Inhibition of AMPK and Krebs Cycle Gene Expression Drives Metabolic Remodeling of Pten-Deficient Preneoplastic Thyroid Cells

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Abstract

Rapidly proliferating and neoplastically transformed cells generate the energy required to support rapid cell division by increasing glycolysis and decreasing flux through the oxidative phosphorylation (OXPHOS) pathway, usually without alterations in mitochondrial function. In contrast, little is known of the metabolic alterations, if any, which occur in cells harboring mutations that prime their neoplastic transformation. To address this question, we used a Pten-deficient mouse model to examine thyroid cells where a mild hyperplasia progresses slowly to follicular thyroid carcinoma. Using this model, we report that constitutive phosphoinositide 3-kinase (PI3K) activation caused by PTEN deficiency in nontransformed thyrocytes results in a global downregulation of Krebs cycle and OXPHOS gene expression, defective mitochondria, reduced respiration, and an enhancement in compensatory glycolysis. We found that this process does not involve any of the pathways classically associated with the Warburg effect. Moreover, this process was independent of proliferation but contributed directly to thyroid hyperplasia. Our findings define a novel metabolic switch to glycolysis driven by PI3K-dependent AMPK inactivation with a consequent repression in the expression of key metabolic transcription regulators. Cancer Res; 73(17): 1–14. ©2013 AACR.

Introduction

Highly proliferative conditions, such as embryonic development, tissue regeneration, lymphocyte activation, and neoplastic transformation, require a complex reorganization of energy metabolism to feed cell growth and division (1). A major feature of this metabolic remodeling is the switch from oxidative phosphorylation (OXPHOS) to aerobic glycolysis, a phenomenon that was first observed by Warburg (2). As a consequence, highly proliferating cells exhibit an increase in the amount of lactate production regardless of oxygen availability (3).

It is widely accepted that several oncogenes, including MYC, hypoxia-inducible factor (HIF)-1α, and AKT (4–6), drive this switch by increasing the expression and activity of glycolytic genes, including hexokinase II, lactate dehydrogenase A, pyruvate kinase M2, and pyruvate dehydrogenase kinase 1 (7–9).

The reduced flux through the tricarboxylic acid (TCA) cycle and OXPHOS is not accompanied by compromised mitochondrial function, and is not associated with alterations in the expression levels of genes involved in oxidative metabolism. One exception to this notion is represented by those tumors in which FH, SDH, or IDH are mutated (10).

To characterize the molecular changes resulting from constitutive activation of the phosphoinositide 3-kinase (PI3K) pathway in tissues where this genetic alteration is causally linked to neoplastic transformation, we have generated a mouse model in which loss of the Pten tumor suppressor gene is targeted to the follicular epithelium of the thyroid gland (11). Thyrocyte-specific deletion of Pten constitutively activates the PI3K signaling cascade, leading to hyperplastic thyroid glands at birth, to the development of thyroid follicular adenomas by 6 to 8 months of age (11), and of metastatic follicular carcinomas after 1 year of age (12). The progression to neoplastic transformation is dramatically accelerated by simultaneous Cdkn1b deletion (12), activation of Kras (13), or Tp53 deletion (14).

By analyzing the molecular and metabolic alterations found in thyroids from young, tumor-free Pten<sup>fl/fl</sup> mice, we have discovered a novel mechanism responsible for the active repression of TCA cycle and OXPHOS in preneoplastic thyrocytes. This pathway is independent of both proliferation and of the known pathways classically associated with the Warburg effect. We show in fact that PI3K activation induces, through the inactivation of AMPK, a coordinated repression of the expression of TCA cycle and respiratory genes.
genes, which favors aerobic glycolysis at the expense of OXPHOS.

Materials and Methods

Animals

The Pten^+/+ and TPO-Cre strains have been previously described (11). PDK2/L-2K mice were kindly provided by Dr. Dario Alessi (University of Dundee) (15). All strains were backcrossed in the 129Sv background for at least 10 generations, and littermates were used as controls. RAD001 (Evelolimus; kindly provided by Novartis Institutes for Biomedical Research, Basel, Switzerland) was given daily by oral gavage at a dose of 10 mg/kg body weight for 2 weeks, starting at age 4 weeks. 5-Aminomimidazole-4-carboxamide-1-β-4-ribofuranoside (AICAR; TRC) was injected intraperitoneally (i.p.) at 400 mg/kg/d, for 4 weeks, starting at age 4 weeks.

Measurement of glucose uptake by PET

Wild-type (WT) and Pten^+/−/− mice were fasted overnight before a tail vein injection of 2[^18F]fluoro-2-deoxy-D-glucose (FDG; 300 μCi). One hour after injection, mice were subjected to positron emission tomography (PET) scanning with the Concorde Microsystems R4 microPET Scanner. Animals were imaged while anesthetized with isoflurane. Image acquisition was done using the MicroPET Manager with the ASPIRO-dedicated software.

