Divergent dysregulation of gene expression in murine models of fragile X syndrome and tuberous sclerosis

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Abstract

Background: Fragile X syndrome and tuberous sclerosis are genetic syndromes that both have a high rate of comorbidity with autism spectrum disorder (ASD). Several lines of evidence suggest that these two monogenic disorders may converge at a molecular level through the dysfunction of activity-dependent synaptic plasticity.

Methods: To explore the characteristics of transcriptomic changes in these monogenic disorders, we profiled genome-wide gene expression levels in cerebellum and blood from murine models of fragile X syndrome and tuberous sclerosis.

Results: Differentially expressed genes and enriched pathways were distinct for the two murine models examined, with the exception of immune response-related pathways. In the cerebellum of the Fmr1 knockout (Fmr1-KO) model, the neuroactive ligand receptor interaction pathway and gene sets associated with synaptic plasticity such as long-term potentiation, gap junction, and axon guidance were the most significantly perturbed pathways. The phosphatidylinositol signaling pathway was significantly dysregulated in both cerebellum and blood of Fmr1-KO mice. In Tsc2 heterozygous (+/−) mice, immune system-related pathways, genes encoding ribosomal proteins, and glycolipid metabolism pathways were significantly changed in both tissues.

Conclusions: Our data suggest that distinct molecular pathways may be involved in ASD with known but different genetic causes and that blood gene expression profiles of Fmr1-KO and Tsc2+/− mice mirror some, but not all, of the perturbed molecular pathways in the brain.

Keywords: Fragile X syndrome, Tuberous sclerosis, Autism, Cerebellum, Blood, Gene expression, Murine model

Background

Autism spectrum disorder (ASD) manifests significant heterogeneity in part because of the interaction of underlying genetic [1-3], neurobiological, and environmental factors [4,5] during early brain development. This heterogeneity presents one of the main obstacles to the development of effective treatments for ASD. The complex genetics of ASD suggest that it is a large set of related disorders with diverse mechanisms; however, many of the etiologies implicated in ASD may converge on a few common pathways. Further research on single gene disorders associated with ASD such as tuberous sclerosis complex (TSC) and fragile X syndrome (FXS) may lead to an understanding of common dysfunction at the cellular or circuit level for a majority of ASD. In a recent survey of over 14,000 individuals under age 35 with ASD in a Boston area hospital, Kohane and colleagues reported that the prevalence of genetic disorders of FXS and TSC in individuals with ASD were 0.5% and 0.8% [6]. Conversely, 30% and 50-61% of patients with FXS and TSC present ASD core symptoms, respectively [7,8]. If such shared pathophysiology exists, then treatments developed for a target in one disorder might be applicable to others. Mouse models for ASD serve an increasingly important role in...
providing a pre-clinical test of promising pharmacological therapeutics [9,10]. Inactivating mutation in Tsc2 (Tsc2+/- mice) showed defects in axon guidance [11] and cognitive deficits such as impaired water maze performance [12], and mice with Fmr1-knockout (KO) presented impairments in long-term depression, hyperactivity, anxiety-like, and unusual social behaviors [13]. Therefore, determining the degree to which there are shared molecular mechanisms in these models will inform clinical trials, particularly those that address populations with genetically heterogeneous causes of ASD.

Although several cellular mechanisms may be implicated (reviewed in Fatemi et al. [14]), accumulating data support a role for the PI3K-mTOR signaling cascade in several genetic causes of ASD. Evidence for the PI3K-mTOR pathway first emerged from TSC [15,16] and mutations in the PTEN gene associated with ASD and macrocephaly [17-19]. Later, investigation of copy number variants (CNV) in autistic individuals identified that PI3K-mTOR pathway-related genes were located in CNV hotspots [20]. These findings have led to the hypothesis that overactivation of the mTOR pathway could lead to abnormal synaptic function owing to an excess of protein synthesis at the synapse [21]. Genetic evidence that directly implicates a translation initiating factor, EIF4E, which is a downstream target of mTOR, in ASD has provided further support for this hypothesis [22]. Interestingly, exposure to teratogens such as valproate in utero can lead to ASD in children [23], and valproate can also modulate this signaling pathway [24], suggesting that environmental factors associated with ASD can also play a role in PI3K-mTOR pathway regulation [25]. More recently, studies have found that PI3K-mTOR signaling is upregulated in mouse models of FXS, one of the most common genetic causes of ASD [26-28].

