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Evidence for evolutionary divergence of activity-dependent gene expression in developing neurons

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Abstract
Evolutionary differences in gene regulation between humans and lower mammalian experimental systems are incompletely understood, a potential translational obstacle that is challenging to surmount in neurons, where primary tissue availability is poor. Rodent-based studies show that activity-dependent transcriptional programs mediate myriad functions in neuronal development, but the extent of their conservation in human neurons is unknown. We compared activity-dependent transcriptional responses in developing human stem cell-derived cortical neurons with those induced in developing primary- or stem cell-derived mouse cortical neurons. While activity-dependent gene-responsiveness showed little dependence on developmental stage or origin (primary tissue vs. stem cell), notable species-dependent differences were observed. Moreover, differential species-specific gene ortholog regulation was recapitulated in aneuploid mouse neurons carrying human chromosome-21, implicating promoter/enhancer sequence divergence as a factor, including human-specific activity-responsive AP-1 sites. These findings support the use of human neuronal systems for probing transcriptional responses to physiological stimuli or indeed pharmaceutical agents.

Introduction
The last common ancestor of mice and humans existed around 80 million years ago, sufficient time for divergence in signal-dependent gene regulation \(^1\). However, the conservation or divergence of dynamic signal-dependent programs of gene expression is incompletely understood, particularly in the nervous system. A fundamental transcriptional program of this sort is that elicited in neurons by electrical activity, triggered via Ca\(^{2+}\) influx through ligand- and/or voltage-gated Ca\(^{2+}\) channels and activating genes containing Ca\(^{2+}\)-responsive transcription factor binding sites in their promoters \(^2\)\(^-\)\(^4\). These changes in gene expression are critical to a neuron’s functional response to electrical activity in development and maturity \(^5\)\(^-\)\(^7\), and are totally distinct from the toxic sequelae of excitotoxic Ca\(^{2+}\) influx \(^8\)\(^,\)\(^9\). For example, rodent studies have shown that activity-dependent gene expression programs direct myriad processes in developing neurons, including neuroprotection, dendritic arborization, and synaptic plasticity \(^5\)\(^-\)\(^7\). Moreover, certain neurodevelopmental disorders are associated with defects in activity-dependent transcriptional mechanisms \(^6\), making it important to fully understand transcriptional programs that are triggered by electrical activity in developing human neurons.

Comparing the influence of neuronal electrical activity on mouse-human orthologs, and identifying the basis for any differences, requires a combination of approaches, given the inability to reproducibly study primary human developing neurons. Embryonic stem cell (ESC)-based technology enables the generation of glutamatergic cortical-patterned neurons from human
embryonic stem cells (hESC CORT-neurons) of sufficient homogeneity and electrical maturity \textsuperscript{10, 11} to enable the study of activity-dependent gene expression. Such responses can then be compared to both primary mouse cortical neurons (of differing developmental stages) as well as those derived from mouse ESCs, to distinguish species-specific differences from those dependent on developmental stage or origin (primary tissue vs. stem cell line). Moreover, it is in theory possible to study mouse-human ortholog regulation in the same neuron by exploiting the aneuploid Tc1 mouse which carries a freely segregating copy of human Chromosome 21, to identify whether any differences are independent of cellular environment (i.e. are due to DNA sequence divergence). A combination of these approaches has been employed to reveal strong conservation of the neuronal activity-dependent transcriptome onto which a significant degree of divergence is overlaid.

Results and Discussion

We generated dissociated glutamatergic cortical-patterned neurons from human embryonic stem cells (hESC CORT-neurons), whose characterisation is described fully elsewhere \textsuperscript{10, 11}. These cells have a combination of homogeneity and electrical maturity that is hard to achieve with classical human stem cell-derived neurosphere-based preparations, which have a poor dynamic range in terms of gene regulation \textsuperscript{12} compared to primary mouse neuronal preparations \textsuperscript{13}

We studied transcriptional responses to L-type Ca\textsuperscript{2+} channel activation, an important mediator of activity-dependent gene regulation \textsuperscript{6, 14-18}. To do this, hESC CORT-neurons were subject to KCl-induced membrane depolarization in the presence of the L-type Ca\textsuperscript{2+} channel agonist FPL64176 (KCl/FPL), which leads to robust and uniform Ca\textsuperscript{2+} responses (\textsuperscript{10}, Figure 1-figure supplement 1a, Figure_1_Source_Data_2) similar to rodent neurons \textsuperscript{19}. QPCR analysis revealed robust induction of classical early-response activity-regulated genes (ARGs) FOS, FOSEL2 (FRA-2), and FOSB, and neurobiologically important ARGs BDNF, ARC, and NPAS4 (Figure 1-figure supplement 1b, Figure_1_Source_Data_3) whose regulation has hitherto been studied primarily in rodent neurons.

We then performed RNA-seq analysis of KCl/FPL-induced gene expression changes in hESC CORT-neurons as well as days-in-vitro (DIV)10 mouse primary cortical neurons (Mus-PRIM CORT-neurons, predominantly excitatory, like Hum-ESC CORT-neurons), focusing on a 4h timepoint (Fig. 1a,b, Figure_1_Source_Data_1). QPCR validation of fold-induction of selected ARGs revealed a tight correlation with the RNA-seq data (r >0.99 for Hum-ESC CORT-neurons and mouse neurons respectively, Figure 1-figure supplement 1c, 1d, Figure_1_Source_Data_4), supporting the reliability of the RNA-seq data set. Our interspecies comparisons were restricted to 1:1 orthologous pairs whose average expression level met a minimum threshold (11,302 genes expressed >0.5 FPKM on average, in all cell types). Genes induced >5-fold in both Hum-ESC CORT- and DIV10 Mus-PRIM CORT-neurons according to RNA-seq included neurobiologically important genes such as PER1, EGR2, EGR4 and ATF3 (Fig. 1c). Global comparison of Log\textsubscript{2} (gene fold-
change) in orthologous pairs revealed a correlation between Hum-ESC\textsuperscript{CORT}-neurons and DIV10 Mus-PRIM\textsuperscript{CORT}-neurons (\(r=0.480,\) 95% CI: 0.465 to 0.494, \(P<0.0001,\) Fig. 1d, Figure_1_Source_Data_1), pointing to a significant, but incomplete, degree of conservation. For the 11,302 ortholog pairs, we calculated the "differential regulation index" (DRI), defined as the fold-change (Hum-ESC\textsuperscript{CORT}-neurons) divided by the fold-change (Mus-PRIM\textsuperscript{CORT}-neurons). The log\(_2\) of the DRIs are normally distributed about zero with a standard deviation of 0.68 (Figure 1-figure supplement 1e, Figure_1_Source_Data_5).