Primary cultures

Thyroid glands were minced and resuspended in Ham’s F12/10% FBS with 100 U/mL type I collagenase (Sigma) and 1 U/mL dispase (Roche). Enzymatic digestion was carried out for 90 minutes at 37°C. After digestion, follicles were seeded in Ham’s F12 containing 40% Nu-Serum IV (Collaborative Biomedical), gly-his-lys (10 ng/mL; Sigma), and somatostatin (10 ng/mL; Sigma) and allowed to spread for 24 to 36 hours before carrying out the experiments.

Proliferation analysis

Anti-Ki67–stained thyroid sections were photographed at x400 magnification and analyzed using the Image J software. Between 1,500 and 3,000 cells per slide were analyzed. For bromodeoxyuridine (BrdUrd) incorporation experiments, mice were injected i.p. with BrdUrd (10 mg/kg; Sigma) and dissolved in PBS 2 hours before sacrifice. Anti-BrdUrd–stained sections were analyzed as above.

Lactate assay

Lactate levels were assayed using a commercially available kit (Biovision). Lactate levels were normalized to the amount of DNA or proteins extracted from each tissue fragment or primary culture.

Oxygen consumption rate and extracellular acidification rate

Freshly isolated thyroid follicles from pools of 12 WT or 3 mutant mice were plated as described in “Primary cultures,” and oxygen consumption rate (OCR) was measured using the Seahorse XF24 instrument (Seahorse Biosciences) under basal conditions, in the presence of the mitochondrial inhibitor oligomycin (1 μg/mL), the mitochondrial uncoupler carbonyl cyanide 4-(trifluoromethoxy)phenylhydrazone (FCCP; 1 μmol/L), and the respiratory chain inhibitors antimycin A (2 mmol/L) and rotenone (0.1 μmol/L).

Flow cytometry analysis of mitochondrial membrane potential

Primary thyroid cells were treated with solvent or 20 μmol/L FCCP for 10 minutes before staining with 100 nmol/L tetramethylrhodamine, ethyl ester (TMRE; Invitrogen) for 20 minutes. Cells were rinsed in Dulbecco’s PBS (DPBS), trypsinized, and analyzed for TMRE staining by flow cytometry.

Western blot analysis

Thyroids and cells were homogenized on ice in radioimmunoprecipitation assay (RIPA) buffer supplemented with Complete Protease Inhibitor Tablet (Roche Diagnostics). Western blot analysis was conducted using 20 to 40 μg proteins using antibodies from Cell Signaling Technology, except for aconitate 2, isocitrate dehydrogenase 3a, HIF-1α, and succinate dehydrogenase b (Santa Cruz Biotechnology) and β-actin (Sigma-Aldrich).

Cell lines and drug treatments

FTC-133 cells (kindly donated by Dr. Matthew Ringel, Ohio State University, Columbus, OH) were maintained in Dulbecco’s Modified Eagle Medium (DMEM) with 10% FBS at 37°C in 5% CO₂. 8505c cells (kindly donated by Dr. Sareh Parangi, Massachusetts General Hospital, Boston, MA) and THJ16T cells (kindly donated by Dr. John A. Copland, Mayo Clinic, Jacksonville, FL) were grown in RPMI medium with 10% FBS. All cell lines were validated before starting the experiments by amplifying and sequencing genomic fragments encompassing their known mutations (FTC133: Pten^+/+; BRAF^R130H; 8505c: BRAF^V600E, THJ16T: PIK3CA^E545K).

Pharmacologic inhibitors of protein kinase A (PKA; H89; Cell Signaling Technology), and PI3K (BKM120; Selleck Chemicals), or AMP analog (AICAR; TRC) were added 24 hours after plating. After 30 minutes, samples were collected and prepared for Western blot analysis. For luciferase and quantitative PCR (qPCR) experiments, samples were collected after 72 hours.

Real-time PCR

Total RNA was extracted with TRIzol and reverse transcribed using the ThermoScript Kit (Life Technologies). Quantitative real-time PCR (qRT-PCR) was conducted on a StepOne Plus apparatus using the Absolute Blue qPCR Rox Mix (Thermo Scientific) and TaqMan expression assays (Applied Biosystems). Each sample was run in triplicate and glyceraldehyde 3-phosphate dehydrogenase (GusB) or β-actin was used to control the input RNA. Data analysis was based on the Ct method and experiments were repeated at least three times using at least two independent thyroid pools (at least five mice/pool).

AMP, ADP, ATP level determination

AMP, ADP, and ATP levels were assayed using three independent thyroid pools (10 mice/pool). Approximately, 30 mg of
tissue was extracted in 210 μL of extraction solvent, 40%/40%/20% acetonitrile/methanol/0.1% formamide in water containing a 15N_AMP extraction standard. The column [Sequent ZIC-cHILIC (3.5 μm, 100 mm × 2.1 mm, inner diameter)] was run with a gradient of 90% acetonitrile/10% water containing 10 mmol/L ammonium formate, pH 3.0, at 0.2 mL/min using an ACQUITY ultra performance liquid chromatography attached to a Waters Xevo triple quadruple mass spectrometry.