Together, the aforementioned findings suggest that an upregulated PI3K-mTOR signaling cascade might be a common mechanism in ASD and therefore would potentially be a promising drug target. Indeed, clinical trials using inhibitors of mTOR are already in progress in patients with TSC. We hypothesized that if the PI3K-mTOR signaling pathway is dysregulated in various causes of ASD, then these disorders should present with a similar gene expression profile signature. We chose to analyze TSC and FXS, two Mendelian disorders highly associated with ASD. Better understanding of similarities and differences of the cellular and molecular defects leading to abnormal neurological function in these two disorders is essential to the development of new therapies for ASD. Here, we used mouse models available for both genetic disorders to investigate the similarities and differences between gene expression profiles in the brain and blood cells.

Methods

Murine models of fragile X syndrome and tuberous sclerosis

To identify molecular signatures of each mouse model of ASD, we performed gene expression profiling on cerebellum and peripheral blood collected from two mouse models and compared to wild-type (WT) controls. We used cerebellum where the most consistent abnormalities were reported in the patients with ASD [14]. Post-mortem studies have shown a reduced number of Purkinje cells (PC), and several neuroimaging studies reported enlarged cerebella in ASD [29,30]. The cerebellum is also implicated in social interaction [31], and the loss of Tsc1 from cerebellar PC was associated with autistic-like behaviors [32]. Additionally, we profiled whole blood from the same individual mouse to compare with the gene expression changes in cerebellum.

All male C57BL/6 congeneric Fmr1-KO mice and Tsc2+/- mice with mixed 129/SvJae-C57BL/6 J background have been previously described [33,34]. We profiled Tsc2+/- mice since homozygous Tsc2 KO was embryonic lethal. The mice were killed at 8–10 weeks of age following the institutional animal care and use committee (IACUC) euthanasia criteria (the Boston Children’s Hospital IACUC animal protocol no. 12-07-2227R). For the Fmr1-KO model, 5 KO and 5 WT mice were profiled, and for the Tsc2+/- model 3 transgenic and 3 WT mice were profiled. Paired blood and cerebellum samples were prepared for gene expression profiling.

Genome-wide gene expression profiling using microarrays

A total of 250 ng RNA was processed using established Affymetrix protocols for the generation of biotin-labeled cRNA, and the hybridization, staining, and scanning of arrays were performed. Briefly, total RNA was converted to double-stranded cDNA using a T7 primer and biotin-labeled cRNA was then generated from the cDNA by in vitro transcription. The cRNA was quantified (using A260) and fragmented. Fragmented cRNA was hybridized to the Affymetrix Mouse Gene ST 1.0 array and scanned on an Affymetrix GeneChip scanner 3000 at 2.5 μm resolution [35]. Microarray data are available at the Gene Expression Omnibus database (GSE40630).

Validation of gene expression changes using quantitative RT-PCR

Total RNA was extracted using TRIzol according to the manufacturer’s instruction. The RNA amount was measured using the Nanodrop (Thermo Scientific); 100 ng of total RNA was reverse transcribed using a cDNA reverse transcription kit with random primers (Applied Biosystems). SyBr Green PCR Master Mix (Applied Biosystems) was used to amplify and detect signals from
cDNA with 1 mM gene-specific primers. Amplified signals were collected by the 7300HT Fast Real-Time System (Applied Biosystems) and normalized to Gapdh. Primer sequences used for this study were Gapdh forward 5’-tgtgctctgtggatctga-3’, Fmr1 forward 5’-gtctgaaccttcttgcttgacgtgtgacaggg-3’, reverse 5’-aagtgctcttgcttgacgatag-3’, Tsc2 forward 5’-cagctgtgcac cagctgtcctt-3’, reverse 5’-tcacgctgctgctgtccttgcttgacgatag-3’, Eps811 forward 5’-cagctacaacacgagaagcg-3’, reverse 5’-ctgaaacctgggtgtgaggt-3’, and reverse 5’-aatgctgttcccacacaac-3’. Primer sequences used for this study were Gapdh forward 5’-ctgaaacctgggtgtgaggt-3’, Eps811 forward 5’-cagctacaacacgagaagcg-3’, reverse 5’-ctgaaacctgggtgtgaggt-3’, and reverse 5’-aatgctgttcccacacaac-3’.

