PUBLISHED VERSION

Kai-Leng Tan, King-Hwa Ling, Chelsee A. Hewitt, Pike-See Cheah, Ken Simpson, Lavinia Gordon, Melanie A. Pritchard, Gordon K. Smyth, Tim Thomas, Hamish S. Scott **Transcriptional profiling of the postnatal brain of the Ts1Cje mouse model of Down syndrome** Genomics Data, 2014; 2:314-317

© 2014 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/3.0/).

Originally published at: http://doi.org/10.1016/j.gdata.2014.09.009

PERMISSIONS	
http://creativecommons.org/licenses/by-nc-nd/3.0/	
Attribution-NonCommercial-NoDerive 3.0 Unported (cc. BY NC ND 3.0)	
This is a human-readable summary of (and not a substitute for) the <u>license</u> . Disclaimer	
You are free to:	
Share — copy and redistribute the material in any medium or format	
The licensor cannot revoke these freedoms as long as you follow the license terms.	
Under the following terms:	
Attribution — You must give <u>appropriate credit</u> , provide a link to the license, and <u>indicate if changes were made</u> . You may do so in any reasonable manner, but not in any way that suggests the licensor endorses you or your use.	
NonCommercial — You may not use the material for <u>commercial purposes</u> .	
NoDerivatives — If you remix, transform, or build upon the material, you may not distribute the modified material.	
No additional restrictions — You may not apply legal terms or technological measures that legally restrict othe from doing anything the license permits.	rs

Contents lists available at ScienceDirect

Genomics Data

journal homepage: http://www.journals.elsevier.com/genomics-data/

Transcriptional profiling of the postnatal brain of the Ts1Cje mouse model of Down syndrome

Kai-Leng Tan ^{a,b,1}, King-Hwa Ling ^{a,c,d,*,1}, Chelsee A. Hewitt ^{c,e,1}, Pike-See Cheah ^{a,b}, Ken Simpson ^c, Lavinia Gordon ^c, Melanie A. Pritchard ^f, Gordon K. Smyth ^{c,g}, Tim Thomas ^{c,h}, Hamish S. Scott ^{c,i,j,k,**}

^a Neurobiology and Genetics Group, GRMRC-Medical Genetics Laboratory, Faculty of Medicine and Health Sciences, Universiti Putra Malaysia, Selangor, Malaysia

^b Department of Human Anatomy, Faculty of Medicine and Health Sciences, Universiti Putra Malaysia, Selangor, Malaysia

^c Walter and Eliza Hall Institute of Medical Research, Victoria, Australia

^f Department of Biochemistry and Molecular Biology, Monash University, Melbourne, Victoria, Australia

^g Department of Mathematics and Statistics, The University of Melbourne, Parkville, Victoria, Australia

^h Department of Medical Biology, University of Melbourne, Melbourne, Victoria, Australia

ⁱ Centre for Cancer Biology, University of South Australia, Department of Molecular Pathology, SA Pathology, Adelaide, Australia

^j School of Molecular and Biomedical Bioscience, University of Adelaide, Adelaide, Australia

^k School of Medicine, Faculty of Health Sciences, University of Adelaide, Adelaide, Australia

ARTICLE INFO

Article history: Received 13 September 2014 Accepted 22 September 2014 Available online 2 October 2014

Keywords: Down syndrome Microarray Postnatal brain Gene expression

ABSTRACT

The Ts1Cje mouse model of Down syndrome (DS) has partial trisomy of mouse chromosome 16 (MMU16), which is syntenic to human chromosome 21 (HSA21). It develops various neuropathological features demonstrated by DS patients such as reduced cerebellar volume [1] and altered hippocampus-dependent learning and memory [2,3]. To understand the global gene expression effect of the partially triplicated MMU16 segment on mouse brain development, we performed the spatiotemporal transcriptome analysis of Ts1Cje and disomic control cerebral cortex, cerebellum and hippocampus harvested at four developmental time-points: postnatal day (P)1, P15, P30 and P84. Here, we provide a detailed description of the experimental and analysis procedures of the microarray dataset, which has been deposited in the Gene Expression Omnibus (GSE49050) database.

