Transcriptome profiling of mouse brains with qkl-deficient oligodendrocytes reveals major alternative splicing defects including self-splicing

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The qkl gene encodes a family of RNA binding proteins alternatively spliced at its 3′ end, giving rise to three major spliced isoforms: QKI-5, QKI-6 and QKI-7. Their expression is tightly regulated during brain development with nuclear QKI-5 being the most abundant during embryogenesis followed by QKI-6 and QKI-7 that peak during myelination. Previously, we generated a mouse conditional qkl allele where exon 2 is excised using Olig2-Cre resulting in QKI-deficient oligodendrocytes (OLs). These mice have dysmyelination and die at the third post-natal week. Herein, we performed a transcriptomic analysis of P14 mouse brains of QKI-proficient (QKIFL/FL; -) and QKI-deficient (QKIFL/FL; Olig2-Cre) OLs.

QKI deficiency results in major global changes of gene expression and RNA processing with >1,800 differentially expressed genes with the top categories being axon ensheathment and myelination. Specific downregulated genes included major myelin proteins, suggesting that the QKI proteins are key regulators of RNA metabolism in OLs. We also identify 810 alternatively spliced genes including known QKI targets, MBP and Nfasc. Interestingly, we observe in QKIFL/FL; Olig2-Cre a switch in exon 2-deficient qkl mRNAs favoring the expression of the qkI-5 rather than the qkI-6 and qkI-7. These findings define QKI as regulators of alternative splicing in OLs including self-splicing.

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nuclear localization signal in its unique C-terminal sequence and is exclusively nuclear. Thus QKI-5 participates in nuclear roles such as in the regulation of alternative splicing. QKI-6 is distributed throughout the cell and forms heterodimers with QKI-5 and may perform both nuclear and cytoplasmic functions. QKI-7 is predominantly cytoplasmic and both QKI-6 and QKI-7 are thought to function in mRNA export, mRNA stability and protein synthesis. During embryonic brain development, QKI-5 isoform is the most abundant and its expression declines after the first two post-natal weeks, while the QKI-6 and QKI-7 isoforms are expressed later in embryogenesis and peak during myelination at the third post-natal week. What regulates the qki gene and its alternative splicing is unknown.

The QKI isoforms all share identical KH domains flanked by QUA1 and QUA2 sequences that mediate sequence-specific recognition of an RNA element termed the QKI response element (QRE) with the following sequence ACUAAY-(N1–20)-UAAY. Transcriptome-wide CLIP (crosslinking and immunoprecipitation) demonstrate that the QKI isoforms in cellulo bind the QREs and also showed that QKI associates predominantly with intronic sequences implying that alternative splicing regulation is one of its main nuclear functions. The QKI isoforms were shown to regulate pre-mRNA splicing of myelin components, and later this function of QKI was shown not to be restricted to brain, as QKI-5 also fulfills this role in vascular smooth muscle cells, monocytes and skeletal muscle.

Herein, we report a transcriptomic analysis by RNA-Seq of entire brains of QKIFL/FL.Olig2-Cre mice compared to QKIFL/FL- mice. The results are consistent with the hypomyelination phenotype of the mice and show that mRNAs encoding myelin components are severely downregulated. We also identify alternative splicing defects especially in genes encoding myelin components. Interestingly, we noted that the absence of the QKI proteins also altered the alternative splicing of the qki gene. Deletion of the qki exon 2 generates a null allele, but switches the production of the qki1 and qki2 isoforms to being mainly qki-6 and qki-7 in wild type P14 brains to qki-5 in the brains of QKIFL/FL.Olig2-Cre mice. This work defines a self-splicing mechanism by which the qki gene autoregulates its own alternative splicing in oligodendrocytes.

Results
QKI-depleted oligodendrocytes downregulate genes involved in myelination. To define the genome-wide alterations in gene expression and alternative splicing patterns in the absence of the QKI proteins, we isolated RNA from brains of QKIFL/FL- and QKIFL/FL.Olig2-Cre mice (n = 3 females/genotype) and performed RNA-sequencing at an average of 133 million reads/sample (Supplementary Table S1). Sequencing and quality control metrics showed high quality reads, with a mean alignment rate greater than 90%. More than 95% of reads mapped to genes and there was low ribosomal and mitochondrial gene content (Supplementary Table S1). We first assessed the global impact of oligodendrocyte (OL)-specific depletion of the QKI RNA binding proteins by unsupervised clustering analysis of samples based on expression profiles. QKIFL/FL- and QKIFL/FL.Olig2-Cre mice consistently formed distinct, robust clusters (Fig. 1a). Indeed, Principal Component Analysis (PCA) based on gene expression data showed that the first component (PC1) clearly separates QKIFL/FL- and QKIFL/FL.Olig2-Cre mice, and explained 75% of the variance (Fig. 1a). Bootstrapped hierarchical clustering also separated the two groups with high bootstrap values (Supplementary Fig. S1a). The clustering is very robust, and is not affected by the number of genes or the algorithm used (Supplementary Fig. S1b). Taken together, these results indicate that QKI-depletion in oligodendrocytes induces major changes in gene expression in the brain. In fact, differential expression analysis showed statistically significant differences in 1,899 genes, representing 10.55% of expressed genes (n = 18,004). Of these, 229 had an absolute fold change greater than 2 (Fig. 1b and Supplementary Table S2). A high proportion (70%) of the significant genes is downregulated, in line with a scenario where a well-defined population of cells is depleted in the brain samples. Quantitative RT-PCR analysis of 19 genes (Supplementary Fig. S1c) showed strong correlation between RNA-seq fold change and RT-qPCR data (r² = 0.8049, Fig. 1c). Gene Ontology (GO) analysis using the conservative list of 229 differentially expressed genes (with absolute fold change greater than 2) suggests that the downregulated biological processes are enriched for genes implicated in myelination and ensheathment of neurons (Fig. 2a,b), consistent with the hypomyelination phenotype of these mice. The upregulated genes, on the other hand, are enriched for biological processes involving extracellular matrix organization and cell adhesion, and include a number of genes related to neuronal biology, such as Ntn4, or genes associated with neurological phenotypes, such as Col5a3 (Supplementary Table S2).

