 samples throughout the run to assess instrument stability. Samples were run randomized and blinded. Data were processed using the freeware XCMS^{16,17} using standard parameters. The output consisted of a matrix of metabolite feature m/z, retention time and intensity values, which was imported into SIMCA-P for multivariate analysis to check data quality and sample outliers. Data were normalized, filtered and checked for normality prior statistical analysis. P-values were determined using Wilcoxon's signed-rank test and corrected to control for multiple hypothesis testing (Benjamini-Hochberg).¹⁸ Ratios were calculated for each analyte as the mean intensity values of $Df(16)A^{+/-}$ mice divided by those of wild-type littermates.

Proteomics

For more detailed information, please refer to Supplementary Methods.

Sample preparation and study design

PFC and HPC brain tissues from $Df(16)A^{+/-}$ and wild-type mice were prepared using total lysis protein extraction^{19,20} combined with either insolution (two independent cohorts; cohort 1 (PFC and HPC): 11 Df(16)A^{+/} vs 10 *wild-type* mice, cohort 2 (PFC): 13 $Df(16)A^{+/-}$ vs 13 *wild-type* mice) or in-gel tryptic digestion ((PFC): 20 $Df(16)A^{+/-}$ and 21 *wild-type* mice, combined from both cohorts to increase statistical power, proteome coverage and avoid sample extraction biases). Samples were subsequently analysed using label-free LC-MS^E mass spectrometry, enabling unbiased protein identification and guantification. Mass spectrometrical sample preparation, label-free LC-MS^E analysis and selected reaction monitoring (SRM) analysis were performed blinded and randomized. Sample size was chosen based on previous experiments using FC and HPC brain tissues and proteomic guidelines. In all, 8-10 animals per group are sufficient for the proteomic shotgun discovery phase. Power calculations using MSstats showed that to achieve a fold change of 15% with 0.8 power 10 sample are needed. The validity of the results was further proven in 20 $Df(16)A^{+/}$ and 20 wild-type mice using highly sensitive and robust label-based SRM mass spectrometry.

Label-free LC-MS^E analysis of PFC and HPC tissue

Brain tissue samples were analysed individually in technical duplicates using a splitless UPLC (10 kpsi nanoAcquity; Waters) coupled online to a Waters Q-TOF Premier mass spectrometer. Data were acquired in expression mode (MS^E). The procedure, quality assessment and data processing were performed as described previously.²¹ LC-MS^E data were processed using the ProteinLynx Global Server (PLGS) v.2.4. (Waters) and Rosetta Elucidator v.3.3 (Rosetta Biosoftware, Seattle, WA, USA) was used for time and mass/charge alignment of mass spectrometric data. The Swiss-Prot mouse reference (March, 2013) proteome was used for protein identification searches. Only peptides detected in both replicates and in >80% of samples were included in further analysis. Protein abundance changes were determined using the MSstats package²² based on linear mixed-effects models, which views peptides mapped to the same protein as replicate measurements of protein abundance, following log₂ transformation and exclusion of peptide intensity values that were >3 s.d. from the mean of each group. The MSstats package included guality assessment and PCA on the raw data was carried out to detect data outliers. The P-values were adjusted to control the falsediscovery rate at a cutoff of 0.05 following the Benjamini-Hochberg procedure.23

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Label-based SRM mass spectrometry

Digested FC and HPC proteomes (20 $Df(16)A^{+/-}$ and 20 wild-type mice), prepared using in-solution digestion from total lysates, were analysed using targeted SRM mass spectrometry on a Xevo TQ-S mass spectrometer (Waters) coupled to a nanoAcquity UPLC system (Waters) as described previously.^{13,20,24} Multiplex SRM assays were developed using a highthroughput strategy.²⁵ Physiochemical criteria for selecting tryptic peptides were based on peptide count, uniqueness and quality of transitions. Transitions were selected based on software internal predictions, discovery proteomics data and spectral data (NIST spectral libraries²⁶) and calculated using Skyline version 1.2.0.3425.²⁷ Quantitative SRM measurements comparing abundance levels of 40 proteins between $Df(A)16^{+/-}$ mice and *wild-type* littermates were performed in scheduled SRM acquisition mode. Heavy isotope labelled peptide versions (JPT Peptide Technologies, Berlin, Germany) were spiked in the peptide mixture for accurate quantification and identification. For each peptide, at least three transitions were monitored for the heavy and light versions. The final transitions can be found in Supplementary Table S2. Samples were run blinded, randomized and blocked²⁸ in triplicates, and blanks and quality-control peptide injections (yeast alcohol dehydrogenase)²⁴ were run alternating after each biological replicate. Resulting SRM data were analysed using Skyline and statistical analysis, testing for differential abundance among $Df(A)16^{+/-}$ and wild-type mice, was conducted using SRMstats²³ with the 'expanded technical replication' setting. Data quality was checked using quality plots in the MSstats package and by employing PCA on the raw data. P-values were corrected to control for multiple hypothesis testing (Benjamini–Hochberg).¹⁷

RESULTS

Serum characterization—quantitative serum immunoassay profiling

Immunodeficiency is one of the key features of the 22q11.2DS, but the degree and spectrum of severity varies widely. A large number of studies have provided evidence suggestive of an immunological component of schizophrenia²⁹ and other psychiatric disorders.^{30,31} We hypothesized that immune dysfunction, as reflected in the levels of immune and metabolic circulation markers, may contribute to the elevated risk of schizophrenia in the 22q11.2DS.³² Analysis of 58 such markers (Supplementary Table S1) in the serum of the $Df(16)A^{+/-}$ mouse using a multiplex immunoassay platform resulted in the identification of a significant decrease in matrix metalloproteinase 9 (MMP-9; ratio = 0.74, $P = 0.039, P^* = 0.20$) in the $Df(16)A^{+/-}$ (n = 11) compared with wildtype (n = 10) mice.

Brain characterization-quantitative proteomics

In the proteomic discovery phase, shotgun profiling of PFC and HPC tissue was carried out to identify proteins differentially expressed in the $Df(16)A^{+/-}$ mouse model compared with wild-type mice. The main aim was to identify affected protein networks that might contribute to the development of psychiatric and cognitive phenotypes in the 22q11.2DS, as well as novel miR-185 targets. We focused on the PFC and HPC because these brain regions have previously been characterized at the transcriptomic level^{5,11} and are implicated in the pathology of various psychiatric disorders. Special emphasis was placed on the PFC as the most extensively characterized brain region in the $Df(16)A^{+/-}$ mouse.^{5,11} Total lysis extracts from PFC tissues of two independent cohorts of $Df(16)A^{+/-}$ mice (11 $Df(16)A^{+/-}$ vs 10 wild-type / 13 Df $(16)A^{+/-}$ vs 13 wild-type) were analysed, as well as from the HPC of one of the cohorts (11 $Df(16)A^{+/-}$ vs 10 wild-type). Additionally, PFC tissue of both cohorts were combined (n = 40) and prepared with a different gel-based procedure to increase proteome coverage and statistical power. Shotgun profiling analysis resulted in the identification of 578 distinct proteins (9605 peptides) in the PFC (cohort 1) of which 59 proteins showed significant genotypic differences (P* < 0.05). Likewise, 715 distinct proteins (14 334 peptides) (cohort 2) were identified in cohort 2, with 186 proteins