Microarray analysis
All microarrays were normalized together at the probe-level using a quantile method, and the Affymetrix Probe Logarithmic Intensity ERror (PLIER) model was used to calculate the absolute gene expression levels as previously described [35]. We fitted a linear model of the tissue (i.e., blood vs. cerebellum) and treatment (i.e., transgenic vs. WT) as predicting variables to each probe set. Two sample t-tests were used separately as different background strains were used. Differentially expressed genes in each murine model were compared to the differentially expressed genes between the wild types of two models. The false discovery rate (FDR) was calculated using Storey’s method [36]. We did not use non-parametric tests such as the Wilcoxon rank sum test because of the granularity of the test statistics with a small number of samples per group.

We identified enriched pathways using the Gene Set Enrichment Analysis (GSEA) [37]. The Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways of size 15–500 were used for pathway analysis. Due to the relatively small number of samples in each group, we randomly sampled gene sets of equal size for each KEGG pathway to calculate the background distribution of enrichment scores. This procedure was done with 2,000 random drawings, thus the minimum permutation p-value was 0.0005. We used permutation p-value 0.05 as the significance threshold for GSEA, and corresponding FDRs were described. To identify the core set of genes that accounts for the gene set’s enrichment signal, we used leading edge analysis as described in Subramanian et al. [37] where a leading edge subset was defined as the subset of genes that gave the maximum enrichment score. These genes were the topmost correlated genes with phenotype in a gene set.

To compare differentially expressed genes with the list of known ASD candidate genes as curated in the SFARI-genes 2.0 database (http://gene.sfari.org/) [38], we mapped mouse genes to human homologs using the Mouse Genome Informatics (MGI) Web database (http://www.informatics.jax.org) and performed hypergeometric tests to check the significance of overlap.

Results
Distinct gene expression changes define Fmr1 and Tsc2 transgenic models
A total of 107 and 115 probe sets were significantly changed in Fmr1-KO and Tsc2+/- mice compared to corresponding WT littermates, respectively (uncorrected p-value < 0.01). Not surprisingly, Fmr1 was the most significantly downregulated gene in Fmr1-KO (p-value 8.85 x 10^-6, corresponding FDR 0.29). We used nominal p-values estimated from a linear model less than 0.01 to rank significant probe sets because multiple testing correction procedures did not make any gene significant. The expression levels of 16 out of 107 significant probe sets (15.0%) in Fmr1-KO mice and 58 out of 115 significant probe sets (50.4%) in Tsc2+/- mice did not show a significant difference between blood and brain. We used an agglomerative hierarchical clustering with the significant probe sets for each model to explore the similarity of gene expression profiles in two genotypes and across tissue types. Samples were clearly separated by tissue type and then by genotype (Figure 1A and 1B). Interestingly, 71% of significant probe sets in Fmr1-KO mice were highly expressed in cerebellum compared to blood (Figure 1A). For the Tsc2+/- model, the average expression levels of 63 out of 115 probe sets (57.5%) were higher in cerebellum (Figure 1B). Differentially expressed genes with statistical scores are listed in Additional file 1: Table S1 (Fmr1-KO vs. WT) and Additional file 1: Table S2 (Tsc2+/- vs. WT).

We compared the gene expression profiles of WT mice since Tsc2+/- mice had 129/SvJae-C57BL/6 J backgrounds compared to C57BL/6 congenic backgrounds of Fmr1-KO mice. A total of 1,486 probes sets were differentially expressed between two WT strains. Seven probe sets, representing five genes, Cog7, Cc2d1a, Smurf1, Sec31a, and AU040320, overlapped with the differentially expressed genes in Fmr1-KO vs. WT comparison. For the differentially expressed genes in Tsc2+/- vs. WT comparison, 13 probe sets (representing 9 genes: Fhad1, Pgam5, Cts8, Piwil4, Tmem101, Heatr7a, Mucl1, 2210404f11Rik, and Fam181b) were significantly different between two WT mice strains.

Epidermal growth factor (EGF) receptor pathway substrate 8-like 1 (Eps811) (Affymetrix probe set ID: 10549655) was the only gene that was significant in both murine models. Eps811 was upregulated in Tsc2+/- mice, but downregulated in Fmr1-KO mice. Post hoc two-group comparison of the transgenic model to WT for each tissue showed that Eps811 was upregulated in blood (Welch’s t-test p-value 0.07) and brain (Welch’s t-test p-value 0.06) of Tsc2+/- mice, while it was downregulated in blood (Welch’s t-test p-value 0.015) of Fmr1-KO mice. Downregulation of Eps811 in brain of Fmr1-KO was not significant (Welch’s t-test p-value 0.38). The homolog
of Eps8l1 in human, EPS8L1 is a member of EPS8-related proteins that play an important role in actin remodeling in response to EGF [39], and EPS8 is one of the reported targets of FMRP [40]. Interestingly, Stamatakou and colleagues reported that Eps8 loss of function impaired the structural and functional plasticity of synapses induced by long-term potentiation in primary rat hippocampal neurons [41]. Phenotypically, Eps8-KO mice have impaired learning and memory, and excessive synaptic growth and abnormal spine morphology were observed in the hippocampus [42]. In human samples, the average expression of EPS8 in fusiform gyri was significantly lower among the patients with ASD compared to controls [42]. Although Eps8 itself was not significantly changed in our study, EPS8 family genes are interesting candidates for further investigation. Thus, we performed qRT-PCR of Eps8 in brain samples of two murine models.