While this imperfect conservation could involve species-specific gene responsiveness, it could be due to other factors: i) non-developmental equivalency in the human and mouse neuronal preparations combined with a dependency of the responsiveness of certain genes on developmental stage could underlie differences; ii) if gene-responsiveness were sensitive to the origin of the neurons studied (embryonic stem cell line (human) vs. primary tissue (mouse)) this could also lead to differences, and an erroneous exaggeration of apparent evolutionary divergence.

To address (i) we compared KCl/FPL-induced gene regulation in DIV10 Mus-PRIM\textsuperscript{CORT}-neurons with that in synaptically immature DIV4 Mus-PRIM\textsuperscript{CORT}-neurons, which revealed a strong correlation (\(P<0.0001, r=0.834, 95\% \) CI: 0.833 to 0.844, Fig. 1e, Figure_1_Source_Data_1), and a log\(_2\) DRI (DIV4:DIV10) distribution curve (Figure 1-figure supplement 1f, Figure_1_Source_Data_5) substantially narrower than that observed in Figure 1-figure supplement 1e. Taken together, this suggests that developmental stage does not have a large effect on the activity-responsiveness of genes at this time point and is unlikely to account for the human-mouse differences observed in Fig. 1d.

To address (ii) (i.e. whether stem cell- vs. primary tissue origin causes differential gene responsiveness) we generated cortical-patterned neurons from mouse embryonic stem cells (Mus-ESC\textsuperscript{CORT}-neurons) using established protocols \(^20\), which exhibit spontaneous firing and synaptic activity (Fig. 2a-d), similar to our Hum-ESC\textsuperscript{CORT}-neurons \(^10, 11\). The Mus-ESC\textsuperscript{CORT}-neurons were subject to the same 4h KCl/FPL stimulation and RNA-seq analysis (Figure 2-figure supplement 1a). Same-species comparison of gene fold-change was made between Mus-PRIM\textsuperscript{CORT}-neurons (DIV4 and DIV10) and Mus-ESC\textsuperscript{CORT}-neurons in the 11,302 genes compared in Fig. 1e,f. A strong correlation was observed in gene-responsiveness in Mus-ESC\textsuperscript{CORT}-neurons vs. Mus-PRIM\textsuperscript{CORT}-neurons at either DIV10 or DIV4 (\(P<0.0001, r=0.748\) (DIV10), Fig. 2e; \(r=0.788\), (DIV4), Fig. 2f), significantly stronger than between Mus-ESC\textsuperscript{CORT}- and Hum-ESC\textsuperscript{CORT}-neurons (\(r=0.596,\) Fig. 2g, see also Figure_2_Source_Data_1). Indeed, a comparison of gene fold-change between all biological replicates in all data sets revealed that all mouse neuronal preparations (primary DIV10, primary DIV4, and ESC-derived) correlated far more strongly with each other than with the human ESC\textsuperscript{CORT}-neurons. (Fig. 2h). This suggests that the weaker correlation with the hESC\textsuperscript{CORT}-neurons may indeed be due to some species-specific differences in neuronal gene activity-responsiveness, rather than differences in developmental stage or cellular origin.
As a final genome-wide comparison we wanted to determine whether activity-dependent gene responsiveness showed a higher or lower degree of divergence than basal gene expression levels, and whether differences in basal gene expression were any predictor of differential activity-dependent gene responsiveness. Comparison of basal expression levels between Hum-ESC<sup>CORT</sup>-neurons and mouse neurons (DIV10 Mus-PRIM<sup>CORT</sup>-neurons, DIV4 Mus-PRIM<sup>CORT</sup>-neurons, and Mus-ESC<sup>CORT</sup>-neurons) revealed correlation coefficients of 0.714, 0.711, and 0.710 (Figure 2-figure supplement 1b-d, Figure_2_Source_Data_3). This correlation is substantially stronger than the that observed when comparing gene fold-change after KCl stimulation (0.480, 0.526, .595, Fig. 1d, 2g, Figure 1-figure supplement 1g). Moreover, we performed a similar clustering analysis as in Fig. 2h which illustrates this graphically: i.e. while the basal expression profile of Hum-ESC<sup>CORT</sup>-neurons does not cluster as closely as the three mouse neuronal populations do to each other, the degree of difference is less than that observed in Fig. 2h (Fig. 2i). Therefore basal neuronal gene expression shows less divergence than the responsiveness of genes to depolarisation. We also investigated whether gene differential responsiveness to KCl (DRI) in human vs. mouse neurons has any relationship with the relative basal expression of that gene in human vs. mouse neurons.

For each of the 11,302 orthologous pairs, we plotted the Log<sub>2</sub>(DRI) Hum-ESC<sup>CORT</sup>- vs. DIV10 Mus-PRIM<sup>CORT</sup>-neurons (i.e. DRIs from Figure 1-figure supplement 1e) against the Log<sub>2</sub>(DBEI), where DBI (differential basal expression index) is the ratio of basal expression in Hum-ESC<sup>CORT</sup>- vs. DIV10 Mus-PRIM<sup>CORT</sup>-neurons (Figure 2-figure supplement 1e, Figure_2_Source_Data_2). As can be seen, there is no link between a gene’s relative responsiveness to depolarisation in human vs. mouse neurons, and its relative basal expression levels in human vs. mouse neurons. Moreover, if we consider the 657 genes where Log<sub>2</sub>(DRI)>1, the standard deviation of their respective Log<sub>2</sub>(DBEI), 1.45, is similar to the standard deviation of Log<sub>2</sub>(DBEI) across all 11,302 genes (1.39), suggesting that there is no dramatic change in divergence of basal gene expression regardless of direction, in genes where DRI>1. Thus, evolutionary differences in activity-dependent gene responsiveness are not substantially attributable to differences in basal expression.