In order to identify robust alterations of $Df(16)A^{+/-}$ hemizyaosity, we cross-compared all proteomic profiling studies among each other and to previous transcriptomic profiling studies^{5,11} of the $Df(16)A^{+/-}$ mice (Figure 1a). Most consistently, we found an average decrease of 20% in septin 5 levels (Figure 1d), a protein encoded in the 22g11.2DS critical region. In the case of other 22g11.2DS encoded proteins, we detected a reduction in the levels of protein DGCR14 (ES2EL) (ratio = 0.85, $P = 8.2 \times 10^{-6}$, 1 peptide) in one of the proteomic cohorts. We further detected a 30% increase in the levels of the hypoxanthine-guanine phosphoribosyltransferase (HPRT) in the $Df(16)A^{+/}$ [–] mice. Two mini-cassettes of the Hprt gene are part of the viral transfection vector where they serve as a positive selection marker for successful recombination and deletion of the 22q11.2 critical region. The identified changes in these proteins substantiate the technical robustness of our proteomic platform. All other 22q11.2DS-encoded proteins, which could serve as positive controls, were below the detection limit, probably owing to low or absent expression in the brain.³¹

Other identified proteins, which showed alterations in abundance, are either primary or secondary effects of abnormal miRNA metabolism or hemizygosity of 22q11.2 genes. For example, we identified a decrease in glutathione *S*-transferase Mu7 (GSTM7) in three of the proteomic studies, which could be a secondary effect of the *Txnrd2* (thioredoxin reductase 2) hemizygosity. Both proteins have been implicated in the regulation of the intracellular redox environment.^{34–37} Furthermore, we discovered robust decreases of mitochondrial proteins (cytochrome *c* oxidase subunit 5B, mitochondrial (COX5B), NADH-ubiquinone oxidoreductase chain 4 (NU4M)). A possible reason for these alterations is the involvement of several 22q11.2 genes (*Prodh, Comt, Txtp, Slc25a1*) in mitochondrial function. Indeed, using the IPA software, we confirmed that 'mitochondrial dysfunction' was a common pathway affected across all profiling studies (Figure 1c).

In order to identify proteins that might be upregulated owing to miR-185 hemizygosity, we analysed all upregulated detected proteins using bioinformatic miRNA target prediction tools (MirDB, Targetscan, MirBase, DIANA), which were based on thermodynamics (DIANA), seed complementarity (Targetscan) or support vector machines (MirDB). Potential miR-185 targets are disk large homolog 2 (DLG2), AP2-associated protein kinase 1 (AAK1), sodium/potassium-transporting ATPase subunit alpha-3 (AT1A3), O-GlcNAc transferase subunit p110 (OGT1) and protein bassoon (BSN), which were found to be upregulated at the proteomic and transcriptomic levels.⁵

We were not able to identify a significant increase in sarcoplasmatic/endoplasmic reticulum calcium ATPase 2 (SERCA2 *—Atp2a2*), another recently proposed target of 22q11.2DS-associated miRNA dysregulation,³⁸ in any of our proteomic screens. This is consistent with Fenelon *et al.*,¹¹ who failed to confirm a change in the transcript and protein levels of *Atp2a2*.

Remarkably, the differentially expressed proteins showed an overall increase in abundance levels across all FC proteomic studies (Figure 1b), which is likely owing to a decrease in about 20% of the miRNAs⁵ (including miR-185) and subsequent upregulation of targets. This finding is consistent with the transcriptomic profiling study by Fenelon *et al.*¹¹ On the other hand, the HPC proteome showed a trend towards downregulation of differentially expressed proteins. Because of the lack of replication studies, further confirmation of the opposing changes in the HPC is needed. Nevertheless, the results are consistent with reported miRNA changes in PFC and HPC identified through