Of the differentially expressed genes, several were also found in the expert curated database of autism candidate genes. Of these, Chd7, Fmr1, and Tmlhe were differentially expressed in Fmr1-KO mice, and Oxtr and Taf1c were significantly changed in Tsc2+/- mice. The differentially expressed genes were not significantly enriched for known ASD candidate genes in human (hypergeometric test p-value 0.48 for Fmr1-KO and 0.70 for Tsc2+/-).

Validation of differentially expressed genes using quantitative RT-PCR

Quantitative RT-PCR for individual genes was used to further confirm the results of the expression profiling in the brain samples used for the initial analysis. As expected, Fmr1 gene expression was significantly decreased in Fmr1-KO mice as compared to controls (average 5.24-fold downregulated, Welch's t-test p-value 0.025), while Fmr1 was unchanged in Tsc2+/- mice (Welch's t-test p-value 0.70). Tsc2 gene expression showed a trend of decreased expression in both Tsc2+/- and Fmr1-KO mice as compared to WT mice that did not reach significance (average fold change 1.12 and 1.31 downregulation, Welch's t-test p-value 0.55 and 0.27, respectively). We previously found
a significant reduction of Tsc2 protein in cortical neurons of Tsc2+/- mice [11]; however, Tsc2 mRNA expression was not significantly downregulated. Further, our results confirm that Eps8l1 exhibited a three-fold increase in Tsc2+/- animals as compared to WT mice (average 3.15-fold upregulation, Welch’s t-test p-value 0.06). However, Eps8l1 expression shows a decreased trend—an average 1.26 fold downregulation—in Fmr1-KO compared to WT similar to that observed in the expression profiling but not significantly changed (Welch’s t-test p-value 0.74).

We observed significant downregulation of subtype 3a of the N-methyl-D-aspartate receptor gene (Grin3a) in the blood of Fmr1-KO mice (uncorrected p-value 0.0098). Grin3a was not significant in brain with microarray data; however, a quantitative RT-PCR analysis of the same samples showed a significant downregulation of this gene in Fmr1-KO brain (5.69-fold downregulated, Welch’s t-test p-value 0.046).

**Enriched biological pathways in blood and brain of the two mouse models**

We explored whether similar sets of biological pathways were perturbed in both models using GSEA [37] as there was only one overlapping gene—Eps8l1—between the two lists of differentially expressed genes in Fmr1-KO and Tsc2+/- mice. The genes that contributed to making a pathway significant were identified using leading edge analysis. First, all genes were ranked by a per-gene signal-to-noise ratio that was defined as mean difference divided by the sum of standard deviation of each group. Then a running sum was calculated for each gene set. Beginning with the top-ranked gene, the running sum increased when a gene in a gene set was found and decreased otherwise. The enrichment score (ES) was defined to be the largest value of the running sum, and the genes that maximized ES were defined as the leading edge subset.

Two pathways related to cytokine and complement-mediated signaling, cytokine-cytokine receptor interaction (HSA04060, FDR < 0.0005) and complement and coagulation cascades (HSA04610, FDR < 0.0005), were the most significantly enriched pathways in Fmr1-KO brain (Table 1). Neuroactive ligand receptor interaction (HSA04080, FDR < 0.0005), long-term potentiation (HSA04720, FDR 0.0023), gap junction (HSA04540, FDR 0.0429), and axon guidance (HSA04360, FDR 0.0480) were

