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Proteomic analysis of pRb loss highlights a signature of decreased mitochondrial oxidative phosphorylation

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The retinoblastoma tumor suppressor (pRb) is functionally inactivated in many human cancers [Weinstein et al. 2013; Zack et al. 2013; Leiserson et al. 2015]. This fact is often attributed to pRb’s role as a repressor of E2F-dependent transcription. pRb is primarily a nuclear and chromatin-associated protein, and loss of pRb function is associated with deregulated expression of genes that provide important functions during cell proliferation [Nevins 2001; Hanahan and Weinberg 2011]. In addition to its interactions with E2F, pRb associates with a variety of transcription factors and chromatin-associated proteins [Morris and Dyson 2001]. Indeed, pRb has been implicated in the assembly of regulatory complexes that can either repress or activate transcription [Markey et al. 2002; Lee et al. 2006; Hutcheson et al. 2014]. The direct targets of pRb are diverse. In addition to cell proliferation genes, well-studied targets function in cell differentiation [Calo et al. 2010], cell metabolism [Nicolay and Dyson 2013], and apoptosis [Ianari et al. 2009]. Such transcriptional changes may help to explain why mouse Rb1−/− tissues have defects in differentiation [Jacks et al. 2001].

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sues shortly after proteomics. We generated proteomic profiles of mouse tissues in mass spectrometry-based quantitative analysis shows that carbon metabolism. The collective effect of this metabolic changes that are strikingly different from the RNA stress (Nicolay et al. 2013). An unexpected aspect of these results give a global perspective of the cellular consequences of acute Rb/RB1 loss and indicate, surprisingly, that the most prominent proteomic change common to RbKO cells involves a decrease in mitochondrial oxidative phosphorylation (OXPHOS) function.

Results

Rb ablation alters RNA and protein levels

We examined the consequences of pRb loss in an in vivo model. Mice were bred to introduce a tamoxifen-inducible form of Cre recombinase [ROSA26SorERt2(cre/Esr1)Tyj] into the genetic background of either Rb wild-type (Rb+/+) or Rb-floxed (Rb+/−) alleles. These animals were injected with tamoxifen, ablating Rb in Rb+/− mice and generating RbKO tissues. pRb was readily apparent in Rb+/− tissues but was undetectable by immunohistochemistry in RbKO tissues (Supplemental Fig. 1A–D). The effects of Rb ablation were followed by quantitative PCR (qPCR) analysis of a set of well-characterized pRb/E2F-regulated transcripts (Fig. 1A,B). These RNAs were elevated in lung and colon tissues after 3 d of tamoxifen dosing (time point = 0 h) [Fig. 1A,B]. This change was escalated and sustained in both tissues 120 h after the final tamoxifen dose and correlated with the loss of pRb [Supplemental Fig. 1A–D].

Previous work has shown that Rb ablation leads to the appearance of ectopic, actively cycling cells in the mouse colon but not in the lung (Meuwissen et al. 2003; Haigis et al. 2006). Although E2F targets are elevated in both tissues, the varied cell cycle effects indicate that the impact of Rb ablation differs between well-differentiated lung tissue and the less-differentiated population of cells in colonic crypts. Consistent with previously published data, RbKO colons contained many ectopic mKi67-positive cells, a marker of actively cycling cells, within 96 h of the final tamoxifen exposure [Fig. 1C,E]. In contrast, the percentage of mKi67-positive cells in RbKO lung tissue remained very small 120 h after tamoxifen dosing [Fig. 1D,F].

qPCR analysis showed elevated RNA transcripts from MCM2 (Fig. 1A,B), a known E2F target gene, 24 h post-tamoxifen in both tissues, and a correlative, significant increase in protein was detectable by immunohistochemistry 96 h post-tamoxifen [Fig. 1G,H]. This is consistent with the established fact that loss of pRb elevates E2F-dependent transcription. Despite the fact that mKi67 and MCM2 mRNAs were elevated in both tissues [Supplemental File 1], mKi67 protein was elevated only in the RbKO colon. This distinction shows that transcriptional changes resulting from Rb deletion are not necessarily
accompanied by similar changes in proteins. To learn whether this a general feature or a unique property of mKi67, we performed quantitative proteomic and RNA sequencing analyses on the \( Rb^{+/-} \) and \( Rb^{KO} \) tissue samples and obtained a more extensive picture of the effects of pRb loss.

The effects of Rb loss on the transcriptome

RNA sequencing revealed both tissue-specific RNA changes and changes evident in both \( Rb^{KO} \) tissues [Supplemental File 1]. We focused on changes that were common between \( Rb^{KO} \) tissues because these are more likely to represent broadly relevant features of Rb function [Table 1; Supplemental File 1]. As expected, gene signature enrichment analysis [GSEA] showed that the RNAs elevated in both \( Rb^{KO} \) colon and \( Rb^{KO} \) lung samples encode proteins associated with DNA replication and cell cycle progression [Table 1, Supplemental File 1; Subramanian et al. 2005]. Leading-edge RNAs were overlaid with publicly available pRb chromatin immunoprecipitation [ChIP]-chip data [Chicas et al. 2010], and this confirmed that many of the enriched up-regulated RNAs were from direct pRb/E2F targets [Supplemental File 2]. As expected, RNAs associated with actively cycling cells, such as \( mKi67, E2f1, CDK2, MCM3, MCM2, \) and \( MCM6 \), were significantly elevated in \( Rb^{KO} \) colons. Although \( Rb^{KO} \) lung tissue contains few actively cycling cells, these same RNAs were also significantly elevated, consistent with the idea that ablation of Rb deregulates the transcription of genes that promote proliferation. Because similar changes were seen in both \( Rb^{KO} \) lungs and colons, these transcriptional events appear to reflect pRb loss rather than the proliferative index of the tissue.