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				Proteomics									Transcrip	otomics					
			Gene	PFC -Cohort 1 PFC- Cohort 2 I			PFC- Combined HPC						Df(1	5)A+/-	Prediction				
Uniprot Ac	cc.No	Protein name	name	PC	Ratio	FDR p	РС	Ratio	FDR p	РС	Ratio	FDR p	РС	Ratio	FDR p	Ratio	PFC	HPC	miRNA-185
SEPT5 Q9	9Z2Q6	Septin-5	Pnutl1	8	0.92	2.2E-03	14	0.82	<e-15< td=""><td>2</td><td>0.79</td><td><e-15< td=""><td>2</td><td>0.74</td><td>1E-07</td><td>0.82</td><td>▼</td><td>•</td><td>1of4</td></e-15<></td></e-15<>	2	0.79	<e-15< td=""><td>2</td><td>0.74</td><td>1E-07</td><td>0.82</td><td>▼</td><td>•</td><td>1of4</td></e-15<>	2	0.74	1E-07	0.82	▼	•	1of4
H2B1B Q6	64475	Histone H2B type 1-B	Hist1h2bb	2	1.16	3.5E-03	4	1.11	7.3E-03	3	1.12	3.6E-03		not dete	ected	1.13			-
PP2BC P4	8455	Calmodulin-dependent calcineurin A y isoform	Ppp3cc	3	1.16	1.2E-02	1	1.17	8.7E-04	1	1.06	4.6E-04	14	0.97	0.2976	1.13			-
GSTM7 Q8	30W21	Glutathione S-transferase Mu 7 (EC 2.5.1.18)	Gstm7	2	1.03	7.2E-01	2	0.93	5.7E-03	6	0.88	7.1E-03	7	0.9	0.0248	0.90			D
COX5B P1	.9536	Cytochrome c oxidase subunit 5B, mitochondrial	Cox5b	2	0.95	2.7E-02	4	0.80	1.9E-07		not dete	ected	1	1.04	0.0002	0.87		•	D
DLG2 Q9	91XM9	Disks large homolog 2	Dlg2	2	1.11	4.1E-02	7	1.33	7.4E-04		not dete	ected		not dete	ected	1.22		A	4of4
NU4M PO	3911	NADH-ubiquinone oxidoreductase chain 4 (EC 1.6.5.3)	Mtnd4	1	1.15	3.1E-07	16	1.07	2.0E-02		not dete	ected	1	1.02	0.7437	1.11			-
H2A1H Q8	BCGP6	Histone H2A type 1-H	Hist1h2ah	1	1.11	3.7E-02	2	1.10	1.3E-04		not dete	ected	1	1.2	0.0871	1.10			-
AAK1 Q3	3UHJO	AP2-associated protein kinase 1 (EC 2.7.11.1)	Aak1	1	1.19	2.8E-05	2	1.51	9.1E-06	4	0.99	7.1E-01		not dete	ected	1.35			1of4
NRDC Q8	BBHG1	Nardilysin (EC 3.4.24.61)	Nrd1	n	ot deteo	cted	2	0.90	4.0E-02	1	0.9	4.3E-03		not dete	ected	0.90			D
GNAS2 P6	3094	Guanine nucleotide-binding protein G(s) su α isoform	Gnas	8	0.97	3.3E-01	16	1.08	1.9E-03	3	1.14	3.0E-04	6	0.86	0.229	1.11			-
PURA P4	2669	Transcriptional activator protein Pur-alpha	Pura	4	1.03	5.7E-01	7	0.89	1.9E-03	1	0.85	6.2E-08	2	0.93	0.4909	0.87		•	D
PALM Q9	9ZOP4	Paralemmin-1	Palm	5	1.01	7.5E-01	3	1.68	3.1E-08	1	1.15	1.3E-03	2	0.97	0.7648	1.42			2of4
STIP1 Q6	50864	Stress-induced-phosphoprotein 1	Stip1	4	1.03	8.4E-01	13	0.91	1.8E-04	1	0.85	1.2E-02	6	1	0.9122	0.88		•	D
HPRT PO	0493	Hypoxanthine-guanine phosphoribosyltransferase	Hprt1	1	1.24	6.2E-11	r	not dete	cted	3	1.43	<e-15< td=""><td></td><td>not dete</td><td>ected</td><td>1.33</td><td></td><td></td><td>PC</td></e-15<>		not dete	ected	1.33			PC
NFL PO	8551	Neurofilament light polypeptide (NF-L)	Nefl	17	1.11	3.4E-05	r	not dete	cted	29	1.09	3.3E-12	11	0.98	0.6312	1.10			-
AINX P4	6660	Alpha-internexin	Ina	7	1.12	9.2E-04	r	not dete	cted	29	1.15	<e-15< td=""><td>16</td><td>0.96</td><td>0.0623</td><td>1.14</td><td></td><td></td><td>-</td></e-15<>	16	0.96	0.0623	1.14			-
CPLX2 P8	4086	Complexin-2	Cplx2	9	1.10	5.4E-04	14	0.98	4.0E-01	1	1.11	1.1E-07	5	0.97	0.7674	1.10			1of4
H14 P4	3274	Histone H1.4	Hist1h1e	2	1	0.4494	3	0.91	0.0254		not dete	ected	10	0.86	0.03	0.88			D
H2B3A Q9	9D2U9	Histone H2B type 3-A	Hist3h2ba	7	1	0.9874	4	0.74	6.5E-06	4	1.04	0.235	3	0.94	0.05	0.84			D
VAMP2 P6	3044	Vesicle-associated membrane protein 2 (VAMP-2)	Vamp2	10	1	0.892	23	1.07	2.2E-03	14	1.02	0.268	8	1.08	5.5E-03	1.08			3of4
EIF3H Q9	91WK2	Eukaryotic translation initiation factor 3 subunit 3	Eif3h	n	ot deteo	cted	1	0.88	0.0091		not dete	ected	1	0.91	<e-15< td=""><td>0.90</td><td></td><td></td><td>D</td></e-15<>	0.90			D
NCOA5 Q9	91W39	Nuclear receptor coactivator 5	Ncoa5	n	ot deteo	cted	3	0.82	0.0414		not dete	ected	1	0.87	0.01	0.85			1of4
NSG1 Q6	52092	Neuron-specific protein family member 1	Nsg1	n	ot deteo	cted	2	0.87	0.0209		not dete	ected	1	0.75	0.01	0.81			D
Proteins de	etectec	l only in one cohort*																	
OGT1 Q8	BCGY8	O-GlcNAc transferase subunit p110	Ogt	2	1.58	2.3E-08	r	not dete	cted		not dete	ected		not dete	ected				1of4
KIF1B Q6	50575	Kinesin-like protein KIF1B	Kif1b	4	1.15	0.0203	r	not dete	cted		not dete	ected		not dete	ected				4of4
BSN 08	88737	Protein bassoon	Bsn	n	ot deteo	cted	r	not dete	cted	18	1.05	8.4E-03		not dete	ected				3of4
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Figure 1. (a) Venn diagrams of the overlap of all identified proteins (left) and of all significantly altered proteins (right) identified in label-free LC-MS^E (liquid chromatography-mass spectrometry) proteomic profiling studies of $Df(16)A^{+/-}$ mice compared with *wild-type* mice (Supplementary Table S4 for full list). (b) Percentages of significantly increased (red) and decreased (green) proteins in $Df(16)A^{+/-}$ mice across all proteomic studies. Mean percentages show significantly increased protein levels in the prefrontal cortex (PFC; p < 0.05, *t*-test). PFC combined are samples from both cohorts combined and in-gel digested (see Material and methods). As the hippocampal (HPC) was only investigated from one cohort, significant levels could not be established. (c) Top ingenuity pathway analysis: altered pathways in the PFC and HPC. (d) Detailed information of the overlapping proteins identified as significantly changing across all proteomic screens. Proteins were sorted by the number of proteomic studies in which they have been detected as significantly altered. Only proteins with a mean ratio of > 1.1 or < 0.9 across all identified studies are displayed. *P*-values were determined using MSstats and corrected to control for multiple hypothesis testing after Benjamini–Hochberg.¹¹⁴ (Supplementary Table S3 for full information). Proteins identified by one peptide (italic) were included if they were overlapping between the proteomic screens. Results were compared with the transcriptomic results using Affymetrix chips published by Stark *et al.*⁵ and Feneleon *et al.*¹¹ Stark reported 716 transcripts altered in the FC and 85 transcripts in the HPC (false-discovery rate = 0.01). Significantly changed transcripts are marked with an **4** and **7** according to their reported fold change direction in comparison to *wild-type* mice. Four different bioinformatic miRNA target prediction tools were used to assess if the upregulated proteins are potential Mir-185 targets. Xof4, predicted in X of the four p

transcriptomic profiling, as nearly twice the number of miRNAs were found to be decreased in the PFC (59 miRNAs) than in the HPC (30 miRNAs).⁵

In silico analysis revealed an overrepresentation of developmentally regulated pathways related to nucleosome and chromatin assembly and chromatin-level regulation across all PFC proteomic studies (Figure 2). Further pathway analysis using the IPA software allowed us to identify overlaps between canonical pathways. The proteomic alterations were associated with the canonical pathways 'glycolysis I', 'gluconeogenesis' and 'mitochondrial dysfunction' (Figure 1c) in the PFC and 'Rho signalling' in the HPC, which was also reflected in the enrichment analysis.