© 2014 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND licenses (http://creativecommons.org/licenses/by-nc-nd/3.0/).

Experimental design, materials and methods

Direct link to deposited data

Experimental approach

Specifications

Organism/cell line/tissue	Mus musculus/cerebral cortex, cerebellum and hippocampus
Strain(s)	Ts1Cje (on a C56BL/6 background) & disomic littermates (assumed to be C56BL/6)
Sex	Female
Sequencer or array type	Affymetrix Gene-Chip® Mouse Genome 430 2.0 arrays
Data format	Normalised data
Experimental	Ts1Cje vs disomic littermates; age (P1, P15, P30 and P84);
factors	brain regions (cerebral cortex, cerebellum and hippocampus)
Experimental	72 microarray datasets encompass Ts1Cje vs disomic
features	littermates, four postnatal time-points and three brain regions
Consent	n/a
Sample source	n/a
location	

^{*} Correspondence to: K.-H. Ling, Neurobiology and Genetics Group, GRMRC-Medical Genetics Laboratory, Faculty of Medicine and Health Sciences, Universiti Putra Malaysia, Selangor, Malaysia.

¹ Equal contributors.

brain regions from 3 Ts1Cje and 3 disomic littermate control were compared at each of the following time-points: P1, P15, P30 and P84.

compared at each of the following time-points: P1, P15, P30 and P84. The tissue samples were randomised prior to RNA extraction, quantitation of total RNA and quality/integrity, cRNA preparation and microarray hybridisation steps (Table 1). Fig. 1(A) is a simplified diagram of the experimental design and data processing flow/criteria used for the study.

Three main brain regions including the cerebral cortex, cerebellum and hippocampus were targeted in the study. Transcriptomes of these

http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE49050.

Ts1Cje mouse breeding, ethics statement and genotyping

Ts1Cje and disomic mice were generated by mating Ts1Cje males (originally obtained from The Jackson Laboratory, Bar Harbour, USA)

http://dx.doi.org/10.1016/j.gdata.2014.09.009

2213-5960/© 2014 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/3.0/).





Data in Brief



^d Department of Obstetrics and Gynaecology, Faculty of Medicine and Health Sciences, Universiti Putra Malaysia, Selangor, Malaysia

^e Pathology Department, The Peter MacCallum Cancer Centre, East Melbourne, Victoria, Australia

^{**} Correspondence to: H.S. Scott, Centre for Cancer Biology, University of South Australia, and Department of Molecular Pathology, SA Pathology, Adelaide, Australia.

Та	ble	1

Sample randomisation prior to RNA extraction, determination of total RNA sample quality/ integrity. cRNA preparation and microarray hybridization steps.