Loss of QKI did not lead to major changes in transcript levels of genes involved in oligodendrocyte lineage specification and precursor proliferation, including Olig2, Nkx6.1, Id2, Id4, and Sox5/6 (Supplementary Table S2), consistent with maintenance of PDGFIR+ OLS in QKIFL/FL.Olig2-Cre mice. QKI did appear to regulate RNA metabolism of transcription factors involved in OL differentiation and expression of myelin genes, including Sox10, Myrf, or myelin genes, including Mag, Plp, Mbp, Molp, and Sirt2 (Fig. 1c), consistent with the hypomyelination phenotype of the mice. The sequencing of the entire brain would reveal secondary events occurring, for example in neurons, as it is well-known that OLs tightly regulate the contacts and metabolism of neurons. However, genes involved in neuronal specification and differentiation, including Neurogenin proneural genes, Achaete scute-like 1, as well as neurogenic differentiation, were not affected by the loss of QKI. Therefore, loss of QKI in OLs exerts changes in gene expression that are specific to OL lineage transcripts.

Alternative splicing changes observed in QKIFL/FL.Olig2-Cre mice. A previous search of alternative splicing events (ASE) within a defined subset of genes allowed us to identify 31 QKI-regulated ASEs in oligodendrocytes. Here, we expanded the search genome-wide to identify novel events and to assess the extent of AS regulation by QKI, using two different, complementary methods to infer splicing information from RNA-Seq data (DEXSeq and rMATs). Genome-wide loss of QKI resulted in widespread AS changes, with 1,438 ASEs
detected in 810 genes (Supplementary Tables S3–S5). Of the 31 identified with the candidate approach, 19 events were confirmed by RNA-seq including ASEs of Nfasc, Magi, Lhfp13, Mpzl1, Cadm1, Bin1, Vldlr, Plp1, Phka1, Mbp, Mobp, Grm5, Nhsl1, and Sirt2 (Supplementary Table S6). These data indicate that our RNA-seq analysis identified bona fide QKI RNA targets and that the majority of AS exons are novel events that have not been

**Figure 1.** Gene expression profiles comparing brains of QKIFL/FL;Olig2-Cre and QKIFL/FL- mice. (a) Global effects of QKIFL/FL;Olig2-Cre on cells were evaluated by unsupervised clustering of samples based on expression profiles of 1,000 most variant genes. QKIFL/FL;Olig2-Cre and QKIFL/FL- mice form robust, distinct clusters, both with hierarchical clustering (right) and Principal Component Analysis (left). (b) Volcano plot representing the results of differential expression analysis. Genes with adjusted p-value < 0.05, absolute fold change >2 and base mean >100 are colored in red. (c) Validation of RNA-seq data by qPCR with 19 transcripts. RNA-seq data strongly correlated with qPCR data ($r^2 = 0.8049$, $p < 0.0001$, Pearson correlation coefficient).

**Figure 2.** Gene ontology analysis. Gene Ontology analysis shows enriched Biological Processes in the lists of upregulated genes (a) and downregulated genes (b) in QKIFL/FL;Olig2-Cre mice. Genes with adjusted p-value < 0.05, absolute fold change >2 and a base mean >100 were used for the analysis.
previously reported in QKI<sup>FL/FL;Olig2-Cre</sup> mice. A stringent list of ASEs, detected by both methods is presented in Supplementary Table S3. The majority of the observed events were skipped exons (SE), with only a handful of alternative 5′ or 3′ splice sites detected (Fig. 3a, Supplementary Tables S3 and S5), suggesting that loss of QKI may result in significant changes at the proteome level. Importantly, most SE events corresponded to skipped cassette exons (Fig. 3a). Among AS events identified by DEXSeq only, approximately half affected first or last exons (Fig. 3a). Since alternative promoter or terminator usage is generally not regulated via the splicing machinery, we excluded them from further analyses and concentrated on cassette exons only. We showed previously that the QKI isoforms bind optimally to QKI response elements (QREs) with a sequence of ACUAAY<sup>(N1-20) UAAY17</sup>. Therefore, we searched for the core sequence ACUAA as a minimal sequence and whether it is

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Figure 3. Brain alternative splicing patterns of mice with QKI-deficient oligodendrocytes. (a) Types of alternative splicing events identified in QKI<sup>FL/FL;Olig2-Cre</sup> mice. For skipped exons (SE), the type of exon affected is also indicated. Exons were considered to be “cassette exons” if they were not the first or last exon in any of the gene isoforms or “mixed exons” if they were not the first or last exon in at least one isoform. Abbreviations: SE: skipped exon, A5SS: alternative 5′ splice site, A3SS: alternative 3′ splice site, RI: retained intron. (b,c) Motif enrichment analysis. Percentage of sequences with the motif ACUAA in introns neighboring SE events compared to a set of background sequences for the stringent SE candidates called by both methods (b) or for all SE candidates (c). The occurrence of the motif was counted in a random set of non-alternatively spliced control sequences of the same size of the set of SE events. This was repeated 1,000 times to get the distribution of the motif occurrence in control sequences. The p-value corresponds to the empirical p-value of the enrichment for the set of SE events compared to the control distribution. (d) Gene Ontology analysis shows enriched molecular function, biological processes and cellular component in the list of genes where the stringent alternative splicing events were found. For Molecular Function, all GO terms with q-value < 0.01 are represented. For Biological Process and Cellular Component, GO terms with q-value < 0.01 and enrichment > 5 are represented.
enriched upstream and/or downstream of the skipped cassette exons (SCE) events (Figs 3b,c and S2). More than 40% of SCE events from the stringent list had the exact sequence motif ACUAA in the neighbouring regions (Supplementary Table S7). In particular, this was the case for almost 50% of SCE events with lower inclusion level in QKI FL/FL; Olig2-Cre mice. In both cases, this represents statistically significant enrichment (p-value < 0.001) over the distribution of the ACUAA motif frequency in a random set of control sequences (Figs 3b and S2b).