Brain characterization—quantitative metabonomics

We next carried out metabonomic profiling of PFC and HPC tissues from the $Df(16)A^{+/-}$ mouse in order to determine whether the above changes in proteins could be linked to alterations in metabolites. Therefore, after the initial identification of a metabolite candidate, the sample was re-run and the peaks were subjected to fragmentation in an MS/MS experiment. Identification of a specific metabolite was accomplished though comparison of the fragmentation pattern of a candidate metabolite with the known MS/MS fragmentation pattern in public and in-house databases. This analysis showed significant changes in 21 m/z-retention time pairs in the PFC, and we were able to identify 7 of

Pre-Frontal Cortex

Pre-Frontal Cortex nbined

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GO-Term		р	GO-Term	com
0065004	protein-DNA complex assembly	0.0016	0040029	regulation of gene exp
0031497	chromatin assembly	0.0016	0009112	nucleobase metabolic
0006334	nucleosome assembly	0.0016	0045044	Pos. regulation of tran
0071103	DNA conformation change	0.003	0045944	polymerase II promot
0071824	protein-DNA complex	0.0033	0006260	DNA replication
0071024	organization	0.0055	0006257	Regu. of transcription
0034728	nucleosome organization	0.0033	0006557	promoter
0006333	chromatin (dis)assembly	0.0033	0006366	transcription from RN
0006323	DNA packaging	0.0033	0000300	promoter
0031648	protein destabilization	0.0121	0045034	negative regulation of
0090304	nucleic acid metabolic process	0.0123	0043334	compound metabolic
0051276	chromosome organization	0.0338	0006305	DNA alkylation
0009058	biosynthetic process	0.0355	0072520	pyrimidine-containing
0016070	RNA metabolic process	0.0360	0072529	process
0019751	polyol metabolic process	0.0365	0043550	regulation of lipid kina
0006106	fumarate metabolic process	0.0365	0006304	DNA modification
			0044728	DNA methylation or d

0040029	regulation of gene expression, epigenetic	0.0025
0009112	nucleobase metabolic process	0.0025
0045944	Pos. regulation of transcription RNA polymerase II promoter	0.0037
0006260	DNA replication	0.0039
0006357	Regu. of transcription from RNA polymerase II promoter	0.0078
0006366	transcription from RNA polymerase II promoter	0.0078
0045934	negative regulation of nucleobase-containing compound metabolic process	0.0078
0006305	DNA alkylation	0.0084
0072529	pyrimidine-containing compound catabolic process	0.0084
0043550	regulation of lipid kinase activity	0.0084
0006304	DNA modification	0.0084
0044728	DNA methylation or demethylation	0.0084
0046113	nucleobase catabolic process	0.0084
0006306	DNA methylation	0.0084
0043551	regulation of PI3K activity	0.0084

Hippocampus

Pre-Frontal Cortex

	Cohort 2			Cohort 1	
GO-Term	CONDICE	р	GO-Term		р
0006333	chromatin assembly or disassembly	0.0375	0007266	Rho protein signal transduction	0.0017
0006325	chromatin organization	0.0394	0007229	integrin-mediated signaling pathway	0.0087
0034612	response to tumor necrosis factor	0.0464	0023021	termination of signal transduction	0.0098
0006359	Reg. of transcription RNA polymerase III promoter	0.0464	0038032	termination of G-protein coupled receptor signaling pathway	0.0098
0006122	mitochondrial electron transport,	0.0464	0007265	Ras protein signal transduction	0.0164
0010E10	Pos. regulation of phospholipase	0.0464	0035023	regulation of Rho protein signal transduction	0.0331
0010318	activity	0.0404	0045454	cell redox homeostasis	0.0331
0070995	NADPH oxidation	0.0464	0016337	cell-cell adhesion	0.0370
0002087	see legend	0.0464	0007272	ensheathment of neurons	0.0421
0006383	transcription RNA polymerase III		0045921	positive regulation of exocytosis	0.0421
0000303	promoter	0.0464	0008366	axon ensheathment	0.0421
0060193	positive regulation of lipase activity	0.0464	0045744	Neg. Reg of G-protein coupled receptor	0 0 4 2 1
0071103	DNA conformation change	0.0525	0045744	protein signaling pathway	0.0421
0008610	lipid biosynthetic process	0.0525	0045044	positive regulation of transcription from RNA	0 0 4 2 9
0051276	chromosome organization	0.0533	0045944	polymerase II promoter	0.0428
0006334	nucleosome assembly	0.0592	0009072	aromatic amino acid family metabolic process	0.0461
0034728	nucleosome organization	0.0592	0060070	canonical Wnt receptor signaling pathway	0.0461

Figure 2. Significantly enriched pathways identified by gene set enrichment analysis using GOstats (proteins per pathway > 3). The top 15 significant Gene Ontotlogy (GO) terms are displayed. GO terms falling into the category of chromosomal regulation are highlighted in grey. A full colour version of this figure is available at the *Molecular Psychiatry* journal online.

these metabolites via MS/MS fragmentation. In the HPC, we detected 27 significantly changed *m*/*z*-retention time pairs, which we could match to 12 metabolites using metabolome databases and identified 4 via MS/MS fragmentation (Table 1, Supplementary Table S5). The most significant change in the PFC was a decrease in ceramide phosphoethanolamine (SP0302), which is a sphingomyelin analog. SP0302 may be linked to the hemizygosity of the ZDHHC8 palmitoyltransferase in the $Df(16)A^{+/-}$ mouse, as palmitoyl-CoA is a precursor in the ceramide synthesis. Alongside, further ceramides were found to be decreased in the PFC and S-palmitoyl-L-cysteine was decreased in the HPC. In the HPC, we also found that sphingomyelin (SM(d17:1/24:1) was the most significantly increased metabolite (ratio = 1.72, P = 0.031). We were able to validate sphingomyelin's databank identification using tandem MS/MS. We detected changes in further lipids such as phosphatidylcholines and glycerolipids, which reflect the

changes in lipid metabolism found in the proteomic pathway analysis of $Df(16)A^{+/-}$ mice (Figure 2). In addition, several carnitines, which have a role in fatty acid metabolism, were found to be decreased (Table 1). Furthermore, proteomic abnormalities in aromatic amino acids could be confirmed via metabonomic profiling, detecting changes in different tyrosine derivates in both regions (N-stearoyl tyrosine and metyrosine). Although we were not able to validate the identification of these metabolites using MS/MS, their changes were reflected in the protein set enrichment analysis of the HPC 'aromatic amino-acid family metabolic process'. A decrease in panthothenic acid-essential in the synthesis of coenzyme-A (CoA)—was also reflected in the proteomic pathway analysis with CoA biosynthesis being affected in cohort 1, where we found increased levels of two of the key enzymes in this pathway.