Day	Week	Tube	Mouse ID	Strain	Age	Tissue	Accession ID
1	1	1	10.1.1	Disomic	P1	Hippocampus	GSM1193014
1	1	2	A40	Ts1Cje	P84	Cerebral cortex	GSM1193015
1	1	3	A104	Ts1Cje	P30	Cerebral cortex	GSM1193016
1	1	4	10.1.3	Ts1Cje	P1	Cerebellum	GSM1193017
1	1	5	A104	Ts1Cje	P30	Hippocampus	GSM1193018
1	1	6	A63	Disomic	P15	Cerebral cortex	GSM1193019
2	1	7	A40	Ts1Cje	P84	Cerebellum	GSM1193020
2	1	8	10.1.1	Disomic	P1	Cerebellum	GSM1193021
2	1	9	A41	Disomic	P84	Hippocampus	GSM1193022
2	1	10	A60	IsiCje	P15	Cerebral cortex	GSM1193023
2	1	11	A40	TsTCJe	P84	Hippocampus	GSIVI1193024
2	1	12	10.1.3	Diagonia	P1 020	Hippocampus	GSIM1193025
2	1	13	A105	Disomic	P30	Coroballum	GSIVIT195020
2	1	14	A63	Disomic	P04 D15	Hippocampus	CSM1103027
3	1	16	A41	Disomic	P84	Cerebral cortex	GSM1193029
3	1	17	A60	Ts1Cie	P15	Hippocampus	GSM1193030
3	1	18	A104	Ts1Cie	P30	Cerebellum	GSM1193031
4	1	19	A103	Disomic	P30	Cerebellum	GSM1193032
4	1	20	10.1.3	Ts1Cie	P1	Cerebral cortex	GSM1193033
4	1	21	A60	Ts1Cje	P15	Cerebellum	GSM1193034
4	1	22	A103	Disomic	P30	Cerebral cortex	GSM1193035
4	1	23	A63	Disomic	P15	Cerebellum	GSM1193036
4	1	24	10.1.1	Disomic	P1	Cerebral cortex	GSM1193037
5	2	25	A44	Ts1Cje	P84	Hippocampus	GSM1193038
5	2	26	10.1.5	Ts1Cje	P1	Cerebellum	GSM1193039
5	2	27	A91	Disomic	P15	Hippocampus	GSM1193041
5	2	28	105	Disomic	P30	Cerebral cortex	GSM1193044
5	2	29	105	Disomic	P30	Hippocampus	GSM1193048
5	2	30	10.1.5	Ts1Cje	P1	Hippocampus	GSM1193049
6	2	31	A91	Disomic	P15	Cerebellum	GSM1193050
6	2	32	A33 3.2.4	Ts1Cje	P30	Cerebellum	GSM1193051
6	2	33	10.1.2	Disomic	P1	Cerebral cortex	GSM1193052
6	2	34	10.1.5	Ts1Cje	P1	Cerebral cortex	GSM1193053
6	2	35	10.1.2	Disomic	P1	Cerebellum	GSM1193054
6	2	36	A92	IsiCje	P15	Cerebral cortex	GSM1193055
7	2	3/	10.1.2	DISOIIIIC Te1Cio	P1 D20	Hippocampus	GSIVIT193050
7	2	20	A33 3.2.4	Disomic	P30 DQ4	Corobollum	GSIVIT195057
7	2	40	ΔQ1	Disomic	P04 D15	Cerebral cortex	CSM1103050
7	2	40	A42/43	Disomic	P84	Cerebral cortex	CSM1193060
, 7	2	42	A92	Ts1Cie	P15	Cerebellum	GSM1193061
8	2	43	105	Disomic	P30	Cerebellum	GSM1193062
8	2	44	A42/43	Disomic	P84	Hippocampus	GSM1193063
8	2	45	A92	Ts1Cje	P15	Hippocampus	GSM1193064
8	2	46	A44	Ts1Cje	P84	Cerebral cortex	GSM1193065
8	2	47	A33 3.2.4	Ts1Cje	P30	Cerebral cortex	GSM1193066
8	2	48	A44	Ts1Cje	P84	Cerebellum	GSM1193067
9	3	49	A76	Ts1Cje	P15	Cerebellum	GSM1193068
9	3	50	A34 3.2.5	Ts1Cje	P30	Hippocampus	GSM1193069
9	3	51	10.1.9	Ts1Cje	P1	Cerebellum	GSM1193070
9	3	52	106	Disomic	P30	Cerebellum	GSM1193071
9	3	53	A49	Ts1Cje	P84	Hippocampus	GSM1193072
9	3	54	A76	Ts1Cje	P15	Hippocampus	GSM1193073
10	3	55	A76	Ts1Cje	P15	Cerebral cortex	GSM1193074
10	3	56	10.1.7	Disomic	PI	Hippocampus	GSM1193075
10	3	57	106	DISOIIIIC	P30	Cerebral cortex	GSIMI 193076
10	2	50	A49	Disomia	P04	Lippocompus	GSIVIT195077
10	3	59	AJ0	Te1Cia	F 04 DQ/	Cerebral cortex	CSM1103070
11	3	61	A50	Disomic	P84	Cerebellum	CSM1193080
11	3	62	1019	Ts1Cie	P1	Cerebral cortex	GSM1193081
11	3	63	A34 3.2 5	Ts1Cie	P30	Cerebral cortex	GSM1193082
11	3	64	10.1.7	Disomic	P1	Cerebral cortex	GSM1193083
11	3	65	A34 3.2.5	Ts1Cie	P30	Cerebellum	GSM1193084
11	3	66	A75	Disomic	P15	Cerebral cortex	GSM1193085
12	3	67	A75	Disomic	P15	Hippocampus	GSM1193086
12	3	68	10.1.7	Disomic	P1	Cerebellum	GSM1193087
12	3	69	A75	Disomic	P15	Cerebellum	GSM1193088
12	3	70	10.1.9	Ts1Cje	P1	Hippocampus	GSM1193089
12	3	71	106	Disomic	P30	Hippocampus	GSM1193090
12	3	72	A50	Disomic	P84	Cerebral cortex	GSM1193091

