

# Reactive astrocytes in ALS display diminished intron retention

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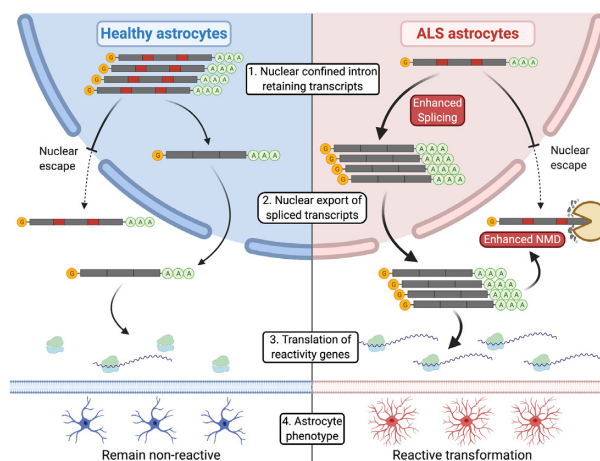
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## ABSTRACT

Reactive astrocytes are implicated in amyotrophic lateral sclerosis (ALS), although the mechanisms controlling reactive transformation are unknown. We show that decreased intron retention (IR) is common to human-induced pluripotent stem cell (hiPSC)-derived astrocytes carrying ALS-causing mutations in *VCP*, *SOD1* and *C9orf72*. Notably, transcripts with decreased IR and increased expression are overrepresented in reactivity processes including cell adhesion, stress response and immune activation. This was recapitulated in public-datasets for (i) hiPSC-derived astrocytes stimulated with cytokines to undergo reactive transformation and (ii) *in vivo* astrocytes following selective deletion of TDP-43. We also re-examined public transcriptome sequencing (TRAP-seq) of astrocytes from a *SOD1* mouse model, which revealed that transcripts upregulated in translation significantly overlap with transcripts exhibiting decreased IR. Using nucleocytoplasmic fractionation of *VCP* mutant astrocytes coupled with mRNA sequencing and proteomics, we identify that decreased IR in nuclear transcripts is associated with enhanced nonsense mediated decay and increased cytoplasmic expression of transcripts and proteins regulating reactive transformation. These findings are consistent with a molecular model for reactive transformation in astrocytes whereby poised nuclear reactivity-related IR transcripts are spliced, undergo nuclear-to-cytoplasmic translocation and translation. Our

study therefore provides new insights into the molecular regulation of reactive transformation in astrocytes.

## GRAPHICAL ABSTRACT



## INTRODUCTION

Amyotrophic lateral sclerosis (ALS) is a rapidly progressive and incurable neurodegenerative disease. The pathological hallmark of ALS is cytoplasmic aggregation of the RNA-binding protein, transactive response DNA-binding protein 43 (TDP-43, encoded by *TARDBP*) accompanied by loss of nuclear TDP-43, in the motor neurons of >97% of ALS cases (1). Since the identification of the *SOD1* gene mutation in ALS, which accounts for ~15% of familial cases, ~30 gene mutations have been discovered of

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which *C9orf72* is the most common (~35% of familial cases) (2,3). Valosin-containing protein (*VCP*) is a transitional endoplasmic reticulum ATPase essential for maturation of ubiquitin-containing autophagosomes, and accounts for ~2% of familial ALS (4). Although motor neuron degeneration was traditionally regarded as the cellular substrate of ALS pathogenesis, accumulating evidence implicates astrocytes, which play vital roles in providing neuronal support including metabolite provision and modulating the immune response (5–7). ALS astrocytes can non-cell autonomously impair motor neuron function through loss of supportive and/or gain of toxic factors, and recent studies additionally revealed cell autonomous astrocyte phenotypes (8–17). ALS postmortem studies consistently report neuroinflammation and reactive gliosis (18–23). Neuroinflammation induces quiescent astrocytes to undergo dramatic changes in gene expression, morphology and function, called ‘reactive transformation’ (24–26). Using inflammatory cues to mimic activated microglia, Liddel *et al.* profiled astrocyte gene expression changes accompanying reactive transformation and categorized them into pan-reactive, A1 (harmful) and A2 (protective) responses (24,27–28).

Although gene expression changes accompanying reactive responses have been extensively studied, the regulating molecular mechanisms are unknown (24,26–27,29). Alternative splicing, particularly intron retention (IR), is recognized as a major regulator of gene expression during both differentiation and activation of several cell types (30–36). Although other modes of alternative splicing (exon skipping and alternative splice site usage) typically enhance protein synthesis, IR can serve to suppress translation—introns often contain premature termination codons promoting degradation via nonsense-mediated decay (NMD) (36,37). IR can also regulate RNA localization by promoting nuclear confinement, further suppressing translation (31–32,38–40). We recently reported aberrant IR in human-induced pluripotent stem cell (hiPSC)-derived ALS motor neuron cultures undergoing differentiation (41,42). IR has also been implicated in the function of other neuronal and immune cells—orchestrated IR regulates activation of macrophages, granulocytes and lymphocytes—however, its role in astrocytes is unknown (30,33–34,36,42–47).

We sought to determine if alternative splicing, and specifically IR, plays a role in astrocyte reactivity in ALS. Using RNA sequencing from a variety of sources (Supplementary Table S1), including enriched hiPSC-derived astrocytes harbouring mutations in *VCP*, *SOD1* and *C9orf72*, we found the loss of thousands of IR transcripts that correlates with increased abundance of their cognate spliced transcripts and are enriched for astrocyte reactivity processes. This decreased IR signature was recapitulated in both astrocytes stimulated with inflammatory cues to undergo reactive transformation as well as *in vivo* astrocytes following selective deletion of *TDP-43*, which itself triggered reactive transformation. We additionally found that a significant number of reactivity genes with reduced IR in ALS astrocytes also display increased translation. Using nucleocytoplasmic fractionation, we revealed that *VCP* mutant astrocytes have fewer IR transcripts in the nucleus, coupled with enhanced NMD of cytoplasmic IR transcripts. This was associated with increased cytoplasmic spliced mRNA and

protein that are reactivity-related, suggesting nuclear-to-cytoplasmic translocation and translation of spliced reactivity transcripts. Cumulatively this work identifies a physiological and orchestrated IR programme in healthy astrocytes, which is lost in ALS and coincides with an increased abundance of reactivity regulators, thus potentially contributing to astrocyte reactive transformation.

## MATERIALS AND METHODS

### Human-induced pluripotent stem cell derived astrocytes

hiPSCs were maintained using standard protocols and were differentiated into astrocytes as described previously, generating highly enriched (>90%) populations of astrocytes (Supplementary Figure S1A) (8,11,48–51). hiPSCs were maintained on Geltrex (Life Technologies) with serum-free Essential 8 Medium media (Life Technologies), and passaged using EDTA. After neural conversion (7 days in a chemically defined medium containing 1  $\mu$ M Dorso-morphin (Millipore), 2  $\mu$ M SB431542 (Tocris Bioscience) and 3.3  $\mu$ M CHIR99021 (Miltienyi Biotec), neural precursors were patterned for 7 days with 0.5  $\mu$ M retinoic acid and 1  $\mu$ M purmorphamine, followed by a 4-day treatment with 0.1  $\mu$ M purmorphamine. After a propagation phase (60–120 days) with 10 ng/ml FGF-2 (Peprotech), astrocytes were terminally differentiated in presence of BMP4 (10 ng/ml, R&D) and LIF (10 ng/ml, Sigma-Aldrich) for 30 days. Informed consent was obtained from all patients and healthy controls in this study. Experimental protocols were all carried out according to approved regulations and guidelines by UCLH’s National Hospital for Neurology and Neurosurgery and UCL’s Institute of Neurology joint research ethics committee (09/0272).

### Nuclear and cytoplasmic RNA purification

Subcellular fractionation was achieved using the Ambion PARIS kit (ThermoFisher Scientific) following the manufacturer’s protocol. The cytosolic fraction was obtained by lysing cells in ice-cold cell fractionation buffer for 5 min, disrupting plasma membranes while leaving nuclear membranes intact. Lysates were centrifuged for 3 min at 500  $\times$  g at 4°C. Supernatant was further centrifuged at maximum speed at 4°C for 1 min, and the resulting supernatant was processed as the cytosolic fraction. Nuclear pellets from the first centrifugation step were gently washed with cell fractionation buffer and then lysed on ice for 30 min in 8 M Urea Nuclear Lysis Buffer. The resulting nuclear fraction was homogenized using a QIAshredder (Qiagen) to shred chromatin and reduce viscosity. Both lysis buffers were supplemented with 0.1 U/ $\mu$ l RiboLock RNase Inhibitor (ThermoFisher Scientific) and HALT Protease Inhibitor Complex (ThermoFisher Scientific).

### RNA isolation and RT-qPCR

The Promega Maxwell RSC simplyRNA cells kit including DNase treatment, alongside the Maxwell RSC instrument, was used for RNA extractions. The Nanodrop was used to assess RNA concentration and the 260/280 ratio, and the Agilent TapeStation was used to assess quality. All RNA

samples have RIN value > 8.4. RT-qPCR was performed on cDNA generated from 50 ng DNaseI-treated total RNA using SuperScript® IV First-Strand Synthesis System (Invitrogen) and random hexamers, according to the manufacturers' instructions. RT-qPCR reactions were performed in 10 µl volumes containing 1× SYBR® Green Mastermix (Bio-Rad) and 0.5 µM of the respective forward and reverse primers. Samples were amplified and analysed using the CFX96™ Real Time PCR Machine (Bio-Rad). Cycling conditions were: 50°C for 2 min, 95°C for 2 min, followed by 40 cycles at 95°C for 15 s, then 60°C for 60s. Samples were run in triplicate and all programs contained a melt curve and a no template control. The absence of contaminating gDNA was confirmed by PCR on negative RT samples. Fold change was calculated using the  $C_T$  method.