mz_Rt pair	lon mode	HMDB/Chempub ID	Name	Class/function	Exact mass	Delta (mz – exact mass)	MS/MS	Ratio	P-value
Frontal cortex									
645.5_1323	+	LMSP03020039	PE-Cer(d14:2(4E,6E)/19:0) PE-Cer(d15:2(4E,6E)/18:0)	Ceramide phosphoethanolamines (SP0302)	644.4893	6.5E-05	ID	0.64	0.0001
426.4_681	+	HMDB06464 HMDB05065	Elaidic carnitine Oleoylcarnitine	Acyl carnitine	425.3505	4.5E-03	ID	0.59	0.0101
232.2 270	+	HMDB02013	Butvrvlcarnitine	Acvl carnitine	231.147	4.4E-03	ID	0.33	0.0133
448.3 667	+	74380333	N-docosanoyl taurine	N-acyl amine	447.3382	4.0E-04	NP	0.70	0.0133
		123060515	N-stearovl tyrosine	N-acyl amine	447.3349	3.7E-03			
765.6 1179	+	123064881	PG(O-20:0/16:0)/PG(O-16:0/20:0)	Glycerophosphoglycerol	764.5931	1.6E-03	ID	0.69	0.0220
367.1 609	+	8439	Salicin 6-phosphate	Glycoside phosphate	366.0716	3.0F-03	NP	1.44	0.0350
650.4 1019	+	135642574	PC(16:0/9:0(CHO))	Phophatidylcholine	649 4319	4 5E-03	NP	0.71	0.0350
530 5 1146	+	-	Water loss from $#16 (5485, 1146sec)$					0.58	0.0350
130 25	+	HMDR33561	2-Acetyl-4 5-dibydrothiazole	_	129 0248	1 4F-03	NP	0.50	0.0550
184 1 1095		HMDB33141	Fragment of a phosphatidylcholine		183 0796	5.0E-04		1 3 3	0.0435
550 6 1035	-		Cor(m18.1(AE)/18.0)	Coramida/sphingolipid	540 5485	0.00013		0.55	0.0435
631 6 1302	т +	LMSP03020003	$PE_Cor(d14.1(AE)/18.1(07))/PE_Cor(d14.2)$	Ceramide phosphoethanolamines	630 4737	0.00013	חו	0.55	0.0435
031.0_1302	Ŧ	LIVI3F03020003	(4E,6E)/18:0)/PE-Cer(d16:2(4E,6E)/16:0)	Ceramide phosphoethanolamines	030.4737	0.00022	D	0.08	0.0455
252.1_526	+	HMDB00101	Deoxyadenosine	Purine nucleosides and analogues	251.101	0.00412	NP	2.62	0.0435
643.5_1269	+	HMDB07312	DG(18:3(9Z,12Z,15Z)/20:2(11Z,14Z)/0:0) (iso2) (several)	Diacylglycerol	642.5223	9.8E-03	NP	0.60	0.0435
Hippocampus									
382.3_619	+	4266008 24701448	Prostaglandin F2 alpha dimethyl amide 5,6-DiHETrE-EA (several)	Prostaglandin Endocannabinoid	381.2879	5.8E-03	NP	0.39	0.0033
771.6 1472	+	123068775	SM(d18:2/21:0)	Ceramide posphocholine	770.6302	4.5E-03	NP	0.57	0.0229
_		123067209	PA(O-20:0/22:2(13Z,16Z))	Glycerophosphate	770.6189	6.8E-03	NP		
359.2 661	+	HMDB12983	Kinetensin 1-3	Peptide	358.2329	5.9E-03	NP	0.56	0.031
799.7 1372	+	123068785	SM(d17:1/24:1) or SM(d18:2/23:0)	Sphingomyelin	798.6615	7.8E-03	ID	1.72	0.031
360.3 662	+	_	S-palmitovl-L-cvsteine	Palmitovlated residue	359.2494	5.7E-03	NP	0.59	0.031
140.1 1645	+	207572	4-Amino-5-hydroxymethyl-2-	Thiamin metabolism	139.0746	1.7E-04	NP	0.35	0.042
		5035	methylpyrimidine L-Histidinal	Histidin metabolism					
545.3_725	+	_	PC (no database match, Isotope of 544.3 (PC 20:4))	Phophatidylcholine	—	—	Т	0.67	0.042
882.6_1302	+	123061384	PC(22:5(4Z,7Z,10Z,13Z,16Z)/22:5 (4Z,7Z,10Z,13Z,16Z))	Phophatidylcholine	881.5935	7.7E-03	NP	0.69	0.042
479.3 760	_	_	Phospholipid—ID cannot be confirmed	_	_	_	т	0.55	3E-05
218.1 300	_	149588	Pantothenic acid	Vitamine	219.1107	4.6E-05	NP	0.80	0.019
194.1 632	_	HMDB14903	Metvrosine	Tyrosine	195.0895	2.3E-04	NP	0.61	0.023
12111_032		HMDR29217	Tyrosine methylester		122.0023			0.01	0.020
566 3 758	_	I MGP01050056	PC(22:6(47 77 107 137 167 197)/0:0)	Glycerophosphocholine	567 3325	3 2F-05	ID	1 30	0.023
567.3 758	_	LMGP01050056	Isotope of 566.3 758	arycerophosphoenonne	507.5525	J.22 0J		1.50	0.025
480.3 746	_	LMGP01060010	PC(O-16:0/0:0)	Phosphatidylcholine	481 3532	-7.8F-05	ID	1.43	0.043
F22.2 015		LMSD02010055	Cor(d14.2)(AE 6E)/20.1(117))	N-acylsphingosines (ceramides)	533 / 808	-0.0003		0.66	0.042

Abbreviations: HFC, hippocampus; ID, identified; MS/MS, tandem mass spectrometry; NP, cannot (dis)prove ID; PFC, prefrontal cortex; T, tentative, class indicated but not database match. Databank searches in METLIN, Lipidmaps, HMDB and MS/MS fragmentation were employed for metabolite identification.

Proteomic validation

We developed highly specific and sensitive targeted SRM assays in order to follow-up and validate alterations identified in the proteomic discovery phase. We assayed for potential miRNA targets (OGT1, ADA10) as well as altered proteins identified in the LC-MS^E profiling approach (PP2BC, NRDC). The inclusion of the miR-185 targets was based on LC-MS^E analysis, bioinformatic predictions and the shotgun proteomics analysis-derived in silico pathway analysis. We further included 6 of the 27 22g11.2 gene products to determine whether or not the mRNA changes are reflected at the protein level and could serve as potential positive controls. The vesicular glutamate transporter 1 (VGLU1) was included based on evidence obtained from the transcriptomic screen where VGLU1 was one of the three transcripts found to be significantly upregulated in both PFC and HPC.⁵ Unfortunately, we were not able to develop assays for DLG2 and BSN owing to assay design issues. We further extended the validation screen to include proteins of pathways that have been implicated in some of the major symptoms observed in 22q11.2DS. This included the mammalian target of rapamycin (mTOR) signalling pathway, which has been linked to autism-like behaviour, cognitive function and glutamatergic signalling. Details of the composition of the multiplex assay can be found in Table 2.