We hypothesized that species-specific gene-responsiveness to neuronal activity could be in part due to evolutionary divergence in genetic sequence, such as the promoters and distal enhancers that mediate activity-dependent changes in gene expression<sup>6, 21</sup>. To investigate whether genetic sequence could be sufficient to quantitatively influence species-specific gene inducibility, we utilized neurons cultured from the Tc1 transchromosomal mouse strain which carries a copy of human chromosome 21 (Hsa21), albeit with approximately 10% deleted<sup>22, 23</sup>. Thus, regulation of Hsa21 genes, plus their mouse 1:1 ortholog, could be studied side-by-side in the same cellular environment of a mouse primary neuron. We cultured cortical neurons from Tc1 embryos to DIV10, stimulated ± KCL/FPL, performed RNA-seq, and developed a workflow that distinguished between human and mouse RNA-seq reads, discarding any reads that were ambiguous. Once the initial criterion of a perfect sequence match of the RNA-seq read to either Hsa21 or the mouse genome was met, only a further 0.055% of reads were discarded due to
100% sequence conservation between species. Thus, expression level of human and mouse
genes carried in the Tc1 mouse can be analysed ± stimulation (Fig. 3a). We first verified that the
mouse genes in Tc1 neurons responded similarly to wild-type mouse neurons of the same age
(Figure 3-figure supplement 1a, Figure_3_Source_Data_1, r=0.900).

72 orthologous pairs were then focussed on, where the Hsa21 ortholog met an expression
cut-off of >100 reads. For each pair, we calculated the "Tc1 DRI" (fold-change (human
ortholog)/fold-change (mouse ortholog)) as a measure of whether responsiveness of the orthologs
was different or similar, and plotted it against the gene’s DRI calculated from fold-induction in
separate human/mouse neuronal preparations: fold-change (Hum-ESC\textsuperscript{CORT}-neurons) / fold-change
(DIV10 Mus-PRIM\textsuperscript{CORT}-neurons). Importantly, we observed a significant correlation between these
DRIs ($P<0.0001$, r=0.730, Fig. 3b, Figure_3_Source_Data_2), meaning that differences in gene
regulation observed in separate human and mouse cortical neuron preparations ("separate DRI")
are broadly recapitulated when the orthologs are studied side-by side in the same cellular
environment ("Tc1 DRI"). This points to the DNA sequence as a contributor to quantitative species-
specific gene responsiveness. An example of such a gene is \textit{ETS2}, the human ortholog of which is
induced more strongly and rapidly than the mouse ortholog (Fig. 3c,d, Figure_3_Source_Data_3).
Interestingly, a region of its proximal promoter contains three human-specific AP-1 sites (Fig. 3e,
right) and was found to confer activity inducibility on a luciferase reporter, unlike the corresponding
region of the mouse \textit{Ets2} promoter (Fig. 3e, left). Moreover, expression of inhibitory AP-1 mutant
TAM\textit{67} \textsuperscript{24} inhibited Hum-\textit{ETS2}-luciferase reporter induction, and mutation of the three AP-1 sites to
the corresponding mouse sequence abolished inducibility (Fig. 3e). Thus, the \textit{ETS2} proximal
promoter contains functional human-specific AP-1 sites, potentially contributing to the stronger and
more rapid induction of the human ortholog in neurons.

Although the focus of this study is understanding the extent and causes of differences in
gene activity-responsiveness in human vs. mouse neurons, it should be noted that qualitative gene
up/down regulation is well conserved. For example, of the top 100 most strongly induced human
genes in Hum-ESC\textsuperscript{CORT}-neurons (FC>3.5 fold, Padj<0.05), 85 are also induced in Mus-ESC\textsuperscript{CORT}-
neurons (FC>1.5-fold, Padj<0.05). Of the remaining 15 genes, 6 are induced in DIV4 and/or DIV10
Mus-PRIM\textsuperscript{CORT}-neurons, taking the total to 91/100. Conserved inducibility remains high even at a
lower fold-change cut-off: e.g. 409 out of 475 genes induced >2-fold (Padj<0.05) in Hum-ESC\textsuperscript{CORT}-
neurons are also induced (FC>1.5-fold, Padj<0.05) in one or more of the mouse neuronal
preparations. Although we confirmed examples of genes regulated both up and down at 4h in
Hum-ESC\textsuperscript{CORT}-neurons but not in mouse neurons, by performing a timecourse in Hum-ESC\textsuperscript{CORT}-
and DIV10-Mus-PRIM\textsuperscript{CORT} neurons (Fig. 3f), we hesitate to label these as definitively human-
specific activity-response genes: there may be time-points, stimulation paradigms, or
developmental stages that do expose activity-dependency on mouse neurons.

Nevertheless, our study points to quantitative differences in activity-dependent gene-
responsiveness in developing human neurons, compared to their mouse counterpart, mediated at
least in part by DNA sequence divergence. It is possible that these differences have a functional impact on the physiological response of human neurons to electrical activity. More generally, given that diverse signal dependent transcriptional programs mediate cellular responses to physiological stimuli, as well as the effects (and side-effects) of pharmaceutical agents, this study points to the value of employing human neurons as a tool to help bridge the translational gap between rodent studies, and human neurophysiology.

Materials and Methods

Generation of cortical-patterned neurons from human ES cells. A detailed description of the derivation of hESC-CORT-neurons, including immunohistochemical validation of cortical markers as well as electrophysiological characterization can be found in recent studies from the authors. Briefly, the hESC line H9 was used, and obtained from WiCell (Madison, WI) under full ethical / IRB approval of the University of Edinburgh. hESCs were maintained on CF-1 irradiated mouse embryonic fibroblasts, neurally converted to anterior-patterned neural precursor cells (NPCs) in suspension in chemically defined medium as described, whereupon neural rosettes were mechanically isolated, dissociated and plated for proliferative expansion. For differentiation of anterior NPCs, they were plated in default media (A-DMEM/F12, 1% P/S, 0.5% Glutamax, 0.5% N2, 0.2% B27, 2 μg/mL Heparin (Sigma)) on poly-D-lysine (Sigma), laminin (Sigma), fibronectin (Sigma) and matrigel coated coverslips for differentiation and fed twice a week. Default media was supplemented with 10 μM forskolin (Tocris) in weeks 2 and 3. From week 4 onwards forskolin was removed and default media was supplemented with 5ng/mL BDNF and 5 ng/mL GDNF. Experiments were performed 5 weeks after the differentiation of anterior NPCs was commenced.