### RNA sequencing

Poly(A)+ selected reverse stranded RNA sequencing libraries were prepared from whole astrocyte and their nuclear and cytoplasmic fractions obtained from astrocyte differentiation from three control and two VCP mutant cell lines using the KAPA mRNA HyperPrep Library kit for Illumina®, with 50 ng of total RNA as input. Libraries were sequenced on a HiSeq 4000 platform at the Francis Crick Institute. One control sample failed quality control and was discarded. A total of 129 million 100 bp-long paired-end strand-specific reads were sequenced per sample split over five lanes. All libraries generated in this study had <1% rRNA, <1% mtDNA, >90% strandedness and >70% exonic reads (Supplementary Table S1).

### Proteomics

Protein samples from nuclear and cytoplasmic fractions were reduced, alkylated and acetone precipitated overnight. Each protein pellet was resuspended in 1 M guanidine hydrochloride and 100 mM HEPES. Proteins were tryptically digested overnight at 37°C with mixing. Digested samples were acidified then stored at −80°C. Each sample was split into triplicates and loaded onto prepared Evotips. Samples were analysed using Evosep 15 cm column and an Orbitrap Fusion mass spectrometer operating in data-dependent acquisition mode. A 44-minute universal (OT/IT) method was used. Raw files were analysed with MaxQuant v1.6.12.0 using the LFQ algorithm against a 2020 SwissProt *Homo sapiens* protein database.

### Western blotting

Protein lysates were loaded onto NuPAGE, 4–12%, Bis-Tris protein gels (Invitrogen) and subjected to electrophoresis before being transferred onto nitrocellulose membrane (Biorad). Blocking was performed in PBS, 0.1% tween, 5% bovine serum albumin for 1 h at room temperature and incubated overnight with mouse anti  $\alpha$ -Tubulin (1:1000, Sigma T5168) and rabbit anti-histone 3 (1:1000, Abcam ab179). Following washes, membranes were incubated for 1 h at room temperature with fluorescent secondary anti-rabbit and anti-mouse antibody (1:10000, LI-COR IRDye). Membranes were imaged using Odyssey Fc Imaging System (LI-COR).

### Statistical analyses

Raw mRNA sequencing reads were aligned to hg38 human reference genome using splice-aware aligner, STAR v2.6.1 (52). Aligned reads were quantified with HTSeq v0.12.4 (53) using intersection-strict mode to enable counting of spliced transcripts at gene-level based on Ensembl GRCh 38.99 annotation. Detailed quality control of the raw RNAseq was assessed utilizing the nf-core/rna-seq pipeline (54) (Supplementary Tables S1, S3). Differential gene expression was measured using DESeq2 (55) in R v4.0.3. Results were generated by comparing the mutant (or treated, or cytoplasmic fraction) versus control (or untreated, or nuclear fraction) and a gene was considered significantly differentially expressed if false discovery rate (FDR) < 0.05.

Alternative splicing was quantified using both a coverage based tool, *VAST-TOOLS* (56) and a splice-junction tool, *SGSeq* (57). IR focused analysis was quantified using *IRFinder*, (58) which measures IR levels using the IR ratio metric (intron read depth divided by the sum of the intron and flanking exon read depth, Figure 1B). *IRFinder* addresses specific peculiarities in IR quantification including: low complexity regions, non-poly(A)+ RNA and DNA contamination, and overlapping exons. To improve consistency in the quantification of IR only 'clean' events are included, filtering out spurious IR events with reliability warnings LowCover (spliced reads mapping across the 3' and 5' flanking exons + intron reads > 10) and LowSplicing (> 4 reads from across the 3' and 5' flanking exons) in the *IRFinder-IR-dir.txt* output. Differential IR was quantified using *analysisWithLowReplicates.pl* from *IRFinder* between mutant and control, using the Audic and Claverie test and threshold of  $P < 0.05$  (uncorrected for multiple testing). To assess the relationship between differential gene expression and IR, we assigned the  $\log_2$ foldchange in its gene expression (mutant versus control) to the  $\Delta$ IR ratio for each reliable differential IR event. To identify enriched functional pathways, Gene Ontology (GO) enrichment was performed using *g:Profiler2* (59). Significantly over-represented (FDR < 0.05) up and downregulated IR genes were grouped according to their differential gene expression directional change and these subsets were used as input, with all measured genes used as background. In the bar charts the top significant GO terms were manually curated by removing redundant terms. Gene set enrichment analysis (GSEA) was performed using the *fgsea* R package (60) on the transcriptomic signature gene sets for NMD components (61) and Liddelow *et al.* reactivity genes (24). Overlapping gene lists between datasets were tested using the Fisher's exact test. Similarly, Liddelow *et al.* reactivity genes with reduced IR were tested for overlap with genes increased in expression using the Fisher's exact test. For Liddelow *et al.* genes with multiple reliable  $\Delta$ IR ratios, we calculated the genes mean  $\Delta$ IR ratio, normalizing for each retained introns length.

Features of retained and spliced introns (length, GC content, PhastCons conservation score) were analysed as reported previously (36,42). To identify RBPs that bind to aberrantly retained introns, we examined iCLIP data for 21 RBPs (62), and eCLIP data from HepG2 cells for 112 RBPs available from ENCODE (63). Relative enrichment for each



of the RBPs was obtained by calculating the proportion of crosslink events mapping to retained introns compared with non-retained introns of the same genes. Maximum Entropy splice site scoring software (MaxEntScan) was used to predict the splice site strength for the 5' splice sites (5' ss) and 3' splice sites (3' ss) (64). The 5' ss score uses 9-bases (last 3 bases of the upstream exon and first 6 bases of the intron), whereas the 3' ss score uses 23-bases (last 20 bases of intron and first 3 bases of the downstream exon). We determined the MaxEntScan score for retained introns, using all 5' ss (exon–intron junctions) and 3' ss (intron–exon junctions) of annotated introns, using the *GencoDymo* R package. NMD probability was predicted by searching for premature termination codons in the open reading frame of transcripts containing retained introns, using *GeneStructureTools* and *not-NMD* bioconductor packages. Differential transcript-level isoform expression analysis was performed with Kallisto (65) and transcript biotypes were annotated using the Ensembl biotype classification to enable comparison between NMD and protein coding annotated transcripts.

For comparison with *SOD1* and *C9orf72* mutant hiPSC astrocytes, we downloaded RNAseq data from Tyzack *et al.* (8) (GSE102902 and GSE99843) and Birger *et al.* (66) (GSE142730). *SOD1*<sup>G37R</sup> mouse spinal cord astrocyte TRAP-Seq was downloaded from Sun *et al.* (67) (GSE74724). For comparison with hiPSC stimulated reactive astrocyte dataset, we downloaded bulk RNAseq from Barbar *et al.* (27) (synapse ID: 21861229). For TDP-43 deleted mouse astrocytes, we accessed bulk RNAseq from Peng *et al.* (68) (GSE156542). RNAseq from these studies were processed using the same pipeline as described above and quality control metrics are reported in Supplementary Table S1. Sequencing depth was on average ~50 million reads per sample, which ranged from 10 million in *SOD1* to 129 million in *VCP* mutant astrocytes. Lower sequencing depth limits the ability to detect IR; however, using the stable IRFinder algorithm, we were able to identify reliable IR events even at 10 million albeit at lower absolute numbers (Supplementary Figure S3) (8,69). Mass spectrometry proteomics data were analysed using DEP package (v1.11.0) (70) on MaxQuant results. Data were filtered, normalized and imputed using default parameters. Differential enrichment analysis was performed using protein-wise linear models combined with Bayes statistics that utilizes *limma*. A protein was considered significantly differentially expressed when  $P < 0.05$  (uncorrected for multiple comparisons). Schematics were created with *BioRender.com*. All error bars shown represent either the standard error of the mean (SEM) or 1.5 times the interquartile range, from independent experiments.