**Transient transfection assays**

Cells were transfected with 1 μg PGC-1α WT or T177A/S538A plasmid (#1026, #18093; Addgene). After 24 hours, cells were treated with 1 mmol/L AICAR (TRC) for 48 hours and collected for RNA isolation.

**Dual luciferase assay**

Cells were cotransfected with 10 ng phRG-TK plasmid (Promega) and 2 μg PGC-1α promoter luciferase plasmid (#8887; Addgene). After 24 hours, cells were treated with inhibitors for 48 hours and collected for dual-luciferase reporter assay (Promega).

**Transmission electron microscopy**

Thyroid glands were fixed with 2% paraformaldehyde and 2.5% glutaraldehyde in 0.1 mol/L sodium cacodylate buffer, postfixed with 1% osmium tetroxide followed by 1% uranyl acetate, dehydrated through a graded series of ethanol and embedded in LX112 resin (LADD Research Industries). Ultrathin (80 nm) sections were cut on a Reichert Ultracut UCT, stained with uranyl acetate followed by lead citrate, and viewed on a JEOL 1200EX transmission electron microscope at 80 kV.

**Proteomic analysis**

Sets of thyroid extracts (100 μg) from 3-month-old WT and mutant mice were subjected to two-dimensional (2D) gel electrophoresis (pH 4-11) and Sypro Ruby gel staining as described in ref. 16. About 30 differentially expressed spots were selected for cutting and in-gel tryptic digestion. Matrix-assisted laser desorption/ionization–time-of-flight (MALDI-TOF) peptide mass fingerprinting was conducted as described in ref. 16. Liquid chromatography/tandem mass spectrometry (LC/MS-MS) peptide sequencing was conducted as described in ref. 17. All analyses were conducted by the Fox Chase Cancer Center Proteomics Facility, Philadelphia, PA.

**Statistical analysis**

Experiments were carried out at least three times. Data were analyzed using the Prism software package. Differences with P values less than 0.05 were considered statistically significant.

**Results**

**Constitutive PI3K activity results in transcriptional repression of TCA cycle and OXPHOS genes**

The thyroids of Pten<sup>Bay<sup>−/−</sup></sup> mice are enlarged from birth due to an increase in the thyrocytes' proliferative index associated with constitutive PI3K signaling. However, these glands do not display any functional alteration or signs of neoplastic transformation until the mice are at least 10 months of age (12).

To identify early molecular changes that may favor or contribute to the development of thyroid tumors in aging mice, we have initially used a small-scale proteomic approach to detect proteins differentially expressed in thyroids from 3-month-old WT and Pten<sup>Bay</sup>−/−<sup>−</sup> mice. Several spots, with significantly different intensity between normal and mutant thyroids, were selected and excised from 2D gels, and 35 proteins were identified using MALDI-TOF and LC/MS-MS. Surprisingly, 7 of 28 proteins found downregulated in mutant thyroids were enzymes involved in the TCA cycle (Table 1). To validate these findings, we measured by RT-PCR the mRNA levels of the genes encoding these proteins and found that Cs, Aco2, Idh3a, Dld, Pdhb, and SdhA expression in mutant thyroids was reproducibly 20% to 50% lower compared with that of WT glands, whereas the expression of the glucose transporter Glut1 was slightly increased (Fig. 1A). Downregulation of these genes was also confirmed by Western blot analysis using antibodies specific for aconitase 2, isocitrate dehydrogenase 3a, and succinate dehydrogenase b (Fig. 1B).

To determine to what extent metabolic genes are transcriptionally deregulated upon PI3K activation, we interrogated our Affymetrix gene expression dataset (14) to extend the analysis to all the genes directly involved in the TCA cycle and glycolysis. Interestingly, while none of the glycolytic enzymes was transcriptionally deregulated in the mutant glands (with the exception of a modest increase in the Glucokinase mRNA