Generation of cortical-patterned neurons from mouse ES cells. Mouse ES cells (E14tg2α) were maintained in GMEM (Sigma) supplemented with 2-mercaptoethanol (Gibco), non-essential amino acids (Gibco), glutamine, pyruvate, 10% foetal calf serum (Gibco) and 100 units/ml LIF (made in-house) on gelatinised tissue culture flasks Monolayer neural differentiation is described elsewhere and modified here to include cyclopamine. Briefly, ES cells were washed to remove all traces of serum and then plated at a density of 1x10⁴ cells/cm² onto gelatin-coated tissue culture plastic in N2B27 serum-free medium. N2B27 consists of a 1:1 ratio of DMEM/F12 and Neurobasal media (Gibco) supplemented with 0.5% modified N2 (made in house as described, 1% B27 (Gibco), 2-mercaptoethanol (Gibco), 2mM L-Glutamine, and 1uM Cyclopamine (Sigma). Medium was changed every day. Cells were cultured for 9 days under these conditions before transferring to terminal differentiation conditions. Terminal differentiation of mouse NPCs into dorsal forebrain (cortical) neurons was achieved as described elsewhere by withdrawal of cyclopamine from day 9 NPCs dissociated into single cells, with Accutase (Life technologies), and re-plated on Matrigel (SLS, 354230; 1 in 100 dilution), Fibronectin 20mg/ml (F2006 - Sigma-Aldrich), Laminin 10 mg/ml
Coated coverslips at a density of 30000 cells per 0.3 cm². Addition of 5 µM AraC on day 10, for 24h, was used to remove residual proliferating cells. Differentiating cortical neurons were maintained for a further 12-14 days, with media changes every 2-3 days, in Advanced DMEM/F12 (Invitrogen) containing: 1% N-2 supplement (Invitrogen), 1% B27 supplement (Invitrogen), 1% penicillin/streptomycin (Invitrogen), 0.5% GlutaMAX (Invitrogen), and 5 mg/ml heparin (Sigma), supplemented with 1 mM N-acetyl cysteine (Sigma) and 10 ng/ml BDNF (R&D Systems) and finally processed for RNAseq and electrophysiological studies. For cell culture IHC, cells were fixed in 4% formaldehyde for 20 min at room temperature, washed with PBS and permeabilized with the detergent NP40 (Life Technologies). Cells were subsequently incubated in primary antibodies over night at 4°C. The next day, cells were washed with PBS and incubated with the appropriate secondary antibody at room temperature for 2h. Cells were then mounted using the mounting medium Vectashield (Vector Labs).

Mouse primary cortical neuronal culture from wild-type and Tc1 mice. Cortical mouse neurons were cultured from either wild-type CD1 mice or Tc1 mice described at a density of between 9-13 x 10⁴ neurons per cm² from E17.5 mice with Neurobasal growth medium supplemented with B27 (Invitrogen, Paisley, UK). Stimulations of cultured neurons were done after a culturing period of 9-11 days during which neurons develop a network of processes, express functional NMDA-type and AMPA/kainate-type glutamate receptors, and form synaptic contacts. For Tc1 mouse genotyping, DNA was isolated from cerebellum using Qiagen QIAamp DNA Mini Kit following manufacturer’s instructions. Genotyping reactions were performed using the following primers: D21S55 Forward = 5’-GGT TTG AGG GAA CAC AAA GCT TAA CTC CCA-3’, reverse = 5’-ACA GAG CTA CAG CCT CTG ACA CTA TGA ACT-3’; Myo Forward = 5’-TTA CGT CCA TCG TGG TGG ACA GCAT-3’, reverse = 5’- TGG GCT GGG TGT TAG TCT TAT-3’. D21S55 recognizes the Tc1 allele (product = 208 bp) whereas Myo recognizes both Tc1 and WT alleles (product = 245 bp).

Stimulation of mouse cortical and hESC-CORT-neurons. Prior to stimulation, neurons were first placed overnight into a minimal defined medium containing 10% MEM (Invitrogen), 90% Salt-Glucose-Glycine (SGG) medium (SSG: 114 mM NaCl, 0.219 % NaHCO₃, 5.292 mM KCl, 1 mM MgCl₂, 2 mM CaCl₂, 10 mM HEPES, 1 mM Glycine, 30 mM Glucose, 0.5 mM sodium pyruvate, 0.1 % Phenol Red; osmolarity 325 mosm/l). KCl/FPL stimulations were performed as described previously. KCl/FPL stimulation involved neurons being exposed to 50 mM KCl by adding 0.41 volumes of KCl depolarization solution (10 mM HEPES, pH 7.2, 170 mM KCl, 1 mM MgCl₂, 2 mM CaCl₂) in the presence of 5 µM FPL64176 (Tocris), plus an NMDA receptor antagonist (MK-801, 5 µM) to prevent any excitotoxicity. As described recently, hESC-CORT-neurons respond strongly and uniformly to KCl/FPL treatment, with over 80% of hESC-CORT-neurons exhibiting a >10-fold increase in [Ca²⁺], and the remainder showing at least a 5-fold increase.
Calcium imaging. Ca\textsuperscript{2+} imaging was performed as described\textsuperscript{34,35} at 37 °C in aCSF (150 mM NaCl, 3 mM KCl, 10 mM HEPES, 2 mM CaCl\textsubscript{2}, 1 mM MgCl\textsubscript{2}, 1 mM glucose). Briefly, cells were loaded with 11 \(\mu\)M Fluo-3 AM (from a stock solution of 2.2 mM Fluo-3 dissolved in anhydrous DMSO containing 20% (w/v) Pluronic detergent) for 30 min at 37 °C. Fluo-3 fluorescence images (excitation 472 ± 15 nm, emission 520 ± 15 nm) were taken at one frame per 5 sec using a Leica AF6000 LX imaging system, with a DFC350 FX digital camera. To calibrate images, Fluo-3 was saturated by adding 50 \(\mu\)M ionomycin to the perfusion chamber (to obtain \(F\text{max}\)) and quenched with 10 mM MnCl\textsubscript{2} + 50 \(\mu\)M ionomycin to levels corresponding to 100 nM Ca\textsuperscript{2+}\textsuperscript{36}, which was in turn used to calculate \(F\text{min}\). Free Ca\textsuperscript{2+} concentrations were calculated from fluorescence signal (\(F\)) according to the equation \([\text{Ca}^2+] = K_d(F - F\text{min})/(F\text{max} - F)\), and expressed as a multiple of the \(K_d\) of Fluo-3 (which is approximately 315 nM).