## RESULTS

### Intron retention is decreased in ALS astrocytes across *VCP*, *SOD1* and *C9orf72* mutants

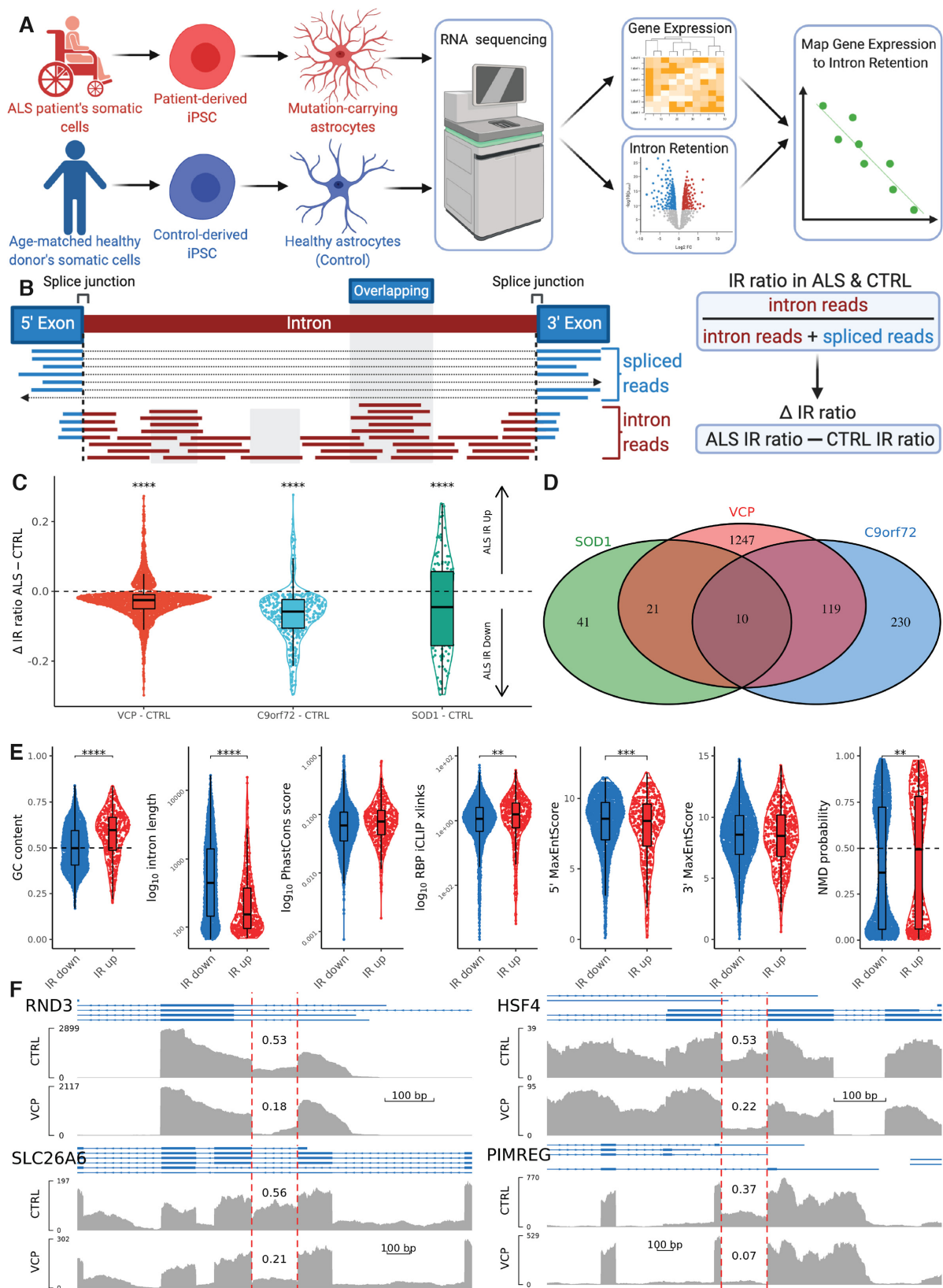
To identify transcriptome-wide changes in ALS astrocytes, we performed RNA sequencing on poly(A)<sup>+</sup> selected mRNA libraries isolated from highly enriched and functional hiPSC-derived astrocytes using our previously established platform (Supplementary Figure S1A) (8,11,49–50,71). Specifically, we first examined RNA sequencing

from two lines derived from two patients with ALS-causing *VCP* gene mutations (p.R155C and p.R191Q) and two healthy control lines (Figure 1A) (8,11,49). To improve confidence in detecting differential splicing, libraries were deeply sequenced to ~130 million reads per sample (Supplementary Table S2). Principal component analysis and unsupervised hierarchical clustering demonstrated that samples segregated based on genotype (Supplementary Figure S1B–D). We first defined differences between *VCP* mutant and control astrocyte gene expression patterns and found 170 differentially expressed genes (FDR < 0.05), which is 11-fold more than what we previously established in terminally differentiated *VCP* mutant motor neurons (41). Differentially expressed genes were enriched for developmental, structural and nervous system processes (Supplementary Figure S1E and Supplementary Table S3).

We sought to examine alternative splicing in hiPSC-derived astrocytes, comparing those carrying *VCP* mutations to control counterparts. After examining all modes of alternative splicing and finding that only IR was significantly different between *VCP* mutant and control astrocytes (Supplementary Figure S2A and B), we focused further on IR quantification (Figure 1B) (58). Across both conditions, we found 16,317 reliable unique retained introns (IR ratio > 0.1 i.e. intron abundance is  $\geq 10\%$  of the sum of intron + spliced abundance); however, there were 14.1% fewer retained introns in *VCP* mutant compared with control (12,169 versus 14,162, respectively, Supplementary Figure S3A). Differential IR comparison revealed 2,309 introns with reliable coverage from 1,699 genes that were retained at significantly different levels ( $P < 0.05$ , Audic and Claverie test, Supplementary Figure S3B), with 1,885 (81.6%) displaying decreased IR in *VCP* (Figure 1C, 1F;  $P = 2.39 \times 10^{-68}$ ), confirming reduced IR in *VCP* mutant astrocytes.

To assess whether IR was also reduced in other ALS mutant astrocytes, we analysed published *C9orf72* (66) and *SOD1* (8) mutant astrocyte RNA sequencing datasets (*C9orf72* dataset: two mutant and two control lines; *SOD1* dataset: one mutant and two control lines) (Supplementary Table S1). In both datasets we identified dramatically less retained introns in ALS mutants compared with control (*C9orf72*: 7,925 versus 14,937 [47% less, Supplementary Figure S3C]; *SOD1*: 3,602 versus 7,842 [54% less, Supplementary Figure S3E] respectively, Figure 1C). In *C9orf72* astrocytes, 448 introns in 376 genes were significantly differentially retained, of which 93% (416/448) were decreased in *C9orf72* relative to control (Supplementary Figure S3D). In *SOD1* mutant astrocytes, 139 introns in 131 genes showed significant differential retention of which 59% (82/139) were reduced in *SOD1* mutants compared with control (Supplementary Figure S3F). These findings from across five ALS mutant and six control lines, when taken together, strongly indicate that decreased IR in astrocytes is common across diverse genetic forms of ALS.

To investigate whether ALS mutant astrocytes exhibit decreased IR within the same genes, we overlapped genes with IR events between *VCP*, *C9orf72* and *SOD1* mutants. Comparing genes exhibiting decreased IR between *VCP* and *C9orf72* datasets, revealed a significant overlap of 129 / 347 (37.2%,  $P = 2.0 \times 10^{-118}$ , Fisher's exact test; Figure 1D). Likewise, *VCP* and *SOD1* mutants, shared 31 / 76 (40.1%,



**Figure 1.** RNA-sequencing reveals loss of intron retention in ALS astrocytes. (A) Schematic depicting human-induced pluripotent stem cells (hiPSC) astrocyte differentiation and fractionation in ALS (red) and control (blue). Astrocytes were subjected to RNA-sequencing and were analysed for differential gene expression and alternative splicing, with a focus on intron retention. We then mapped the spliced-transcript expression and intron retention values of each gene. See Supplementary Figure S1A for stepwise differentiation strategy of iPSC-derived astrocytes. Two hiPSC lines were obtained from two ALS

$P = 9.0 \times 10^{-31}$ ) genes with decreased IR (Supplementary Table S4). Ten genes with decreased IR were common to all three mutations (chi-squared  $P = 0.005$ ), of which six are relevant to reactivity: *FLNA*, *PLOD3*, *MRC2*, *ACTN4*, *GPS2* and *MCAM* (28,72). This significant overlap between independent ALS datasets recapitulates our observation in *VCP* mutant astrocytes, suggesting that diminished IR of astrocyte reactivity genes is generalizable across ALS mutations.

Consistent with prior studies, retained introns in all three datasets (i.e. *VCP*, *C9orf72* and *SOD1* mutant ALS astrocytes) exhibited significantly shorter lengths, higher GC content, higher conservation score, higher predicted binding affinities to RNA-binding proteins (RBPs) and lower splice site strength compared with spliced introns (Supplementary Figure S4A) (36,73–74). Introns with significantly decreased retention in any of the three ALS mutants exhibited longer lengths, lower GC content, and lower RBP-binding affinities but similar conservation scores compared with gained introns (Figure 1E) (63,75). Although 3' splice site strengths were similar between increased and decreased IR events, the 5' splice site strengths were higher in decreased IR events, consistent with increased activity of the spliceosome machinery in ALS astrocytes (64). Scanning for premature termination codons (PTC) within the open reading frame of intronic sequences revealed that the probability of NMD was significantly lower in IR transcripts decreased in ALS compared to IR transcripts increased in ALS. These observations were generally consistent between *VCP*, *C9orf72* and *SOD1* mutants (Supplementary Figure S4B) and confirm previous studies demonstrating that *cis* features define an 'IR code' (36,42,45).

### Decreased intron retention and increased expression of astrocyte reactivity regulators is generalizable across ALS astrocytes