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Abbreviation: N/A, not applicable.
Figure 1. Metabolic reprogramming in Pten<sup>thyr</sup><sup>−/−</sup> mice. A, relative expression of the indicated genes in WT and mutant thyroids. Bars represent mean ± SD of triplicate measurements. * indicates significant (P < 0.05) differences. B, Western blot analysis showing downregulation of representative TCA cycle enzymes in mutant glands. C, TCA cycle diagram showing the genes downregulated in Pten<sup>thyr</sup><sup>−/−</sup> thyroids. The color scale reflects changes calculated from the Affymetrix data. Genes circled in red have been validated by qPCR. D, heatmap showing the repression of TCA cycle genes in Pten<sup>thyr</sup><sup>−/−</sup> thyroids and in follicular carcinomas (FTC) arising in older mice. E, DNA content-based assessment of mitochondria number in 3-month-old WT and mutant mice (top). Expression levels of selected mitochondrial genes in the thyroids of WT and mutant mice (bottom). Bars represent mean ± SD of triplicate measurements. * indicates significant (P < 0.05) differences. F, mitochondrial damage (swelling, clarification, and cristae disruption) in Pten<sup>−/−</sup> glands detected by transmission electron microscopy. G, OCR in control and mutant primary thyrocytes cells in response to 1 µg/mL oligomycin, 1 µmol/L FCCP, or 2 µmol/L antimycin A + 2 µmol/L rotenone. H, ECAR in control and mutant primary thyrocytes cells. P = 0.004. I, mitochondrial membrane polarization was measured in primary thyrocytes by flow cytometry using TMRE. FCCP pretreatment of WT cells was used to determine the baseline.
levels), the expression of 22 of 28 genes encoding enzymes associated with the TCA cycle was significantly repressed in \( Pten^{−/−} \) thyroids (Fig. 1C; data not shown). About 60% of these genes were still repressed in the metastatic follicular carcinomas developing in older \( Pten^{lov−/−} \) mice (Fig. 1D).

Furthermore, when we quantitated citrate and isocitrate in the thyroids of control and \( Pten^{thr−/−} \) mice by gas chromatography-mass spectroscopy (GC-MS), we found significantly reduced levels of these metabolites in the mutant glands (citrate, 2,577 ± 525 pmol/mg in the WT and 985 ± 302 in the mutants; isocitrate, 217 ± 42 pmol/mg in the WT and 80 ± 26 in the mutants).

The tight connection existing between TCA cycle, OXPHOS, and mitochondrial function prompted us to test whether this global downregulation of TCA cycle genes is accompanied by alterations of the expression of mitochondrial-encoded OXPHOS genes, as well as by changes in the number of mitochondria. We first used RT-PCR to measure the relative mitochondrial genome copy number by amplifying four different mitochondrially encoded genes and one reference nuclear gene from total (genomic and mitochondrial) DNA isolated from WT and mutant thyroids. No significant differences were found between control and mutants, suggesting that constitutive PI3K activation does not alter mitochondrial mass in the mouse thyroid (Fig. 1E). Conversely, the expression of most mitochondrial-encoded genes was heavily downregulated in \( Pten^{−/−} \) thyroids (Fig. 1E). Furthermore, electron microscopy analysis of control and mutant thyroid sections showed that \( Pten \) loss caused profound morphologic defects in the mitochondria, including swelling, matrix clarification, and disruption of the cristae (Fig. 1F). All these features are commonly associated with reduced mitochondrial oxidative capacity (18).

To test whether the aforementioned findings reflect an impairment in mitochondrial function, we measured OCR as well as extracellular acidification rate (ECAR) in primary cultures of control and \( Pten^{−/−} \) thyocytes. Strikingly, \( Pten^{−/−} \) thyocytes exhibited significantly reduced spare respiratory capacity, that is, lower maximal rate of respiration (Fig. 1G) and increased ECAR (Fig. 1H). Finally, we used flow cytometry to compare the mitochondrial membrane potential in primary cultures of control and \( Pten^{−/−} \) thyocytes using the cationic potential-sensitive dye, TMRE, which accumulates in intact mitochondria. The main population of mutant cells displayed reduced mitochondrial membrane potential compared with control thyrocytes (about 50% of WT levels, note the logarithmic scale; Fig. 1I), whereas a second population of mutant cells was characterized by even lower potential, similar to that of cells treated with the uncoupler FCCP.

Taken together, these data show that constitutive PI3K activation in the mouse thyroid results in the coordinated downregulation of the expression of genes encoding members of both the TCA cycle and OXPHOS pathways, in extensive mitochondrial damage, and in severe impairment of mitochondrial function.

**A glycolytic switch in preneoplastic \( Pten^{−/−} \) thyrocytes**

Because the expression changes described earlier result in reduced efficiency of the TCA cycle/OXPHOS metabolic pathway, the mutant cells might have to increase their glycolytic rate to meet their energetic needs.

As a proxy for the thyrocytes' glycolytic rate, we assessed the amount of lactate present in thyroids from 3-month-old WT and mutant mice. Strikingly, \( Pten^{−/−} \) thyroids displayed a 3-fold increase in normalized lactate content, suggesting a higher metabolic flux through glycolysis (Fig. 2A). Along the same line, we measured the lactate secretion rate in primary thyrocyte cultures. Mutant cells produced 3- to 5-fold more lactate than their WT counterparts (Fig. 2B), strongly indicating that glycolysis is indeed upregulated to compensate the PI3K-mediated repression of the TCA cycle/OXPHOS pathway. Finally, to validate this hypothesis, we conducted FDG-PET imaging on control and \( Pten^{lov−/−} \) mice and found that the hyperplastic \( Pten^{−/−} \) thyroids showed strong FDG uptake, whereas no uptake was detected in WT mice (Fig. 2C).