Electrophysiology. The whole-cell patch configuration was used to record membrane currents and voltage deflections as previously described\textsuperscript{10,37}. Membrane potential data are corrected for liquid junction potential (+14 mV). Current and voltage measurements were typically low-pass filtered online at 2 kHz, digitized at 10 kHz and recorded to computer using the WinEDR V2.7.6 Electrophysiology Data Recorder (J. Dempster, Department of Physiology and Pharmacology, University of Strathclyde, UK).

RNA sequencing. Neurons were treated with KCl/FPL as described above. Cells were lysed and RNA extracted at 4h post-stimulation. For both mouse and human-based experiments, three independent biological replicates were performed, and analysed in parallel. Three replicates were deemed sufficient based on previous transcriptome studies on activity-dependent gene expression in which we successfully identified differentially regulated genes with this number of replicates\textsuperscript{38}. Total RNA was assessed for quality (Agilent Bionalyzer) and quantity (Invitrogen Qubit) before library preparation. Illumina libraries were prepared from 1 \(\mu\)g of total RNA using TruSeq RNA Sample Prep Kit v2 with a 10 cycle enrichment step as per the manufacturer’s recommendations. Final libraries were pooled in equimolar proportions before Illumina sequencing on a HiSeq 2500 platform using 75 base paired-end reads. Raw reads were processed using RTA 1.17.21.3 and Casava 1.8.2 (Illumina). Reads were mapped to the mouse (mm10) and human (hg38) reference genomes using version 2.4.0i of the STAR RNA-seq aligner\textsuperscript{39}. A table of per-gene read counts was generated from the mapped reads with featureCounts version 1.4.6-p2\textsuperscript{40}, using gene annotations from Ensembl version 82. Differential expression analysis was then performed using DESeq2 (R package version 1.10.0)\textsuperscript{41}. Raw data will be deposited in the European Nucleotide Archive in advance of publication.

Species-specific sorting of Tc1 neuron RNA-seq data: Given a set of RNA-seq reads that have derived from transcripts of both the mouse genome and human chromosome-21, we implemented a sorting procedure to assign reads to their true species of origin. In the balance
between precision and recall, our strategy is conservative, in that it aims foremost to minimise the number of reads allocated to the incorrect species; but we simultaneously seek to maximise the number of reads that can be unambiguously assigned to the correct species. The sorting criteria detailed below were chosen to effect this goal. In this species-specific sorting (SSS) procedure, reads are first mapped to the genomes of each species with the STAR RNA-seq aligner. At this stage multi-mapping alignments are allowed (--outFilterMultimapNmax 10000), but only those with an alignment score equal to the maximum (--outFilterMultimapScoreRange 0). Subsequently, for each RNA-seq read the alignments of that read to each genome are compared. If the read has alignments to the mouse genome, but no alignments to the human chromosome-21 exist, the read is provisionally assigned to the mouse; note, however, the further requirements given below for a read to be finally allocated to this genome. Similarly, if alignments to human chromosome-21 exist, but there are no alignments to the mouse genome, the read is provisionally assigned to the human. If alignments to both genomes exist, these alignments are examined in more detail. Firstly, any read which aligns multiple times to either genome is discarded, since in this initial conservative strategy their exclusion removes a potential source of ambiguity. Next, the number of mismatched bases between the mapped read and each genome is examined. If the number of mismatches is smaller for one species, the read is provisionally allocated to that genome. If the number of mismatches for each species is equal, a further check is made on the structure of the alignments; a successful alignment is required to span the full length of the read (without any clipping of bases), and, if the alignment spans an intron, at least 5 bases are required to align to the exons on either side of the boundary. If these criteria are satisfied for the alignment to one genome, but not to the other, then the read is provisionally assigned to the first genome. If the criteria are satisfied for the alignments to both genomes, the read is rejected as ambiguous; it cannot be assigned with confidence to one species rather than the other. The read is also rejected if the structural criteria are not satisfied for the alignments to either genome.

At this stage, a read has either been provisionally allocated to one species, or has been rejected. In the former case, a final set of checks are made on the provisional alignment. In this conservative strategy, these are that there should be no mismatches between the mapped read and the genome, and that the structural criteria outlined above are satisfied (if this has not already been confirmed). The outcome of this procedure is that all reads successfully assigned to one species or the other have a single, full-length alignment to that species' genome, with no mismatched bases. Subsequently, for each sample, per-gene read counts were summarised using featureCounts. Relative expression levels of genes are expressed as fragments per million reads per kilobase of message (FPKM). Within the SSS-workflow, only reads that are unambiguously attributed to a particular species are used as the denominator in the FPKM calculation. The value for the length of message for a particular gene refers to the maximum transcript length. Where gene length data is given, this refers to the number of nucleotides contained within the union of all
exons of all transcripts of the gene, including 5' and 3' UTRs. Differential expression analysis on
data sets was performed using DESeq2.