Furthermore, over 30% of the complete set of 842 detected SCE events had the sequence motif in neighbouring sequences (Supplementary Table S8), again representing a significant enrichment over the control distribution (Figs 3c and S2c,d). In addition, we determined whether the motif was present upstream or downstream of the AS exon. Similar to previous reports 3, we observed that the QRE core sequence tends to be located more often upstream of ASEs that are repressed by QKI and downstream of ASEs that are activated by QKI (Supplementary Tables S7, S8 and Supplementary Fig. 2g).

We performed GO analysis on the stringent list of candidates. Interestingly, the top enriched GO terms for biological processes included regulation of gliogenesis and myelination with myelin structural constituents and paranodal junction being one of the top molecular functions and cellular components, respectively (Fig. 3d). Moreover, many of the top AS candidates are associated with neurological disease(s) and/or involved in important brain function, with the top 15 represented in this table. (Table 1). In some cases, we found almost complete exon skipping in QKI FL/FL; Olig2-Cre mice, such as for exon 7 of Bcas1, exon 18 of Sema6a and exon 9 of Capzb (Fig. 4a,b,c, left panels, sashimi plots are shown in Supplementary Fig. S4a,b,c), which we have validated by semi-quantitative PCR showing complete exon exclusion in the QKIFL/FL; Olig2-Cre mice (Fig. 4a,b,c, middle panels).

Table 1. A large number of novel AS events observed in QKI FL/FL; Olig2-Cre mice are located in genes associated with neurological diseases and/or involved in important brain function, with the top 15 represented in this table. Abbreviations: SE: skipped exon, A5SS: alternative 5′ splice site, A3SS: alternative 3′ splice site, RI: retained intron.

| Gene Symbol | Gene Description | Splicing inclusion change in QKI FL/FL;Olig2-Cre | Event Type | Associated neurological diseases/brain function | Reference (PMID) |
|-------------|------------------|-----------------------------------------------|------------|-----------------------------------------------|-----------------|
| Vcan        | versican         | inclusion                                      | SE         | Neurite outgrowth                              | 24955366        |
| Gpm6b       | glycoprotein m6b | exclusion                                      | A3SS       | Oligodendrocyte differentiation and myelination | 23322581        |
| Pprr21      | protein tyrosine phosphatase, receptor type Z, polypeptide 1 | inclusion/exclusion                   | SE         | Oligodendrocyte differentiation and myelination | 23144976        |
| Mag         | myelin-associated glycoprotein | inclusion                                     | SE         | Associated with demyelinating leukodystrophy   | 26179919        |
| Sema6a      | semaphorin 6A    | exclusion                                      | SE         | Oligodendrocyte differentiation and myelination | 2277942         |
| Ndirg2      | N-myc downstream regulated gene 2 | inclusion                                    | SE         | Mainly expressed in astrocytes, regulates neurogenesis, associated with neurodegeneration | 26341979        |
| Ank3        | ankyrin 3, epithelial | exclusion                                    | SE         | Genetic risk factor for psychiatric disorders; cytoskeletal scaffolding and synaptic protein | 25374361        |
| Daam2       | dishevelled associated activator of morphogenesis 2 | inclusion                             | SE         | Suppresses oligodendrocyte differentiation during development, elevated in Guillain-Barré syndrome | 25754822 26293489 |
| Epb4.1L2    | erythrocyte protein band 4.1-like 2 | inclusion                                | SE         | Organization of internodes in peripheral myelin | 22291039        |
| Map4k4      | mitogen-activated protein kinase kinase kinase kinase 4 | inclusion                             | SE         | Neurite outgrowth and retraction               | 18007665        |
| Myo6        | myosin VI        | exclusion                                      | SE         | Hippocampal synaptic transmission              | 16819522        |
| Map4        | microtubule-associated protein 4 | inclusion                                   | SE         | Microtubule-stabilizing activity in oligodendrocytes and astrocytes | 15642108        |
| Bcas1       | breast carcinoma-amplified sequence 1 | exclusion                               | SE         | Among most abundant transcripts in myelin      | 27173133        |
| Fgfr3       | fibroblast growth factor receptor 3 | inclusion                               | RI         | Involved in neurogenesis, associated with hypochondroplasia, medial temporal lobe dysgenesis and focal epilepsy | 26180198 24630288 |
| Map2        | microtubule-associated protein 2 | exclusion                               | SE         | Microtubule-stabilizing activity and regulation of microtubule networks in dendrites | 15642108        |

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We have also quantified the exon inclusion levels normalized over a constitutive exon present in all isoforms for each of those three genes and a significant reduction was observed ($p < 0.001$, Fig. 4a,b,c).

