

Conservation of Regional Gene Expression in Mouse and Human Brain

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Many neurodegenerative diseases have a hallmark regional and cellular pathology. Gene expression analysis of healthy tissues may provide clues to the differences that distinguish resistant and sensitive tissues and cell types. Comparative analysis of gene expression in healthy mouse and human brain provides a framework to explore the ability of mice to model diseases of the human brain. It may also aid in understanding brain evolution and the basis for higher order cognitive abilities. Here we compare gene expression profiles of human motor cortex, caudate nucleus, and cerebellum to one another and identify genes that are more highly expressed in one region relative to another. We separately perform identical analysis on corresponding brain regions from mice. Within each species, we find that the different brain regions have distinctly different expression profiles. Contrasting between the two species shows that regionally enriched genes in one species are generally regionally enriched genes in the other species. Thus, even when considering thousands of genes, the expression ratios in two regions from one species are significantly correlated with expression ratios in the other species. Finally, genes whose expression is higher in one area of the brain relative to the other areas, in other words genes with patterned expression, tend to have greater conservation of nucleotide sequence than more widely expressed genes. Together these observations suggest that region-specific genes have been conserved in the mammalian brain at both the sequence and gene expression levels. Given the general similarity between patterns of gene expression in healthy human and mouse brains, we believe it is reasonable to expect a high degree of concordance between microarray phenotypes of human neurodegenerative diseases and their mouse models. Finally, these data on very divergent species provide context for studies in more closely related species that address questions such as the origins of cognitive differences.

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Introduction

Here we compare and contrast gene expression in three different regions of the human brain, motor cortex, caudate, and cerebellum, to identify genes that are differentially expressed between the regions. In other words, we seek to identify genes that show patterned expression. Knowledge of such regionally enriched genes may provide insight into the development and biochemistry of different brain structures. This information may also hold potential biomedical implications. Many neurodegenerative diseases, such as Huntington's disease, have a hallmark regional and cellular pathology affecting one or another of these regions while sparing the others. It is reasonable to assume that unique susceptibilities in disease may relate to distinctive brain gene expression patterns in health.

We also perform a parallel analysis on the functionally and anatomically corresponding regions of the mouse brain, anterior cortex, striatum, and cerebellum. This allows us to begin to compare and contrast patterns of gene expression in these tissues across these two species. Our motivation for this cross-species analysis also has a biomedical consideration. In recent years, mice have become the most important model organism for human neurological and neurodegenerative

diseases. The brains of humans and mice are clearly different with respect to size, complexity, and cognitive abilities. The belief that mice can accurately model human neurodegenerative or neurological diseases rests on assumptions about deeper biological similarities between mouse and human brains that have not been systematically examined. While it is impossible to directly compare mouse arrays to human arrays, one may compare patterns of gene expression in several corresponding brain regions. Comparing gene expression patterns is one way to obtain objective and global information on how similar the brains of humans and mice are. A practical use for this comparative cross-species gene

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Abbreviations: Brodmann area 4, BA4; dN, number of nonsynonymous substitutions/number of nonsynonymous sites; dS, number of synonymous substitutions/number of synonymous sites; GO, gene ontology; RMA, Robust Multiple-array Average

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Author Summary

Animal models of human neurodegenerative and psychiatric disorders, particularly mouse models, have assumed a central role in biomedical research aimed at discovering the causes of disease and generating novel, mechanism-based treatments. But to what degree can a mouse brain serve as a model for a human brain? Here we begin to address this question by looking at patterns of gene expression across three corresponding regions of mouse and human brains. We find that within each species, the different regions (motor cortex, striatum, and cerebellum) have very distinct gene expression profiles. It is likely that these differences reflect distinctions in regional neurochemistry and function. We then show that genes that are enriched in one of the three areas relative to the other two in mice have the same pattern of expression in humans. Looking at the relationship between conservation of expression and amino acid sequence, we find that genes showing patterned expression generally have been more conserved than more uniformly expressed genes. This suggests that in the brain, constraints on the evolution of DNA sequence and gene expression can also be particularly high for genes with regional or tissue-specific expression.

expression information is as a baseline for the comparison of microarray phenotypes of human diseases and their mouse models. For example, if expression changes in a human disease and its mouse model have a global correlation of $r = 0.5$, is this the best possible correlation that can be expected or might we reasonably expect more? Obviously, to answer this question it is useful to know what sort of correlation array data from mice and human brains have initially.

Comparative cross-species information of brain gene expression would also seem to have natural implications for studies that address the origin of cognitive differences between humans and other species. It is generally believed that with respect to other species, even our closest relatives the chimpanzees, humans have unique abilities pertaining to language and higher order cognitive functions. Sequencing projects have revealed that the human and chimpanzee genomes are ~96% identical, that their typical protein amino acid sequences are ~99% identical, and that both species have essentially the same number of genes [1]. Since an increase in genomic complexity seems inadequate to explain the apparent mental differences between humans and chimpanzees, the idea that these differences may be due to changes in gene expression has attracted new attention [2].

Recent microarray studies have sought to identify general trends in brain gene expression that distinguish humans from other primates [3–10]. The conclusions reached by these studies have been somewhat discordant. One study found more expression differences between human and chimpanzee liver than prefrontal cortex, but by using orangutan liver and cortex as out groups, concluded there had been an accelerated rate of change in brain gene expression during human evolution [3]. A reanalysis of this data supported this interpretation of rapid and recent human evolution [6]. Other investigations have found that elevated levels of gene expression further distinguish the human brain from that of other primates [4–6]. Several studies have cast doubt upon these findings, attributing them to improper array normalization or to hybridization artifacts rooted in measuring

nonhuman primate expression with arrays designed for human sequences [7–10].

Since we are comparing human and mouse expression, we cannot make definitive statements regarding differential expression between humans and chimpanzees. However, examination of global similarity of expression patterns in species as divergent as mice and humans can provide useful context for studies that aim to correlate expression changes with cognitive differences between more closely related species. Presumably, if gene expression patterns distinguish human brains from the brains of other primates, then changes due to recent human evolution may be even more apparent when comparing humans to mice.

Results

Absolute and Relative Gene Expression Levels in Equivalent Human and Mouse Brain Structures

The first part of our analysis focuses on expression in different regions of nondiseased brain. We determined absolute and relative gene expression in three anatomically distinct regions of human brain: motor cortex (Brodmann area 4, BA4), caudate nucleus, and cerebellum. The data consisted of samples of all three tissues from 12 donors. These samples constituted a portion of the control group in a study comparing gene expression in Huntington's disease and non-Huntington's brain [11]. For the present reanalysis, the 36 arrays were normalized using Robust Multiple-array Average (RMA) [12]. To assess differential expression between brain regions, three sets of paired t-tests were performed; caudate-to-cerebellum, BA4-to-cerebellum, and BA4-to-caudate using the Bioconductor package LIMMA (<http://www.bioconductor.org/packages/1.9/bioc/html/limma.html>) [13,14]. To confirm the primary human data, we used a second set of caudate and cerebellum samples from nine different donors [11]. Because of the original study's design, there were no motor cortex samples from these nine donors. The absolute and differential expression analysis of the human samples is provided in Datasets S1 and S2. A key that associates samples with GEO accession numbers, age, gender, post mortem delay, and other covariables can be found in Table S1.

Comparing the \log_2 (fold change) (i.e., log ratio) for the replicate caudate-to-cerebellum comparisons indicated that these independent data were highly correlated, with Pearson's correlation coefficient $r = 0.93$ (Figure S1). In the primary caudate-to-cerebellum comparison, 9,088 probesets met $p < 0.001$ with respect to differential expression. In the smaller secondary dataset, 8,074 probesets met $p < 0.001$. Of these, 82% (6,589/8,074) met $p < 0.001$ in both comparisons, and only four probesets showed discordant directions of change. These results demonstrate that the caudate and cerebellum have quite distinct gene expression profiles. They also show that the relative differences between the regions were robust and reproducible in these post mortem human samples. This is consistent with results from a detailed analysis of the relationship between prehybridization variables and posthybridization assessments of data quality, which found little negative contribution from post mortem interval to data quality in these samples [15].