| Table 1 Enriched pathways in blood and brain of Fmr1 knockout mice |
|-------------------------|-----------------|---|---|---|---|
| **KEGG categories** | **Name** | **SIZE** | **NES** | **NOM p-val** | **FDR q-val** |
| **Brain** | | | | | |
| Nervous system | HSA04080 Neuroactive ligand receptor interaction | 223 | 2.34 | < 0.0005 | 0.0000 |
| | HSA04720 Long-term potentiation | 62 | -2.15 | < 0.0005 | 0.0023 |
| | HSA04540 Gap junction | 77 | -1.75 | < 0.0005 | 0.0429 |
| | HSA04360 Axon guidance | 125 | -1.81 | < 0.0005 | 0.0480 |
| Immune system | HSA04060 Cytokine-cytokine receptor interaction | 212 | 2.72 | < 0.0005 | < 0.0005 |
| | HSA04610 Complement and coagulation cascades | 47 | 2.32 | < 0.0005 | < 0.0005 |
| | HSA04650 Natural killer cell mediated cytotoxicity | 95 | 1.70 | 0.0015 | 0.0579 |
| | HSA04612 Antigen processing and presentation | 42 | 1.67 | 0.0049 | 0.0397 |
| | HSA04640 Hematopoietic cell lineage | 66 | 2.09 | < 0.0005 | 0.0004 |
| Signaling pathways | HSA01430 Cell communication | 118 | 2.14 | < 0.0005 | 0.0004 |
| | HSA04630 Jak-STAT signaling pathway | 137 | 2.07 | < 0.0005 | 0.0005 |
| | HSA04070 Phosphatidylinositol signaling system | 65 | 1.77 | 0.0029 | 0.0420 |
| | HSA04910 Insulin signaling pathway | 126 | 1.80 | < 0.0005 | 0.0455 |
| Metabolism | HSA00150 Androgen and estrogen metabolism | 33 | 1.84 | 0.0043 | 0.0118 |
| | HSA005590 Arachidonic acid metabolism | 39 | 1.81 | 0.0017 | 0.0147 |
| | HSA00361 y-hexachlorocyclohexane degradation | 17 | 1.69 | 0.0118 | 0.0370 |
| | HSA00020 Citrate cycle | 25 | 1.93 | 0.0023 | 0.0375 |
| | HSA00592 a-linolenic acid metabolism | 15 | 1.70 | 0.0127 | 0.0392 |
| | HSA00251 Glutamate metabolism | 30 | 1.87 | 0.0038 | 0.0430 |
| Folding, sorting, and degradation | HSA04120 Ubiquitin mediated proteolysis | 36 | 1.78 | 0.0039 | 0.0453 |
| | HSA04130 Snare interactions in vesicular transport | 29 | 1.84 | < 0.0005 | 0.0450 |
| **Blood** | | | | | |
| Signaling pathways | HSA04070 Phosphatidylinositol signaling system | 65 | 1.78 | < 0.0005 | 0.0500 |
also enriched in \textit{Fmr1}-KO brain. Among the signaling pathways, the PI3K signaling pathway (HSA04070) was changed in both brain (FDR 0.042) and blood (FDR 0.05) of \textit{Fmr1}-KO mice. Eight genes were in the leading edges of the two tissues (see Methods). These were Dgkb, Dgkg, Dgkh, Inpp4b, Inpp5a, Itpr3, Plcb4, and Prkca. Glutamate metabolism (HSA00251) was also enriched in \textit{Fmr1}-KO brain (FDR 0.043).

Eight pathways were significantly enriched in \textit{Tsc2+/-} brain, while 11 pathways were enriched in \textit{Tsc2+/-} blood (Table 2). Pathways associated with the immune system were significantly enriched in both blood and brain. Ribosome (HSA03010, FDR < 0.0005), cytokine-cytokine receptor interaction (HSA04060, FDR < 0.0005), and oxidative phosphorylation (HSA00190, FDR < 0.0005) were the most significant pathways in the brain, and the ribosome pathway was also significantly deregulated in \textit{Tsc2+/-} blood (FDR 0.0468). In the \textit{Tsc2+/-} brain, cytokine-cytokine receptor interaction (FDR < 0.0005) and hematopoietic cell lineage (HSA04640, FDR 0.0389) were significant, while the Toll-like receptor signaling pathway (HSA04620, FDR 0.0016) and B-cell receptor signaling pathway (HSA04662, FDR 0.0072) were significant in \textit{Tsc2+/-} blood. None of the differentially expressed genes between two WT strains was a member of significant pathways that we identified above.