**Quaking regulates the alternative splicing of its pre-mRNA.** Remarkably, among the most significant events detected were exons of *qkl* itself, indicating a switch of isoform usage in *QKI*FL/FL,*-Olig2-Cre* mice. The alternative splicing of exons 7 and 8 in the 3′ UTR of the *qkl* gene generates the different QKI isoforms. In fact, exons 7c and 8 as well as the 3′ UTR of the *qkl*-5 isoform were significantly upregulated, while exons 7a, 7b and the 3′ UTRs of *qkl*-6 and *qkl*-7 isoforms were significantly downregulated in *QKI*FL/FL,*-Olig2-Cre* mice compared to *QKI*FL/FL*-* mice (Fig. 5a, sashimi plots are shown in Supplementary Fig. S4d). We also searched the *qkl* gene for QREs and we were able to identify many that mapped in intronic or exonic sequences. We identified 24 QREs present in the 3′ alternatively spliced region of *qkl* that harbour either a perfect QRE or differ by 1 mismatch (Fig. 5b and Supplementary Table S9). We next designed primer pairs aimed at amplifying the different *qkl* isoforms with their locations and expected products shown in Supplementary Table S10. We validated the alternative splicing of *qkl* pre-mRNA by RT-PCR using RNA from brains of three mice each for *QKI*FL/FL,*-Olig2-Cre* and *QKI*FL/FL*-* mice. Since exon 6 of *qkl* mRNA is common to all the isoforms, we designed a primer (Fig. 6a; primer #1) that is located in this exon and performed PCR with primers that map to either *qkl*-5 (Fig. 6a; primer#4), *qkl*-6 (Fig. 6a; primer#3) or *qkl*-7 (Fig. 6a; primer#2) specific exon 7 as well as in *qkl*-5 specific exon 8 (Fig. 6a; primer#5) and 3′-UTR (Fig. 6a; primer#6). *qkl*-5, *qkl*-6 and *qkl*-7 share overlapping sequences in the 3′-end and

**Figure 4.** Alternative splicing event in Bcas1, Sema6a and Capzb in *QKI*FL/FL,*-Olig2-Cre* and *QKI*FL/FL*-* mice. (a,b,c) Left: IGV views of the exon skipping events in *Bcas1*, *Sema6a* and *Capzb*. All samples are represented on the same scale for each screenshot. Black arrows represent the primers used for validation by RT-PCR. Right: Validation of exon skipping events by semi-quantitative PCR in $n = 3$ mice per genotype. The constitutive exon is present in all isoforms of the gene and is not alternatively spliced. The pixel densities of PCR bands from 3 independent mice brain RNA samples were quantified using ImageJ software, normalized over wild-type mice and represented as mean ± SEM (Student $t$-test, **$p < 0.001$).
therefore these primers amplify regions in each of these transcripts with the size of the PCR product indicating which qkI isoform is being amplified. In QKI FL/FL;Olig2-Cre; mice, we observed a decrease in the qkI-7 isoform as evident by the lower PCR product obtained from primers #1 and #2 that corresponds to exon 6; exon 7a (Fig. 6b). The upper band in Fig. 6b corresponds to a product if the intron was retained (unspliced pre-mRNA) and shows a non-significant increase in QKI FL/FL;Olig2-Cre; mice. Primers #1 and #3 show a decrease in the qkI-6 isoform (the lower PCR product in Fig. 6c that corresponds to exon 6; exon 7b). The upper band corresponds to the 3′-UTR of qkI-7 which shows a significant decrease in the QKI FL/FL;Olig2-Cre; mice. Primers #1 and #4 show a significant increase in qkI-5 isoform (Fig. 6d, lower band, exon 6; exon 7c) with a significant decrease in the 3′-UTR of qkI-6 (Fig. 6d, upper band). A significant increase in qkI-5 isoform is shown using primers #1 and #5 (Fig. 6e, exon 6; exon 7c; exon 8) as well as an increase in the 3′-UTR of qkI-5 using primers #1 and #6 (exon 6; exon 7c; exon 8; 3′-UTR). We have also quantified the exon inclusion/exclusion levels normalized over a constitutive exon of qkI (in exon 1) shown on the right of each semi-quantitative PCR (Fig. 6b,c,d,e). The observed decrease in the inclusion of exons 7a (qkI-7) and 7b (qkI-6) and the increase in the inclusion of exon 7c of qkI-5 were confirmed by RT-qPCR (Supplementary Fig. S3a). The deletion of exon 2 generates a frame-shift resulting in a null allele and therefore a complete lack of QKI proteins as previously shown2. The loss of QKI-6 and QKI-7 specific exons and 3′-UTRs in QKI FL/FL;Olig2-Cre; brains indicates that expression of the nuclear QKI-5 isoform is required to promote the inclusion of exons 7a (qkI-7) and 7b (qkI-6) as well as their 3′-UTRs. To confirm this, we transfected plasmids expressing green fluorescent protein (GFP) alone or GFP-QKI-5, -QKI-6, -QKI-7, mutated versions of QKI-5 that either cannot dimerize (GFP-QKI-5:E148G), or a version that cannot bind RNA (GFP-QKI-5:V157E) in HEK293T cells and assessed the inclusion of exon 7b of human QKI-7 by RT-PCR. Note, in humans the qkI-7 exon is 7b and not 7a as in mice (Fig. 7a). We observed an increase in the Hqk-7 isoform (exon 6; exon 7b) only when GFP-QKI-5 was transfected, but not when GFP-QKI-6, GFP-QKI-7 or the mutated versions of QKI-5 were transfected (Fig. 7a,b). We have also quantified exon 7b (Hqk-7) inclusion levels normalized over a constitutive exon of qkI (exon 1), and observed a significant increase in exon 7b inclusion when QKI-5 was transfected (Fig. 7b, right). The increase observed when GFP-QKI-7 was transfected is due to the primer being in exon 7b of the cDNA in pGFP-QKI-7, recognizing the ectopically expressed isoform. Equivalent QKI protein expression was confirmed by immunoblotting (Fig. 7c). These findings show that QKI-5 promotes the inclusion of exon 7b of Hqk-7 in cellulo.