We next performed identical comparative analyses of anterior cortex, striatum, and cerebellum samples from six five-week old wild type C57BL/6 mice. It is generally accepted

Table 1. Different Regions of the Brain Show Many Statistically Significant Differentially Expressed Genes

Organism	Tissue 1	Tissue 2	$p < 0.001$ (%)	Percentage 1 > 2	Mean ABS log ratio
Human	Striatum	Cerebellum	9,088 (41)	49	0.65
	Motor cortex	Striatum	5,992 (27)	45	0.52
	Motor cortex	Cerebellum	9,880 (44)	53	0.60
Mouse	Striatum	Cerebellum	7,844 (35)	53	0.87
	Anterior cortex	Striatum	5,926 (26)	46	0.66
	Anterior cortex	Cerebellum	7,920 (35)	48	0.84

Probeset counts and percentages for differential expression ($p < 0.001$) in human and mouse regional comparisons are shown, along with the fraction of the differentially expressed probes increasing in one tissue relative to the other. The mean of the absolute log ratios in the differentially expressed probesets is also shown.
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that these mouse brain regions are anatomically and functionally homologous to human motor cortex, caudate, and cerebellum respectively. We used young mice since very often identifying the earliest changes in a mouse neurological disease model is of primary experimental interest. The complete mouse RMA and regional comparison data are provided in Datasets S3 and S4. Counts of probesets meeting $p < 0.001$ for differential expression in one region relative to the others for both the mouse and primary human data are shown in Table 1. As was found with the human analysis, the three mouse brain regions examined had very distinct gene expression profiles with many statistically significant changes (Table 1).

To provide additional verification of the data, we queried the human and mouse caudate/striatum-to-cortex and caudate/striatum-to-cerebellum comparisons against a published list of 54 striatum-enriched mouse genes (Table S2) [16]. Table 2 shows both the human and mouse array data to be consistent with known mouse striatal genes ($p \approx 10$).

Table 3 shows the 30 named genes with the highest regional scores (see Materials and Methods) in each species. At their most extreme, the pattern of expression for these genes is “on” in one region and “off” in the other two regions. Table 3 represents only a small subset of regionally enriched genes, and the complete data pertaining to differential expression between brain regions can be found in Datasets S2 and S4. Several genes appear in both the mouse and human lists. Even considering these short lists of top regional genes, the intersections between human and mouse gene lists are highly statistically significant ($p < 10^{-7}$ for each intersection).

Gene Expression Variation between Tissues and Individuals

While the primary interest of this study was in regional differences, many factors such as age, gender, tissue heterogeneity, post mortem interval, medication, and cause of death may influence gene expression in the brain and contribute to differences between individuals. To examine the effect of individual variability of the gene expression on the profiles, the between-tissue and within-tissue variances for each probeset were computed from the human RMA signals. This was repeated for the mouse probesets. As post mortem delay was not a concern with the mouse samples, and all of the mice

Table 2. Mouse and Human Microarray Data Are Consistent with Previously Identified Striatal Genes

Criterion	Human		Mouse	
	Striatum: Cerebellum	Striatum: Cortex	Striatum: Cerebellum	Striatum: Cortex
Matching and $p < 0.001$	31	30	39	36
Indeterminant	8	9	3	7
Mismatch and $p < 0.001$	1	1	1	0

The probesets corresponding to genes in a list of striatally enriched mouse genes were identified and scored as enriched in striatum/caudate relative to cerebellum or cortex. A gene was scored as matching the striatum list if any one of its representative probesets met $p < 0.001$, otherwise it was scored as indeterminate. If a gene met $p < 0.001$ but was less abundant in striatum/caudate than the other tissues, it was scored as a mismatch.
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were the same age, mouse individual variability might reasonably be expected to be smaller than human variability. We found that the between-tissue variability was greater for 89% of the human probesets and 85% of the mouse probesets. This suggested that human individual variability in gene expression and factors such as tissue heterogeneity and post mortem interval were not obscuring or significantly contributing to regional differences. It also suggested that compared to expression dictated by regional identity, age and gender appear to have effects of small magnitude or of large magnitude on a small fraction of genes, even in humans. Some evidence for this can also be inferred from the two independent human caudate-to-cerebellum comparisons. In these comparisons, age and gender were not balanced between the groups, yet relative expression levels were highly correlated and the slope of the regression line was 0.967 (Figure S1).

Cross-Species Comparison of Regional Gene Expression

To explore the gene expression of each tissue in more depth, we used gene ontology (GO) [17]. GO provides means of objectively identifying functional themes in large groups of genes, in this case the genes that were differentially expressed in the pair-wise regional comparisons within each species. A significantly high number of overrepresented GO categories were found for both human and mouse in all three types of regional comparisons. This was true whether considering increased or decreased probesets separately or together.

While GO is not intended for rigorous assessment of evolutionary relationships, GO nomenclature is standardized. This allowed us to compare and contrast the regionally enriched functions in the homologous mouse and human brain regions. Of the hundreds of GO categories differentially represented in one region relative to another, many were common between the corresponding human and mouse comparisons (Tables 4 and S3). Permutation testing showed these intersections to be significantly greater than would be expected by chance ($p < 0.0001$) (Table S4).

Both the functional GO analysis and the intersections among top regional marker genes hinted that relative gene expression levels across brain regions have been conserved between mice and humans. To examine this in depth, it was