Leading-edge analysis of the RNAs down-regulated in both \( Rb^{KO} \) tissues shows an enrichment for genes encoding ribosomal proteins and components of lysosomal pathways [Table 1]. Few RNAs that decreased in this manner were transcribed from previously identified pRb/E2F-bound promoters. However, those that were from direct pRb/E2F targets were also significantly enriched for ribosomal and lysosomal functions [Supplemental File 2]. Several studies have described transcriptional signatures associated with the loss of pRb [Muller et al. 2001; Ren et al. 2002; Witkiewicz et al. 2012], and our results are consistent with these reports: pRb loss increased the transcription of a set of E2F targets and proliferation-related genes and decreased a less-well-characterized group of transcripts.
Global analysis of the effects of Rb loss on the proteome in vivo

In parallel to the RNA analysis, we used multiplexed quantitative proteomics and tandem mass tag (TMT) technology to identify proteome changes caused by Rb ablation (Supplemental Material; Thompson et al. 2003; Ting et al. 2011; McAlister et al. 2014). Proteome profiles were generated using colon and lung samples from six separate mice per genotype. After peptide alignment, we identified 8063 proteins [present in both colon and lung samples] for which we had corresponding RNA sequencing data [Supplemental File 3]. Next, we calculated a log2 fold change ratio \( \frac{\text{RbKO}}{\text{RbWT}} \) for protein and RNA in each tissue and compared these data [Fig. 2B,C; Supplemental File 3].

Figure 2. Loss of Rb leads to distinct changes in proteins and transcripts. (A) Key of the plots in B–J. (B–J) RNA and protein levels were normalized, and a fold change of \( \frac{\text{RbKO}}{\text{RbWT}} \) was taken and log2 transformed (0 value = ratio of 1/1). Eight-thousand-sixty-three gene products (RNA and protein) were correlated from the effects of Rb loss. The data shown are from four fold change ratios of biological replicates. Proteomics were from six mice. Little correlation between RNA and protein was detected in either the lung (B) or the colon (C). (D,E) E2F targets [blue] show increased RNA but little correlation in protein in the lung. (F) Mitochondria proteins decrease in the RbKO lung. (G,H) E2F targets [blue] show increased RNA and a correlation in protein in the colon. (I) Mitochondria proteins decrease in the RbKO colon. (J) Representative ribosome fractionation plot from the RbWT lung. (K,L) qPCR analysis of ribosomal fractions of specific genes from the colon and lung upon loss of Rb. Error bars are the 95% confidence intervals. Statistical significances are shown as follows: (*) \( P < 0.05 \); (**) \( P < 0.02 \). Statistical differences are between RbKO and RbWT animals.
This integrated data set gives a detailed picture of the consequences of Rb loss. A striking feature is that, while many of the protein:RNA changes in RbKO tissues show a nearly 1:1 ratio (log2 value of 0), a substantial number of changes seen in the RNA levels were not evident in the protein levels, and vice versa [Fig. 2B,C]. These disparities were present in both tissues [Fig. 2B,C]. The RNA sequencing and proteomic data were highly reproducible [Pearson’s correlation of >0.75 of biological replicates] (Supplemental Fig. 2), but the overall correlation between the changes in RNA and protein following Rb loss was much lower [Pearson’s correlation of 0.20 of biological replicates] [Fig. 2B,C, Supplemental Fig. 2]. Within these data, mKi67 and MCM2 showed the changes predicted from our initial experiments [mKi67 and MCM2 RNAs were elevated in both RbKO tissues, and MCM2 protein was elevated in both RbKO tissues, but mKi67 protein was elevated only in the RbKO colon] [Supplemental File 3].

Analysis of E2F/pRb targets

Since E2F regulation is the best-known function of pRb, we first assessed the overall impact of Rb loss on E2F/pRb targets. We made a list of 358 pRb/E2F-dependent RNAs (“E2Fs”) [Supplemental File 3] from reported ChIP and expression array data in the GSEA database [Subramanian et al. 2005]. mRNAs from multiple pRb/E2F targets were elevated in both RbKO lungs and RbKO colons [Fig. 2D,G, right quadrant, X-axis], but only a subset increased in both tissues [Supplemental File 3]. This may reflect a context-dependent regulation of E2F-regulated transcripts and is consistent with previous work showing that pRb is rate-limiting at only a small subset of E2F target genes [Hurford et al. 1997]. We identified 27 pRb/E2F targets that were transcriptionally up-regulated in both tissues, but, as with mKi67 and MCM2, this had variable impacts on the proteins. These proteins were more significantly increased in RbKO colons [Fig. 2E,H, Supplemental Fig. 3A,B].