Plasmids and reporter assays. A region of the human ETS2 promoter (NM_001256295 -1080 to -
19, or NM_005239.5, positions -1622 to -542) was generated by PCR amplification using
Stratagene Ultrall Fusion HS DNA Polymerase kit and subcloned into pGL4.10. All other promoter
sequences (both wild-type and mutant) were obtained as synthetic clones from Life Technologies
(Geneart) and then subcloned into pGL4.10. This included the corresponding mouse Ets2
promoter sequence (ENSMUST00000023612 -1055 to -65, or NM_011809.3 -1387 to -397) and a
mutated form of the human ETS2 promoter with the putative AP-1 sites mutated to the
corresponding mouse sequence (ETS2(ΔAP1)) was synthesized. The following plasmids have
been described previously: pcDNA3-cJun(3-122) (TAM-67) 24. Firefly luciferase-based reporter
gene constructs were transfected (Lipofectamine 2000) into DIV8 mouse cortical neurons as along
with a Renilla expression vector (pTK-RL) control. The number of biological replicates were
performed based on previous studies where site-directed mutagenesis was performed on reporter
constructs to assess activity-dependent promoter regulation 38. Neurons were stimulated with
KCl/FPL (where appropriate) 24 h after transfection. Luciferase assays were performed using the
Dual Glo assay kit (Promega) with Firefly luciferase-based reporter gene activity normalized to the
Renilla control (pTK-RL plasmid) in all cases.

RNA extraction and qPCR. For RNA-seq and qPCR, RNA was isolated using the Roche HP RNA
Isolation kit according to manufacturer’s instructions. For qPCR, cDNA was synthesized from 1– 3
μg RNA using Roche Transcriptor cDNA Synth kit. Briefly, RNA was diluted in RNase-free water
and mixed on ice with 1 μl oligo-dT primers (50pmol/μl), 2 μl random hexamers (600 pmol/μl), 4 μl
Transcriptor RT Reaction Buffer (5x), 0.5 μl Protector RNase Inhibitor (40 U/μl), 2 μl
Deoxynucleotide Mix (10 mM each), 0.5 μl Transcriptor Reverse Transcriptase and made up to 20
μl with RNAase-free water. Reaction mixtures were incubated for 2 min at 25°C, 40 min at 42°C
and 5 min at 95°C. Resultant cDNA was stored at -20°C.

qPCR was performed in an Mx3000P QPCR System (Stratagene) using Roche FS
Universal SYBR Green MasterRox. Briefly, cDNA (equivalent to 6ng of starting RNA) was mixed
with concentrated master mix (2x) and appropriate forward and reverse primers (20nM; Table S6)
and made up to a final volume of 15 μl with RNase-free water. Technical replicates as well as no
template and no RT controls were included in each run. Gene expression levels were normalized
in all cases to Gapdh housekeeping control. Primer sequences used are as follows:

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
</tr>
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<tr>
<td>Human-ARC</td>
<td>F: 5'-CTGAGCCACCTAGAGGATCT-3'. R: 5'-AACTCCACCCAGTTCTTCAGG-3'</td>
</tr>
<tr>
<td>Mouse-Arc</td>
<td>F: 5'-GCTGGGAAGAAATCCCATCAAGGC-3'. R: 5'-ACCTCTCCAGACGTTAGAAGAC-3'</td>
</tr>
<tr>
<td>Gene</td>
<td>Human Forward</td>
</tr>
<tr>
<td>----------</td>
<td>---------------</td>
</tr>
<tr>
<td>BDNF</td>
<td>5’-AGCTGAGCGTG GTGACAGT-3’</td>
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<tr>
<td>CCNH</td>
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</tr>
<tr>
<td>ETS2</td>
<td>5’-CAACTCCTTTTCAGAGATCAGG-3’</td>
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<tr>
<td>FOS</td>
<td>5’-CTACCACTCAACCCGAGACT-3’</td>
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<td>FOSB</td>
<td>5’-GCCGGGAACGAAATAAAACTA-3’</td>
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<td>FOSL2</td>
<td>5’-ACACCCCTGTTTCTCTCCG-3’</td>
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<tr>
<td>GAPDH</td>
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<tr>
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<td>5’-CCTCAAGAGGGCGACAGGTG-3’</td>
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<td>5’-TGCCCTTGGCACCCTGCAACCAG-3’</td>
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<td>PTSS1</td>
<td>5’-GAGGATGGAGTTCTGATCTTGTGGTTGC-3’</td>
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<tr>
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<tr>
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<td>Kcd1</td>
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<tr>
<td>Lmbrd2</td>
<td>5’-CTGGAGAGCGTGGATAGGTA-3’</td>
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</table>

**Statistical analysis, equipment and settings.** Statistical testing of the RNA-seq data is described in that section. Other testing involved a 2-tailed paired Student's t-test, or a one- or two-way ANOVA followed by Sidak’s post-hoc test, as indicated in the legends. Correlation coefficients were calculated assuming a Gaussian distribution of the data. Throughout the manuscript, independent biological replicates are defined as independently performed experiments on material derived from different animals.

**Acknowledgements**

This work was supported by the MRC, the Wellcome Trust, the BBSRC and the Royal Society. SL is a Wellcome Trust Senior Research Fellow, GEH is a MRC Senior Non-Clinical Research Fellow. We thank Eleanor Coffey for supplying us with the TAM-67 vector.
List of Supplemental Files

- Figure_1_Figure_Supplement1.pdf
- Figure_2_Figure_Supplement1.pdf
- Figure_3_Figure_Supplement1.pdf
- Figure_1_Source_Data_1.xlsx
- Figure_1_Source_Data_2.xlsx
- Figure_1_Source_Data_3.xlsx
- Figure_1_Source_Data_4.xlsx
- Figure_1_Source_Data_5.xlsx
- Figure_2_Source_Data_1.xlsx
- Figure_2_Source_Data_2.xlsx
- Figure_2_Source_Data_3.xlsx
- Figure_3_Source_Data_1.xlsx
- Figure_3_Source_Data_2.xlsx
- Figure_3_Source_Data_3.xlsx

References

Correlation of KCl/FPL-induced fold-change in 11,302 ortholog pairs in DIV10 Mus-PRIM CORT-<0.05) induced in Hum-ESC CORT-neurons and DIV10 Mus-PRIM CORT-neurons respectively. Pairs where both are strongly (>5 fold) and significantly (Benjamini-Hochberg-adjusted P-value <0.05, calculated within DESeq2) are highlighted in red.