Table 3. Selected Regionally Enriched Genes in Human and Mouse Brain Tissues

Tissue	Human			Mouse			
	Probeset ID	Gene Title	Gene Symbol	Probeset ID	Gene Title	Gene Symbol	
Human cerebellum/ mouse cerebellum	207182_at	gamma-aminobutyric acid (GABA) A receptor, alpha 6	GABRA6	1417121_at	gamma-aminobutyric acid (GABA-A) receptor, alpha 6	Gabra6	
	206914_at	class-I MHC-restricted T-cell associated molecule	CRTAM	1419085_at	Purkinje cell protein 2 (L7)	Pcp2	
	206282_at	neurogenic differentiation 1	NEUROD1	1423287_at	cerebellin 1 precursor protein	Cbln1	
	206373_at	Zic family member 1 (odd-paired homolog, <i>Drosophila</i>)	ZIC1	1422911_at	cerebellin 3 precursor protein	Cbln3	
	208153_s_at	FAT tumor suppressor homolog 2 (<i>Drosophila</i>)	FAT2	1449903_at	cytotoxic and regulatory T cell molecule	Crtam	
	205747_at	cerebellin 1 precursor	CBLN1	1426413_at	neurogenic differentiation 1	Neurod1	
	206163_at	mab-21-like 1 (<i>Caenorhabditis elegans</i>)	MAB21L1	1424958_at	carbonic anhydrase 8	Car8	
	211343_s_at	collagen, type XIII, alpha 1	COL13A1	1451129_at	calbindin 2	Calb2	
	203706_s_at	frizzled homolog 7 (<i>Drosophila</i>)	FZD7	1450079_at	Nik related kinase	Nrk	
	204431_at	transducin-like enhancer of split 2 homolog (<i>Drosophila</i>)	TLE2	1418868_at	engrailed 2	En2	
	219423_x_at	tumor necrosis factor receptor superfamily, member 25	TNFRSF25	1424679_at	mab-21-like 1 (<i>C. elegans</i>)	Mab21l1	
	221911_at	ets variant gene 1	ETV1	1418933_at	solute carrier family 1 member 6	Slc1a6	
	210770_s_at	calcium channel voltage-dependent P/Q type, alpha 1A	CACNA1A	1417391_a_at	interleukin 16	Il16	
	200920_s_at	B-cell translocation gene 1, antiproliferative	BTG1	1450428_at	LIM homeobox protein 1	Lhx1	
	203895_at	phospholipase C, beta 4	PLCB4	1418983_at	InaD-like (<i>Drosophila</i>)	Inadl	
	219778_at	zinc finger protein, multitype 2	ZFPM2	1423477_at	zinc finger protein of the cerebellum 1	Zic1	
	212825_at	PAX interacting protein 1	PAXIP1	1427624_s_at	interleukin 22	Il22	
	219825_at	cytochrome P450, family 26, subfamily B, polypeptide 1	CYP26B1	1424007_at	growth differentiation factor 10	Gdf10	
	214734_at	exophilin 5	EXPH5	1420709_s_at	D-amino acid oxidase 1	Dao1	
	207110_at	potassium inwardly rectifying channel, subfamily J, member 12	KCNJ12	1434719_at	alpha-2-macroglobulin	A2m	
	205336_at	Parvalbumin	PVALB	1428574_a_at	chimerin (chimaerin) 2	Chn2	
	214936_at	leucine-rich repeats and calponin homology containing 1	LRCH1	1431829_a_at	ral guanine nucleotide dissociation stimulator-like 3	Rgl3	
	206557_at	zinc finger protein 702	FLJ12985	1421435_at	glutamate receptor, ionotropic, delta 2	Grid2	
	214705_at	InaD-like (<i>Drosophila</i>)	INADL	1419271_at	paired box gene 6	Pax6	
	205646_s_at	paired box gene 6 (aniridia, keratitis)	PAX6	1456140_at	Opr	Zic5	
	212224_at	aldehyde dehydrogenase 1 family, member A1	ALDH1A1	1452650_at	tripartite motif-containing 62	Trim62	
	219572_at	Ca ²⁺ -dependent activator protein for secretion 2	CADPS2	1424859_at	homer homolog 3 (<i>Drosophila</i>)	Homer3	
	207060_at	engrailed homolog 2	EN2	1422929_s_at	atonal homolog 7 (<i>Drosophila</i>)	Atoh7	
	205923_at	Reelin	RELN	1417639_at	solute carrier family 22, member 4	Slc22a4	
	205380_at	PDZ domain containing 1	PDZK1	1417995_at	protein tyrosine phosphatase, non-receptor type 22	Ptpn22	
	Human motor cortex/ mouse anterior cortex	205827_at	Cholecystokinin	CCK	1427017_at	special AT-rich sequence binding protein 2	Satb2
		204338_s_at	regulator of G-protein signaling 4	RGS4	1428664_at	vasoactive intestinal polypeptide	Vip
		201340_s_at	ectodermal-neural cortex (with BTB-like domain)	ENC1	1418047_at	neurogenic differentiation 6	Neurod6
220551_at		solute carrier family 17, member 6	SLC17A6	1422580_at	myosin, light polypeptide 4	Myl4	
205825_at		proprotein convertase subtilisin/kexin type 1	PCSK1	1448366_at	syntaxin 1A (brain)	Stx1a	
209200_at		myocyte enhancer factor 2C	MEF2C	1418370_at	troponin C, cardiac/slow skeletal	Tnnc1	
213435_at		SATB family member 2	SATB2	1416711_at	T-box brain gene 1	Tbr1	
210181_s_at		calcium binding protein 1 (calbrain)	CABP1	1451620_at	C1q-like 3	C1ql3	
211685_s_at		neurocalcin delta	NCALD	1426851_a_at	nephroblastoma overexpressed gene	Nov	
206481_s_at		LIM domain binding 2	LDB2	1416846_a_at	PDZ domain containing RING finger 3	Pdzrn3	
33767_at		neurofilament, heavy polypeptide 200kDa	NEFH	1419473_a_at	cholecystokinin	Cck	
213326_at		vesicle-associated membrane protein 1 (synaptobrevin 1)	VAMP1	1416658_at	frizzled-related protein	Frzb	
213745_at		attractin-like 1	ATRNL1	1451507_at	myocyte enhancer factor 2C	Mef2c	
213479_at		neuronal pentraxin II	NPTX2	1419230_at	keratin complex 1, acidic, gene 12	Krt112	
219032_x_at		opsin 3 (encephalopsin, panopsin)	OPN3	1418317_at	LIM homeobox protein 2	Lhx2	
212922_s_at		SET and MYND domain containing 2	SMYD2	1419517_at	cornichon homolog 3 (<i>Drosophila</i>)	Cnih3	

Table 3. Continued.

Tissue	Human			Mouse		
	Probeset ID	Gene Title	Gene Symbol	Probeset ID	Gene Title	Gene Symbol
	211616_s_at	5-hydroxytryptamine (serotonin) receptor 2A	HTR2A	1454770_at	cholecystokinin B receptor	Cckbr
	221805_at	neurofilament, light polypeptide 68kDa	NEFL	1455893_at	R-spondin 2 homolog (<i>Xenopus laevis</i>)	Rspo2
	205635_at	kalirin, RhoGEF kinase	KALRN	1435551_at	formin-family protein FHOS2	FHOS2
	206051_at	ELAV (embryonic lethal, abnormal vision)-like 4 (Hu antigen D)	ELAVL4	1450061_at	ectodermal-neural cortex 1	Enc1
	205352_at	serpin peptidase inhibitor, clade I (neuroserpin), member 1	SERPINI1	1433909_at	Synaptotagmin XVII (Sy17), mRNA	Bk
	210121_at	UDP-Gal:betaGlcNAc beta 1,3-galactosyltransferase, polypep 2	B3GALT2	1418790_at	zinc finger protein 312	Zfp312
	212976_at	leucine rich repeat containing 8 family, member B	LRRC8B	1427351_s_at	immunoglobulin heavy chain 6 (heavy chain of IgM)	Igh6
	213920_at	cut-like 2 (<i>Drosophila</i>)	CUTL2	1428572_at	brain abundant, membrane attached signal protein 1	Basp1
	209205_s_at	LIM domain only 4	LMO4	1449298_a_at	phosphodiesterase 1A, calmodulin-dependent	Pde1a
	205113_at	neurofilament 3 (150kDa medium)	NEF3	1453317_a_at	KH domain containing RNA binding signal transduction asc.3	Khdrbs3
	209485_s_at	oxysterol binding protein-like 1A	OSBPL1A	1417262_at	prostaglandin-endoperoxide synthase 2	Ptgs2
	204697_s_at	chromogranin A (parathyroid secretory protein 1)	CHGA	1428118_at	leucine rich repeat neuronal 6A	Lrrn6a
	205591_at	olfactomedin 1	OLFM1	1436066_at	kalirin, RhoGEF kinase	Kalrn
	219736_at	tripartite motif-containing 36	TRIM36	1420388_at	protease, serine, 12 neurotrypsin (motopsin)	Prss12
Human caudate/mouse striatum	220313_at	G-protein coupled receptor 88	GPR88	1423171_at	G-protein coupled receptor 88	Gpr88
	207307_at	5-hydroxytryptamine (serotonin) receptor 2C	HTR2C	1416783_at	tachykinin 1	Tac1
	215241_at	transmembrane protein 16C	TMEM16C	1420437_at	indoleamine-pyrrole 2,3 dioxygenase	Indo
	206552_s_at	tachykinin, precursor 1	TAC1	1418950_at	dopamine receptor 2	Drd2
	220359_s_at	cyclic AMP-regulated phosphoprotein, 21 kDa	ARPP21	1417804_at	similar to Ca and DAG-regulated g-nucleotide exchange fact.	Rasgrp2
	205229_s_at	coagulation factor C homolog, cochlin (<i>Limulus polyphemus</i>)	COCH	1418691_at	regulator of G-protein signaling 9	Rgs9
	213791_at	proenkephalin	PENK	1450339_a_at	B-cell leukemia/lymphoma 11B	Bcl11b
	214655_at	G protein-coupled receptor 6	GPR6	1417129_a_at	myeloid ecotropic viral integration site-related gene 1	Mrg1
	215867_x_at	carbonic anhydrase XII	CA12	1418782_at	retinoid X receptor gamma	Rxrg
	205013_s_at	adenosine A2a receptor	ADORA2A	1419390_at	phosphodiesterase 10A	Pde10a
	206355_at	G-nucleotide BP, alpha activating activity polypeptide, olfactory type	GNAL	1454906_at	retinoic acid receptor, beta	Rarb
	206803_at	prodynorphin	PDYN	1455961_at	Membrane metallo endopeptidase	Mme
	214071_at	metallophosphoesterase 1	MPPE1	1450723_at	ISL1 transcription factor, LIM/homeodomain (islet 1)	Isl1
	213338_at	Ras-induced senescence 1	RIS1	1427344_s_at	RASD family, member 2	Rasd2
	221008_s_at	alanine-glyoxylate aminotransferase 2-like 1	AGXT2L1	1449420_at	phosphodiesterase 1B, Ca2+-calmodulin dependent	Pde1b
	205454_at	hippocalcin	HPCA	1427519_at	adenosine A2a receptor	Adora2a
	205478_at	protein phosphatase 1, regulatory (inhibitor) subunit 1A	PPP1R1A	1416776_at	crystallin, mu	Crym
	210372_s_at	tumor protein D52-like 1	TPD52L1	1451280_at	cyclic AMP-regulated phosphoprotein, 21	Arpp21
	204712_at	WNT inhibitory factor 1	WIF1	1448327_at	actinin alpha 2	Actn2
	206518_s_at	regulator of G-protein signalling 9	RGS9	1415904_at	lipoprotein lipase	Lpl
	203548_s_at	lipoprotein lipase	LPL	1418881_at	EF hand calcium binding protein 2	Efcfbp2
	205330_at	meningioma (disrupted in balanced translocation) 1	MN1	1427038_at	preproenkephalin 1	Penk1
	215506_s_at	DIRAS family, GTP-binding RAS-like 3	DIRAS3	1417680_at	K voltage-gated channel, shaker-related subfamily, member 5	Kcna5
	214652_at	dopamine receptor D1	DRD1	1425870_a_at	Kv channel-interacting protein 2	Kcnip2
	211021_s_at	regulator of G-protein signalling 14	RGS14	1423544_at	protein tyrosine phosphatase, nonreceptor type 5	Ptpn5
	214608_s_at	eyes absent homolog 1 (<i>Drosophila</i>)	EYA1	1451331_at	protein phosphatase 1, regulatory (inhibitor) subunit 1B	Ppp1r1b
	201842_s_at	EGF-containing fibulin-like extracellular matrix protein 1	EFEMP1	1427300_at	LIM homeobox protein 8	Lhx8