To test whether pRb loss affects translation of individual mRNAs, ribosome fractionation analysis was performed on RbKO and control tissues. We examined the distribution of mRNA for CDK2 [increased RNA and protein in RbKO tissues] and MCM3 [increased RNA and no increase in protein in either RbKO tissue]. Consistently, CDK2 mRNA was more enriched within the polyribosome fractions of both RbKO tissues compared with Rb+/+ tissues [Fig. 2J–L], suggesting that the loss of pRb increased the proportion of CDK2 mRNA that was actively translated. Conversely, MCM3 transcripts were distributed between both monosome and polyribosome fractions of control tissues but switched to being largely present in the polyribosome fractions in RbKO tissues [Fig. 2J–L]. These changes show that Rb loss affects not only the level of RNAs but also the extent to which specific transcripts are actively translated. This agrees with previous work showing that translational regulation of specific E2F-driven mRNAs can prevent the aberrant appearance of the protein [Miles et al. 2014].

Taken together, these results support the idea that Rb mutation increases the levels of E2F-regulated mRNAs and that this occurs in both proliferating and nonproliferating cells. However, many of the classic E2F targets that had increased RNA in both the lung and colon displayed a corresponding increase in protein in the RbKO colon but not in the RbKO lung. In part, this distinction may reflect differences in mRNA translation and raise the possibility that the increased levels of cell cycle proteins seen in RbKO tissues may often be a consequence, rather than solely a cause, of increased cell proliferation.

A proteomic signature of Rb loss highlights changes in mitochondrial proteins

Given that increased protein production from the classic E2F-regulated transcripts was not a universal consequence of Rb loss, we looked for a proteomic signature of Rb inactivation that was present in both RbKO tissues. Functional classification of proteins elevated twofold or decreased onefold in RbKO tissues was carried out using DAVID [Supplemental Material]. Several categories of proteins were modestly enriched in the lists of proteins showing a twofold increase in both RbKO colons and lungs [Table 2]; however, the most striking pattern was observed among the proteins that decreased in both RbKO tissues. This list showed a strong enrichment for multiple categories of proteins that function in mitochondria [enriched terms included mitochondrial inner membrane, organelle inner membrane, mitochondrion, oxidative phosphorylation, mitochondrial envelope, mitochondrial membrane, Table 2. DAVID enrichment analysis of protein changes common to both RbKO tissues

<table>
<thead>
<tr>
<th>Enrichment analysis of increased proteins</th>
<th>FDR Q-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>GOBP_Protein transport</td>
<td>0.0023</td>
</tr>
<tr>
<td>GOBP_Establishment of protein localization</td>
<td>0.0026</td>
</tr>
<tr>
<td>GOBP_Protein localization</td>
<td>0.0053</td>
</tr>
<tr>
<td>GOBP_Cell cycle</td>
<td>0.01</td>
</tr>
<tr>
<td>GOBP_DNA metabolic process</td>
<td>0.028</td>
</tr>
<tr>
<td>KEGG_RNA polymerase</td>
<td>0.024</td>
</tr>
<tr>
<td>KEGG_Pyrimidine metabolism</td>
<td>0.041</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Enrichment analysis of decreased proteins</th>
<th>FDR Q-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>KEGG_Parkinson’s disease</td>
<td>0.00029</td>
</tr>
<tr>
<td>GOCC_Mitochondrial inner membrane</td>
<td>0.00062</td>
</tr>
<tr>
<td>GO_CC_Oranelle INNER MEMBRANE</td>
<td>0.00014</td>
</tr>
<tr>
<td>GO_CC_mitochondrial</td>
<td>0.00018</td>
</tr>
<tr>
<td>KEGG_Alzheimer’s disease</td>
<td>0.00019</td>
</tr>
<tr>
<td>GO_CC_Mitochondrial part</td>
<td>0.00062</td>
</tr>
<tr>
<td>KEGG_Oxidative phosphorylation</td>
<td>0.00069</td>
</tr>
<tr>
<td>GO_CC_Mitochondrial envelope</td>
<td>0.0011</td>
</tr>
<tr>
<td>GO_CC_Mitochondrial membrane</td>
<td>0.0014</td>
</tr>
<tr>
<td>GO_CC_Oranelle envelope</td>
<td>0.0026</td>
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<tr>
<td>GOCC_Respiratory chain</td>
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<tr>
<td>GO_CC_Organelle membrane</td>
<td>0.0058</td>
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<tr>
<td>GO_CC_Secretory granule</td>
<td>0.011</td>
</tr>
<tr>
<td>KEGG_Huntington’s disease</td>
<td>0.014</td>
</tr>
</tbody>
</table>