These data show that, in response to constitutive PI3K activation, thyroid epithelial cells undergo a glycolytic switch, which is functionally reminiscent of the Warburg effect observed in cancer cells, as well as in highly proliferating cells such as activated lymphocytes.

The Warburg effect is usually driven by a set of "master genes," including c-MYC, AKT, mTOR, and HIF-1a, which control the expression, localization, and function of both glucose transporters and glycolytic enzymes (5, 19). Upregulation of hexokinase 2, pyruvate kinase M2, and lactate dehydrogenases A and B are hallmarks of this glycolytic switch.

The establishment of a glycolytic switch in slowly proliferating, non-neoplastic (i.e., nontransformed) cells, and the absence of glycolytic genes upregulation in our expression profiling data would argue against a mechanism involving the classical Warburg effect–related master regulators. In fact, we did not detect significant alterations in the expression levels of Hif-1a, c-Myc, Hk2, Pkm2, Ldha, or Ldhb both at the RNA and protein level (Fig. 2D and E; data not shown).

These data suggest that PI3K activation in nontransformed thyroid epithelial cells induces a glycolytic switch through a novel molecular mechanism.

**Pdk1 drives TCA cycle/OXPHOS gene repression downstream of PI3K, independent of mTOR**

Pdk1 is an essential member of the PI3K cascade, as it phosphorylates Akt on T308 and S6k1 on T229, allowing their downstream activation, independent of mTOR.

To test to what extent the TCA cycle gene expression changes described earlier result in reduced efficiency of the TCA cycle/OXPHOS metabolic pathway, the mutant cells might have to increase their glycolytic rate to meet their energetic needs.

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Pdk1 is an essential member of the PI3K cascade, as it phosphorylates Akt on T308 and S6k1 on T229, allowing their full activation (20). To test to what extent the TCA cycle gene repression observed in \( Pten^{−/−} \) thyroids depends on the Pdk1/Akt/mTOR cascade, we first crossed \( Pten^{lov−/−} \) mice to \( Pdk1^{lov−/−} \) conditional mutants (21), obtaining simultaneous thyroid-specific deletion of these two genes. Combined loss of \( Pten \) and \( Pdk1 \) rescued both the repression of the TCA cycle genes (Fig. 3A) and that of mitochondrially encoded OXPHOS genes (Fig. 3B). Interestingly, mitochondrial DNA (mtDNA)-encoded gene expression in the compound mutants was higher than in WT controls. Furthermore, simultaneous inactivation of \( Pten \) and \( Pdk1 \) restored lactate production in the thyroid to WT levels (Fig. 3C).

Constitutive PI3K activation in the thyroid follicular cells results in mTOR activation (22), and mTOR activity has been proposed to play a key role in establishing the glycolytic switch.
To test the role of the PI3K/Akt/mTOR axis in the metabolic phenotype of \( \text{Pten}^{+/+} \) mice, we treated 3-month-old WT and mutant mice \((n = 5/\text{group})\) daily for 2 weeks with the mTOR inhibitor RAD001 or placebo (Fig. 3D). As previously shown by our group, this treatment is sufficient to inhibit mTOR activity and drastically reduce proliferation in mutant thyroids (Fig. 3D and E; ref. 22). Strikingly, despite the clear inhibition of the thyrocytes' proliferation rate, RAD001 treatment had no effect on the repression of TCA cycle genes (Fig. 3F), on the repression of mitochondrial genes (Fig. 3G), or on thyroid lactate production (Fig. 3H). These data strongly suggest that PI3K activation in thyroid cells induces a glycolytic switch independent of mTOR activity. Furthermore, they show that the increase in glycolysis in \( \text{Pten}^{+/+} \) mice is not a direct consequence of the increased proliferative rate observed in the hyperplastic glands.

**PI3K/AKT inhibit AMPK, leading to decreased OXPHOS metabolism in preneoplastic thyrocytes**

The expression of nucleus- and mitochondrial-encoded respiratory proteins is controlled by a number of nucleus-encoded transcription factors and coactivators that modulate mitochondrial function in response to extra- and intracellular signals. To identify the mechanism through which loss of \( \text{Pten} \) results in a coordinated reprogramming of the expression of respiratory genes, we initially measured in thyroids from 3-month-old WT and \( \text{Pten}^{+/+} \) mice the expression levels of the transcription factors and cofactors involved in the regulation of TCA cycle/OXPHOS. Although the expression levels of ERR\( \beta \), PRC, NRF-1 and -2, and TFAM were not altered by constitutive PI3K activation, \( \text{Pten}^{+/+} \) thyroids had significantly lower expression of ERR\( \gamma \), PGC-1\( \alpha \) and -\( \beta \), and, to a lesser extent, ERR\( \alpha \) (Fig. 4A). These data suggest that PI3K activation might alter the expression or activity of a common master regulatory gene.