Table 3. Continued.

Tissue	Human			Mouse		
	Probeset ID	Gene Title	Gene Symbol	Probeset ID	Gene Title	Gene Symbol
	216086_at	synaptic vesicle glycoprotein 2C	SV2C	1448269_a_at	kelch-like 13 (<i>Drosophila</i>)	<i>Klh13</i>
	220331_at	cytochrome P450, family 46, subfamily A, polypeptide 1	<i>CYP46A1</i>	1449979_a_at	sparc/osteonectin, cwcv and kazal-like proteoglycan 3	<i>Spock3</i>
	207174_at	glypican 5	<i>GPCS</i>	1425503_at	glucosaminyl (N-acetyl) transferase 2, l-branching enzyme	<i>Gcnt2</i>

Criteria for inclusion were $p < 0.001$ and $\log \text{ratio} > 1$ in both relevant tissue comparisons. Genes were sorted by the sum of the log ratios and the top 30 named genes are shown. Genes in red appear in corresponding mouse and human regional lists: (a) 30 human and 30 mouse genes whose expression is relatively restricted to cerebellum when compared to the caudate/striatum and motor cortex/anterior cortex; (b) Human motor cortex and mouse anterior cortex enriched genes; (c) Human caudate and mouse striatum enriched genes. doi:10.1371/journal.pgen.0030059.t003

necessary to contrast expression ratios on a gene-by-gene basis across the human and mouse arrays. This was complicated by the fact that genes are often represented by more than one probeset on each array. To lessen this complication for our initial analyses, genes were identified where only one probeset existed on each array. We also arbitrarily required that human and mouse gene symbols were identical, since it was a clean and simple way to identify genes that have met widely accepted criteria for being orthologous pairs. Using these criteria, 2,998 one-to-one orthologous pairs were found on the mouse and human arrays.

Taking this set of genes, we first asked whether genes with high variance of expression across the brain regions of one species would cluster the entire set of samples sensibly. This was motivated by our earlier observation that the largest component of a gene's expression variance was due to tissue specificity. Thus, high variance implied patterned expression across the three brain regions. Figure 1 shows that both the mouse and human genes with the largest variance in the one-to-one gene set cluster all of the samples perfectly, first by tissue, then by species. In other words, for these three brain regions, the equivalent human and mouse regions are more alike than different regions within a species or individual. We also note that while we selected the genes based on variability

of expression across regions within a species and not conservation between species, there were 43 genes in common on the two lists of 125 most variable one-to-one genes ($p \approx 0$). The heat maps of normalized expression indicated that relatively few genes in corresponding brain regions were on opposite sides of their mean signal within a species. All of these observations suggest a high degree of similarity in the genes with patterned expression in mouse and human brain.

Using all genes in the one-to-one set, we next examined relatedness of regional gene expression within and between species by computing normalized Euclidian distances between all possible nonself pairs of tissues (Table 5). The similarity of corresponding tissues between the species was apparent by their consistently having the minimal between-species distance. The pattern of distances between regions within the human brain was essentially identical to the pattern of distances within the mouse brain, suggesting that no single region of the human brain had diverged from the other two regions any more than regions in the mouse brain had diverged from each other.

To expand our analysis beyond the one-to-one subset of genes, ENSEMBL (<http://www.ensembl.org>) information was used to identify a more complete set of mouse-human orthologs. Where more than one probeset represented a gene, we retained only information pertaining to the probeset with the highest mean RMA signal. This collapsed the arrays to a nonredundant set of 8,499 genes common to both array types (Dataset S5). We then correlated log ratios in the appropriate pairs of tissue comparisons over all the genes. The correlation coefficient of the mouse and human caudate-to-cerebellum log ratio was $r = 0.47$. For the cortex-to-cerebellum comparisons and for the cortex-to-caudate comparisons, $r = 0.45$.