Figure 5. Alternative splicing patterns of qkI in QKI FL/FL;Olig2-Cre; and QKI FL/FL; mice. (a) IGV view of the alternative splicing events in qkI. All samples are represented on the same scale for each screenshot. (b) A schematic of the 3′-UTR of mouse qkI gene showing the location of Quaking Response Elements (QREs) identified. QREs are either a perfect match to consensus QRE or differ by one mismatch.

Figure 6. Alternative splicing patterns of qkI in QKI FL/FL;Olig2-Cre; and QKI FL/FL; mice. (a) Genomic coordinates (- strand) for qkI and Exons 6 and 7. (b) Exons and introns of qkI in QKI FL/FL;Olig2-Cre; mice. (c) Exons and introns of qkI in QKI FL/FL; mice. (d) Exons and introns of qkI in QKI FL/FL;Olig2-Cre; mice. (e) Exons and introns of qkI in QKI FL/FL; mice.
Figure 6. QKI proteins regulate self-splicing in mice. (a) A schematic of the 3′-UTR of mouse qki gene. Arrows indicate location of primers used for splicing analysis. Primer 1 is in exon 6 that is common to all QKI isoforms. The lower band corresponds to qki-7 isoform (exon 6; 7a) whereas the upper band corresponds to the unspliced pre-mRNA. (Right) The pixel densities of PCR bands from 3 independent mice brain RNA samples were quantified using ImageJ software, normalized over constitutive exon, and then normalized over wild-type mice and represented as mean ± SEM (Student t-test, **p < 0.001). (b) (Left) RT-PCR analysis using primers #1 and #2 from n = 3 mice brain RNA/genotype. The lower band corresponds to qki-7 3′-UTR. (Right) The pixel densities of PCR bands from 3 independent mice brain RNA samples were quantified using ImageJ software, normalized over constitutive exon and then normalized over wild-type mice and represented as mean ± SEM (Student t-test, *p < 0.05). (c) (Left) RT-PCR analysis using primers #1 and #3 from n = 3 mice brain RNA/genotype. The lower band corresponds to qki-6 isoform (exon 6; 7b) whereas the upper band corresponds to qki-7 3′-UTR. (Right) The pixel densities of PCR bands from 3 independent mice brain RNA samples were quantified using ImageJ software, normalized over constitutive exon and then normalized over wild-type mice and represented as mean ± SEM (Student t-test, *p < 0.05). (d) (Left) RT-PCR analysis using primers #1 and #4 from n = 3 mice brain RNA/genotype. The lower band corresponds to qki-5 isoform (exon 6; 7c) whereas the upper band corresponds to qki-6 3′-UTR. (Right) The pixel densities of PCR bands from 3 independent mice brain RNA samples were quantified using ImageJ software, normalized over constitutive exon and then normalized over wild-type mice and represented as mean ± SEM (Student t-test, *p < 0.05, **p > 0.01). (e) (Left) RT-PCR analysis using primers #1 and #5 (qki-5 isoform: exon 6; 7c; exon 8), primers #1 and #6 (qki-5 isoform: exon 6; 7c; exon 8; 3′-UTR), constitutive qki exon 1, and GAPDH from n = 3 mice brain RNA/genotype. (Right) The pixel densities of PCR bands from 3 independent mice brain RNA samples were quantified using ImageJ software, normalized over constitutive exon and then normalized over wild-type mice and represented as mean ± SEM (Student t-test, *p < 0.05).
The presence of a QKI response element in >1400 mRNAs suggested that the QKI isoforms regulate many genes. In the present manuscript, we perform a transcriptomic analysis using whole brain RNA from QKI FL/FL and QKIFL/FL;Olig2-Cre mice and we identify >1,800 genes that are significantly differentially expressed of which 229 genes show an absolute fold change greater than 2. More than 1400 alternative splicing candidates were identified using two methods, DEXseq and rMATS. Gene ontology analysis of the 229 genes has identified genes involved in glial cell differentiation (OLs), axon ensheathment, and myelination to be the most downregulated. Surprisingly, we identified qkI as the top alternatively spliced event by DEXseq and we show that QKI-5 regulates the alternative splicing of the qkI-6 and qkI-7 isoforms. Inspection of the qkI pre-mRNA reveals the presence of many QREs distributed across the intronic regions of these pre-mRNAs. Our findings define the QKI isoforms as master regulators of RNA metabolism in oligodendrocytes and show that QKI-5 isoform autoregulates alternative splicing of its own gene.

Northern blot analysis shows that the qkI-5 mRNA is highly expressed during embryogenesis and maintained until approximately P14 at which point its expression declines thereafter. In contrast, the mRNA expression of qkI-6 and qkI-7 begins embryonically and peaks at P14 and is maintained into adulthood. Thus our observation in P14 wild type mice (QKIP/FL) that qkI-6 and qkI-7 are the main mRNA isoforms observed is expected. What is unexpected is that qkI-5 mRNA is the major isoform observed at P14 in the brains of QKIP/FL;Olig2-Cre mice. These findings suggest that the complete absence of the QKI proteins favors the splicing of the qkI-5 isoform. It also suggests that the partial qkI promoter deletion observed in the quaking viable mice may decrease the transcriptional output of the qkI gene such that the levels of qkI-5 are insufficient to drive the subsequent pre-mRNA splicing of qkI-6 and qkI-7 and their expression. Therefore, it is the qkI isoform imbalance that blocks cellular differentiation and these findings support the tumor suppressor function of the QKI proteins, as their deletion or isoform imbalance favors cell proliferation. Furthermore, the haploinsufficient expression of the human qkI gene is enough to block cellular differentiation, as observed with its monocyte to macrophage differentiation defect.