DAVID analysis that used gene ontology (GO) and KEGG pathway enrichment was performed to monitor the effects of Rb1 loss on RNAs from both the lungs and colons. FDR Q-value is a P-value corrected for multiple hypothesis testing.
and respiratory chain) [Fig. 2F,I; Supplemental Fig. 3A,B; Supplemental File 3]. Importantly, unlike the E2F targets and cell cycle proteins, the relative changes in mitochondrial proteins were similar between 
\( Rb^{K0} \) colons and lungs [Fig. 2F,I; Supplemental Fig. 3A,B]. pRb and E2F proteins have been shown to bind to the promoters of several genes encoding proteins that function in mitochondria [Cam et al. 2004]. Although there was a consistent decrease in many mitochondrial proteins in the proteomic data, strikingly, this was not accompanied by a uniform change in RNA levels, with some transcripts being increased in 
\( Rb^{K0} \) tissues, while others decreased [Fig. 2F, I]. This indicates that transcriptional regulation is unlikely to be the sole cause of the mitochondrial changes. Ribosome fractionation profiles showed no significant change in the distribution of 
\( TOMM20 \) mRNA even though levels of this mitochondrial protein marker decreased in both the 
\( Rb^{K0} \) colon and lung [Fig. 2J–L]. This suggests that changes in the levels of mitochondrial proteins likely involve post-translational regulation. Interestingly, several studies have linked pRb to mitochondria, but, confusingly, loss of pRb has been described to have both a positive and cell cycle proteins, the relative changes in mitochondrial mass. Indeed, TOMM20 and VDAC1, two mitochondrial proteins decreased in both 
\( Rb^{K0} \) RPE cells (Fig. 3A). All subcellular changes could be the result of an overall drop in mitochondrial mass per cell. This indicates that transcriptional regulation is unlikely to be the sole cause of the mitochondrial changes.

**Rb loss reduces mitochondria proteins and leads to a general decrease in mitochondrial mass**

Many mitochondria proteins decrease in both the 
\( Rb^{K0} \) colon and lung tissues [Supplemental File 3]. Such extensive changes could be the result of an overall drop in mitochondria mass. Indeed, TOMM20 and VDAC1, two proteins that are often used to approximate mitochondria mass, were reduced in both 
\( Rb^{K0} \) tissues [Supplemental File 3]. However, not all mitochondrial proteins were reduced, and the enrichment of proteins involved in OXPHOS suggests that changes in this process may represent a specific mitochondrial phenotype.

To assess the effects of pRb loss in cultured cells, genetic knockouts of 
\( RB1 \) [\( Rb^{K0} \)] were made by CRISPR/Cas9 technology in the human hTERT-RPE1 cell line because of its use in vivo growth conditions by modifying the DMEM [Cold Spring Harbor Laboratory Press on October 30, 2015 - Published by genesdev.cshlp.org Downloaded from genesdev.cshlp.org on October 30, 2015 - Published by Cold Spring Harbor Laboratory Press]. We began our analysis by assessing the effects of 
\( RB1 \) loss on mitochondria mass using three approaches to avoid ambiguity. First, immunoblot analysis confirmed that, as in the mouse tissues, TOMM20 and VDAC1 were decreased in the 
\( Rb^{K0} \) RPE cell lines [Fig. 3B]. Second, live cells were stained with MitoTrackerGreenFM, a cell-permeable dye that enters polarized mitochondria and fluoresces independently of shifts in membrane potential. Loss of pRb significantly decreased the signal of green fluorescence per cell. [Fig. 3B]. Finally, four unique amplicons from the mitochondrial genome were used to calculate the ratio of mitochondria DNA [mtDNA] to nuclear DNA [NucDNA]. This ratio was also significantly decreased in 
\( Rb^{K0} \) RPE cells [Fig. 3B]. These different assays led to the same conclusion: Ablation of 
\( RB1 \) reduces mitochondrial mass per cell.

The decreased OXPHOS proteins in the proteomic profiles of 
\( Rb^{K0} \) mouse tissues included components of multiple ETC complexes [CI–CV]. Examples included ATP5A (CV), UQCRCC2 (CIII), COX4 (CIV), and NDUFB8 (CI). Similar decreases were evident in 
\( Rb^{K0} \) RPE cells [Fig. 3C], confirming that these effects are conserved between mouse and human cells. Immunoblot analysis provided additional examples of ETC proteins that decreased in 
\( Rb^{K0} \) RPE cells [COX1 (CIV) and SDHB (CII)] [Fig. 3C]. However, the levels of SDHA (CII) and cytochrome c (CIII/CIV) were unaffected [Fig. 3C], indicating some degree of specificity. Accordingly, other mitochondrial proteins [HSP60, PGAM5, and UCP1] were also unaffected by pRb loss in RPE cells [Fig. 3C].

Time-course experiments were performed to examine the kinetics of these changes [Fig. 3D,E]. shRNAs induced knockdown of pRb in both the RPE cells and hTERT-Bj fibroblasts. In both cell lines, acute knockdown of pRb reduced mitochondrial proteins and increased transcription of Cyclin E, an E2F target gene [Fig. 3D,E]. TOMM20 protein decreased in both cell lines within 24 h of shRb hairpin induction and showed a continued decline [Fig. 3D]. Other mitochondrial proteins also decreased but with different kinetics, suggesting a progressive process [Fig. 3D]. Although changes in RNAs were apparent, the initial decrease of TOMM20 protein occurred as TOMM20 mRNA transiently increased. This indicated that the initial loss of TOMM20 protein is unlikely to be transcriptionally linked. We infer that the mitochondrial protein changes cannot be described simply as an “early” or “late” event. Within the resolution of our experiments, the earliest changes to TOMM20 occurred at a time frame similar to the transcriptional up-regulation of Cyclin E. Collectively, these results show that the loss of pRb causes both a reduction in mitochondria mass and decreases of specific OXPHOS proteins.