Relationships between Tissue-Specific Expression, Conservation of Sequence, and Conservation of Expression

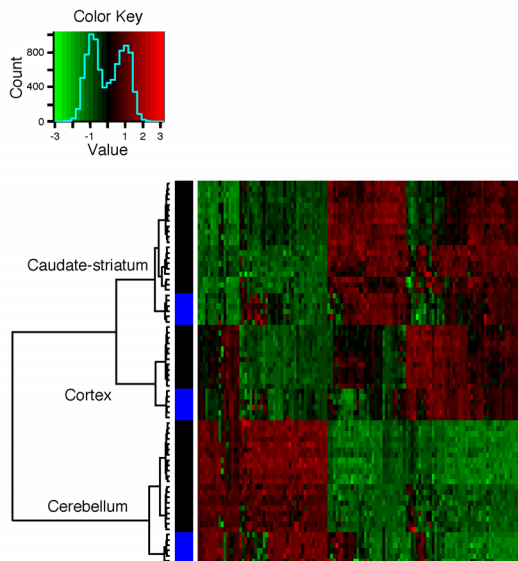
We explored the hypothesis that genes whose sequences had been under stabilizing selective pressure would also be constrained in their pattern of expression. Information about nonsynonymous and synonymous amino acid substitution ratios (dN/dS) and percent nucleotide identity for mouse and human orthologs were retrieved from the ENSEMBL database. The set of 8,499 orthologous genes was ranked by each

Table 4. Mouse and Human Brain Regions Share a Higher Number of Overrepresented Functional Groups than Would Be Expected by Chance

Criterion	Tissue 1	Tissue 2	Human	Mouse	Intersection
Tissue 1 > 2	Striatum	Cerebellum	170	241	42
	Motor cortex	Striatum	150	200	47
	Motor cortex	Cerebellum	132	258	47
Tissue 2 > 1	Striatum	Cerebellum	110	230	23
	Motor cortex	Striatum	116	209	23
	Motor cortex	Cerebellum	98	230	25

The numbers of GO processes found to be overrepresented in one tissue relative to another in mouse and human brain are shown for specified pairs of tissues. The first three rows show the numbers of GO processes overrepresented in tissue 1 relative to tissue 2 for human and mouse plus their intersections. The last three rows show the numbers of GO processes overrepresented in tissue 2 relative to tissue 1 plus their intersections. doi:10.1371/journal.pgen.0030059.t004

A



B

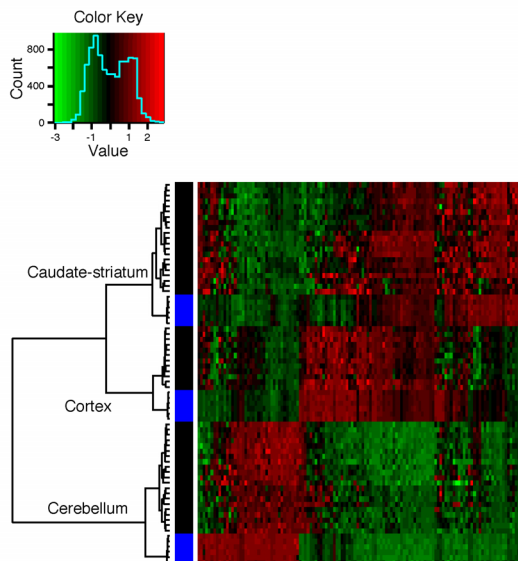


Figure 1. The Most Variable Genes in One Species Accurately Cluster the Brain Regions of the Other Species, and Orthologous Structures Cluster Together

(A) Hierarchical clustering of regions and species using the 125 human genes in the one-to-one set with the highest variance across the 36 primary human arrays is shown.

(B) Hierarchical clustering of regions and species using the 125 mouse genes in the one-to-one set with the highest variance across the 18 mouse arrays is shown. While variable genes were identified using only the primary human data, clustering included the secondary set of human data. The mouse, primary, and secondary human data were normalized separately. Expression levels are colored by standard deviation from the within-group mean. Columns represent genes. Rows represent samples with black bars on the left signifying human samples, and blue signifying mouse samples.

doi:10.1371/journal.pgen.0030059.g001

of these metrics, and a correlation coefficient between appropriate human and mouse log ratios was computed for each quartile of genes. The quartile-based correlation coefficients are plotted for each class of tissue comparison in Figure 2. This shows that there is a positive relationship

Table 5. Orthologous Brain Regions between Species Are More Similar to Each Other than to Different Regions within a Species

Tissue	Human			Mouse	
	Motor Cortex	Caudate	Cerebellum	Anterior Cortex	Striatum
Human caudate	1.20	—	—	—	—
Human cerebellum	1.67	1.73	—	—	—
Mouse anterior cortex	1.00	1.27	1.46	—	—
Mouse striatum	1.26	1.00	1.54	1.44	—
Mouse cerebellum	1.43	1.50	1.29	1.75	1.81

Using the 2,998 orthologous genes that were represented just once on the human and mouse arrays, Euclidian distances for gene expression in tissues between all possible pairs of nonself tissues were calculated. These were then normalized to the smallest distance. In each row or column of distances, the minimum distance is found at the pairing of corresponding mouse and human brain regions.

doi:10.1371/journal.pgen.0030059.t005

between conservation of sequence and conservation of expression.

It seems natural to assign greater confidence in a pairing between two genes that are 95% conserved at the sequence level than between two genes that are 75% conserved. Furthermore, as homology thresholds decrease, the number of potential ortholog pairings increases. Because of these factors, we assume that our rate of incorrectly pairing orthologs may increase as percent nucleotide identity decreases. Pairing errors also likely reduce correlation, thus any bias in the error rate of pairing may introduce a false positive relationship between homology and gene expression. To avoid this potential bias, we examined the relationship between variability of expression across tissues and sequence conservation. Results presented above suggest that in mouse and human brain, the genes with the greatest variability of expression in the three examined brain regions were similar (Figure 1). We also showed that expression variance was most strongly dependent upon tissue specificity rather than variability between individuals. Variance within a species can be determined in the absence of homology information, so we examined the within-species variance of bins of genes with integral percent identities. Figure 3 shows that there is a clear tendency in both species for genes with higher expression variance across brain regions to have higher identity with their orthologs. Since expression variance is a surrogate for tissue specificity, this indicates that region-specific genes in the brain tend to have greater homology with their orthologs than more widely expressed brain genes. This is consistent with the idea that functional constraints have applied selective pressure on brain gene expression since the mouse and human lineages diverged some 80 million years ago.

Discussion

Our data indicate that expression patterns across comparable regions of human and mouse brains have generally been conserved since the two lineages diverged. This is consistent with classical comparative neuroanatomy, which has long indicated general conservation of gross mammalian