We next compared our mouse alternative splice analysis with two other RNA-seq analyses that were performed in human cells. The qkI haploinsufficient patient exhibits intellectual disabilities, hypotonia, seizures, brain anomalies and specific dysmorphic features. She carries a balanced reciprocal translocation (t(5; 6) (q23.1; q26), where a break point in one of her qkI alleles causes reduced expression of QKI mRNA and protein.

**Discussion**

The presence of a QKI response element in >1400 mRNAs suggested that the QKI isoforms regulate many genes. In the present manuscript, we perform a transcriptomic analysis using whole brain RNA from QKIP/FL and QKIP/FL;Olig2-Cre mice and we identify >1,800 genes that are significantly differentially expressed of which 229 genes show an absolute fold change greater than 2. More than 1400 alternative splicing candidates were identified using two methods, DEXseq and rMATS. Gene ontology analysis of the 229 genes has identified genes involved in glial cell differentiation (OLs), axon ensheathment, and myelination to be the most downregulated. Surprisingly, we identified qkI as the top alternatively spliced event by DEXseq and we show that QKI-5 regulates the alternative splicing of the qkI-6 and qkI-7 isoforms. Inspection of the qkI pre-mRNA reveals the presence of many QREs distributed across the intronic regions of these pre-mRNAs. Our findings define the QKI isoforms as master regulators of RNA metabolism in oligodendrocytes and show that QKI-5 isoform autoregulates alternative splicing of its own gene.

Northern blot analysis shows that the qkI-5 mRNA is highly expressed during embryogenesis and maintained until approximately P14 at which point its expression declines thereafter. In contrast, the mRNA expression of qkI-6 and qkI-7 begins embryonically and peaks at P14 and is maintained into adulthood. Thus our observation in P14 wild type mice (QKIP/FL) that qkI-6 and qkI-7 are the main mRNA isoforms observed is expected. What is unexpected is that qkI-5 mRNA is the major isoform observed at P14 in the brains of QKIP/FL;Olig2-Cre mice. These findings suggest that the complete absence of the QKI proteins favors the splicing of the qkI-5 isoform. It also suggests that the partial qkI promoter deletion observed in the quaking viable mice may decrease the transcriptional output of the qkI gene such that the levels of qkI-5 are insufficient to drive the subsequent pre-mRNA splicing of qkI-6 and qkI-7 and their expression. Therefore, it is the qkI isoform imbalance that blocks cellular differentiation and these findings support the tumor suppressor function of the QKI proteins, as their deletion or isoform imbalance favors cell proliferation. Furthermore, the haploinsufficient expression of the human qkI gene is enough to block cellular differentiation, as observed with its monocyte to macrophage differentiation defect.

We next compared our mouse alternative splice analysis with two other RNA-seq analyses that were performed in human cells. The qkI haploinsufficient patient exhibits intellectual disabilities, hypotonia, seizures, brain anomalies and specific dysmorphic features. She carries a balanced reciprocal translocation (t(5; 6) (q23.1; q26), where a break point in one of her qkI alleles causes reduced expression of QKI mRNA and protein.

**Figure 7.** QKI-5 regulates self-splicing in vitro in human cells. (a) A schematic of the 3′-UTR of human Hqk gene. Arrows indicate location of primers used for RT-PCR/splicing analysis. (b) (Left) HEK293T cells were transfected with expression vectors encoding green fluorescent protein (GFP) alone or expression vectors for GFP-QKI-5, -QKI-6, -QKI-7, -QKI-5:V157E or -QKI-5:E148G. Total RNA was isolated 72 h later and analyzed by RT-PCR (n = 3). The pixel densities of PCR bands from 3 independent transfections were quantified using ImageJ software, normalized over Hqk constitutive exon 1 and represented as mean ± SEM (One-way ANOVA, **p < 0.01, ***p < 0.001). (c) Protein expression of transfected plasmids in HEK293T cells is determined by immunoblotting using the indicated QKI antibodies or β-actin antibody for equivalent loading. The molecular mass markers are shown on the left in kDa and the migration of the GFP-QKI isoforms indicated on the right.
RNA-seq analysis of her monocyes and macrophages compared to her control sibling revealed 1,513 alternative splice events. These spliced events were indeed enriched for QREs next to these regulated exons, suggesting these events are mainly direct targets of the QKI isoforms. Low QKI levels is a prognostic marker for poor survival for patients with NSCLC. The human lung/bronchus epithelial cell line, BEAS-2B, was depleted of the QKI isoforms using shRNAs. RNA-seq was performed and 451 ASE were identified of which 81 were validated by RT-qPCR. Recently, transcriptomic profiling and alternative splicing analysis was performed in wild type versus qki-deficient neural stem cells either in the presence or absence of Pten and Trp53 identifying 388 and 215 genes that are differentially regulated and alternatively spliced, respectively.

Our RNA-seq analysis identified a total of 1438 ASEs (Tables S3 to S5) with 190 events (Supplementary Table S3) that we termed 'stringent', as they were identified by both DEXSeq and rMATS. Comparing our stringent list with Zong et al. and Shingu et al., we have 96, 93, and 46 common gene names, respectively. If we compare all ASEs from all 4 gene sets, we can assemble a common signature of 4 genes in QKI-depleted cells (ADD3, Bin1, MAP4K4 and WDFY3). Interestingly, this list of 4 gene names spans mouse, human, oligodendrocytes, neural stem cells, monocyte/macrophages and lung epithelial cells. The other alternative splice targets are, therefore, more tissue- or cell-specific. Other genes commonly identified in three out of four gene sets include H1P1BP3, MAP3K7, SBF1, SPAG9 and VLDLR.