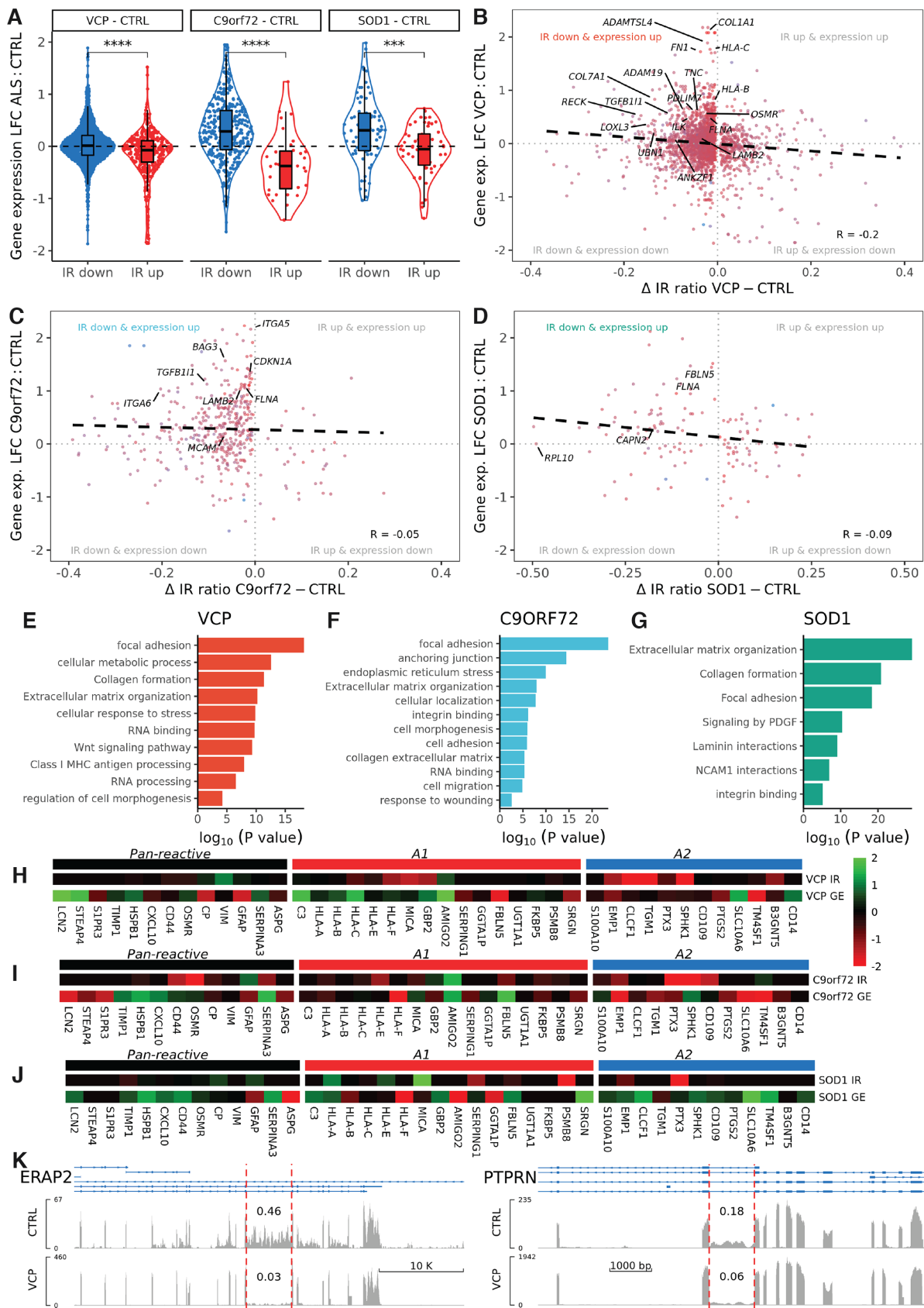
To understand the consequences of decreased IR in ALS astrocytes on gene expression, for each gene we mapped levels of IR to expression (Figure 1A). Previous studies indicate that decreased IR can lead to increased gene expression through avoiding NMD (30,76–77). Comparing changes in IR with changes in gene expression, we found that genes with decreased IR had higher expression compared with

genes with increased IR in *VCP* mutant versus control astrocytes (*t*-test  $P = 2.6 \times 10^{-11}$ ; Figure 2A). Correlating changes in IR with changes in gene expression between *VCP* mutant and control revealed a negative association (Pearson  $R = -0.20$ , Figure 2B), consistent with a relationship between transcripts with decreased IR and increased expression of their spliced isoforms. Of 817 genes with increased expression and exhibiting a reliable IR event, 702 (85.9%) exhibited decreased IR in *VCP* mutant astrocytes. Restricting to genes with significantly increased expression in *VCP* mutant astrocytes (FDR < 0.05) and a reliable IR event, revealed that all four were also significantly decreased in IR (*ERAP2*, *PTPRN*, *IL20RB*, and *KCNE4*; Figure 2K). Using gene ontology analysis, we found genes with reduced IR and increased expression in *VCP* mutant astrocytes were over-represented in pathways associated with cell adhesion (e.g. *TGFB11*, *COL7A1*, *TNC*, *RECK*, *ERAP2*), stress response (e.g. *STX4*, *PTPRN*, *SLC26A6*, *NUP199*, *HSF4*) and immune activation (e.g. *LOXL3*, *NFKB2*, *HLA-B* and *HLA-C*) - processes involved in astrocyte reactivity (Figure 2E) (78–80). Conversely, genes with increased IR and increased expression in *VCP* mutant astrocytes were enriched for enzyme activity rather than reactivity related terms (Supplementary Figure S5A). Of the Liddelou *et al.* (24) astrocyte reactivity genes with a reliable IR event, 21/31 were decreased in IR and 15/31 were increased in expression, while 10/31 exhibited both (32%; overlap  $P = 0.056$ ; Figure 2H). Taken together, these data suggest an inverse relationship between IR and expression, with *VCP* mutant astrocytes exhibiting decreased IR and increased expression of genes regulating astrocyte reactivity.

Similar to *VCP* mutant astrocytes, for both *C9orf72* and *SOD1* mutants, genes with decreased IR exhibited significantly increased gene expression, whereas genes with increased IR correlated with decreased expression (*C9orf72*:  $P = 4.5 \times 10^{-8}$ ; *SOD1*:  $P = 1.5 \times 10^{-4}$ ; Figure 2A). Additionally, both *C9orf72* and *SOD1* mutants exhibited a negative correlation between changes in IR and changes in gene expression (*C9orf72*  $R = -0.05$ ; *SOD1*  $R = -0.09$ ; Figure 2C and D). Gene ontology analysis revealed transcripts with decreased IR and increased expression were also over-represented in astrocyte reactivity pathways (Figure 2F and G; Supplementary Figure S5B and C). Of the Liddelou *et al.*

patients with *VCP* mutations (R155C and R191Q) and two healthy controls. (B) Schematic of IR quantification and differential IR calculation between mutant and control. Median intron coverage is calculated after excluding non-unique multi-mapping reads, overlapping exons (blue) and outliers (highest and lowest 30%) depicted by the shaded regions. Exon and intron read abundance is normalized for feature length. IR ratio is calculated as intron abundance divided by the sum of intron and spliced abundance. Reliable IR event expression is defined as (1) spliced reads mapping across the 3' and 5' flanking exons + intron reads > 10 and (2) > 4 reads from across the 3' and 5' flanking exons. Delta IR ratio is calculated as mutant minus control IR ratio. Adapted from Wong *et al.* (36) and Middleton *et al.* (58). (C) Violin plot showing delta IR ratio in ALS mutants control across differential intron retention events. Intron retention was significantly downregulated in *VCP* (red,  $n = 2309$ ), *C9orf72* (light blue,  $n = 448$ ) and *SOD1* (green,  $n = 139$ ) astrocytes versus CTRL (one-sample *t*-test, \*\*\*\* represents  $P < 0.0001$ ). (D) Venn diagram showing the number of overlapping genes with decreased intron retention events in *VCP* (red), *C9orf72* (blue) and *SOD1* (green). Gene overlap exhibiting decreased IR between *SOD1* and *VCP* datasets was 31/76 (40.1%,  $P = 1.8 \times 10^{-18}$ , hypergeometric test). *C9orf72* and *VCP* overlap was 129 / 347 (37.2%,  $P = 1.8 \times 10^{-67}$ ). Overlapping all three mutants chi-squared  $P = 0.005$ . (E) Violin plots showing intron characteristics (GC content; intron length; phastCons conservation score; iCLIP crosslinking with RBPs; 5' and 3' Maximum Entropy Scores); and nonsense mediated decay (NMD) probability of upregulated versus downregulated retained introns in ALS mutants (*VCP*, *C9orf72* and *SOD1*) versus control astrocytes (IR down,  $n = 2339$ ; IR up,  $n = 490$ ; \*\*\*\* represents adjusted  $P$  values from Wilcoxon test < 0.0001, \*\*\* < 0.001, \*\*  $P < 0.01$ ). For intron characteristics of individual ALS mutants see Supplementary Figure S4B. (F) Coverage plots of reads per million from *VCP* mRNA sequencing reads for *RND3*, *SLC26A6*, *HSF4* and *PIMREG* genes. Within each are alignment tracks for CTRL (top) and *VCP* (bottom) astrocytes. Ensembl transcript annotation tracks are shown above the alignment tracks with exons (thick blue boxes) and introns (thin blue arrowed lines). Vertical red dashed lines show the retained intron of interest and the IR ratio is labelled above the retained intron. Y-axis limits for each track are auto-adjusted to the peak exon coverage so that the target feature fills the view and enables easier visual comparison of IR ratios between conditions.





**Figure 2.** Intron retention is associated with increased expression of genes involved with astrocyte reactivity. (A) Violin plots showing log2 fold change in gene expression (Y-axis) in downregulated IR events (blue) and upregulated IR events (red) for VCP (IR down,  $n = 1885$ ; IR up,  $n = 424$ ), C9orf72

*al.* (24) astrocyte reactivity genes exhibiting a reliable IR event, we observed decreased IR and increased expression in 44% (11/25, overlap  $P = 0.0002$ ) for *C9orf72* mutants and 42% (8/19, overlap  $P = 0.0002$ ) for *SOD1* mutants (Figure 2I and J). Comparing genes exhibiting decreased IR and increased expression in *SOD1* mutants with *VCP* mutants revealed a significant overlap of 18/55 (33%,  $P = 1.1 \times 10^{-21}$ , hypergeometric test; Supplementary Figure S5D). Likewise, of the 251 genes exhibiting decreased IR and increased expression in *C9orf72*, 70 (28%) were also found in *VCP* mutants ( $P = 2.7 \times 10^{-75}$ ; Supplementary Table S5). Six genes with decreased IR and increased expression were common to all three mutations (chi-squared  $P = 0.008$ ), of which four are directly relevant to reactivity: *FLNA*, *PLOD3*, *MRC2* and *ACTN4* (28,72). This significant overlap between independent ALS datasets recapitulates our observation in *VCP* mutant astrocytes, demonstrating that diminished IR and enhanced expression of astrocyte reactivity genes are generalizable across ALS mutations.