Expression was significantly altered by KCl/FPL treatment (Benjamini-Hochberg-adjusted p-value 0.05). Figure 1. Conservation and divergence in gene regulation in neurons of human and mouse origin. All data can be found in Supplemental Table 1. A,B) Analysis of gene expression changes induced by KCl/FPL in Hum-ESC CORT-neurons (A) and DIV10 Mus-PRIM CORT-neurons (B). Normalised RNA-seq read density (FPKM) mapping to each gene in RNA extracted from control vs. KCl/FPL-treated neurons is shown (n=3 independent biological replicates). Genes whose expression was significantly altered by KCl/FPL treatment (Benjamini-Hochberg-adjusted p-value <0.05, calculated within DESeq2) are highlighted in red. C) Cohort of human:mouse orthologous pairs where both are strongly (>5 fold) and significantly (Benjamini-Hochberg-adjusted P-value <0.05) induced in Hum-ESC CORT-neurons and DIV10 Mus-PRIM CORT-neurons respectively. D) Correlation of KCl/FPL-induced fold-change in 11,302 ortholog pairs in DIV10 Mus-PRIM CORT-
neurons vs. DIV10-Hum-ESC<sup>CORT</sup>-neurons. **E** Correlation of KCl/FPL-induced fold-change in the same 11,302 genes as in (D) in DIV10 vs. DIV4 Mus-PRIM<sup>CORT</sup>-neurons.

**Figure 2. Stem cell origin does not substantially impact on activity-dependent gene responsiveness.** **A** Example immunofluorescence pictures of Mus-ESC<sup>CORT</sup>-neurons stained for neuronal markers TuJ1 (upper) and Reelin (lower). Note also absence of Nestin staining (upper), a marker of undifferentiated neural precursor cells. Scale=20 µm. **B** Example trace of a burst of action potentials (APs) induced in Mus-ESC<sup>CORT</sup>-neurons by current injection (see methods). **C** Example trace illustrating spontaneous TTX-sensitive AP firing. **D** Example trace illustrating spontaneous TTX-sensitive EPSCs, as well as TTX-insensitive, CNQX-sensitive miniature EPSCs (also see inset; scale bar: 20 pA, 5 ms). Activity returned upon wash out of TTX and CNQX (not shown). **E,F** Correlation of KCl/FPL-induced fold-change in the same 11,302 genes as in Fig. 1d,e in Mus-ESC<sup>CORT</sup>-neurons vs. DIV10 (E) or DIV4 (F) Mus-PRIM<sup>CORT</sup>-neurons. **G** Correlation of KCl/FPL-induced fold-change in 11,302 ortholog pairs in Mus-ESC<sup>CORT</sup>-neurons vs. Hum-ESC<sup>CORT</sup>-neurons. **H** A connection map generated in Cytoscape illustrating the relative determination coefficients (R<sup>2</sup>) between the fold-inductions of the 11,302 genes studied in each of the three biological replicates of the experiments performed in each of the four different neuronal preparations. The thickness of the connecting line and the attractive force of the connecting nodes are both directly proportional to R<sup>2</sup>, against a background of constant inter-node repulsion. Note that all mouse neurons of differing developmental stage and origin (primary vs. ES cell) cluster strongly together, with the Hum-ESC<sup>CORT</sup>-neuronal replicates clustering away from them. **I** A connection map generated as for (H) but illustrating the relative determination coefficients (R<sup>2</sup>) between the basal expression levels (FPKM) of the 11,302 genes studied in each of the three biological replicates of the experiments performed in each of the four different neuronal preparations.

**Fig. 3. DNA sequence is a contributing factor to species-dependent gene responsiveness to neuronal activity.** **A** Analysis of gene expression changes induced by KCl/FPL in mouse Tc1 neurons. Mouse genes are shown in grey, human chromosome-21 (Hsa-21) genes in red. **B** The graph concerns 72 orthologous pairs whose human ortholog is on Hsa-21 and carried by the Tc1 mouse strain, and which meets expression cut-off (100 reads). For each pair, the differential responsiveness index (DRI) was calculated from the species-separated Tc1 neuron RNA-seq data(see text), and plotted against the DRI calculated from separate neuronal preparations (Hum-ESC<sup>CORT</sup>-neurons vs. DIV10 Mus-PRIM<sup>CORT</sup>-neurons, as was done in Figure 1-figure supplement 1e). **C** KCl/FPL-induced fold induction of the human (red) and mouse (blue) orthologs of Hsa-21 gene ETS2, analysed side-by-side in Tc1 neurons. *P*=0.035. unpaired t-test; # indicates P=0.0464, 0.0026 (left to right) vs. unstimulated control (4h, n=7; 24h, n=3). **D** Kinetics of KCl/FPL-induced ETS2 induction in Hum-ESC<sup>CORT</sup>-neurons vs. DIV10 Mus-PRIM<sup>CORT</sup>-neurons. *P*=0.0017, 0.004 (left to right), unpaired 1-way ANOVA vs. control (Hum-ESC<sup>CORT</sup>-neurons: n=3 (2h, 8h), n=6 (4h), P=; DIV10 Mus-PRIM<sup>CORT</sup>-neurons: n=4, P. **E** The indicated firefly luciferase reportersbased on human (Hsa) or mouse (Mmu) ETS2 promoters (See (Right) for schematic) plus a pTK-renilla control were transfected into mouse cortical neurons, treated ± KCl/FPL after which firefly:renilla luciferase ratio was measured, and fold-induction calculated. For comparing the effect of TAM67, a control vector (encoding β-globin) was used. *P=0.0015, 0.0015, 0.0015, 2-tailed paired t-test (n=3). **F** Kinetics of gene regulation of 6 human:mouse ortholog pairs that show quantitative differences in activity-dependent regulation in Hum-ESC<sup>CORT</sup>-neurons vs. all mouse neuronal preparations from prior RNA-seq analyses. Analysis was performed in Hum-ESC<sup>CORT</sup>-neurons vs. DIV10 Mus-PRIM<sup>CORT</sup>-neurons (n=3,4 respectively). At the timepoints studied, three of these pairs exhibit quantitatively higher induction in human neurons, and three stronger repression in human neurons. *P=<0.0001, 0.0004, 0.0005, <0.0001, <0.0001, 0.0073 (top to bottom, 2-way ANOVA, P-value corresponds to the main species effect).
Figure 1-figure supplement 1. A) Example fluo-3 Ca^{2+} imaging trace of KCl/FPL-treated Hum-ESC_CORT-neurons. B) qPCR analysis of the fold induction of the indicated genes in Hum-ESC_CORT-neurons after 4h of KCl/FPL treatment. *P=0.0001, 0.0013, 6E-05, 0.0005, 0.0086, 0.0010, 2-tailed \(t\)-test (n=3). C,D) A comparison of the magnitude of gene induction of a selection of human (C) and mouse (D) genes, as assayed by RNA-seq and by qRT-PCR. In both cases identical sets of 3 independent replicates were assayed and the mean calculated. E) For the 11,302 orthologous pairs, the "differential regulation index" (DRI) was calculated, defined as the fold-change (Hum-ESC_CORT-neurons) divided by the fold-change (DIV10 Mus-PRIM_CORT-neurons). The distribution of the log2 of the DRIs is shown. F) As for (E) except the DRI was calculated for DIV4 vs. DIV10 Mus-PRIM_CORT-neurons. G) Correlation of KCl/FPL-induced fold-change in 11,302 ortholog pairs in DIV4 Mus-PRIM_CORT-neurons vs. Hum-ESC_CORT-neurons.