The SRM analysis (Table 2) validated the expected decrease in all the five tested 22g11.2 gene products and an increase of the HPRT peptides from the vector cassette, showing the technical robustness of the developed method. Importantly, SRM confirmed an increase in the miR-185-target OGT1 in PFC and HPC tissue. Alterations of OGT1 have been identified in the transcriptomics⁵ and in proteomics (Figure 1) high-throughput screens of the PFC. The OGT1 enzyme exhibits protein N-acetylglucosaminyltransferase activity, which leads to attachment of O-GlcNAc onto intracellular proteins.³⁹ Furthermore, we were able to validate an increase in levels of the glycogen synthase kinase-3 beta (GSK3b) and kinesin heavy chain isoform 5A (KIF5A), a microtubule-dependent molecular motor that is important for neuronal functions especially for GABA_A receptor transport.⁴⁰ Similar to the transcriptomic and PROTEOMIC profiling analyses, we did not find altered levels of the miR-185 target transforming protein RhoA (RHOA). This might be due to secondary regulation caused by the hemizygous genes or downregulated miRNA that might involve alterations in transcriptional levels.

Further validated proteins were serine/threonine protein phosphatase 2B catalytic subunit gamma isoform (PP2BC), the neurofilament light polypeptide (NFL) and VGLU1. The latter was one of the only three proteins found in the transcriptomic screen in both PFC and HPC. The extracellular signal-regulated kinase (ERK) pathway was most significantly affected in the PFC, while abnormal glutamatergic signalling was predominately found in the HPC. The latter finding is consistent with published transcriptomic profiling results from the $Df(16)A^{+/-}$ mice.⁵

DISCUSSION

To the best of our knowledge, this study represents the first comprehensive quantitative proteomic and metabonomic characterization of a 22q11.2DS mouse model. We assume that the phenotype of this disease model is due to cumulative effects of abnormal miRNA regulation and hemizygosity of genes residing within the 22q11.2 chromosome region, which requires analysis on the systems level rather than by targeting the effects of individual genes.^{41–43}

Multiplex immunoassay profiling analysis identified altered serum levels of only one protein, the endopeptidase MMP-9, which has been implicated in many pathological conditions such as cancer, cardiovascular disease,⁴⁴ amyotrophic lateral sclerosis⁴⁵ and stroke.^{46,47} Recently, MMP-9 has been reported to have a role in the plasticity of the central nervous system⁴⁸ by regulating the activity of PFC.⁴⁹ A functional polymorphism of the MMP-9 gene in schizophrenia⁵⁰ and bipolar disorder type II⁵¹ has also been reported. The identified alterations might serve as surrogate readouts for the psychiatric comorbidities in the 22q11.2DS and help to elucidate the molecular mechanisms of the immunodeficiency in 22q11.2DS patients.

The proteomic brain profiling analysis revealed PFC and HPC changes in various molecular pathways associated with chromatin remodelling and RNA transcription, indicative of an epigenetic component of 22q11.2DS. Recently, it has been shown that developmentally regulated genes involved in chromatin remodelling are overrepresented among targets of *de novo* mutations in schizophrenia.^{52,53} Therefore, this study highlights a possible contribution of chromatin remodelling in the manifestation of the psychiatric and cognitive phenotype in 22q11.2DS. Other associated pathways were glycolysis, gluconeogenesis, lipid biosynthesis and the mitochondrial transport chain. Effects on the latter two pathways were also reflected in the transcriptomic enrichment analysis of the $Df(16)A^{+/-}$ mouse⁵ and together with lipid metabolism have frequently been linked to psychiatric conditions.^{54,55} Altered levels of sphingomyelin and ceramide have been reported in erythrocytes and postmortem brain tissues of schizophrenic patients.56

Based on the proteomic and metabonomic findings, we created a schematic model summarizing the most prominent molecular network findings in the $Df(16)A^{+/-}$ mouse. Interestingly, all implicated pathways can be linked through one of the most consistent and strongest proteomic candidates, (OGT1), which is a predicted miR-185 target (Figure 3).

OGT1 adds single terminal *N*-actylglucosamine (O-GlcNAc) residues to serine/threonine residues of specific target proteins.^{39,57} This represents a crucial signalling modification relevant for cellular functions, such as epigenetic regulation, transcription activation processes, ^{58,59} chromatin remodelling,⁶⁰ translation, proteasomal degradation, signal transduction and stress responses.⁶¹ Cellular substrates recognized by OGT1 are mostly present in macromolecular complexes, such as nuclear pores, 62-65 the transcriptional machinery 66-68 and vesicle docking components.^{69–72} Consistently, we identified abnormalities in biological functions associated with these complexes in the Df $(16)A^{+/}$ brain and most of these were found in the PFC. Interestingly, we identified robust alterations of several histone proteins in the $Df(16)A^{+/-}$ mouse model (Figure 1d). O-GlcNAcylation occurs on all four core-nucleosomal histone proteins (H2A, H2B, H3 and H4).⁷³ However, such chromatin and transcriptionallevel abnormalities could also be due to the hemizygosity for four transcription factors (for example, Tbx1, Hira) in the $Df(16)A^{+/}$ mouse. Abnormalities in glycosylation have already been postulated to occur in the $Df(16)A^{+/-}$ mouse by a bioinformatic enrichment analysis, which found coordinated dysregulation of Golgi-related putative miR-185 targets in the $Df(16)A^{+7-}$ mouse.⁹ Furthermore, the miR-185 target Mirta22 has been localized to the Golgi apparatus. This is the first study to suggest that changes in OGT activity may be linked to the above glycosylation alterations. This is also supported by the fact that in silico analysis of the 25 identified human OGT interacting proteins showed an enrichment of intra Golgi-vesicle-mediated transport and vitamin biosynthetic processes.⁷⁴ Also, bioinformatic analysis of an OGT-interactome substantiated the role of OGT in transcriptional regulation.⁷⁴ A potential role for O-GlcNAcylation in modifying synaptic efficacy and cognition has previously been suggested.⁷⁵ O-GlcNAcylation of AMPA receptors GluA2 has been linked to long-term depression in HPC synapses.⁷⁶ OGT1 is also involved in mitochondrial function,⁷⁷ consistent with the finding of alterations in the levels of mitochondrial proteins in this study. Overexpression of OGT1 leads to decreases in proteins involved in the respiratory chain and the TCA cycle,⁷⁷ as well as altered mitochondrial morphology. The majority of O-GlcNAc-modified proteins are soluble nuclear or

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Table 2. Significantly changed proteins identified using label-based LC-SRM (targeted proteomics) in the PFC and HPC of the $Df(16)A^{+/-}$ mouse compared with *wild-type* mice (PFC = 20 vs 20, HPC = 11 vs 10)