### Stimulating reactive transformation using inflammatory cues recapitulates decreased intron retention and increased expression of reactivity genes

Having established that astrocytes with a range of familial ALS mutations exhibit reduced IR, we sought to determine if artificially stimulating astrocyte reactivity with inflammatory cytokines (TNF $\alpha$ , IL-1 $\alpha$  and C1q) was sufficient to induce the same effect (Figure 3A) (27). By analysing RNA sequencing from Barbar *et al.* hiPSC-derived astrocytes stimulated to undergo reactive transformation (three reactive and three untreated lines) (27), we identified 11% less retained introns in astrocytes stimulated with TNF $\alpha$ , IL-1 $\alpha$  and C1q compared with untreated (basal) astrocytes (2,143 versus 2,402, respectively; Supplementary Figure S3G). Differential IR analysis revealed 578 introns from 430 genes that were retained at significantly different levels, with 304 (53%) being reduced in cytokine-stimulated astrocytes (Figure 3D, S3H; one-sample *t*-test  $P = 0.001$ ), indicating reduced IR in hiPSC-derived reactive astrocytes. Of the 240 genes with decreased IR in cytokine-stimulated astrocytes, 86 (35.8%) also exhibited decreased IR in the ALS astrocytes (overlap  $P = 3.8 \times 10^{-71}$ ; Supplementary Table S4). Taken together, we find that IR is prevalent in healthy quiescent hiPSC-derived astrocytes but that ALS astrocytes (with *VCP*, *SOD1* and *C9orf72* mutations) as well as astro-

cytes stimulated using TNF $\alpha$ , IL-1 $\alpha$ , and C1q, share a common decreased IR signature.

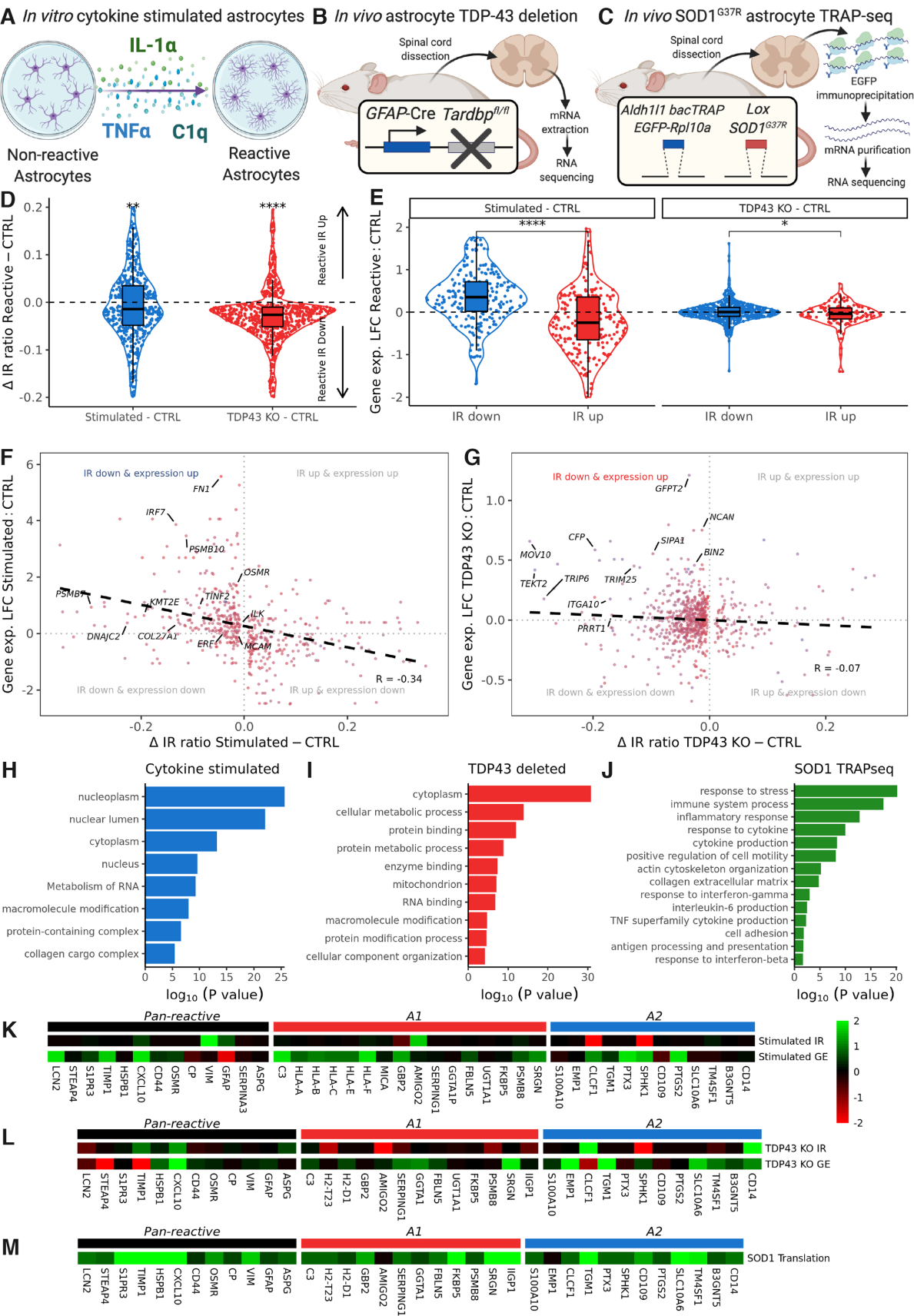
By comparing gene expression with IR after cytokine-stimulation in control astrocytes, we observed that genes with decreased IR in stimulated astrocytes had significantly higher gene expression ( $P < 2.22 \times 10^{-16}$ ; Figure 3E). Furthermore, IR events with greater decreases in the IR ratio exhibited larger increases in gene expression ( $R = -0.34$ , Figure 3F). Enrichment analysis of genes with decreased IR and increased expression reveals that they are over-represented in pathways associated with cellular compartments, RNA metabolism and collagen processing, analogous to processes implicated in ALS astrocytes (Figure 3H). Of the Liddelow *et al.* (24) reactivity genes with a reliable IR event, 11/32 (34%) exhibited both decreased IR and increased expression (overlap  $P = 0.005$ , Figure 3K). Comparing genes with decreased IR and increased expression between stimulated astrocytes and ALS astrocytes revealed an overlap of 45/189 (23.8%, overlap  $P = 4.2 \times 10^{-40}$ ), of which 17 are directly relevant to astrocyte reactivity (Supplementary Table S5) (8,26,81). This indicates that stimulating astrocyte reactivity via established pro-inflammatory cues reproduces reduced IR and increased expression in similar reactivity related genes as ALS astrocytes.

### *In vivo* astrocytes with selective deletion of TDP-43 exhibit decreased intron retention

To further explore the molecular causes associated with decreased IR in ALS astrocytes, we next investigated the role of TDP-43 in this context given its salience as a pathological hallmark in motor neurons. To address this, we leveraged RNA sequencing data from a recent *in vivo* study that demonstrated A1-like reactive transformation in astrocytes upon TDP-43 depletion (68). We sought to determine if this mouse spinal cord astrocyte-specific *TARDBP* knockout (TDP-43 deletion) influenced IR (four knockout and four control mice; Figure 3B). This revealed 11% less retained introns in TDP-43 deleted astrocytes compared with control (12,716 versus 14,338 respectively; Supplementary Figure S3I). Differential IR analysis revealed 700 introns from 643 genes that were retained at significantly different levels ( $P < 0.05$ , Supplementary Figure S3J), with 581 (83%) being decreased (Figure 3D; one-sample *t*-test  $P = 1.10 \times 10^{-20}$ ), indicating that astrocyte TDP-43 deletion induces a global reduction in IR. Comparing genes with decreased IR

(IR down,  $n = 416$ ; IR up,  $n = 32$ ), *SOD1* (IR down,  $n = 82$ ; IR up,  $n = 57$ ). Student's *t*-test was used to determine significance (\*\*\*\* indicates  $P < 0.0001$ , \*\*\*  $P < 0.001$ ). (B–D) Scatterplots of the  $\Delta$  IR ratio (ALS minus control, X-axis) against log<sub>2</sub> fold change in gene expression (Y-axis) in *VCP* (B), *C9orf72* (C) and *SOD1* (D) versus control astrocytes. Points are coloured by mean gene expression across mutant and control (red = high, blue = low expression). Black dashed line indicates linear regression correlation (VCP pearson correlation,  $R = -0.20$ ; *C9orf72*  $R = -0.05$ , *SOD1*  $R = -0.09$ ). Grey dotted lines indicate the points of no difference between ALS versus control astrocytes for IR ratio (X-axis) and gene expression (Y-axis). Reactivity related transcripts with IR down and expression up are labelled. See Figure 2E–G for processes enriched amongst genes in the IR down and expression up quadrant. For the three remaining quadrants see Supplementary Figure S5A–C. (E–G) Bar graphs showing significantly over-represented functional categories (FDR < 0.05) determined by gene ontology analysis of transcripts with reduced IR and increased expression in *VCP* (E), *C9orf72* (F), *SOD1* (G) versus control astrocytes. (H–J) Heatmaps of pan-reactive (black), A1 specific (red), and A2 specific (blue) markers from Liddelow *et al.*, 2017 (24). IR and gene expression (GE) data represent the log<sub>2</sub> fold change in *VCP* (J), *C9orf72* (K), *SOD1* (L) versus control astrocytes. (K) Coverage plots of reads per million from *VCP* mRNA sequencing reads for differentially expressed genes *ERAP2* and *PTPRN* genes. Within each are alignment tracks for CTRL (top) and *VCP* (bottom) astrocytes. Ensembl transcript annotation tracks are shown above the alignment tracks with exons (thick blue boxes) and introns (thin blue arrowed lines). Vertical red dashed lines show the retained intron of interest and the IR ratio is labelled above the retained intron. Y-axis limits for each track are auto-adjusted to the peak exon coverage so that the target feature fills the view and enables easier visual comparison of IR ratios between conditions.