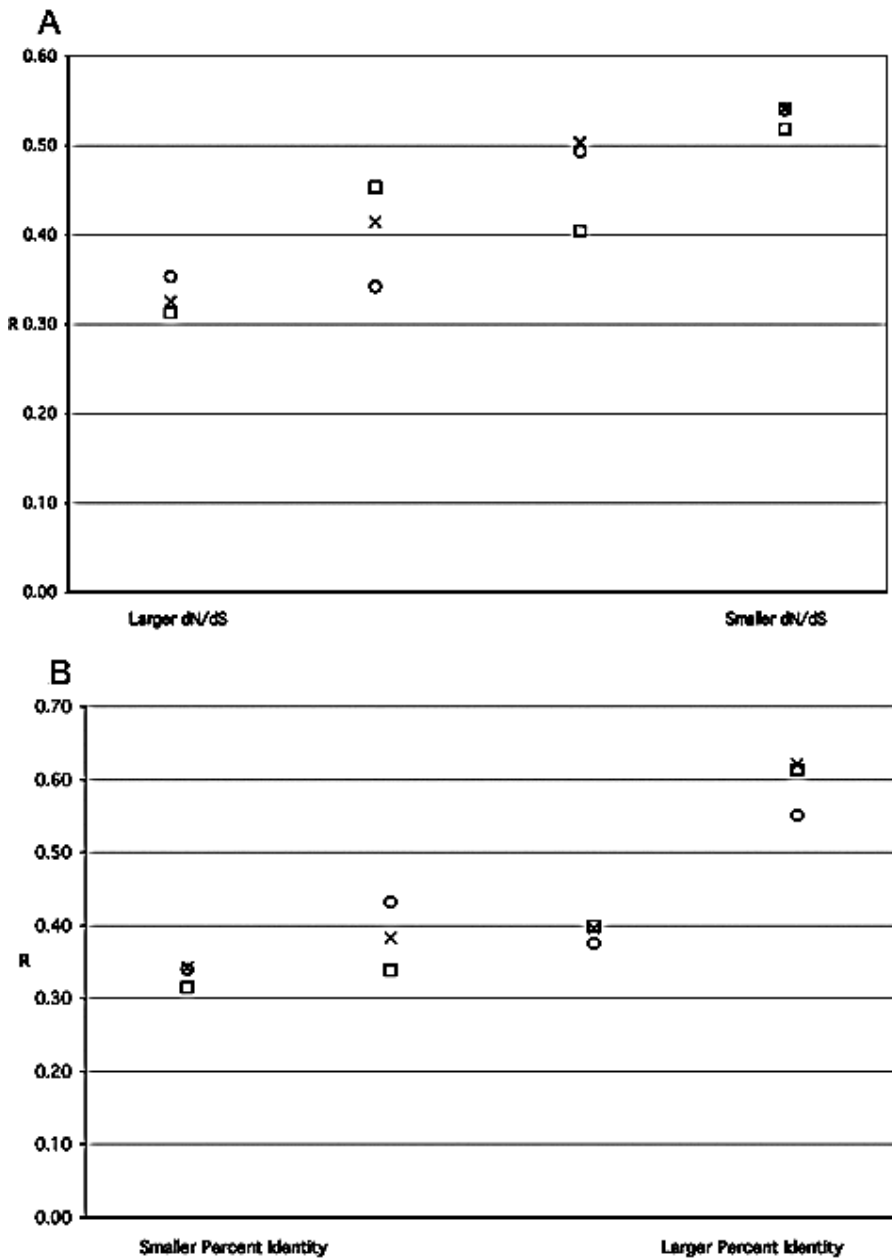


Figure 2. Conservation of Gene Expression Increases with Conservation of Sequence

(A) Orthologous human and mouse genes were ranked by their dN/dS ratios, as given in the ENSEMBL database, from least conserved at the DNA sequence level (larger dN/dS), to most conserved. For each of the three pair-wise regional comparisons, the correlation coefficient between human and mouse log ratios was determined and plotted for each quartile of genes.

(B) Orthologous genes were ranked by their percent nucleotide identity, as given in the ENSEMBL database, and quartile correlation coefficients were determined and plotted for each quartile of genes. X, caudate or striatum-to-cerebellum; Circle, cortex-to-cerebellum; Square, cortex-to-caudate or striatum.

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brain structure and conservation of cell types within comparable regions [18,19]. Conservation of patterned gene expression in the mammalian brain is consistent with standard assumptions of biological uniformity justifying the use of model organisms. Further underscoring conservation of mammalian brain gene expression, we find that in the three brain regions examined, equivalent regions in mouse and human brain are more alike than different regions within a species. This is apparent whether considering the 125 genes with the most variable expression within a species (Figure 1)

or whether considering Euclidian distances based on expression of thousands of orthologous gene pairs (Table 5). Our finding is consistent with other studies contrasting brain gene expression in dogs and humans [20], chimpanzees and humans [7], and mice and humans [10].

We do not mean to suggest, and our findings should not be interpreted to mean that gene expression in human and mouse brains is identical. Here we are mainly concerned with the genes that show an extremely patterned expression across three particular brain regions. Because our study examines

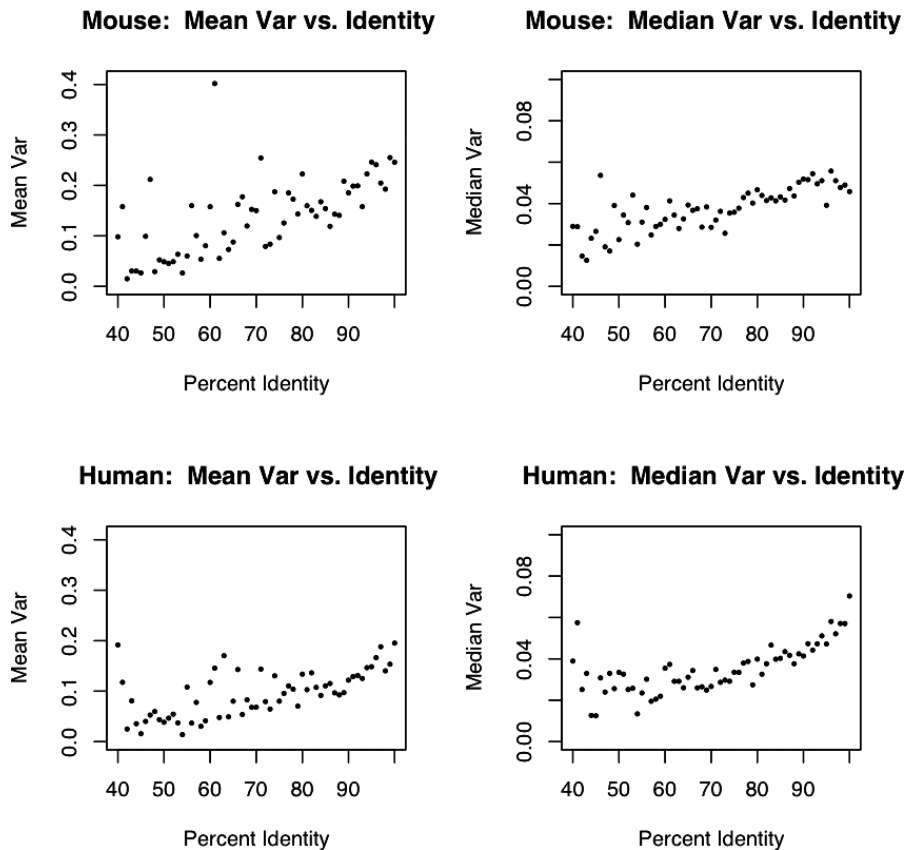


Figure 3. Genes with High Variance across Tissues Have Greater Conservation of Nucleotide Sequence

Orthologous mouse and human gene pairs were binned according to their nearest integral percent identity. The mouse or human mean and median variance of each bin are plotted against the bin's percent identity.

doi:10.1371/journal.pgen.0030059.g003

expression of the tissue, we cannot discern evolutionary changes within specific cell types. However, within the three regions examined here, the overall trend is for regional-marker genes to have been conserved. For example, considering the human motor cortex-to-cerebellum comparison, in the 100 human genes with greatest evidence for differential expression between these regions, there are only nine discordant changes in the 100 mouse orthologs. The overall correlation between human and mouse log ratios in this set of 100 genes is $r = 0.86$. In the top 250 human caudate-to-cerebellum changes, $r = 0.81$, and there are 39 discordant changes. In the top 500 changes, $r = 0.75$, and there are 93 discordant changes. Essentially identical correlations and trends appear in cross-species correlation of the other two regional comparisons (unpublished data). It is very likely that our data somewhat underestimate the true correlation, since factors such as post mortem delay, tissue dissection, and gender ratios were not strictly controlled. Other technical sources of variability include possibly measuring different splice variants in mice and humans, comparing young mice to old humans, and differences in cell-type composition arising from comparing whole mouse tissues to small portions of the human tissues. Finally due to evolution of genomic sequence, Affymetrix (<http://www.affymetrix.com>) must almost always use probes of different sequences to assay human and mouse gene expression. Probe sequence has a profound influence upon the signal detected in a microarray experiment [21].

Overall we observe correlation of relative expression levels in mice and humans on the order of $r = 0.45$. This leaves the proportion of unexplained variance due to technical factors and evolutionary changes as roughly 80% ($1 - 0.45^2 = 0.8$). If an estimate of the variance due to technical factors can be made, in theory it is possible to determine the proportion of unexplained variance due to the evolution of gene expression. From the correlation observed in our independent human regional comparisons, one can arrive at an estimate of 14% of the variance being due to technical noise for a within-species regional comparison ($1 - 0.93^2 = 0.14$). Perhaps twice this or 28% may serve as an estimate of cross-species technical noise. Thus at the high end, our data suggest 52% (80%–28%) of the variance could be due to evolutionary changes. It may be more accurate to suppose that cross-species correlations are subject to the same technical noise effects as within-species correlations on different generations of Affymetrix microarrays. In that case, typically $r = 0.7$ [21]. Therefore, the estimate of variance due to noise is $1 - 0.7^2 = 0.51$, which leaves $0.8 - 0.51 = 0.29$ or 29% as our estimate of the unexplained variance due to evolution of expression in mice and humans. Since we are examining log ratios, evolutionary contributions from both tissues in each species are combined in this number. We presume the true variance due to evolution within each single tissue would be less than 29%.