3							Prefrontal cortex							Hippocampus					
							Proteomics							Proteomics					
						Ĺ	Discovery (D))	Targeted proteomics (SRM)					D	Targeted proteomics (SRM)				
					Transcrip-	10 vs 10	12 vs 10	20 vs 20	20		20 vs 21		Transcrip-	10 vs 10		11 v	s 10		
Uniprot	Acc. no.	Protein	Gene	Label	tonnics				Re	atio	Р	P*	tonnes		Rat	io	Ρ	P*	
22q11.2 critical Sep-05 RANG Q8JZU2 PROD COMT HPRT	l gene prod Q9Z2Q6 P34022 Q8JZU2 Q9WU79 O88587 P00493	ucts+vector construct Septin-5 Ran-specific GTPase-activating protein Protein Slc25a1 Proline dehydrogenase 1, mitochondrial Catechol O-methyltransferase Hypoxanthine-guanine phosphoribosyltransferase	Sep-05 Ranbp1 Slc25a1 Prodh Comt Hprt1	22q11.2 22q11.2 22q11.2 22q11.2 22q11.2 22q11.2 Vector construct	* 4 4 4 4		▼ 	▼ 	V V V V	0.78 0.55 0.79 0.62 0.76 1.56	5.2E-11 < E-15 4.8E-08 3.5E-07 1.4E-06 < E-15	1.1E-09 < E-15 7.5E-07 4.4E-06 1.4E-05 < E-15		▼	* * * *	0.85 0.47 0.64 0.90 0.71 1.77	1.8E-04 < E-15 6.7E-10 0.26 1.6E-03 < E-15	4.9E-04 < E-15 5.9E-09 0.30 3.6E-03 < E-15	
miRNA targets GSK3B OGT1 KIF5A A4 KDM2A RHOA MBOA5 ADA10 ANFY1	Q9WV60 Q8CGY8 P33175 P12023 P59997 Q9QUI0 Q91V01 O35598 Q810B6	Glycogen synthase kinase-3 beta O-GlcNAc transferase subunit p110 Kinesin heavy chain isoform 5A Amyloid beta A4 protein Lysine-specific demethylase 2A Transforming protein RhoA Lysophospholipid acyltransferase 5 Disintegrin metalloproteinase domain- containing protein 10 Ankyrin repeat and FYVE domain- containing protein 1	Gsk3b Ogt Kif5a App Kdm2a Rhoa Lpcat3 Adam10 Ankfy1	miRNA target (2/4) miRNA target (1/4) + Figure 1d miRNA target (1/4) miRNA target (1/4) miRNA target (2/4) miRNA target, Liu <i>et al.</i> , ¹¹⁵ miRNA target (1/4) + Figure 1d miRNA target (3/4) miRNA target (1/4)	☆ NS NS NS NS NS NS NS	NS — NS 	 NS 	▼ ▲ NS —	NS NS NS NS NS NS	1.14 1.22 1.22 1.13 1.02 1.05 1.09 1.03	4E-06 1.3E-02 3.6E-02 6.5E-02 0.77 0.37 0.13 0.59 0.85	3.2E-05 3.5E-02 0.09 0.13 0.86 0.48 0.21 0.73 0.91	NS NS NS NS NS NS NS NS	_ - -	NS NS NS NS NS NS NS	1.16 1.20 1.15 1.11 1.06 1.08 1.04 1.22	6.1E-04 8.9E-03 0.18 0.16 0.71 0.05 0.696 0.05	1.5E-03 1.6E-02 0.22 0.21 0.74 0.08 0.7443 0.07	
Discovery PP2BC	P48455	Serine/threonine-protein phosphatase 2B	Ррр3сс	Figure 1d	NS					1.06	1.1E-02	3.1E-02	NS	NS		1.21	4.7E-10	4.8E-09	
PP2BB	P48453	catalytic γ isoform Serine/threonine-protein phosphatase 2B	Ppp3cb		NS	NS	NS	NS		1.16	5.8E-04	2.4E-03	NS	NS		1.16	8.9E-03	1.6E-02	
NFL AINX NRDC VGLU1	P08551 P46660 Q8BHG1 Q3TXX4	Neurofilament light polypeptide (NF-L) Alpha-internexin (Alpha-Inx) Nardilysin (EC 34.24.61) Vesicular glutamate transporter 1 (VGluT1)	Nefl Ina Nrd1 Slc17a7	Figure 1d Figure 1d Figure 1d Transcriptomics ⁵	NS NS NS 企	NS NS	- 		NS NS	1.09 0.96 1.02 1.16	1.8E-03 0.31 0.60 1.6E-06	6.3E-03 0.42 0.73 1.4E-05	NS NS NS 介	NS NS NS NS	NS	1.15 1.21 0.99 <i>1.19</i>	1.6E-04 3.0E-08 0.89 1.5E-04	4.9E-04 2.3E-07 0.90 4.8E-04	
Implicated path mTOR-autist MTOR MK03 MK01 PEA15 KPCG	hways n spectrum Q9JLN9 Q63844 P63085 Q62048 P63318	disorders and mental retardation Serine/threonine-protein kinase mTOR Mitogen-activated protein kinase 3 (ERK-1) Mitogen-activated protein kinase 1 (ERK-2) Astrocytic phosphoprotein PEA-15 Protein kinase C gamma type	Mtor Mapk3 Mapk1 Pea15 Prkcg		NS NS NS 압		 NS	 NS		1.28 1.16 1.11 1.15 1.11	4.8E-04 2.3E-05 5.9E-05 5.0E-05 1.2E-03	2.1E-03 1.5E-04 3.0E-04 2.8E-04 4.6E-03	NS NS NS NS	NS NS MS NS	NS	1.10 1.23 1.07 0.91 1.21	0.30 3.7E-05 6.3E-03 2.8E-02 6.2E-07	0.34 1.3E-04 1.2E-02 4.4E-02 2.9E-06	
Glutamatergic NMDZ1	signalling— P35438	-schizophrenia and affective disorders Glutamate receptor ionotropic, NMDA 1	Grin1	Hippocampus network ⁵	NS	_	_	_		1.15	0.015	3.8E-02	仓	NS		1.86	1.1E-12	2.3E-11	
GRIA1 GRIA2 GRIA3 KCC2A	P23818 P23819 Q9Z2W9 P11798	(INNU-R1) Glutamate receptor 1 (GluR-1) Glutamate receptor 2 (GluR-2) Glutamate receptor 3 (GluR-3) Calcium/calmodulin-dependent protein binase true II o	Gria1 Gria2 Gria3 Camk2a	Hippocampus network ⁵ Hippocampus network ⁵ Hippocampus network ⁵	爺 NS NS 爺	NS NS	NS NS	(▲) —	NS NS NS	1.08 1.12 1.03 1.05	0.18 4E-04 0.64 0.20	0.26 2E-03 0.76 0.28	☆ NS ☆	NS NS		1.43 1.25 1.30 1.27	2.7E-06 1.9E-12 1.1E-03 1.6E-07	1.0E-05 2.9E-11 2.6E-03 1.1E-06	
KCC2G	Q923T9	Calcium/calmodulin-dependent protein	Camk2g		NS	NS	NS	NS	NS	1.05	0.11	0.20	NS	NS		1.25	1.7E-04	4.9E-04	
SYT1 DLG4 NCDN	P46096 Q62108 Q9Z0E0	Synaptotagmin-1 Disks large homolog 4 Neurochondrin (Norbin)	Syt1 Dlg4 Ncdn	Hippocampus network ⁵ Hippocampus network ⁵	NS 介 介	NS NS	▼ NS	NS	NS NS (▲)	1.05 1.00 1.09	0.07 0.81 0.05	0.13 0.88 0.10	仓 仓 NS	NS NS NS	NS	1.08 1.17 1.27	0.05 7.6E-05 3.0E-03	0.08 2.6E-04 6.2E-03	
Housekeeping RL8 MAP2 RS3A	P62918 O08663 P97351	60S ribosomal protein L8 Methionine aminopeptidase 2 40S ribosomal protein S3a	Rpl8 Metap2 Rps3a		小 NS NS	NS	 	 NS 	NS NS NS	1.05 1.04 1.04	0.35 0.21 0.11	0.46 0.29 0.20	NS NS NS	NS NS NS	NS (▲)	 1.06 1.12	 0.04 0.04	 0.06 0.06	