**Figure 3.** Loss of intron retention and increased expression of astrocyte reactivity genes is reproduced cytokine stimulation of astrocytes *in vitro* and astrocyte TDP-43 deletion *in vivo*, which coincides with increased translation in SOD1<sup>G37R</sup> astrocytes. (A–C) Schematics depicting (A) *in vitro* inducing

between TDP-43 deleted astrocytes and ALS astrocytes revealed an overlap of 139/525 (26.5%,  $P = 1.2 \times 10^{-92}$ ; Supplementary Table S4). These results indicate that astrocyte TDP-43 is required to maintain the abundant physiological IR levels and the loss of TDP-43 may trigger the decrease in IR we have revealed in ALS astrocytes.

By comparing gene expression with IR in TDP-43 deleted astrocytes, we observed that genes with decreased IR had significantly higher gene expression ( $P = 0.01$ , Figure 3E). Consistent with the other datasets, we found a negative correlation between changes in IR and changes in gene expression ( $R = -0.07$ ; Figure 3G). Also similar to both ALS and cytokine-stimulated astrocytes, gene ontology analysis showed that they are enriched in metabolic processes, cellular compartments and RNA binding (Figure 3I). Of the Liddelow *et al.* (24) astrocyte reactivity genes with a reliable IR event, 12/31 (39%) were decreased in IR and increased in expression (overlap  $P = 0.01$ , Figure 3L). Comparing genes with decreased IR and increased expression between the TDP-43 deleted astrocytes and the ALS astrocytes revealed an overlap of 46/269 (17.1%,  $P = 4.6 \times 10^{-34}$ ), of which 19 are directly relevant to astrocyte reactivity (Supplementary Table S5) (26). This suggests that astrocyte TDP-43 depletion recapitulates reduced IR and increased expression in similar reactivity related genes as ALS astrocytes.

### Reactivity genes with decreased IR in ALS astrocytes are translated in a *SOD1* mouse model of ALS

To determine whether decreased IR in ALS astrocytes influences translation of astrocyte reactivity genes, we examined a translating ribosome affinity purified RNA sequencing (TRAP-Seq) dataset from Sun *et al.* (Figure 3C) (67). *SOD1*<sup>G37R</sup> mutant mouse spinal cord astrocytes were isolated and mRNAs bound to ribosome subunits were sequenced enabling interrogation of these translating mRNAs (four *SOD1*<sup>G37R</sup> and six control samples). In concordance with Sun *et al.*, we found that *SOD1*<sup>G37R</sup> astrocytes exhibit upregulation of inflammatory pathways (Figure 3J and Supplementary Figure S5G) as well as established astrocyte reactivity genes (24) (enrichment  $P = 7.3 \times 10^{-10}$ ; Figure 3M). Comparing genes with significantly increased

translation in Sun *et al.* (FDR < 0.05) with those with decreased IR in ALS astrocytes, revealed 44/429 (9%,  $P = 9.2 \times 10^{-14}$ ) overlapping genes, of which 32 are directly relevant to astrocyte reactivity, including *OSMR*, *FBLN5*, *VIM* and *PTX3* (Supplementary Table S4) (26). Additionally, of the genes exhibiting decreased IR in stimulated reactive astrocytes, 6/223 ( $P = 0.01$ ) displayed increased translation in *SOD1*<sup>G37R</sup> mouse astrocytes, of which all eight were decreased in IR in ALS astrocytes. Similarly, in the astrocyte TDP-43 deleted mouse, 17 genes with decreased IR were increased in translation in *SOD1*<sup>G37R</sup> astrocytes ( $P = 4.4 \times 10^{-7}$ ), of which 7 were also decreased in IR in ALS mutants. Collectively, these data suggest that decreased IR in ALS astrocytes is correlated with translation of reactivity genes.

### Decreased nuclear intron retention is associated with increased cytoplasmic spliced reactivity transcripts and protein in *VCP* mutant astrocytes

To gain further mechanistic insight into reactive transformation of astrocytes, we examined the nuclear and cytoplasmic transcriptomes and proteomes in *VCP* mutant astrocytes. To achieve this we performed nuclear–cytoplasmic fractionation of astrocytes, followed by high depth poly(A)+ selected RNA sequencing and mass spectrometry (Figure 4A). Fractionation quality was confirmed by determining that transcripts known to localize to the nucleus (histone *H3*, intronic *GAPDH*, *NEAT*, *MALAT1*) or cytoplasm (exonic *GAPDH*, *Tubulin*) were enriched in the expected compartment, using western blot (Figure 4B), RT-qPCR (Figure 4C), RNA sequencing (Supplementary Figure S6A) and mass spectrometry (Supplementary Figure S6B) of both *VCP* mutant and control astrocytes (40,82). Principal component analysis and unsupervised hierarchical clustering demonstrated that samples segregated based on both fraction (nuclear or cytoplasmic) and genotype (Supplementary Figure S6C and D).

To determine the subcellular location of intron-retaining transcripts, IR focused analysis was performed on nuclear and cytoplasmic isolates. This revealed 5.8-fold more retained introns in nuclear than cytoplasmic fractions (41,138

astrocyte reactivity with inflammatory cues (TNF $\alpha$ , IL-1 $\alpha$ , C1q) in hiPSC-derived control astrocytes, (B) *in vivo* astrocyte-specific TDP-43 deletion using conditional GFAP-Cre recombinase promoter. Spinal cord was dissected and mRNA extracted before RNA sequencing, and (C) ALS *SOD1*<sup>G37R</sup> mutant mouse with astrocyte-specific (Aldh1l1) bacTRAP reporter, which expresses the EGFP-tagged ribosome protein (Rpl10a) within astrocytes. Polyribosome-associated mRNAs undergoing translation are then isolated by EGFP immunoprecipitation before mRNA purification and RNA translome sequencing. (D) Violin plot showing delta IR ratio in reactive astrocytes (blue, left,  $n = 578$ ) and TDP43 deletion (red, right,  $n = 700$ ) minus control across differential intron retention events. Intron retention was significantly downregulated in both cytokine stimulated astrocytes and TDP43 deleted astrocytes (one-sample  $t$ -test, \*\*\*\* represents  $P < 0.0001$  and \*\*  $P < 0.01$ ). (E) Violin plots showing log2 fold change in gene expression (reactive state versus control, y-axis) in downregulated IR events (blue) and upregulated IR events (red) for cytokine stimulated (IR down,  $n = 304$ ; IR up,  $n = 274$ ) and TDP-43 deleted (ko, knockdown) (IR down,  $n = 581$ ; IR up,  $n = 119$ ). Student's  $t$ -test was used to determine significance (\*\*\*\* indicates  $P < 0.0001$ , \*  $P < 0.05$ ). (F and G) Scatterplots of the  $\Delta$  IR ratio (reactive state minus control, x-axis) against log2 fold change in gene expression (y-axis) in (F) cytokine stimulated and (G) TDP43 deleted versus control astrocytes. Points are coloured by mean gene expression across mutant and control (red = high, blue = low expression). Black dashed line indicates linear regression correlation (cytokine stimulated Pearson correlation  $R = -0.34$ ; TDP43 deleted  $R = -0.07$ ). Reactivity related genes with IR down and expression up are labelled. See Figure 3H–I for processes enriched amongst genes in the IR down and expression up quadrant. (H and I) Bar graphs showing significantly over-represented functional categories (FDR < 0.05) determined by gene ontology analysis of genes with reduced IR and increased gene expression in cytokine stimulated (H) and TDP43 deleted (I). See Supplementary Figure S5E and F for the other IR and expression categories. (J) Bar graph showing curated significantly over-represented functional categories (FDR < 0.05) determined by gene ontology analysis of genes with increased translation in *SOD1*<sup>G37R</sup> versus control mouse spinal cord astrocytes. See Supplementary Figure S5G for genes decreased in translation. (K and L) Heatmaps of pan-reactive (black), A1 specific (red) and A2 specific (blue) markers from Liddelow *et al.*, 2017 (24). IR and gene expression (GE) data represent the log2 fold change in cytokine stimulated (K) and TDP43 deleted (L) versus control astrocytes. (M) Heatmap of pan-reactive, A1 specific and A2 specific markers in astrocyte translome data log2 fold change in *SOD1*<sup>G37R</sup> versus control samples.





versus 7,036, respectively, Supplementary Figure S7C and D), consistent with previous reports (38,74,82). Of the 16,317 retained introns identified in whole astrocytes, 92% ( $n = 14,952$ ) were observed in nuclear isolates, 40% ( $n = 6,493$ ) in cytoplasmic isolates, while 37% ( $n = 6,059$ ) were observed in both (Supplementary Figure S7E). Retained introns in both nuclear and cytoplasmic fractions exhibited similar characteristics as in whole astrocytes (shorter in length, higher in GC content, conservation score, RBP binding affinity and lower in splice site strength; Supplementary Figure S8A–F).