**pRb loss reduces mitochondrial function**

Oxygen consumption rate (OCR) is used as a measure of mitochondrial activity and a correlate of mitochondrial health. OCRs were taken to ask whether the decrease in OXPHOS proteins has consequences in 
\( Rb^{K0} \) RPE cells. A reduced basal OCR was observed in 
\( Rb^{K0} \) RPE cells and shRb-expressing RPE cells [Fig. 4A,B; Supplemental Fig. 4A]. Similarly, decreased OCR was seen upon pRb

We began our analysis by assessing the effects of 
\( RB1 \) loss on mitochondria mass using three approaches to

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These results remained evident when OCR was normalized to total mitochondrial mass per cell (Fig. 4C), indicating an intrinsic drop in mitochondria activity.

Decreased OCR may reflect a slower rate of carbon oxidation by the cell. Under cell culture conditions, glucose and glutamine are the two primary sources of oxidation. Glucose is oxidized by way of glycolysis en route to the TCA cycle for further oxidation, and glutamine is well known as an anaplerotic nutrient for TCA cycle oxidation in cell culture. Genetic ablation of RB1 in RPE cells led to decreased oxidation of both glucose and glutamine (Fig. 4E–H). This change correlated with little to no decreases in uptake from the extracellular media of both substrates (Fig. 4D; Supplemental Fig. 4C). Significantly, the loss of pRb reduced ATP levels in RPE cells (Supplemental Fig. 4D), further suggesting that pRb loss negatively impacts energy metabolism. Additionally, we found that, independent of RB status, the RPE cells showed little ability for fatty acid oxidation in our cell culture conditions (Supplemental Fig. 4E). Finally, the impact on OCRs was not due to the use of DMEMphyso, as the basal OCR was still found depleted upon pRb knockdown when cell lines were grown in standard DMEM (Supplemental Fig. 4F,G).

These results suggest that the drop in OCRs in RBKO RPE cells reflects a decrease in mitochondrial function. The ratio between the rate of uncoupled respiration and the OCR after oligomycin treatment is a good indicator of mitochondrial function. This ratiometric parameter (analogous to the respiratory control ratio [RCR]) is independent of mitochondrial mass and is sensitive to changes in substrate utilization and mitochondrial membrane proton leaks but not to ATP turnover (Brand and Nicholls 2011). Strikingly, the loss of RB1 significantly reduced the RCR (Fig. 4I).

To ensure that the changes in the RCR were not due to the slight differences in glucose and glutamine uptake by the RBKO RPE cells, permeabilized cell mitochondrial analysis was carried out. This method allows a defined, mechanistic analysis of mitochondrial OXPHOS with fewer of the challenges associated with mitochondrial isolation (Brand and Nicholls 2011). Additionally, this method makes it possible to directly compare the relative substrate capacities of each ETC complex in RBKO RPE cells.
and control cells. RBKO RPE cells demonstrated an ability to use substrates for CI (pyruvate + malate), CII (succinate), and CIV (TMPD/ascorbate) (Fig. 5A). However, substrate utilization was significantly reduced for CI and CII (Fig. 5A). These results could reflect the decrease in mitochondrial mass in RBKO RPE cells. However, if this were the case, then any differences would be neutralized by normalizing to the CI OCR for each genotype. Interestingly, when the OCR of both RBKO and control RPE cells was normalized to CI activity, CII was still functionally reduced in the RBKO cells. In contrast, CIV was capable of higher substrate capacity in the RBKO RPE cells (Fig. 5B). These results were consistent with the RCR analysis and raise the possibility that decreased OXPHOS function of RBKO mitochondria may be linked to decreased proton movement.

Decreased proton pumping or increased proton leak would reduce not only the OCR but also mitochondria polarization. To measure the amount of membrane potential per unit of mitochondrial mass, intact cells were costained with both MitoTrackerGreenFM (measure of mitochondria mass) and TMRE [a lipophilic dye with fluorescence dependent on membrane potential] and monitored by FACS. Strikingly, RBKO RPE cells had 15%–30% less TMRE fluorescence per unit of mitochondria mass per cell when compared with control cells (Fig. 5C). This indicates that the RBKO mitochondria have decreased membrane potential. We conclude that RBKO mitochondria have reduced respiration, and this is likely due to the reduced mitochondrial mass combined with an increase in mitochondrial proton leak.

RB loss alters TCA cycle activity in vivo

OXPHOS is tightly associated with the activity and function of the TCA cycle. Flux through the TCA cycle contributes electrons for reduction along the ETC. Metabolic isotopic enrichment analysis was performed to test whether changes in flux through the TCA cycle contribute to the reduced OXPHOS in RBKO tissues. Using the same population of mice assayed for RNA and protein analysis, we performed in vivo isotopic analysis of U13C-glucose.