### Discussion

We hypothesized that \textit{Fmr1}-KO and \textit{Tsc2+/-} mice would have similar gene expression profiles. In contrast, our findings indicate that different gene expression signatures define these two monogenic mouse models of

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**Table 2 Enriched pathways in blood and brain of \textit{Tsc2+/-} mice**

<table>
<thead>
<tr>
<th>KEGG categories</th>
<th>Name</th>
<th>SIZE</th>
<th>NES</th>
<th>NOM p-val</th>
<th>FDR q-val</th>
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<td>Brain</td>
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**Common signature of \textit{Fmr1} and \textit{Tsc2} transgenic models**

Cytokine-cytokine receptor interaction pathway, hematopoietic cell lineage, and cell communication were enriched in brain gene expression profiles of both \textit{Fmr1}-KO and \textit{Tsc2+/-} mice (Figure 2). We performed leading edge analysis to find core genes that made a pathway significant, although each gene was not necessarily differentially expressed. For the cytokine-cytokine receptor pathway (N = 212), 105 and 106 genes were the core genes in \textit{Tsc2+/-} and \textit{Fmr1}-KO brain data sets. \textit{Il7r} was the only gene that showed marginal significances in both data sets (post-hoc Welch’s \textit{t}-test \textit{p}-values 0.011 and 0.013 in \textit{Tsc2+/-} and \textit{Fmr1}-KO brain profiles, respectively). The same gene was also the only common significant gene for the hematopoietic cell lineage pathway. Thirty-two genes were in common between the leading edges of cell communication for \textit{Fmr1}-KO and \textit{Tsc2+/-} brain; however, no genes were significantly differentially expressed. Interestingly, \textit{Tsc2} gene expression was downregulated in \textit{Fmr1}-KO blood (post-hoc Welch’s \textit{t}-test \textit{p}-value 0.0045).
ASD. Global expression profiles of the two models examined were distinct, such that only one gene—Eps8l1—was in common. This is a particularly surprising result since translational dysregulation as well as aberrant synaptic protein synthesis associated with both disorders has been proposed as one possible pathway leading to autistic phenotypes, including cognitive impairment [21].

Our data indicate that FXS and TSC may have very distinct brain and blood cellular phenotypes despite the fact that both syndromes result in similar behavioral and cognitive symptoms.

Nonetheless, we did find that the cytokine and complement signaling pathways are differentially regulated in both mouse models although the specific genes

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**Figure 2 Enriched pathways in Fmr1-KO and Tsc2+/− models of ASD, found using the Gene Set Enrichment Analysis (GSEA).** Red (upregulated) and blue (downregulated) squares in the matrix represent enriched pathways for each data set (false discovery rate ≤ 0.05). Two immune system pathways (cytokine-cytokine receptor signaling pathway and hematopoietic cell lineage) and one signaling pathway (cell communication) were significant in the brain gene expression profiles of both mice models.
affected within these pathways were different. The immune system has been implicated in ASD in multiple ways, but the exact mechanism of the interaction between the immune system and genetic disorders that result in an increased risk of autism has not been well studied. Modulation of the immune system may not be completely unexpected since the TSC-mTORC1 pathway regulates inflammatory responses after bacterial stimulation in monocytes, macrophages, and primary dendritic cells [43], and this pathway contributes to cytokine upregulation in response to endotoxins [44]. However, this is of particular interest in light of a recent study that showed that immune activation during gestation can markedly worsen the neurological phenotype of Tsc2−/− mouse pups [45]. Similarly, preliminary observations indicate that plasma protein levels of a number of cytokines differ between individuals with and without FXS. Furthermore, differences in cytokine and other immune-signaling genes were observed between the FXS group with autism and the FXS group without autism [46]. On the other hand, Yuskaitis and colleagues investigated the peripheral immune system of Fmr1-KO mice, but did not find any differences in either the T-cell population at basal and stimulated status or the proinflammatory cytokines TNFα and IFNγ at basal and stimulated status [47]. How the loss of FMRP leads to changes in the immune system, however, remains unclear. Accumulating evidence over the last few years indicates that the TSC and FMRP pathways interact. However, precisely how these pathways interact remains an open question. On the one hand, FMRP can be phosphorylated by S6K, an enzyme downstream of TSC [48]. On the other hand, mTOR, the kinase inhibited by the TSC2 protein, has increased activity in Fmr1-KO neurons [26], and FMRP-deficient cells display increased activity of PI3K, an enzyme upstream of TSC proteins [27]. Recently Auerbach et al. reported that the synaptic dysfunction in the CA1 region of the hippocampus of Tsc2+/− mice was opposite to that of Fmr1-KO mice [49]. In fact, manipulating the mGluR receptors with positive allosteric modulators was sufficient to rescue this defect in Tsc2+/−, while inhibiting the mGluR receptors was necessary in the case of Fmr1-KO mice. Finally, a genetic cross of the Tsc2+/− and Fmr1-KO mice was similar to WT in CA1 synaptic physiology and contextual learning. These results argue that loss of Tsc2 and Fmr1 have some opposite cellular phenotypes, which can be rescued in a double knockout. However, the mechanisms by which Tsc2 and Fmr1 result in opposite synaptic phenotypes is not yet clear. In post-hoc analysis, we found that Tsc2 gene expression was downregulated in Fmr1-KO blood (Welch’s t-test p-value 0.0045). Thus, in the Tsc2+/− Fmr1-KO mice, the Tsc2 expression level may be closer to WT and may contribute to the rescue of the synaptic physiology. This finding points to another level of interaction between the FMRP and Tsc2 functions in the cell. Future studies are required to understand whether Fmr1 loss leads to changes in Tsc2 mRNA via transcriptional or post-transcriptional regulation. Similarities between the proposed roles of TSC1/2 and FMRP proteins in regulation of protein synthesis have led to the popular hypothesis that hyperactive mTOR signaling is pathogenic in both FXS and TSC. Our results indicate the gene expression dysregulation differs markedly in the two conditions. This is consistent with the previously reported changes in neuronal morphology in each of these models. Neurons deficient in Tsc1 or Tsc2 display lower dendritic spine density in contrast to Fmr1-KO neurons [50], which have increased spine density [51]. At the biochemical level there are also some important differences. In Tsc1/2-null cells, mTORC1-dependent negative feedback mechanisms exist to dampen the activation of upstream components of the network such that Akt activation is decreased [52]. However, in Fmr1-KO neurons, Akt activity is enhanced [27]. While mTORC1 may be activated because of loss of either Tsc1/2 or FMRP, the neuronal phenotype and gene expression profiles may be altered by changes in the activation of other signaling pathways. Such differences have implications for targeted treatment options of the two distinct genetic conditions.