Although we observed a reduction in the absolute numbers of retained introns in *VCP* mutant relative to control in both the nucleus (7% less, 34,464 versus 36,943, respectively) and the cytoplasm, this reduction was substantially greater in the cytoplasm (58% less, 2,791 versus 6,607). Comparing IR levels between *VCP* and control, revealed 1.8-fold more significantly differentially retained introns in the cytoplasm (5,138 in 3,065 genes; Supplementary Figure S7B) than the nucleus (2,915 in 2,079 genes; Supplementary Figure S7A). These were decreased in *VCP* mutants in both nuclear (1882/2915, 65%) and cytoplasmic fractions (5089/5138, 99%; Figure 4D, Supplementary Figure S9). Comparing this decrease between the cytoplasm and nucleus confirmed that the cytoplasmic decrease was significantly greater (wilcoxon test,  $P < 2.22 \times 10^{-16}$ ). This excessive cytoplasmic decrease raises the possibility that IR transcripts in *VCP* mutant astrocytes are subject to either (i) increased nuclear confinement or (ii) enhanced cytoplasmic degradation.

To establish whether differentially retained introns in *VCP* mutant astrocytes differed between the nucleus and cytoplasm, we next examined their IR characteristics. Although similar patterns were noted for GC content, we found that nuclear retained introns that were decreased in *VCP* mutants were shorter, higher in conservation and

weaker in RBP binding, compared to introns gained in *VCP* (Supplementary Figure S8G–L). Conversely, cytoplasmic retained introns decreased in *VCP* mutants were longer, lower in conservation and stronger in RBP binding. These opposing nuclear and cytoplasmic IR transcript patterns between *VCP* and control astrocytes support either altered nuclear-to-cytoplasmic transport or differential stability.

To ascertain the degree to which IR transcripts are nuclear confined in *VCP* mutant astrocytes, we compared IR between cytoplasmic and nuclear fractions. Although we noted 10% more differentially retained introns in *VCP* (31,358 events in 9,630 genes) than control (28,447 events in 9,459 genes), the proportion of these that were increased in the nucleus (i.e. nuclear confined) was 99.9% in both conditions (control: 28 410/28 447; *VCP*: 31 330/31 358). Comparing this nuclear confinement between *VCP* and control showed no significant difference (wilcoxon test,  $P = 0.66$ ; Figure 4E). This striking but similar nuclear confinement in both control and *VCP* indicates that astrocytes use IR coupled with nuclear confinement as a strategy to regulate gene expression, consistent with reports in other cell types (30,82).

However, this still leaves unresolved the excessive cytoplasmic decrease in IR within *VCP* mutant astrocytes. To begin to address this, we next examined NMD to establish whether cytoplasmic degradation is responsible for the disproportionate loss of IR. Gene set enrichment analysis of the NMD gene set ( $n = 120$ ) in the cytoplasmic fraction revealed significant upregulation in *VCP* (normalized enrichment score 1.15, enrichment  $P = 0.049$ ), with 94/120 (78%) of the NMD genes being increased (Figure 4F; one-sample wilcoxon test  $P = 2.36 \times 10^{-7}$ ). By performing differential spliced transcript-level isoform expression analysis in *VCP* versus control fractions and comparing transcripts annotated as NMD substrates with those annotated as protein coding biotypes, we found that cyto-

histone 3 (bottom, nuclear marker) and Tubulin (top, cytoplasmic marker) in nuclear (left) and cytoplasmic (right) protein from control (ctrl1, ctrl2) and *VCP* (R155C, R191Q) astrocytes. Gaps indicate non-adjacent samples run on the same blot. (C) RT-qPCR of nuclear markers intronic GAPDH, Malat1, NEAT and cytoplasmic marker exonic GAPDH transcript fold enrichment in nuclear and cytoplasmic fractions from control and *VCP* mutant astrocytes. Data are from two replicates for each sample and show mean  $\pm$  SD. (D) Violin plot showing  $\Delta$  IR ratio across differential intron retention events in nuclear (red,  $n = 5138$ ) and cytoplasmic (blue,  $n = 2915$ ) fractions. One-sample wilcoxon test adjusted  $P = 1.94e-47$  and  $P < 2.00e-145$  respectively, \*\*\*\* represents  $P$  value  $< 0.0001$ . Comparing cytoplasmic and nuclear groups with the wilcoxon test showed significantly greater decreases in IR (*VCP* versus control) in cytoplasmic than nuclear fraction ( $P < 2.2 \times 10^{-16}$ ). (E) Violin plot showing  $\Delta$  IR ratio (cytoplasmic minus nuclear fractions) across differential IR events in *VCP* (blue,  $n = 31\ 358$ ) and CTRL (red,  $n = 28\ 447$ ). One-sample wilcoxon tests  $P < 2.00e-145$  for both. Comparing *VCP* and CTRL groups showed no differences in nuclear confinement (wilcoxon test  $P = 0.66$ ). (F) Nonsense mediated decay (NMD) components ( $n = 120$ ) gene set enrichment analysis of  $\log_2$  fold changes (LFC) in gene expression (*VCP* versus CTRL) in the cytoplasmic fraction. One-sample wilcoxon test  $P = 2.36 \times 10^{-7}$ . (G) Violin plots showing  $\log_2$  fold changes in transcript isoform expression in *VCP* versus CTRL for transcripts annotated in Ensembl as nonsense\_mediated\_decay (NMD tx) and protein\_coding (non-NMD tx) biotypes in cytoplasmic (left, NMD tx  $n = 16\ 370$ , non-NMD tx  $n = 91\ 880$ , wilcoxon  $P < 2.2 \times 10^{-16}$ ) and nuclear fractions (right, NMD tx  $n = 16\ 370$ , non-NMD tx  $n = 91\ 880$ , wilcoxon  $P = 0.11$ ). (H) Violin plot showing  $\log_2$  fold change (*VCP* versus control) in cytoplasmic transcript abundance for upregulated (blue,  $n = 1033$ ) and downregulated (red,  $n = 1882$ ) nuclear IR events (*VCP* minus control). Student's  $t$ -test was used to compare IR up with IR down. (I) Scatterplot of the nuclear  $\Delta$  IR ratio (*VCP* minus control) against  $\log_2$  fold change (LFC) in cytoplasmic spliced transcript abundance in *VCP* versus control. Black dashed line indicates linear regression correlation (Pearson correlation  $R = -0.21$ ). Reactivity related genes with nuclear IR down and cytoplasmic expression up are labelled. (J) Bar graph showing curated overrepresented functional categories (FDR  $< 0.05$ ) by gene ontology analysis of genes with reduced nuclear IR and increased cytoplasmic mRNA abundance, in *VCP* versus control. See Supplementary Figure S7G for GO terms in other IR and expression categories. (K) Violin plot showing log fold change (*VCP* versus control) in cytoplasmic mass spectrometry (protein) intensities for upregulated (blue,  $n = 621$ ) and downregulated (red,  $n = 413$ ) nuclear IR events (*VCP* minus control). Student's  $t$ -test was used to compare IR up with IR down ( $P = 0.14$ ). (L) Scatterplot of the nuclear  $\Delta$  IR ratio (nuclear *VCP* minus nuclear control,  $x$ -axis) against log fold change (LFC) in mass spectrometry ( $y$ -axis) in *VCP* versus control cytoplasmic fractions. Black dashed line indicates linear regression correlation (Pearson correlation,  $R = -0.06$ ). Reactivity related genes with nuclear IR down and cytoplasmic protein up are labelled. (M) Bar graph showing curated overrepresented functional categories (FDR  $< 0.05$ ) by gene ontology analysis of genes with reduced nuclear IR and increased cytoplasmic protein abundance, in *VCP* versus control. See Supplementary Figure S7H for GO terms in other IR and protein categories. (N) Heatmap of astrocyte reactivity markers from Liddelow *et al.*, 2017. Nuclear intron retention (IR), cytoplasmic gene expression (Cyto mRNA) and cytoplasmic mass spectrometry (Cyto protein) data represent the log fold change in *VCP* versus control.

plasmic NMD substrate transcripts exhibited significantly lower expression than protein coding biotypes in *VCP* mutant cytoplasm (wilcoxon  $P < 2.2 \times 10^{-16}$ ). Comparatively there was no significant difference between differential transcript expression between NMD and protein coding biotypes in the nuclear fraction ( $P = 0.11$ ; Figure 4G). This contrast between nuclear and cytoplasmic differential transcript expression according to NMD transcript annotation supports enhanced cytoplasmic NMD activity in *VCP* mutant astrocytes. Taken together, this suggests that increased cytoplasmic NMD activity underlies the excessive cytoplasmic degradation of IR transcripts in *VCP* mutant astrocytes.

To explore whether the nuclear confinement of IR transcripts determines cytoplasmic expression, we next mapped nuclear IR to cytoplasmic spliced transcript abundance. Genes with decreased nuclear IR in *VCP* mutant astrocytes displayed higher cytoplasmic spliced transcript abundance than genes with increased IR ( $P = 1.71 \times 10^{-35}$ ,  $R = -0.21$ ; Figure 4H and I), suggesting that spliced transcripts are more likely to be exported from the nucleus. Genes with reduced nuclear IR and increased cytoplasmic spliced transcript abundance were over-represented in astrocyte reactivity processes: cell adhesion (*MCAM*, *COL1A1*, *ILK*), translation (*EIF2B4*, *EIF5*, *EIF3A*) and immune activation (*HLA-B*, *HLA-C*, *TGFB1*; Figure 4J). NMD was also enriched, indicating NMD component transcripts themselves are subject to enhanced splicing and cytoplasmic translocation. Of the astrocyte reactivity genes identified in Liddelow *et al.* (24) with a reliable IR event, we observed decreased nuclear IR and increased cytoplasmic expression in 11/30 markers (37%, overlap  $P = 0.002$ ; Figure 4N).