It might be reasonable to expect that gene expression

variability would be significantly larger between individual humans than between inbred mice housed in uniform conditions. There is little evidence for this in our profiles. We find the fraction of genes that vary between individuals more than between tissues is roughly the same in the two species. These findings could be unique to the regions examined, or they may be a consequence of the between-region variability being so much larger than individual variability for both humans and inbred mice. A more interesting alternative is that this implies that the constraints on brain gene expression are quite strict and that many commonly presumed sources of individual variability are just not that influential. Gender may be one of the largest contributors to individual gene expression variability. In analyses to identify gender-dependent gene expression differences in human brains, differential expression was limited to a rather small set of genes when the X and Y chromosomes were excluded (L. Jones, unpublished data).

Examining orthologous mouse and human genes, we find that conservation of amino acid and nucleotide sequence is correlated with conservation of regional expression. Since this relationship could have been an artifact of our ability to identify homologous genes, we re-examined this relationship by beginning with genes that showed evidence for regional expression within one or the other species. This showed that the genes with higher variability of expression between brain regions within a species also tended to have greater sequence homology with their orthologs than genes that are expressed in multiple brain regions. This is somewhat surprising if one imagines that evolutionary constraints act additively on genes widely expressed in different tissues. It may be that regional gene expression in the brain is particularly highly constrained since the proper behavior of the organism depends upon each brain region functioning smoothly with the others. Wider surveys of tissue gene expression tend to support constraints on brain gene expression, finding that the brains of humans and chimpanzees show fewer differentially expressed genes than kidney, heart, liver, and testes [3,22].

Particular interest has been devoted to finding differences between humans and chimpanzees. Some studies have concluded that there is a bias for genes to be more highly expressed in human cortex relative to chimpanzee [4,6]. While our data cannot directly address chimpanzee and human gene expression, and this claim was made for a rather small number of genes, we see little evidence that the human cortex has uniquely undergone extensive and rapid evolutionary change. Based on the Euclidian distances shown in Table 5, we find it is the cerebellum that is the outlier tissue both within and between the two species.

It is quite possible that complexity of higher order brain functions relate to splicing or protein modifications that escape microarray analysis, but our data suggest some boundaries on the idea that gene expression differences explain differences in cognitive abilities between species. Few genes appear to have evolved new patterns of regional expression. The minority of genes that do show discordant regional expression between adult mice and humans may indeed be key genes regulating brain functions. Alternatively, since general expression patterns in the adult brain have largely been conserved, perhaps it is gene expression during development that ultimately wields the most influence upon higher brain functions by specifying the complexity of

neuroanatomy. Humans have at least two orders of magnitude greater numbers of neurons and neuronal connections than mice [18,19]. Our data suggest the active genes in those neurons and connections are quite similar in adult mice and humans, species with extremely different cognitive abilities. This similarity should become greater as more closely related species, such as chimpanzees and human, are considered. The most important genes relating to cognitive differences may be genes that specify how the machinery is assembled.

Transgenic mice have become the most common model organism for human neurodegenerative diseases [23]. Scrutiny of models has previously involved comparing histopathological and neurochemical phenotypes, or extrapolating from mouse neurobehavioral tests to human disease signs and symptoms. We suggest that the transcriptional signature of the human disease can be used to objectively and globally assess both genetic and phenotypic models; the assumption being that a model that recapitulates the human disorder should have a similar expression profile. Ideally, such assessment involves reference to a range of expression profiles so that the biological specificity of the disease phenotype can be addressed and to provide outlier groups to place relatedness in context [24]. We believe that contrasting healthy mouse- and human-brain gene expression profiles provides a reasonable context with which to assess likeness between mouse models and human neurodegenerative diseases. The high correlation between regional gene expression in healthy brain suggests that mouse models of human neurodegenerative diseases may quite accurately recapitulate the human microarray phenotype and should be held to a high standard.

Here we have focused on the general similarity rather than specific differences between two species. Using several different methods, we find that regional gene expression in the mouse anterior cortex, striatum, and cerebellum is very similar, respectively, to gene expression in human motor cortex, caudate, and cerebellum. Classical comparative neuroanatomy has identified a general conservation of mammalian brain structure, with differences between species arising from elaboration of ancestral forms. Our data indicate that this general conservation continues down to the gene expression level, and that expression patterns in our brains may be less far removed from ancestral forms than apparent differences in mental abilities might suggest.

Materials and Methods

Human tissue dissection and RNA processing. Post mortem human tissue was gathered with ethical approval and permissions, dissected, and processed as specified [11]. The samples were hybridized to Affymetrix HG-U133A arrays containing 22,283 probesets. The primary dataset consisted of caudate, cerebellum, and motor cortex samples from eight men and four women, whose ages ranged from 36 to 77 with an average age of 58 years. Confirmation of the primary data was performed with an independent second group that consisted of caudate and cerebellum samples from seven men and two women whose ages ranged from 22 to 72 with an average of 49 years. Clustering included all human and mouse samples.

Mouse tissue dissection and RNA processing. Postnatal day-35 C57BL/6 mice, five females and one male, were killed by cervical dislocation. The brain was immediately dissected into ice-cold phosphate-buffered saline. Tissue microdissections were performed at 4 °C on one hemisphere at a time with the brain on a bed of dry ice. The cortex was divided into an anterior and posterior portion with the line of division at the point where the striatum and hippocampus meet. Tissue was collected into 5-ml polypropylene Falcon tubes, submerged in liquid nitrogen, and stored at -80 °C. Total RNA was

isolated by adding 1 ml of Qiazol reagent (Qiagen, <http://www.qiagen.com>) to each frozen sample and homogenizing the tissue with a polytron for 40 s at medium speed. Residual salts and proteins were removed with an RNeasy Lipid Kit per the manufacturer's instructions (Qiagen). RNA concentration was determined with spectrometer. The Affymetrix single-cycle probe synthesis kit was used to generate cRNA probe per the manufacturer's instructions. For the cortical and cerebellar samples, 5 µg of total RNA was used as starting material. For striatal samples, 2 µg of total RNA was used. Biotinylated-cRNA was checked on a bioanalyzer prior to and after the fragmentation reaction. Samples representing tissue from a single mouse were hybridized to MOE_430A_2 chips containing 22,690 probesets, $n = 6$ for each tissue. The raw image data is available at <http://www.hdbase.org>.

Microarray analysis. Primary analysis of microarray data was performed using Bioconductor, a freely available software package designed for the analysis of genomic data (<http://www.bioconductor.org>). We first preprocessed and normalized the CEL files with RMA. The primary and secondary groups of human samples were normalized and analyzed separately. Then we fit a linear model (gene expression \approx donor + tissue type) for each of the three paired comparisons of tissue using the Bioconductor library package LIMMA to calculate log ratios, moderated paired t -statistics, and corresponding p -values. We did not further adjust p -values for multiple testing. Here we primarily used p -values for ordering genes. Additional adjustments, such as a Bonferroni or Benjamini-Hochberg correction, would not affect how we ordered genes since such adjustments are typically monotonic operations.