The mouse models we have explored have been developed as models of ASD with known divergent genetic etiologies. The overall divergence in gene expression dysregulation in these two models does not rule out shared downstream effects, and indeed we observed an overlapping dysregulation of the cytokine signaling pathway. Nonetheless, it does suggest that a multiplicity of therapeutics will have to be developed for the varied mechanisms contributing to the increasingly fine-grained distinctions between the etiologies of ASD.

We could identify similar sets of biological pathways enriched in both tissues. In the Fmr1-KO mice, the PI3K signaling pathway was dysregulated in both blood and brain, while the ribosome pathway was dysregulated in both tissues of Tsc2+/−. With the two mouse models, we also could identify biological pathways that were positively correlated with genetic background in both tissues. Previous studies demonstrated that peripheral blood expression signatures could be used to classify the clinical conditions of brain disorders [35,53]. Likewise, our results suggest that peripheral blood signatures could be used to classify the varied mechanisms contributing to the increasingly fine-grained distinctions between the etiologies of ASD.

The current study is limited by the different background strains of Fmr1-KO and Tsc2+/− mice and by the small sample size. The differentially expressed genes between two WT background strains overlapped with the
significant genes in each murine model – 5 for Fmr1-KO and 9 for Tsc2+/- and Tsc2+/- mice in brain and blood. Due to the small sample size, we did not have enough statistical power for microarray experiments to detect and compare relatively low-expressed genes. Further study using the same background strain and increasing the sample size will be essential to confirm our findings, and using a more sensitive quantification method such as RNA-seq will improve sensitivity for low abundance transcripts. For the Fmr1-KO model, we did not include female mice heterozygous for Fmr1 in this experiment. Heterozygous Fmr1 female mice should exhibit genetic mosaicism due to random X-inactivation of one X chromosome during development. For this reason, most previous studies characterized male Fmr1 KO models. Qin and colleagues performed an interesting comparison of male Fmr1-KO and homozygous and heterozygous KO in female mice [54]. They reported that only the homozygous mice had a deficit on the passive avoidance test, whereas both homozygous and heterozygous female mice exhibited hyperactivity and increased susceptibility to seizures. A follow-up experiment with both sexes and different dosages of Fmr1 in female mice with a larger sample size would be ideal since a gender effect on global gene expression profiles should be considered when both sexes are included in the experiment.