Abbreviations: HFC, hippocampus; NS, not significant. Significantly ($P^* < 0.05$) downregulated and upregulated in proteomic profiling ($\nabla \blacktriangle$) or transcriptomic profiling^{5,8} ($\Uparrow \Downarrow$), in brackets if only P < 0.05; PFC, prefrontal cortex; —, not detected. Findings are compared with label-free LC-MS^E proteomic discovery profiling and transcriptomic profiling results. Italicized cells indicates consistency across the studies. Column 'Label' indicates implication of protein (for example, predicted miRNA target by Xof4 bioinformatic prediction tools). *P*-values were determined using SRM*stats* and corrected for multiple hypothesis testing (P^*).¹⁸

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Figure 3. Schematic model summarizing the findings of the proteomic and metabonomic profiling analyses. Green boxes: altered levels of molecules determined by proteomic or metabonomic profiling. Orange boxes: altered pathways determined by *in silico* pathway analysis using IPKB or gene set enrichment analysis. Increased levels of O-GlcNAc transferase subunit p110 (OGT1) lead to abnormal O-GlcNAcylation, which affects chromatin remodelling and transcriptional regulation. Levels of the OGT1 substrate UDP-GlcNac are generated by glycolysis and gluconeogenesis (both found to be enriched). Both glutamatergic and Ca²⁺ signalling are mainly affected in the hippocampus (HPC). (R) = regulates. IPKB, Ingenuity Protein Knowledge Base; PFC, prefrontal cortex.

cytoplasmic proteins that are modified in response to cellular or environmental cues, such as growth factors, signalling molecules, glucose and other nutrient fluxes and stressors. O-GlcNAcylation has also been implicated in the aetiology of human disorders, including type II-diabetes,⁷⁸ Alzheimer's disease⁷⁹ and cancer.⁸⁰ Notably, O-GlcNAcylation has also been shown to be a negative regulator of insulin signalling.⁸¹ Transgenic mice overexpressing OGT in muscle and fat show elevated insulin levels and insulin resistance.^{81,82} Interestingly, schizophrenia patients show an increased prevalence of diabetes, impaired glucose tolerance⁸³ and metabolic syndrome.⁸⁴ Similarly, molecular analyses of postmortem brain tissues and blood cells have substantiated these findings.^{85–87} We also confirmed changes in GSK3beta and KIF5A, a microtubule-dependent molecular motor that is important for neuronal function, especially for GABA_A receptor transport.⁴⁰ In contrast, we did not find changes in the known miRNA target RHOA in any of the omics experiments. This might be due to the secondary effects of 22q11.2-encoded proteins. Further novel candidate miRNA targets implicated in both this and previous transcriptomic studies^{5,9} include BSN, VAMP2, AAK1 and DLG2.

We were also able to validate the increased levels of PP2BC and NFL in the model. PP2BC is involved in synaptic plasticity and has already been shown to be associated with schizophrenia in genome-wide association studies,^{88,89} via genome-wide DNA methylation analysis⁹⁰ and transcriptomic studies.⁹¹ As PP2BC phosphorylates proteins on serine or threonine residues, it may have a role in the extensive crosstalk between O-GlcNAcylation and phosphorylation involved in cellular signalling.⁹² Interestingly, VGLU1 was increased significantly in our proteomics and also the

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transcriptomic (PFC and HPC) screens. This protein has already been shown to be increased in bipolar disorder and major depressive disorder⁹³ and decreased in schizophrenia.⁹⁴

We also tested proteins that have already been implicated at the pathway level in major psychiatric symptoms observed in 22q11.2DS. The first of these was the mTOR pathway, which has been linked to autism spectrum disorder,^{95–97} cognition^{98,99} and glutamate signalling.^{100–102} The SRM analysis showed that mTOR was specifically affected in the PFC together with changes in the ERK1/2, PKCG and CAMK2 isoforms, which are essential components of NMDAR-related signal transduction. ERK is regulated by the activity of dopamine, serotonin and glutamate receptors.¹⁰³ ERK signalling has also been implicated in the mechanism of action of mood stabilizers¹⁰⁴ and antipsychotics¹⁰⁵ and has been shown to be affected in social behaviour¹⁰⁶ and autism spectrum disorder.^{107–109}

In contrast to the above findings, we found a robust increase in proteins relating to glutamatergic signalling in the HPC. This is consistent with the findings of the transcriptomic profiling study of the $Df(16)A^{+/-}$ mouse, which found upregulation of a multimodal gene interaction network in the HPC associated with DLG4 and glutamatergic synapses.⁵ The HPC has been implicated in the cognitive deficits in 22q11 as alterations in the gross morphology of this brain region have been observed in 22q11.2DS patients,^{110–112} and this positively correlates with cognitive impairment.¹¹² Interestingly, PKCG was also found to be increased in the HPC. Furthermore, O-GlcNAcvlation of AMPA receptor GluA2 is associated with NMDA receptor and PKC-independent long-term depression in Ca3-Ca1 synapses.¹¹³ Therefore, our findings of alterations in the levels of OGT1 support the possibility that disruption of O-GlcNAcylation pathways could be involved in cognitive dysfunction processes.

In conclusion, this is the first system-based study to identify proteomic and metabonomic abnormalities in a 22q11.2DS mouse model. The main effects were brain region specific and involved proteins associated with chromatin modulation pathways, along with alterations in lipid and energy metabolism pathways. As one of the most robust candidates, we were able to identify OGT1, a potential miR-185 target, and postulated that O-GlcNAc might have effects on many of the affected pathways. These results provide further insights into the molecular basis of the synaptic, circuitry and behavioural deficits of the 22q11.1 mouse model and potentially of the human syndrome. A shortcoming of this study is that the employed approaches focus on global changes of FC and HPC tissue, although these brain regions are comprised of different subpopulations of brain cells, which might lead to a dilution of the magnitude of the changes. Further analysis into these brain regions and of the role of OGT1 in these effects might help to elucidate its neuronal functions and increase our understanding of the development and pathogenesis of schizophrenia and other psychiatric disorders in 22q11.2DS and in the wider population.

CONFLICT OF INTEREST

SB is a director of Psynova Neurotech. The remaining authors declare no conflict of interest.

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