In sum, we define the QKI RNA binding proteins as master regulators of RNA metabolism driving oligodendrocyte differentiation. Our RNA-seq analysis defines QKI as regulators of alternative splicing and herein we define many spliced targets including self-regulation.

Materials and Methods

Isolation of brain RNA. All mouse procedures were performed in accordance with McGill University guidelines, which are set by the Canadian Council on Animal Care. Experimental protocols involving animal use were approved by the McGill University Animal Care Committee. QKIP1/FL;Olig2-Cre and QKIP1/FL; mice were generated as described. Exon 2 of the qki gene is flanked with loxP sites and the recombination with Cre results in a null. Total RNA was extracted from brains of QKIP1/FL; and QKIP1/FL;Olig2-Cre P14 mice (n = 3 for each genotype) with appropriate amounts of TRIzol® reagent according to the manufacturer’s instructions (Invitrogen). 30 µg of RNA were DNase I (Promega) treated for 30 mins at 37°C.

RNA-Sequencing analysis. Total RNA from QKIP1/FL; and QKIP1/FL;Olig2-Cre brains (n = 3 females for each genotype) were purified using polyA selection and converted into cDNA by Illumina TruSeq and sequenced on a HiSeq 2000 (Illumina) with paired ends 100 bp in length (Genome Quebec, Canada) at an average of 133 million reads/sample. Illumina sequencing adapters were removed from the reads, and all reads were required to have a length of at least 32 bp. Sequencing runs were processed with Illumina CASAVA software. Trimomatic v0.32 was used to trim reads, including removal of low-quality bases at the end of reads (phred33 < 30), clipping of the first three bases and clipping of Illumina adaptor sequences using the palindrome mode. We performed quality trimming with a sliding window, cutting once the average quality of a window of four bases fell below 30. We discarded reads shorter than 32 base pairs after trimming. Trimmed reads were aligned to the reference genome mm 10 using STAR v2.3.0c. Quality control was performed using metrics obtained with FASTQC v0.11.2, SAMtools v1.3, BEDtools v2.27 and custom scripts. This showed a good proportion of high quality reads and mapping rates higher than 90% in all samples. Bigwig tracks were produced with custom scripts, using BEDtools and UCSC tools. Data were visualized using the Integrative Genomics Viewer (IGV) 22.

Gene expression analysis. Expression levels were estimated by featureCounts v1.5.0 using exonic reads and normalized using DESeq2 24. Global gene expression changes were assessed by unsupervised hierarchical clustering of samples and PCA, as previously described. Differential expression analysis between QKIP1/FL; and QKIP1/FL;Olig2-Cre mice was performed using DESeq2 for all UCSC known genes. Genes were considered differentially expressed if they had an adjusted p-value < 0.05 and a base mean >100. Expressed genes were defined as all genes with the DESeq2 base mean higher than the first expression quartile.

Alternative splicing analysis. We used rMATS v3.2.2 to identify alternative splicing (AS) events. Since rMATS requires reads of equal lengths, we performed the analysis on untrimmed reads after verifying that the adaptor content was low in the FASTQC reports. We ran rMATS with standard parameters and analyzed both outputs: AS events identified using reads mapping to splice junctions and reads on target (JC + ROT), and AS events identified using reads mapping to splice junctions only (JC). The list of putative AS events was obtained by including all events with an adjusted p-value (FDR) <0.05 in either JC + ROT or JC. We also computed differential exon usage with DEXSeq v1.12.2. To generate counts compatible with DEXSeq, we used featureCounts to count reads mapping to all annotated exons in Ensembl GRCm38.44 and converted them to the standard DEXSeq input format using scripts available from Vivek Bhardwaj (Subread_to_DEXSeq, https://github.com/vivekbhr/Subread_to_DEXSeq). Differential exon usage analysis between WT and QKI KO mice was performed using standard DEXSeq parameters. Exons with a minimum length of 15 nucleotides and with an adjusted p-value >0.05 were considered statistically significant.

To obtain a reduced set of stringent, robust ASE candidates, we combined predictions from DEXSeq and rMATS, since the underlying methods and the type of information (read counts, splice junction reads, etc) used by each approach are quite dissimilar. Significant events called both by rMATS (p-value <0.05) and by DEXSeq (adjusted p-value <0.05) were intersected based on genomic location. For this, we first merged the two rMATS outputs (JC + ROT and JC) so that each AS event was represented by one entry. We then used the BEDTools intersect function to identify the genomic regions of overlap between the results from rMATS and DEXSeq, keeping
the information from rMATS (Table S3). Since a number of exons are present as multiple entries in the rMATS outputs (for each isoform they belong to), we then intersected (BEDTools intersect) the results from rMATS and DEXSeq with the GENCODE VM11 annotation to identify all unique AS exons. AS events that were identified by rMATS and DEXSeq but that were not annotated in GENCODE VM11 were then added to the list. Exons with identical genomic coordinates were merged and counted only once. This approach allowed us to count the number of unique AS events without duplicates. To obtain a more comprehensive list of AS exons, we generated the union of the DEXSeq (adjusted p-value < 0.05) and rMATS (adjusted p-value < 0.05) and for both the stringent and the comprehensive lists, we then used the GENCODE annotation to distinguish SE events corresponding to cassette exons compared to first or last exons. Exons were considered to be “cassette exons” if they were not the first or last exon in any of the gene isoforms or “mixed exons” if they were not the first or last exon in at least one isoform. SE events identified by rMATS that were not included in the GENCODE annotation were counted as “mixed” exons. This annotation is indicated in the last columns of Tables S3 and S5. Exons that were first or last exons in all isoforms were excluded from further analyses.