Using mass spectrometry, we detected 1160 proteins in the cytoplasmic fractions of which 580 were increased in protein levels in *VCP* versus control (Supplementary Table S6). Of these 580 proteins, 500 had a reliable IR event, of which 299 (60%) were decreased in nuclear IR and of these, the majority (155/299, 52%) are involved with reactive transformation. Across the entire cytoplasmic proteome, we found no significant difference in protein abundance between those whose cognate transcripts exhibited decreased compared with increased nuclear IR in *VCP* versus control astrocytes (wilcoxon test  $P = 0.46$ ; Figure 4K). However, correlating changes in nuclear IR with changes in cytoplasmic protein abundance revealed a negative correlation ( $R = -0.06$ ; Figure 4L), indicating that decreased nuclear IR is associated with increased cytoplasmic protein. Genes with reduced nuclear IR and increased cytoplasmic protein abundance were over-represented in the infection response, cell adhesion, RNA processing as well as NMD, indicating that an increase in protein levels is a specific phenomenon amongst astrocyte reactivity processes (Figure 4M). Seven of the Liddelow astrocyte reactivity genes were detected in cytoplasmic mass spectrometry of which HSPB1 and HLA-E were both increased in *VCP* mutants, which were also increased in cytoplasmic mRNA (Figure 4N). Comparing cytoplasmic mRNA and protein abundance changes in *VCP* versus control revealed significantly increased mRNA fold changes than that of protein (wilcoxon test  $P = 3.74 \times 10^{-4}$ ; Supplementary Fig-

ure S7F), implicating enhanced cytoplasmic mRNA decay within *VCP* mutant astrocytes. These findings together suggest that under normal circumstances astrocytes employ IR to achieve nuclear confinement of 'poised' reactivity transcripts; however in *VCP* mutant astrocytes increased splicing releases them to the cytoplasm permitting their translation (depicted in graphical abstract).

## DISCUSSION

Recent advances in astrocyte biology have implicated reactive astrocytes in ALS pathogenesis (14,22–23,29,81,83). However, despite this accumulating evidence (5–7) and the increased availability of transcriptomic data (8,27,66), the molecular determinants of astrocyte gene expression changes that drive reactive transformation have remained unknown. Although previous transcriptome-wide analyses exposed the reactive nature of ALS astrocytes, they mostly relied on differential gene expression rather than alternative splicing methods (8–9,23,29,66). Combining expression with splicing analyses of the astrocyte transcriptome, coupled with translome and proteome data, we find that IR is prevalent in healthy quiescent hiPSC-derived astrocytes but that ALS astrocytes share a common decreased IR signature, which is a conserved phenomenon across *VCP*, *C9orf72* and *SOD1* mutations. This is reproduced in publicly available RNA sequencing datasets in both (i) astrocytes stimulated with  $\text{TNF}\alpha$ ,  $\text{IL-1}\alpha$  and  $\text{C1q}$  to undergo reactive transformation (a modest 53% decrease albeit statistically significant) and (ii) astrocytes depleted in TDP-43 *in vivo* (more substantial 83% decrease). Transcripts with decreased IR in ALS and reactive astrocytes displayed increased expression and were over-represented in astrocyte reactivity pathways, supporting IR as a post-transcriptional repressor of reactive transformation (30–32). Furthermore, by re-examining public translational profiling of *SOD1*<sup>G37R</sup> mouse spinal cord astrocytes (67), we observed increased translation of astrocyte reactivity genes that exhibited decreased IR in ALS astrocytes (26). Our study provides insight into the molecular factors regulating reactive transformation of ALS astrocytes, whereby decreased IR might serve to post-transcriptionally enhance astrocyte reactivity genes.

These findings differ substantially from what we and others have established in ALS motor neurons carrying diverse mutations, which exhibit increased IR (41–42,44,46,84–86). Although motor neurons and astrocytes are derived from the same neural precursor cell, they evidently show striking differences in IR regulation during terminal differentiation. Instead, the association between decreased IR and astrocyte reactivity resembles other immune cells, such as lymphocytes and macrophages, where there is an inverse relationship between global IR levels and the cellular activation state (31–34,47). We speculate that the opposing shift in IR between motor neurons and astrocytes is a consequence of distinct cell-type responses to ALS pathogenic processes, such as TDP-43 proteinopathy (71), extrinsic stressors (87), neuronal injury (8,88) or even reactive transformation itself. Although *VCP*, *C9orf72* and *SOD1* mutations are pathologically distinct with divergent effects, a common theme is that they are each linked to perturbed RNA processing and/or

RBP mislocalization, which may be responsible for these ALS cell-type specific alterations in IR (3,41). In support of this, TDP-43 mislocalization triggers neurons to upregulate immune response pathways (NF- $\kappa$ B and type 1 interferon) via the cGAS-STING pathway (89), whereas TDP-43 proteinopathy in astrocytes has been shown to be associated with metabolic dysregulation via cAMP and calcium signalling (90). These divergent responses of astrocytes and motor neurons may differentially affect key splicing factors, provoking deregulation of the spliceosome complex and altering its finely-tuned ability to recognize and act on splicing signals (91–93). Future functional studies with cell-type specific targeted disruption of spliceosome components, as well as splicing enhancers and repressors, would help to elucidate the mechanisms of this differential IR shift and whether it directly contributes to disease pathogenesis (94).

It has been proposed that intron-containing mRNAs may function to encourage their nuclear confinement, preventing nuclear leak and translation (74,82). Thus, an important question is whether transcripts exhibiting decreased IR in ALS astrocytes are able to escape nuclear confinement and undergo nuclear-export. To address this, we used nuclear-cytoplasmic fractionation and found decreased IR in both nuclear and cytoplasmic compartments in *VCP* mutant astrocytes. We identified more than double the number of differential IR events (*VCP* mutant versus control) in the cytoplasm (5,138) than the nucleus (1,882) and although there was an overall decrease in IR in both *VCP* fractions, the proportion lost was dramatically greater in the cytoplasm (99% versus 65%). Although there was no change in the strength of nuclear confinement of IR transcripts, there was a substantial loss of transcripts annotated as NMD substrates from the cytoplasm in *VCP* mutant astrocytes. We further established that NMD components exhibited decreased nuclear IR and increased cytoplasmic abundance in *VCP* mutant astrocytes. This suggests that, together with astrocyte reactivity transcripts, NMD components are subject to enhanced splicing enabling their cytoplasmic translocation. While it is possible that other mechanisms regulating IR transcripts in the cytoplasm are responsible, such as targeted subcellular localization, RBP sequestering or cytoplasmic splicing, the most likely explanation for the excessive cytoplasmic degradation of IR transcripts in *VCP* mutant astrocytes is enhanced NMD activity (42,95–96). This raises the possibility that NMD itself is a key part of reactive transformation, which would be consistent with studies reporting that IR-NMD coupling has additional roles in regulating the stress response, beyond its well established post-transcriptional quality control mechanism serving to degrade PTC-containing IR transcripts (35–36,74,97–101). Indeed, perturbed IR-NMD crosstalk has been shown in neuroinflammation and ALS, where it may act to liberate bound RBPs for further splicing modulation (32,102–105).

Consistent with enhanced NMD in *VCP* mutant astrocytes, we also found that nuclear transcripts with decreased IR manifest higher cytoplasmic spliced transcript abundance than nuclear transcripts with increased IR. We observed a major cluster of astrocyte reactivity related genes in *VCP* mutant astrocytes that are subject to loss of nuclear IR and increased cytoplasmic spliced transcript abundance, including many of the established astrocyte reactiv-

ity genes identified by Liddel et al. (24). Interestingly, of these we found multiple HLA class I genes, which when overexpressed protect motor neurons from reactive astrocyte toxicity (106). Within this cluster we also identified enrichment of other genes involved with the extracellular-matrix (e.g. *COL1A1*, *MMP24*, *ADAM19*), focal adhesion (e.g. *MCAM*, *ILK*, *EPHA2*, *ITGB1*) and immune-activation (e.g. *TGFB1*, *NFKB1*, *NFKB2*, *CD68*, *IL32*), consistent with processes previously identified in ALS astrocytes (8,15–16,29,66). This indicates that transcripts regulating multiple facets of astrocyte reactivity exhibit enhanced nuclear splicing in *VCP* mutant astrocytes that may allow nuclear-export to the cytoplasm for translation, facilitating reactive transformation.

Overall, our study provides new insights into the co-ordinated role of IR coupled with gene expression in astrocytes and its aberration in ALS. Astrocytes employ IR to post-transcriptionally repress reactivity genes; however, ALS astrocytes undergo augmented splicing and lose this homeostatic regulation, which may be an initial compensatory mechanism that may become maladaptive over time. Further investigation into the impact of this phenomenon on neighbouring neurons will enable an improved understanding of ALS, as well as other neurodegenerative diseases characterized by aberrant astrocyte reactivity. Our study raises the prospect of therapeutically targeting astrocyte reactivity through reinstating the physiological IR programme by manipulation of the splicing process, for example using antisense oligonucleotides.

## DATA AVAILABILITY

All raw and processed mRNA sequencing data generated in this study have been deposited in the NCBI Sequence Read Archive (BioProject Gene Expression Omnibus) under accession number GSE160133. RAW Mass Spectrometry data have been deposited to the ProteomeXchange Consortium (<http://proteomecentral.proteomexchange.org>) via the PRIDE partner repository with the dataset identifier PXD022604. Code is available through GitHub [https://github.com/ojziff/ALS\\_astrocyte\\_intron\\_retention](https://github.com/ojziff/ALS_astrocyte_intron_retention). hiPSC-derived astrocytes carrying ALS mutations are available at GSE142730 (C9orf72), GSE102902 and GSE99843 (SOD1 mutants and control respectively). Cytokine-stimulated hiPSC-derived astrocytes are available at syn21861181. *TARDBP* knockout mouse spinal cord astrocyte specific RNA-seq is available at GSE156542. Mouse SOD1 astrocyte TRAP-seq is available at GSE74724.

## SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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