Figure 4. Loss of RB1 leads to decreased respiration and mitochondria activity. (A–C) Loss of pRb decreases OCR independently of mitochondrial mass differences. (B, C) RBKO RPE cells. n = 14, repeated twice. (D) Loss of pRB in RPE cells caused little to no difference in 2-NDBG uptake. (E) Extracellular acidification rate (ECAR) measurement in response to glucose challenge. (E–H) OCR measurements in response to glucose (F), glutamine (G), or glutamine + pyruvate (H). For the ECAR and OCR analysis in E–H, n = 14 replicates, repeated twice. (I) The respiratory control ratio is decreased in RBKO RPE cells. n = 14 replicates, repeated twice. Error bars are the 95% confidence intervals. Statistical significances are as follows: (*) P < 0.05; (**) P < 0.02; (***) P < 0.001. Statistical differences are between the effects from RB+− compared with control cells.
in both the colon and the lung using protocols similar to previously published work [Supplemental Material; Fan et al. 2011; Lane et al. 2011; Yuneva et al. 2012; Sellers et al. 2015]. Mice were given a single bolus of U^{13}C-glucose and were sacrificed after 20 min for tissue isolation. An initial time-course analysis found that, 20 min after injection, the levels of U^{13}C-glucose had peaked within the blood, and \(^{13}\)C enrichments were sustained in downstream intermediates of glycolysis and the TCA cycle within both the lung and the colon (Fig. 6A–D). To determine that the glucose clearance would be similar in both \(Rb^{+/+}\) and \(Rb^{KO}\) mice, glucose tolerance tests (GTT) of the same concentrated bolus of U^{13}C-glucose showed no differences between genotypes 96 h after Cre induction (Fig. 6E). These control experiments demonstrated that qualitative differences in glucose-derived metabolites could be determined using this methodology.

In support of the GTT results, analysis of U^{13}C-glucose uptake from the serum revealed no differences between \(Rb^{+/+}\) and \(Rb^{KO}\) colon or lung tissues [Supplemental Fig. 5A]. In contrast, loss of \(Rb1\) induced a significant enrichment (twofold) of M + 2 citrate in both tissues (Fig. 6F).
No difference was seen in the amount of M + 3 pyruvate or M + 3 lactate produced in either tissue (Fig. 6G,H). Similar to RBKO RPE cells (Fig. 4), no increase in glycolysis was detected following Rb1 loss (as indicated by pyruvate and lactate production) (Supplemental Fig. 5B,C). This curious result suggests that RBKO tissues had increased entry of glucose into the TCA. In agreement with this, a ratiometric that qualitatively measures PDH activity (M + 2 citrate/M + 3 pyruvate) was significantly elevated upon loss of Rb1 (Fig. 6I). Furthermore, we observed a significant enrichment of M + 2 acetyl-CoA in both RBKO tissues as well as a significant increase in total acetyl-CoA in the RBKO lung (Fig. 6J,K). Using this technique, we could not qualitatively measure the TCA cycle activity directly. Increased TCA cycle activity would be expected to increase production of ATP. Strikingly, however, a significant decrease in ATP was found in RBKO lung tissue (Fig. 6L). This result was consistent with our model that the RBKO mitochondria have an increased proton leak, which leads to an uncoupling of TCA cycle activity and ATP production via OXPHOS. In contrast, no difference in ATP was observed in RBKO colons (Fig. 6L). As ATP turnover and production are linked to cellular energetic demands, the lack of a difference in ATP in RBKO colons may reflect the increased percentage of proliferating cells in this tissue. Nevertheless, these results show that the loss of pRb in vivo did not produce a glycolytic phenotype, and, despite increased entry of glucose into the TCA cycle, ATP output decreased under comparable energetic conditions.

To support these in vivo results, a similar analysis was performed using the RPE cell lines. Strikingly, RBKO RPE cells showed a significant increase in the enrichment for
M + 2 citrate (Fig. 7A). This effect again correlated with an elevated ratio of PDH activity (Fig. 7B). These results show that Rb/RB1 ablation similarly affects pyruvate oxidation in the TCA cycle in mice and humans, furthermore, this change is irrespective of the effects of Rb/RB1 loss on cell proliferation. Unlike the in vivo analysis, a 24-h pulse of U^{13}C-glucose in the RPE cells produced a fully labeled fraction of U^{13}C-citrate. From this, we estimated the TCA cycle activity as a ratio between the M + 4 citrate and M + 2 citrate. The M + 4 citrate represents the second pass of U^{13}C-enriched citrate through the TCA cycle (Fig. 6A). Thus, a larger ratio would reflect an accelerated TCA cycle. The loss of Rb1 severely decreased this ratio (Fig. 7C), confirming that the TCA cycle was indeed diminished in these cells. As glutamine is also a source of carbon oxidation for the TCA cycle in vitro, we performed similar fate mapping of U^{13}C-glutamine-derived TCA intermediates. Our respiration assays found glutamine oxidation significantly reduced (Fig. 4); however, the assays using U^{13}C-glutamine showed less consistency in RBKO RPE cells (Fig. 7D). Collectively, our data demonstrate that the loss of pRb negatively affects both the TCA cycle kinetics and OXPHOS energetics.

RB loss renders cells sensitive to mitochondrial stress

Given the reduced mitochondrial capacity associated with RB1 mutation, we tested whether these changes could create a cellular vulnerability. A recent study found that cells with defective OXPHOS are more sensitive to mitochondrial challenge when grown in low glucose (Birsoy et al. 2014). Therefore, we cultured control RPE and RBKO RPE cells for 72 h in DMEMPhysio in which the glucose concentration was reduced from 5 mM to 1 mM. Impressively, this nutrient shift by itself led to a 50% reduction in growth in RBKO RPE cells when compared with control cells (Fig. 7E). Additional treatment with two different OXPHOS inhibitors (rotenone and phenformin, both CI inhibitors) further reduced the viability of RBKO RPE cells compared with similarly treated control cells (Fig. 7F). These results strongly support that the loss of Rb/RB1 impairs OXPHOS capacity and show that the mitochondrial changes have physiological consequences.