Recently published data have directed our attention on AMPK as a candidate for this role. AMPK contributes to the control of respiratory genes in muscle cells (24) and phosphorylates PGC-1\( \alpha \) on T177 and S538 to increase its transcriptional activity, including its ability to transactivate its own promoter (25). In addition, the ESRRA promoter contains a PGC-1\( \alpha \)/ERR\( \alpha \)-responsive regulatory element that can also be
recognized and activated by ERRβ and ERRγ (26). Furthermore, in certain cell types, AKT can phosphorylate AMPK on S485, preventing LKB1 from phosphorylating (on T172) and activating AMPK (27, 28). Indeed, we found that thyroids from Pten+/mice have drastically reduced pT172- and higher pS485-AMPK, compared with WT controls, and reduced levels of phosphorylated ACC (Fig. 4B). Because it is theoretically possible that the reduced levels of pT172-AMPK in the mutant glands might be due to a low AMP/ATP ratio, we measured AMP, ADP, and ATP levels in extracts from WT and mutant glands. AMP and ATP levels in mutant glands were comparable with WT controls, whereas ADP levels were slightly higher on the mutant glands. These data indicate that PI3K inhibits AMPK despite AMP/ATP and ADP/ATP ratios that would stimulate AMPK activation, as in WT glands (Fig. 4C).

To test the hypothesis that PI3K activation represses TCA cycle/OXPHOS gene expression through the inhibition of AMPK, we treated WT and Pten+/mice with the AMPK activator AICAR for 4 weeks, and then isolated their thyroid glands. AICAR treatment increased pT172-AMPK levels not only in WT mice, but also (and to the same levels) in Pten+/mice, despite the persistent AKT activation and inhibitory AMPK phosphorylation on S485 (Fig. 5A). Furthermore, AICAR-activated AMPK restored ACC phosphorylation on Ser79 (Fig. 5D).

We next measured the expression of ERRα, ERRγ, PGC-1α, and PGC-1β in the thyroids of control, mutant, and AICAR-treated mutant mice. Strikingly, AICAR treatment increased the mRNA levels of all four transcription factors and cofactors to near-WT levels (Fig. 5B). To verify that the increase in transcriptional regulator levels translates in a restoration
of normal TCA cycle/OXPHOS gene expression, we measured by qPCR and Western blotting the expression levels of selected TCA cycle genes, and found that AICAR treatment had completely rescued the PI3K-dependent repression of Aco2, Idh3a, and Pdhb (Fig. 5C and D). However, when we tested the expression of the mitochondrial-encoded OXPHOS genes, we could only detect a partial rescue upon AICAR treatment (Fig. 5E). Furthermore, the extent of mitochondrial damage observed in Pten thyr/−/− thyrocytes was only mildly reduced by AICAR treatment (Fig. 5F). Despite the persistence of some mitochondrial damage, enforced AMPK activation reversed the glycolytic switch induced by constitutive PI3K activation, as shown by the reduction of lactate content to WT levels in AICAR-treated Pten thyr/−/− mice (Fig. 5G).

These data strongly suggest that PI3K/AKT–mediated AMPK inhibition is responsible for the repression of oxidative metabolism gene expression observed in Pten thyr/−/− mice.

**AMPK inhibition actively contributes to thyroid hyperplasia**

To test whether the glycolytic switch observed in Pten thyr/−/− mice plays an active role in the hyperproliferative and protu- morigenic phenotype of mutant thyrocytes, we compared the weight and proliferative index of WT, Pten thyr/−/−, and AICAR-treated mutant thyroid glands. During the 4 weeks of enforced AMPK reactivation, mutant glands grew at a much slower rate compared with untreated mutants, so that the weight of the mutant thyroids at the end of the treatment was significantly reduced compared with untreated Pten thyr/−/− mice (−34%; Fig. 5H). This size reduction was associated with decreased proliferation: the thyrocyte proliferation index in AICAR-treated mice, determined by BrdUrd incorporation, was drastically reduced, compared with that of untreated mutant mice (−53%; Fig. 5I). Thus, it is possible that that the glycolytic switch caused by PI3K-mediated AMPK inhibition actively contributes to the development of thyroid hyperplasia in Pten thyr/−/− mice.

**The PI3K/AMPK–mediated repression of oxidative metabolism is maintained in human thyroid cancer cells and involves AMPK-mediated phosphorylation of PGC-1α**

Having established in a relevant in vivo system that sustained PI3K activation redirects energy metabolism toward glycolysis by repressing the expression of TCA cycle genes, we measured by qPCR and Western blotting the expression levels of selected TCA cycle genes, and found that AICAR treatment had completely rescued the PI3K-dependent repression of Aco2, Idh3a, and Pdhb (Fig. 5C and D). However, when we tested the expression of the mitochondrial-encoded OXPHOS genes, we could only detect a partial rescue upon AICAR treatment (Fig. 5E). Furthermore, the extent of mitochondrial damage observed in Pten thyr/−/− thyrocytes was only mildly reduced by AICAR treatment (Fig. 5F). Despite the persistence of some mitochondrial damage, enforced AMPK activation reversed the glycolytic switch induced by constitutive PI3K activation, as shown by the reduction of lactate content to WT levels in AICAR-treated Pten thyr/−/− mice (Fig. 5G).