with C57BL/6 female mice for over 10 generations. All mice were kept in a controlled environment of 12-h light/12-h dark cycle with unlimited access to a standard pellet diet and water. Breeding procedures, husbandry and all experiments were performed under the approval from the Walter and Eliza Hall Institute Animal Ethics Committee (Project numbers 2001.45, 2004.041 and 2007.007). Genomic DNA was extracted from mouse-tails and genotyping was performed using multiplex PCR with primers for neomycin (*neo*) and the glutamate receptor, ionotropic, kainite 1 (*Grik1*) as an internal control as described previously [4].

Tissue procurement

Three female Ts1Cje mice at four time-points (P1, P15, P30 and P84) with sex and age matched disomic littermates were used to avoid the effects of Y-linked genes such as *Sry* (sex-determining region of the Y chromosome), which contribute to neural sexual differentiation of the brain [5]. All mice were euthanized via cervical dislocation. Procurement of the cerebral cortex, cerebellum and hippocampus was conducted according to a method described previously [6].

RNA extraction and microarray hybridisation

The Qiagen RNeasy Micro kit (Qiagen) with a DNase I digestion step was used to extract total RNA from each tissue according to the manufacturer's instructions. All 72 tissues were randomised prior to RNA extraction to avoid biases (Table 1). The quality and quantity of each RNA sample were assessed using an Agilent 2100 Bioanalyzer (Agilent). The RNA Integrity Number (RIN) ranged from 7.0 to 10. Six micrograms of total RNA was used to prepare biotinylated cRNA according to the standard Affymetrix protocol (Expression Analysis Technical Manual, 2001, Affymetrix). Hybridisation of labelled RNA samples onto Affymetrix GeneChip Mouse Genome 430 2.0 Arrays was performed according to the Australian Genome Research Facility (AGRF) protocol. A probe cocktail (cRNA at 0.05 μ g/ μ l), which included 1× Hybridisation Buffer (100 mM MES, 1 M NaCl, 20 mM EDTA, 0.01% Tween-20), 0.1 mg/ml Herring Sperm DNA, 0.5 mg/ml BSA, and 7% DMSO was prepared to a total of 300 µl for each sample and 200 µl was hybridised onto a single GeneChip. The chips were incubated at 45 °C for 16 h in an oven with a rotating wheel at 60 rpm, washed and stained with streptavidin-phycoerythrin (SAPE) using the appropriate fluidics script on the Affymetrix Fluidics Station 450 (Affymetrix). The GeneChips were scanned using a GeneChip Scanner 3000® (Affymetrix) with GeneChip® Operating Software (GCOS). Fig. 1(B) shows a simplified diagram of the sample preparation.

Microarray data normalisation and analysis

The microarray data was analysed using R (www.r-project.org) and Bioconductor (www.bioconductor.org) [7]. The probe-level intensities for the 72 arrays were background corrected, normalised and summarised using the GC Robust Multi-array Average (GC-RMA) algorithm [8] to obtain gene (probe-set) level summaries (see Supplementary File 1 for GC-RMA script used). Differential expression between Ts1Cje and their disomic littermates at different time-points and in different brain regions was assessed using the limma package [9]. A linear model was fitted for multiple contrasts (corresponding to the Ts1Cje vs disomic comparisons) for each gene using the lmFit procedure and differential expression was

Notes to Table 1:

Note: 'Day' and 'Week' refer to the different days in different weeks on which RNA was extracted from these samples. All samples were relabelled with a set of continuous numbers under the 'Tube' column. 'Mouse ID' refers to the internal identification number used in the animal facilities where these mice were bred and maintained. 'Accession ID' refers to GEO sample ID deposited under the GSE49050 data series. Within the 'Strain' column, Ts1Cje denotes model of Down syndrome mice. Under the 'Age' column, P denotes 'Postnatal Development Day'.

assessed using empirical Bayes moderated *t*-statistics [10]. P-values corresponding to the moderated *t*-statistics were adjusted for multiple testing using the false discovery rate (FDR) procedure of Benjamini and Hochberg [11]. Fig. 1(C) shows a simplified diagram of the microarray analysis.

Stringent criteria were applied to identify differentially expressed genes (DEGs) from the datasets, which included *t*-statistic values of \geq 4 or \leq -4 and a FDR of \leq 0.05. As reported in Ling et al. [12], a total number of 317 DEGs were identified from all spatiotemporal comparisons. A top-down screening approach was then used to analyse the 317 DEGs in order to identify any disrupted molecular pathways. Initially, a functional ontology clustering analysis based on all 317 DEGs collectively using the Database for Annotation, Visualisation and

Integrated Discovery (DAVID) [13] was performed. The functional clustering analysis was performed under a stringent classification criteria (a kappa similarity threshold of 0.85, a minimum term overlap of three, two initial and final group membership with a 0.50 multiple linkage threshold and a modified Fisher-exact P-value or enrichment thresholds of 0.05) using the following databases: Biological Biochemical Image Database (BBID), BioCarta database, EC_number, Kyoto Encyclopedia of Genes and Genomes (KEGG) database, PANTHER pathway database and Reactome pathway database [13]. Subsequently, a more refined analysis was carried out involving the DEGs identified from the comparisons that were based on a specific time-point or brain region. Finally, the significant ontologies identified through all analyses were manually curated based on common genes that were found involved in the



Fig. 1. Outline of experimental design (A), sample preparation (B), microarray data analysis (C), and functional clustering analysis using DAVID tools (D). Ts1Cje denotes Down syndrome mice, cRNA denotes complementary RNA, GCRMA denotes Guanine Cytosine Robust Multi-Array Analysis, FDR denotes false discovery rate, DAVID denotes Database for Annotation, Visualization and Integrated Discovery.

ontologies leading to the identification of 7 significant functional clusters. Fig. 1(D) shows a simplified diagram of the functional clustering analysis.

Discussion

Here we provide a detailed description of the generation of a 72 microarray dataset, which is comprised of transcriptome profiling data derived from three brain regions, at four postnatal time-points from the Ts1Cje mouse model of DS and disomic littermates. The strategy used to identify DEGs between the Ts1Cje and disomic littermate data and functional clustering analysis is also described. This comprehensive and well-controlled microarray dataset encompasses postnatal developmental stages from P1 to P84 in the cerebral cortex, cerebellum and hippocampus providing a platform to understand the differences between the Ts1Cje and disomic mouse brain in these regions at a transcriptome level. The analysis of the dataset was fully described and discussed in the study by Ling et al. [12], which demonstrated that the interferon-related pathways were significantly dysregulated in the Ts1Cje brain as compared to their disomic littermates.

Acknowledgement

This work was supported by the National Health and Medical Research Council fellowships (461204 and APP1023059 to HSS); National Health and Medical Research Council Grants 219176, 257501, and 215201 (to HSS and GKS); Sciencefund Grant, MOSTI, Malaysia (02-01-04-SF1306) awarded to P-SC; Research University Grant Scheme, Universiti Putra Malaysia (04-02-12-2102RU) awarded to K-HL; and the APEX Foundation for Research into Intellectual Disability Limited to CAH. K-LT was a recipient of the Malaysian Ministry of Higher Education MyPhD scholarship. The microarrays were performed by the Australian Genome Research Facility, which was established through the Commonwealth-funded Major National Research Facilities programme.

Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx. doi.org/10.1016/j.gdata.2014.09.009.