Discussion

A key goal at the heart of all pRb research is to understand how cells are changed by the loss of pRb. Historically, pRb research has focused on the transcriptional changes associated with the mutation or inactivation of pRb. Here, we describe proteomic changes resulting from pRb ablation. These studies reveal that the effects of Rb loss on the proteome are significantly different from the transcriptional changes. This work provides a new perspective on the cellular consequences of pRb loss, and we highlight two important features of these data.

First, while changes in E2F-driven transcription are the best-known consequence of pRb inactivation, it is striking that proteins encoded by these mRNAs are not the major feature of the proteomic changes. In part, this observation stems from the fact that we looked for common features in two different RbKO tissues: one in which Rb loss causes ectopic cell division, and one in which it does not. Although mRNA from E2F target genes increased similarly in both tissues, altered levels of the encoded proteins were evident only in the proliferative tissue. This suggests

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**Figure 7.** RBKO RPE cells have decreased TCA cycle activity and enhanced sensitivity to mitochondrial stress. (A) RB loss elevates U^{13}C-glucose-derived citrate in RPE cells but has little to no effect on the rates of U^{13}C-glucose-derived pyruvate or lactate. n = 6. (B) The ratio of PDH activity [M + 2 citrate/M + 3 pyruvate]. RB loss elevates PDH activity in RPE cells. (C,D) RBKO RPE cells show reduced TCA cycle oxidation of glucose (C) or glutamine (D). (E,F) RBKO RPE cells are growth-impaired over 72 h when cultured in low-glucose conditions (E) and show significantly enhanced sensitivity to 500 pM rotenone and 10 μM phenformin (F). n = 12 samples per genotype, repeated twice. Error bars are the 95% confidence intervals. Statistical significances are as follows: [*] P < 0.05; [**] P < 0.02; [***] P < 0.001. Statistical differences are between RBKO and RBWT except in F, where it is mock compared with drug treatment.
that there are additional mechanisms to regulate protein synthesis (or accumulation) from canonical E2F targets elevated in nondenuding in $Rb^{-/-}$ tissue. This regulation is not well understood but is clearly of great significance. The miRNA network and RNA-binding proteins that increase in $Rb/RB1$ mutant cells and target cell cycle-regulated transcripts are likely to be involved [Miles et al. 2014]. It may also be relevant that p53, a tumor suppressor commonly inactivated with $Rb$ [Cerami et al. 2012], can regulate translation through modulation of mTOR [Loayza-Puch et al. 2013]. Further studies are needed to investigate these possibilities.

Second, among the proteomic effects of $Rb$ ablation that are common between the lung and colon, the most striking feature is a decrease in multiple mitochondrial proteins. pRb is best known for its effects on important cellular processes such as proliferation, DNA replication, apoptosis, cell differentiation, and senescence. A small number of previous studies have described mitochondrial alterations in $Rb$-deficient cells (Cam et al. 2004; Dali-Youcef et al. 2007; Sankaran et al. 2008; Ciavarra and Zacksenhaus 2010, Blanchet et al. 2011), but these are rarely highlighted in reviews of the pRb literature; given this, it is striking that mitochondrial changes showed the clearest correlation to $Rb$ loss in an unbiased informatics analysis of the proteomic data. Experiments in tissue culture cells confirmed that the effect of $Rb/RB1$ ablation on the levels of mitochondrial proteins is conserved between human and mouse cells.

The mitochondrial changes in $Rb$-deficient cells are a complex phenotype that impacts several processes and has at least two components: a reduction in mitochondrial mass and a change in mitochondrial activity. The drop in mass may reflect decreased mitochondrial biogenesis, increased mitophagy, or a combination of both. Interestingly, independent studies have found that $Rb$ inactivation in erythrocytes [Sankaran et al. 2008] and myoblasts [E. Benevolenskaya, pers. comm.] decreases mitochondrial mass and that increased mitochondrial biogenesis can partially suppress differentiation defects in these cell types. Ciavarra and Zacksenhaus (2010) observed similar mitochondrial defects in $Rb^{-/-}$ myoblasts and found that autophagy inhibitors were sufficient to restore a healthy mitochondrial network and promote differentiation. Collectively, these observations suggest that several of the differentiation defects associated with $Rb$ mutation stem from a failure to generate sufficient mitochondrial function.

In addition to the overall drop in mitochondrial mass per cell, our analysis shows that the OXPHOS capacity of mitochondria is compromised in $Rb^{-/-}$ cells. Metabolic fate mapping of $^{13}$C-glucose and $^{13}$C-glutamine showed that $Rb$ ablation decreased oxidation of downstream derivatives of both carbon sources [Fig. 7]. Additionally, OCRs decreased in $Rb^{-/-}$ cells, and this difference remained even after normalization for mitochondrial mass per cell. We also observed an increased percentage of hypopolarized mitochondria in $RBKO$ cells. The effects on OXPHOS in $RBKO$ cells may stem at least in part from a disruption of the proton gradient. It is intriguing that $Rb^{-/-}$ brown fat cells show increased expression of UCP1, which leads to a hypopolarized state of the mitochondria [Hansen et al. 2004; Nam and Cooper 2015]. While we did not observe this specific change in our systems, such changes are a normal part of thermogenesis. This raises the possibility that pRb may be involved in the thermogenic response such that the inactivation of pRb promotes a hypopolarized mitochondrial network. A second component of this phenotype is suggested by evidence that inactivation of pRb family proteins increases reactive oxygen species [ROS] [Li et al. 2010; Nicolay et al. 2013]. Increased ROS may damage the ETC, decreasing OXPHOS capacity and promoting mitochondrial defects. Further analysis is needed to understand the origins and consequences of the changes in mitochondrial polarization and ROS production in pRb-deficient cells. pRb and E2F proteins are reported to bind directly to the promoters of some genes encoding mitochondrial proteins [Cam et al. 2004]. Although some changes in the levels of mitochondrial proteins were accompanied by changes in transcript levels, approximately half of the changes were not. Time-course experiments revealed that the mitochondrial proteins decrease at very different rates following pRb depletion, with the earliest changes in TOMM20 protein levels occurring prior to any decrease in transcript level. Thus, the mitochondrial phenotype is likely to involve both transcriptional and post-transcriptional regulation and include indirect effects of pRb loss.