These data strongly suggest that PI3K/AKT–mediated AMPK inhibition is responsible for the repression of oxidative metabolism gene expression observed in Pten thyr/−/− mice.

**AMPK inhibition actively contributes to thyroid hyperplasia**

To test whether the glycolytic switch observed in Pten thyr/−/− mice plays an active role in the hyperproliferative and protu-
and OXPHOS genes in mouse thyrocytes, we sought to extend these studies to the human setting. We have initially used the PTEN<sup>−/−</sup> follicular thyroid cancer cell line, FTC-133, untreated, normally growing cells displayed almost undetectable pT172-AMPK, and strong pS485-AMPK. In keeping with the mouse data, AICAR treatment increased pT172-AMPK, as well as pS79-ACC (a direct AMPK target) in a dose-dependent manner (Fig. 6A). Furthermore, enforced...
reactivation of AMPK led to significantly increased expression of the TCA cycle genes *ACO2*, *IDH3A*, and *PDHB* (Fig. 6B).

As mentioned earlier, AMPK might control the expression and function of metabolic transcription factors and enzymes by modulating PGC-1α activity via phosphorylation on PGC-
1αT177 and S538 (25). To test whether this pathway, discovered in skeletal muscle cells, is also active in thyrocytes, we transfected FTC-133 cells with an expression vector encoding WT or phosphorylation-defective PGC-1α (PGC-1α-AA), treated the cells with vehicle or AICAR to reverse the inhibition of AMPK, and extracted mRNA to measure the expression of a group of TCA cycle genes. AMPK reactivation increased the expression of AC02, IDH3A, and PDHB in cells transfected with WT PGC-1α but not, or to a much lower extent, in those cells transfected with the PGC-1α-mutant that cannot be phosphorylated (Fig. 6C). Thus, PGC-1α acts as a conduit for AMPK to control the expression of metabolic genes.

To test whether AMPK repression in human cells is directly associated with P3K activation, we treated FTC-133 cells with the pan-P3K inhibitor BKM120. Surprisingly, P3K inhibition at a BKM concentration that has no off-target effects with the pan-PI3K inhibitor BKM120, treated the cells with vehicle or AICAR to reverse the inhibition of AMPK, and extracted mRNA to measure the expression of a group of TCA cycle genes. AMPK reactivation increased the expression of AC02, IDH3A, and PDHB in cells transfected with WT PGC-1α but not, or to a much lower extent, in those cells transfected with the PGC-1α-mutant that cannot be phosphorylated (Fig. 6C). Thus, PGC-1α acts as a conduit for AMPK to control the expression of metabolic genes.

PKA has been shown to phosphorylate S485-AMPK in the insulin-secreting cell line, INS-1 (29). Furthermore, in mouse adipocytes, PKA can phosphorylate AMPK on S173, inhibiting T172 phosphorylation, likely through steric hindrance (30). Thus, we tested whether PKA inhibition could synergize with P3K inhibition to restore AMPK activity and respiratory gene expression. Strikingly, while FTC-133 treatment with the PKA inhibitor H89 alone did not result in alterations of the phosphorylation status of AMPK, simultaneous inhibition of P3K and PKA abolished pS485-AMPK and drastically increased pT172-AMPK (Fig. 6D). Accordingly, while single-inhibitor treatment of FTC-133 failed to increase the expression of TCA cycle genes, or PGC-1α and PGC-1β, combined treatment significantly increased their mRNA levels (Fig. 6E). Combined inhibition of P3K and PKA was also more effective than either kinase inhibition in increasing the expression of a luciferase reporter driven by the promoter region of PGC-1α, further supporting the role of AMPK in the control of PGC-1α activity and expression (Fig. 6F).

These data suggest that the P3K/AKT axis controls AMPK activity also in PTEN+/− human thyroid cancer cells; however, in these cells, PKA contributes to the metabolic reprogramming through an overlapping pathway.

To test whether this signaling cascade is also active in human anaplastic thyroid cancer, we used two cell lines representative of the most common driver pathways in this aggressive and genetically complex tumor type: THJ16T, harboring an activating PIK3CA mutation (E545K), and 8505c, harboring the BRAF oncogenic allele V600E.

Both cell lines displayed low to undetectable pT172-AMPK, as well as strong pS485-AMPK, suggesting that AMPK inhibition is a common theme in thyroid cancer (Fig. 6G). AICAR treatment restored T172 phosphorylation, more effectively in the cell line harboring the PI3K oncogenic mutation (Fig. 6G).

When we treated these cells with P3K and PKA inhibitors, alone and in combination, we found that simultaneous P3K and PKA inhibition effectively reactivated AMPK and increased the expression of metabolic regulators such as ERRγ and PGC-1β in cells with constitutively active P3K; however, they did not affect AMPK activation or ERRγ and PGC-1β expression in the BrafV600E cell line (Fig. 6H and I). Similar to the in vivo model, AMPK reactivation reduced proliferation of a panel of mouse and human thyroid cancer cell lines carrying P3K-activating mutations (Fig. 6J).