References

- L. Olson, R. Roper, L. Baxter, E. Carlson, C. Epstein, R. Reeves, Down syndrome mouse models Ts65Dn, Ts1Cje, and Ms1Cje/Ts65Dn exhibit variable severity of cerebellar phenotypes. Dev. Dyn. 230 (2004) 581–589.
- [2] P. Belichenko, A. Kleschevnikov, A. Salehi, C. Epstein, W. Mobley, Synaptic and cognitive abnormalities in mouse models of Down syndrome: exploring genotypephenotype relationships. J. Comp. Neurol. 504 (4) (2007) 329–345.
- [3] R. Siarey, A. Villar, C. Epstein, Z. Galdzicki, Abnormal synaptic plasticity in the Ts1Cje segmental trisomy 16 mouse model of Down syndrome. Neuropharmacology 49 (2005) 122–128.
- [4] H. Sago, E.J. Carlson, D.J. Smith, J. Kilbridge, E.M. Rubin, W.C. Mobley, C.J. Epstein, T.T. Huang, Ts1Cje, a partial trisomy 16 mouse model for Down syndrome, exhibits learning and behavioral abnormalities. Proc. Natl. Acad. Sci. U. S. A. 95 (11) (1998) 6256–6261.
- [5] P. Dewing, C.W.K. Chiang, K. Sinchak, H. Sim, P.-O. Fernagut, S. Kelly, M.-F. Chesselet, P.E. Micevych, K.H. Albrecht, V.R. Harley, et al., Direct regulation of adult brain function by the male-specific factor SRY. Curr. Biol. 16 (4) (2006) 415–420.
- [6] K.H. Ling, C.A. Hewitt, T. Beissbarth, L. Hyde, K. Banerjee, P.S. Cheah, P.Z. Cannon, C.N. Hahn, P.Q. Thomas, G.K. Smyth, et al., Molecular networks involved in mouse cerebral corticogenesis and spatio-temporal regulation of Sox4 and Sox11 novel antisense transcripts revealed by transcriptome profiling. Genome Biol. 10 (10) (2009) R104.
- [7] R. Gentleman, V. Carey, D. Bates, B. Bolstad, M. Dettling, S. Dudoit, B. Ellis, L. Gautier, Y. Ge, J. Gentry, et al., Bioconductor: open software development for computational biology and bioinformatics. Genome Biol. 5 (10) (2004) R80.
- [8] Z. Wu, R.A. Irizarry, R. Gentleman, F. Martinez-Murillo, F. Spencer, A model-based background adjustment for oligonucleotide expression arrays. J. Am. Stat. Assoc. 99 (468) (2004) 909–917.
- [9] G.K. Smyth, Limma: linear models for microarray data. in: R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (Eds.), Bioinformatics and Computational Biology Solutions using R and Bioconductor, Springer, New York, 2005, pp. 397–420.
- [10] G. Smyth, Linear models and empirical Bayes methods for assessing differential expression in microarray experiments. Stat. Appl. Genet. Mol. Biol. 3 (2004).
- [11] Y. Benjamini, Y. Hochberg, Controlling the false discovery rate: a practical and powerful approach to multiple testing. J. R. Stat. Soc. Ser. B Methodol. (1995) 289–300.
 [12] K.-H. Ling, C. Hewitt, K.-L. Tan, P.-S. Cheah, S. Vidyadaran, M.-I. Lai, H.-C. Lee, K.
- [12] K.-H. Ling, C. Hewitt, K.-L. Tan, P.-S. Cheah, S. Vidyadaran, M.-I. Lai, H.-C. Lee, K. Simpson, L. Hyde, M. Pritchard, et al., Functional transcriptome analysis of the post-natal brain of the Ts1Cje mouse model for Down syndrome reveals global disruption of interferon-related molecular networks. BMC Genomics 15 (1) (2014) 624.
- [13] G. Dennis, B. Sherman, D. Hosack, J. Yang, W. Gao, H. Lane, R. Lempicki, DAVID: Database for Annotation, Visualization, and Integrated Discovery. Genome Biol. 4 (2003) P3.