Figure 2-figure supplement 1. A) Normalised RNA-seq read density (FPKM) mapping to each gene in RNA extracted from control vs. KCl/FPL-treated Mus-ESC_CORT-neurons is shown (n=3 independent biological replicates). Genes whose expression was significantly altered by KCl/FPL treatment (Benjamini-Hochberg-adjusted p-value <0.05, calculated within DESeq2) are highlighted in red. B-D) Correlation of basal gene expression (Log2(FPKM)) across 11,302 ortholog pairs in Hum-ESC_CORT-neurons vs. DIV4 Mus-PRIM_CORT-neurons (B), DIV4 Mus-PRIM_CORT-neurons (C) and Mus-ESC_CORT-neurons (D). E) For each of the 11,302 orthologous pairs, the Log2(DRI) Hum-ESC_CORT-vs. DIV10 Mus-PRIM_CORT-neurons (i.e. DRIs from Figure 1-figure supplement 1e) were plotted against the Log2(DBEI), where DBI (differential basal expression index) is the ratio of basal expression in Hum-ESC_CORT-vs. DIV10 Mus-PRIM_CORT-neurons.

Figure 3-figure supplement 1. A) Correlation of KCl/FPL-induced fold-change in the same 11,302 mouse genes analysed in Figs 1 and 2, in DIV10 Mus-ESC_CORT-neurons from wild-type vs. Tc1 primary cortical neurons, showing the expected high correlation.

Source Data Legends

Figure_1_Source_Data_1. Data set relating to Figures 1a-e.
Figure_1_Source_Data_2. Data set relating to Figure 1-figure supplement 1a.
Figure_1_Source_Data_3. Data set relating to Figure 1-figure supplement 1b.
Figure_1_Source_Data_4. Data set relating to Figure 1-figure supplement 1c-d.
Figure_1_Source_Data_5. Data set relating to Figure 1-figure supplement 1e-f.
Figure_2_Source_Data_1. Data set relating to Figure 2e-g.
Figure_2_Source_Data_2. Data set relating to Figure 2-figure supplement 1e.
Figure_2_Source_Data_3. Data set relating to Figure 2-figure supplement 1b-d.
Figure_3_Source_Data_1. Data set relating to Figure 3-figure supplement 1a.
Figure_3_Source_Data_2. Data set relating to Figure 3b.
Figure_3_Source_Data_3. Data set relating to Figure 3c-f.
Figure 1-figure supplement 1. A) Example fluo-3 Ca$^{2+}$ imaging trace of KCL/FPL-treated Hum-ESC$^{\text{CORT}}$-neurons. B) qPCR analysis of the fold induction of the indicated genes in Hum-ESC$^{\text{CORT}}$-neurons after 4h of KCl/FPL treatment. *P=0.0001, 0.0013, 6E-05, 0.0005, 0.0086, 0.0010, 2-tailed t-test (n=3). C,D) A comparison of the magnitude of gene induction of a selection of human (C) and mouse (D) genes, as assayed by RNA-seq and by qRT-PCR. In both cases identical sets of 3 independent replicates were assayed and the mean calculated. E) For the 11,302 orthologous pairs, the "differential regulation index" (DRI) was calculated, defined as the fold-change (Hum-ESC$^{\text{CORT}}$-neurons) divided by the fold-change (DIV10 Mus-PRIM$^{\text{CORT}}$-neurons). The distribution of the log$_2$ of the DRIs is shown. F) As for (E) except the DRI was calculated for DIV4 vs. DIV10 Mus-PRIM$^{\text{CORT}}$-neurons. G) Correlation of KCl/FPL-induced fold-change in 11,302 ortholog pairs in DIV4 Mus-PRIM$^{\text{CORT}}$-neurons vs. Hum-ESC$^{\text{CORT}}$-neurons.
Figure 2-figure supplement 1. A) Normalised RNA-seq read density (FPKM) mapping to each gene in RNA extracted from control vs. KCl/FPL-treated Mus-ESC<sup>CORT</sup>-neurons is shown (n=3 independent biological replicates). Genes whose expression was significantly altered by KCl/FPL treatment (Benjamini-Hochberg-adjusted p-value <0.05, calculated within DESeq2) are highlighted in red. B-D) Correlation of basal gene expression (Log<sub>2</sub>(FPKM)) across 11,302 ortholog pairs in Hum-ESC<sup>CORT</sup>-neurons vs. DIV4 Mus-PRIM<sup>CORT</sup>-neurons (B), DIV4 Mus-PRIM<sup>CORT</sup>-neurons (C) and Mus-ESC<sup>CORT</sup>-neurons (D). E) For each of the 11,302 orthologous pairs, the Log<sub>2</sub>(DRI) Hum-ESC<sup>CORT</sup>-vs. DIV10 Mus-PRIM<sup>CORT</sup>-neurons (i.e. DRIs from Figure 1-figure supplement 1e) were plotted against the Log<sub>2</sub>(DBEI), where DBI (differential basal expression index) is the ratio of basal expression in Hum-ESC<sup>CORT</sup>-vs. DIV10 Mus-PRIM<sup>CORT</sup>-neurons.
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