**Gene Ontology.** GO term enrichment analysis was performed GOriilla36. The lists of differentially expressed genes and genes containing AS events were compared to a background list of expressed genes, consisting of all expressed genes in the complete dataset (see Gene expression analysis above). For differentially expressed genes, upregulated and downregulated genes with a base mean higher than 100 and an absolute fold change greater than 2 were used for the analysis. For AS, we used the genes in which stringent AS events (detected by DEXSeq and rMATS, see above) were observed.

**Motif enrichment analysis.** We searched for the motif ACUAA in windows of 200 nucleotides surrounding each side of the identified SE events corresponding to “cassette” or “mixed” exons (see Alternative splicing analysis above). To verify enrichment of this motif in SEs and compute empirical p-values, we generated a set of control exons (n = 88,217) that met the following criteria: “cassette” or “mixed” exons, good coverage at the exon and gene level (coverage >lowest quartile of exon or gene coverage normalized by exon or gene length), not alternatively spliced (DEXSeq adjusted p-value > 0.99 and rMATS p-value < 0.05) and with a minimum length of 15 nucleotides. To compute empirical p-values, we performed 1,000 iterations, where in each iteration we randomly selected a number of control exons equal to the number of observed AS events, and computed the frequency of occurrence of the ACUAA motif within neighbouring windows of 200nt in these controls exons. Motif analyses were performed successively for all SEs, SEs with higher inclusion (QKI-repressed) in KIP1/FL/FLg-Cre mice and SEs with higher exclusion in KIP1/FL/FLg-Cre mice (QKI activated). To determine the position of the core motif relative to the exon, we used the FIMO tool37 (MEME suite) and searched for exact matches to ACUAA.

**Transfections.** HEK293T cells were plated in 6-well plates and transfected with 2 µg of the indicated GFP expression vectors encoding QKI-5, QKI-6, QKI-7, QKI5:148G and QKI5:157E. The common forward primer for RT-PCR detection of qkI alternative splicing was in exon 6 and is the following: 5'-CCT CAC CCA ACT GCT GCA ATA G-3'. The reverse primer to detect exon 7 of QKI-7 inclusion is: 5'-CAT GAC TGG CAT TTC AAT CC-3'. Human constitutive exon: forward, 5'-GTC GGG GAA ATG GAA ACG AAG-3' and reverse, 5'-GGT TGA AGA TCC CGC AGA AG-3'. Human GAPDH forward: 5'-CCG CAT CTT CTT TTG CTT CGT CG-3'. Human GAPDH reverse: 5'-TGG GTG TCT CCG TGT TAA TC-3'.

**Reverse transcript - qPCR.** 1 µg of RNA was reverse transcribed using oligo(dT) primer and M-MLV reverse transcriptase according to the manufacturer's protocol (Promega). cDNAs were then amplified by PCR using the following conditions: 95 °C for 30 seconds, 60 °C for 30 seconds, 72 °C for 1 minute and repeated for 30 cycles. The PCR products were then separated by agarose gel electrophoresis. For real-time PCR, primers were designed and efficiency tested according to the MIQE guidelines. Real-time PCR was performed in triplicates with a 1:4 dilution of cDNA using SyBR Green PCR Mastermix (Qiagen, Valencia, CA). All quantification data were normalized to GAPDH using the ΔΔCt method. All quantification data were normalized to GAPDH using the ΔΔCt method or for the qkI isoforms they were normalized to a constant exon region.

**Primers.** Mouse 1: 5'-TCC TTG AGT ACC CTA TTG AAC CC-3'; Mouse 2: 5'-GGG CTG AAA TAT CAG GCA TGA C-3'; Mouse 3: 5'-TTA GCC TTT CGT TGG GAA AGC-3'; Mouse 4: 5'-GGT CTG CGG TCA CAA TCC TTG G-3'; Mouse 5: 5'-TTG AGT TCC TGG CCG TGG C-3'; Mouse 6: 5'-GGC CTT CTT ACC GTT CG-3'; Mouse constitutive exon (exon 1 in all isoforms): forward, 5'-GTC GGG GAA ATG GAA ACG AAG-3' and reverse, 5'-GGT TGA AGA TCC CGC AGA AG-3'. Mouse GAPDH forward 5'-CCC TCG TCT GCT TCA TCA GG-3'; Mouse GAPDH reverse: 5'-CCA TAC GGC CAA ATC CTT CGT TC-3'. Mouse Bacs1 forward: 5'-GAC ACT GCT TTC ACC TAA CAA G-3'. Mouse Bacs1 reverse: 5'-GTA GCC TTT GTC TCT CCT C-3'. Mouse Bacs2 constitutive forward: 5'-CCG CAT CTT GAT GGC GTT CCT CA-3'. Mouse Bacs1 constitutive reverse: 5'-CCA CGG AGT TTC AGC TTT GG-3'. Mouse Sema6a forward: 5'-GAG ACT GCT ACA ATT CCT CGT C-3'. Mouse Sema6a reverse: 5'-GAA CCA GGT GGT CTG TGC-3'. Mouse Sema6a constitutive forward: 5'-CCA AAC ATG CCA ACA TCG C-3'. Mouse Sema6a constitutive reverse: 5'-TCA CAG TCC TGA GGG TAG GG-3'. Mouse Capzb forward: 5'-CTG GTG GAC ATG GAA AAC-3'. Mouse Capzb reverse: 5'-CTG CTG CTT CCT CCT CTT CAA GG-3'. Mouse Capzb constitutive forward: 5'-TGG GTG GAA GTG CAG GAG-3'. Mouse Capzb constitutive reverse: 5'-AGC CGG ATT TGT TGG TTT GC-3'.
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Additional Information
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