To select sets of genes whose expression was highly enriched in one of the three regions under consideration, we chose as arbitrary criteria that probesets met $p < 0.001$ and log ratio ≥ 1 in both relevant pair-wise comparisons. To rank probesets, the log ratios of the two relevant comparisons were summed in the appropriate fashion to provide a positive regional score. For example, the largest values of $\log_2(\text{BA4/caudate}) + \log_2(\text{BA4/cerebellum})$ would be candidate BA4 genes. Finally, probesets whose summed regional score was > 2 in more than one region were culled from the list.

The variance for a probeset, across n samples, was calculated by

$$\frac{1}{(n-1)} \sum_{i=1}^n (x_i - \bar{x})^2$$

where x_i is the RMA signal for probeset i on array n . After selecting variable genes, to minimize systematic differences of scale between the mouse and human arrays, prior to clustering we separately normalized the mouse and human RMA data to give each probeset zero mean and unit variance. Hierarchical clustering and heat maps using the 125 (an arbitrary number) most variable probesets were generated using Ward's linkage method, which uses an analysis of variance approach to evaluate the distances between clusters. In short, this method attempts to minimize the sum of squares of any two (hypothetical) clusters that can be formed at each step [25]. Heat maps were generated with the Comprehensive R Archive Network (CRAN) package GREGMISC.

Euclidean distances between samples were calculated using RMA signals by

$$\sqrt{(x_1 - y_1)^2 + \dots + (x_g - y_g)^2}$$

where there are g probesets and x and y are any two mouse or human samples. Euclidean distances between regions were calculated using the mean RMA probeset signals for each tissue.

Bioinformatics. To extract ortholog identities, the ENSEMBL database (<http://www.ensembl.org/Multi/martview>) was queried using mouse ENSEMBL identities provided in the Affymetrix annotation. Human ENSEMBL numbers, dN (number of nonsynonymous substitutions/number of nonsynonymous sites), dS (number of synonymous substitutions/number of synonymous sites), dN/dS, and percent identity were retrieved and associated with mouse probesets. dN and dS values were generated using the codeml program included in the PAML package [26,27]. Codeml performs pair-wise Maximum Likelihood calculations of dN and dS for each set of orthologs. We used the F3 \times 4 codon evolution model. This takes into account bias derived from the different probabilities of transition versus transversion mutations and bias due to different nucleotide frequencies at the three codon positions. Incorrect ortholog assignments manifest as anomalously high dS values. We therefore applied a cut off of twice the median dS as the criterion for retaining the dN/dS ratio.

Statistical methods. Since a large but unknown fraction of genes

are coregulated, assumptions of independence are not met. We therefore report extreme statistical significance ($p < 10^{-20}$) as $p \approx 0$, as we do not wish to imply that we believe all assumptions are correct. While additional computation might improve our estimate of p , results when assuming independence are so extreme that our conclusions per statistical significance would not change.

p -Values for the intersections of lists of regional marker genes in Table 3 were calculated assuming a hypergeometric distribution drawing two lists of 30 from a pool of 8,500 genes. p -Values for intersection of most variable mouse and human genes in Figure 1 were calculated assuming a hypergeometric distribution drawing two lists of 125 genes from a pool of 2,998 genes. p -Values for correlation coefficients were calculated with a likelihood ratio test assuming observations are independent realizations from a joint bivariate normal distribution.

Analysis of GO categories in different regions. Categories overrepresented in lists of probes were differentially expressed between different tissue regions (e.g., caudate versus cortex) within species. For the human HG-U133A arrays, 70.6% of the probesets had an assigned GO category. For the mouse MOE430A_2 arrays, 66.2% of the probesets had an assigned GO category. For each GO category, the total number of probes in that category and the number of probes appearing on a list of differentially expressed probes ($p < 0.05$) were calculated. A p -value for overrepresentation of each category was calculated using Fisher's exact test if either the number of probes on the list or the number not on the list was less than ten, otherwise a Pearson chi-square was used. The number of categories achieving a given p -value for overrepresentation was calculated, and its significance assessed by permutation (to account for the overlap in categories). The permutation procedure was as follows: generate a list of differentially expressed probes of equal length to the actual list by sampling probes at random (without replacement); calculate the number of probes on the list for each GO category, and hence a p -value for overrepresentation; count the number of categories with a p -value for overrepresentation less than the specified criterion, and compare to that in the actual data; repeat the process 5,000 times.

Overlaps in overrepresented categories between species for a given regional comparison were examined. These analyses were restricted to the 3,119 GO categories defined for both human and mouse. The number of categories significantly overrepresented ($p < 0.05$) for both mouse and human in the actual data was calculated for each comparison and direction of expression. Significance was again assessed by permutation (to reflect the fact that several probes are in more than one category). A random list of differentially expressed probes of equal length to that observed in human was generated and used to calculate p -values for overrepresentation for the human GO categories, as before. The n most significant categories were selected (n being the number of significantly overrepresented categories in the actual human data), and the overlap between these and the significantly overrepresented mouse categories calculated. The process was repeated 10,000 times. For all three regional comparisons and all expression directions, the number of overlapping categories in the actual data was higher than that obtained in any of the simulated replicates.

Supporting Information

Dataset S1. Complete Data for Human RMA Signals

Found at doi:10.1371/journal.pgen.0030059.sd001 (8.8 MB ZIP).

Dataset S2. Complete Data for Human Regional Comparisons

Found at doi:10.1371/journal.pgen.0030059.sd002 (6.0 MB ZIP).

Dataset S3. Complete Data for Mouse RMA Signals

Found at doi:10.1371/journal.pgen.0030059.sd003 (5.1 MB ZIP).

Dataset S4. Complete Data for Mouse Regional Comparisons

Found at doi:10.1371/journal.pgen.0030059.sd004 (5.0 MB ZIP).

Dataset S5. Alignment of Human and Mouse Orthologous Genes Including Differential Expression Statistics from Regional Comparisons

Found at doi:10.1371/journal.pgen.0030059.sd005 (3.8 MB ZIP).

Figure S1. Relative Gene Expression in Human Brain Tissues Is Robust and Reproducible

The log ratio from the caudate-to-cerebellum comparison in the primary set of human samples is plotted (x-axis) against the caudate-

to-cerebellum log ratio of the independent second set of human samples. The data are highly correlated ($r = 0.93$) despite having unbalanced age and sex ratios between the two groups.

Found at doi:10.1371/journal.pgen.0030059.sg001 (2.1 MB TIF).

Table S1. Key Relating GEO Accession Numbers, Clinical Covariates, and Samples

Found at doi:10.1371/journal.pgen.0030059.st001 (31 KB XLS).

Table S2. Striatal Genes Listed in [16]

Found at doi:10.1371/journal.pgen.0030059.st002 (34 KB XLS).

Table S3. GO Categories Common to Human and Mouse Brain Regions

Found at doi:10.1371/journal.pgen.0030059.st003 (59 KB XLS).

Table S4. Observed and Expected Numbers of GO Categories Reaching Various p -Value Thresholds and $p < 0.05$ False Discovery Rates (FDR)

Found at doi:10.1371/journal.pgen.0030059.st004 (14 KB XLS).

Accession Numbers

The GEO database (<http://www.ncbi.nlm.nih.gov/geo>) accession num-

ber is GSE3790. Affymetrix Web site (<http://www.affymetrix.com>) annotations for human HG-U133A and mouse MOE430_2 are from (<http://www.affymetrix.com/support/technical/byproduct.affx?product=hgu133>) and (<http://www.affymetrix.com/support/technical/byproduct.affx?product=moe430-20>).

Acknowledgments

Author contributions. AH and LJ generated the human microarray data. ZCB and KRJ generated the mouse microarray data. ADS, AKA, and CK analyzed the data and did the supporting statistical analysis. PH and LJ provided GO and supporting statistical analysis. AH and PC extracted homology and ortholog information. ADS, LJ, and JMO wrote the manuscript.

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Competing interests. The authors have declared that no competing interests exist.

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