Conclusions

Contrary to our initial hypothesis that Fmr1-KO and Tsc2+/- mice would share a transcriptional signature, we found that the two mouse models presented distinct sets of differentially expressed genes. In retrospect, this is not surprising as multiple lines of evidence suggest that FXS and TSC are actually driven by opposite molecular phenotypes [49]. Despite these gene-level differences, however, we observed that cytokine signaling, cell communication, and hematopoietic cell lineage genes were differentially expressed in both mouse strains. Second, our results show that blood expression signatures mirror many aspects of the brain transcriptome. Specifically, several pathways were dysregulated in both the brain and blood of the two mouse models studied here. This confirmation is important for the future use of blood tissue to study neurodevelopmental disorders.

Availability of supporting data

The data set supporting the results of this article is available in the Gene Expression Omnibus repository with the accession identifier GSE40630 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE40630).

Additional file

Additional file 1: Table S1. The 107 transcripts were differentially expressed in Fmr1-KO mice compared to wildtype littermates. These probesets were used for the hierarchical clustering analysis as showed in Figure 1A. The transcript that was assigned for uncharacterized gene was designated as ‘NA’ with the Affymetrix probeset identifiers (AffyID). The p-value was estimated from a linear model using genotype and tissue as predicting variables for each probeset. The q-value denotes the false discovery rate that were calculated from the distribution of p-values using Storey and Tibshirani’s (see Methods). Negative values in the fold changes represent down-regulated transcripts in transgenic mice.

Table S2. The 115 transcripts were differentially expressed in Tsc2 +/- mice compared to wildtype littersates. These probesets were used for the hierarchical clustering analysis as showed in Figure 1B. The transcript that was assigned for uncharacterized gene was designated as ‘NA’ with the Affymetrix probeset identifiers (AffyID). The p-value was estimated from a linear model using genotype and tissue as predicting variables for each probeset. The q-value denotes the false discovery rate that were calculated from the distribution of p-values using Storey and Tibshirani’s (see Methods). Negative values in the fold changes represent down-regulated transcripts in transgenic mice.

Abbreviations

Akt: v-Akt murine thymoma viral oncogene; ASD: Autism spectrum disorder; CA1: Cornu ammonis 1; Chd7: Chromodomain-helicase-DNA-binding protein 7; Dgko: Diacylglycerol kinase beta; Dgkg: Diacylglycerol kinase gamma; Dgkgl: Diacylglycerol kinase eta; EGF: Epidermal growth factor; EIF4E: Eukaryotic translation initiation factor 4E; EPSP: Epidermal growth factor receptor pathway substrate 8; EPSP 1: Epidermal growth factor receptor pathway substrate 8 like 1; FDR: False discovery rate; Fmr1: Fragile X mental retardation 1; FMRP: Fragile X mental retardation protein; FXS: Fragile X syndrome; Grin1a: N-methyl-D-aspartate receptor subtype 3a; GSEA: Gene set enrichment analysis; Inpp4b: Inositol polyphosphate-4-phosphatase, type II; Inpp5a: Type I inositol-1,4,5-trisphosphate 5-phosphatase; Inpp5c: Inositol 1,4,5-trisphosphate receptor, type 3; KEGG: Kyoto encyclopedia of genes and genomes; KD: Knockout; mGlur: Metabotropic glutamate receptor; mTOR: Mammalian target of rapamycin; mTORC1: Mammalian target of rapamycin complex 1; Ntr: Nitrilase receptor; Pldb: Phosphatidylinositol 3-kinases; Pldb4: 1-Phosphatidylinositol-4,5-bisphospho-phosphodiesterase beta-4; PLIER: Probe logarithmic Intensity error model; Prka: Protein kinase C, alpha; PTEN: Phosphatase and tensin homolog; RNA-seq: mRNA sequencing; Raf1c: TATA box-binding protein-associated factor RNA polymerase I subunit C; Tmlhe: Trimethyllysine dioxygenase; TSC: Tuberous sclerosis complex; Taf1c: Tuberous sclerosis protein 1; Tsc2: Tuberous sclerosis protein 2; WT: Wild type.

Competing interests

MFB declares a financial interest in Seaside Therapeutics. SWK, MS, CDC, MGC, JL, DK, LMK, and ISK report no biomedical financial interests. The authors declare no competing financial interests.

Authors’ contributions

SWK and CDC collected and analyzed the data, and JDL and MHW performed qRT-PCR validation. SWK and MGC performed statistical analyses. MS provided Tsc2 heterozygous mouse, and DK and MFB provided Fmr1 knockout mouse. ISK and LMK conceived the study, and SWK, MS, and ISK wrote the manuscript. All authors read and approved the final manuscript.

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