These observations illustrate that $Rb$ ablation leads to extensive changes that cannot be predicted or explained by transcriptional profiles alone. The altered properties of pRb-deficient cells in low-glucose medium and in response to mitochondrial poisons indicate that some of these changes have functional consequences.

The notion that mRNAs encoding key cell cycle genes can be elevated in pRb mutant tissue without automatically increasing protein levels suggests one way that $Rb$ inactivation can serve as a priming event for tumorigenesis. Additional events to relieve the proliferative inhibition must presumably be able to compensate for any negative consequence of the mitochondrial changes resulting from $RB1$ inactivation. In most cancers, the mutation of $RB1$ coincides almost universally with concomitant loss of $TP53$ [Weinstein et al. 2013; George et al. 2015; Leiserson et al. 2015]. While the loss of $RB1$ has a negative impact on mitochondrial OXPHOS activity and even causes a slight decrease in glycolysis, the functional loss $TP53$ is known to provide a gain of glycolytic function [Berkers et al. 2013]. In effect, mutation of $TP53$ in $RB1$ mutant cells may jump-start the very pathway that is capable of providing ATP and building blocks in the absence of normal mitochondrial OXPHOS. Indeed, the mitochondrial changes in $RB1$ mutant cells may heighten the selective advantage of $TP53$ mutation. Interestingly, a recent study has shown that OXPHOS deficiency promotes loss of $TP53$, genomic instability, and transformation of neural cells [Bartesaghi et al. 2015]. In addition, genomic instability seen after pRb loss can be partly suppressed by supplementing the culture medium with whole nucleosides [Manning et al. 2014]; we speculate that reduced
mitochondrial OXPHOS capacity may help to promote genomic instability in pRb-deficient cells by reducing macromolecule synthesis.

Importantly, in the absence of TP53 mutation or other similar secondary events, RB mutation renders cells vulnerable to mitochondrial challenges. RBKO RPE cells are more sensitive to mitochondria bioenergetic perturbations generated by changes in the levels of nutrients or by addition of mitochondrial OXPHOS inhibitors [Fig. 7E,F]. Significantly, the sensitivities of the RBKO cells were most evident when cells were cultured with physiological levels of nutrients rather than standard cell culture conditions [Fig. 7E,F]. The fact the RBKO mutant cells are vulnerable to changes in extracellular nutrients may help to explain why Rb mutant mouse embryos are so sensitive to placental defects (Wu et al. 2003). Nutrient sensitivity is an evolutionarily conserved feature, as the loss of pRb orthologs in Drosophila [Nicolay et al. 2013] or Caenorhabditis elegans [Cui et al. 2013] also causes hypersensitivity to fasting. This suggests that the link between pRb and energy sensing is an ancestral feature of pRb function. We speculate that this role may help to modulate pRb function in early G1 and may impact the regulation of cell cycle progression.

Undoubtedly, these proteomic profiles are just a starting point. Further studies are needed to explore each of the novel aspects of the RBKO phenotype and determine how the proteomic changes resulting from Rb/RB1 inactivation vary in different cellular contexts. The proteomic profiles show that Rb/RB1 mutant cells have many features that distinguish them from normal cells, and some of these may provide opportunities for therapeutic targeting. Indeed, these data raise the possibility that the up-regulation of E2F-dependent transcription may not be the most useful or important property of an RB1 mutant cell. Clearly, much remains to be learned about the cellular consequences of Rb/RB1 mutation.

Materials and methods

Cell proliferation assays

Assaying differences in culture recipes Cells were seeded in six-well plates at 1.5 x 10^4 per well and allowed to attach overnight in high-glucose DMEM. Culture medium was then used as specified in the text and figures. In brief, for experiments using nonstandard DMEM, the standard DMEM was aspirated, and cells were washed once with PBS and re-fed with DMEM [5 mM glucose, 0.5 mM L-glutamine, 5% FBS]. Viable cells were counted at the indicated times by Trypan blue (Sigma, T1854) exclusion using a hemocytometer. Each well was counted separately twice, and an average was taken of the four different counts per cell line per condition. After each count, 1.5 x 10^5 cells were reseeded per well in the appropriate medium in new six-well plates.

Additional Materials and Methods are detailed in the Supplemental Material.

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Proteomic analysis of pRb loss highlights a signature of decreased mitochondrial oxidative phosphorylation

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