Thus, in well-differentiated and undifferentiated human thyroid cancer cells, constitutive activation of P3K, in cooperation with PKA, alters the expression of genes involved in respiratory metabolism by inhibiting AMPK.

**TCA cycle gene repression in neoplastic lesions is also observed in other human tissues**

To extend our findings to human tumors other than those arising in the thyroid, we interrogated the Oncomine database for datasets showing significant simultaneous downregulation of both PTEN and TCA cycle genes. Although this approach does not account for other, more common, P3K-activating mechanisms, it represents the only direct way to identify, within large expression datasets, tumors with activated P3K signaling, in the absence of validated "P3K activation" signatures.

Considering the inherent low power of our search, it is remarkable that we found a very strong association between PTEN loss and TCA cycle repression in two datasets, derived from dedifferentiated liposarcoma and from early-stage colon cancer (Supplementary Fig. S1). Thus, other human tumor types display global downregulation of TCA cycle genes in association with PTEN loss and P3K activation, warranting future studies to experimentally validate this association.

**Discussion**

The existence of a tight link between P3K signaling and rerouting of energy metabolism is well established in cancer cells, where activation of P3K contributes to the switch from oxidative to glycolytic pathways through several of its downstream effectors, including AKT, mTOR, and HIF-1α (31). In turn, these effectors increase the expression and/or activity of a variety of glycolytic enzymes, such as hexokinase II (32), phosphofructokinase 2 (33), the GLUT family of glucose transporters (34), lactate dehydrogenases (35, 36), and the M2 isoform of pyruvate kinase (23). Thus, in fully transformed cells, the Warburg effect is achieved through an increase in glycolytic flux, whereas mitochondrial metabolism is unaffected.

Little is known of the mechanism(s) regulating energy metabolism in normal and preneoplastic (i.e., carrying single mutations and not yet transformed) cells. There is a consensus that in normal cells the glycolytic switch is linked to high proliferation rates, as shown for embryonic tissues and activated lymphocytes (37). Although an elegant study has recently established that activated lymphocytes rely mainly on MYC to transcriptionally reprogram their metabolism (38), much less is known when it comes to epithelial cells.

Our data, generated in a relevant in vivo model, support the hypothesis that, in response to P3K activation, nontransformed thyroid epithelial cells increase metabolic flux through...
glycolysis, as indicated by the dramatic increase in lactate production observed in mutant cells; however, this metabolic switch is independent of thyrocyte proliferation. Metabolic reprogramming of thyroid epithelial cells is achieved through a previously unknown mechanism, involving the coordinated downregulation of the expression of TCA cycle and OXPHOS genes, and leading to dysfunctional mitochondria and reduced ability to conduct respiratory metabolism. Although we did not observe any significant increase in the expression of glycolytic genes (with the exception of Glk) upon Pten loss, we found a reproducible upregulation of the Glut1 transporter. Accordingly, a strong connection between loss of Pten expression and increased expression of GLUT1 has been recently reported in thyroid tumors discovered during unrelated FDG-PET scans (39).

Our data are in partial agreement with those recently reported by Garcia-Cao and colleagues using a mouse model with enforced whole-body overexpression of Pten (40). Mouse embryonic fibroblasts (MEF) from these mice display a metabolic shift toward OXPHOS, decreased lactate production, and increased expression of PGC-1α-target genes. However, in striking contrast with our model, cells overexpressing Pten have increased mitochondria number and upregulate Pkm2 through mTOR. Although these differences might be linked to tissue-specificity of some of the controlling pathways, they might also be the consequence of overexpressing Pten, which completely abolishes PI3K signaling. Notably, the expression of OXPHOS genes in Pten−/− mice (in which PI3K signaling is drastically repressed, like in the Pten transgenic strain) was higher than in WT control, supporting the latter hypothesis.

From a mechanistic standpoint, our data support a model in which PI3K activation initiates the remodeling of energy metabolism through the phosphorylation of AMPK on S485 and its consequent inactivation. AMPK is a kinase known to play critical roles in growth, metabolism, autophagy, and cell polarity (41, 42). Its activation requires phosphorylation on T172 by LKB1 (41). Such phosphorylation can be inhibited by PKA-mediated phosphorylation of S173 (30). A recent report has shown that also S6K1 can inhibit AMPK by phosphorylating S485 (44). However, the fact that RAD001 treatment of mutant mice does not rescue any of the metabolic phenotypes despite the complete ablation of S6K activity argues against a role for S6K1 in the phosphorylation of AMPK, at least in the thyroid. For the same reason, although mTOR was found to be central to the glycolytic switch in Tsc2−/− mouse kidney tumors and in Pten−/− MEFs (23), our in vivo data clearly show that mTOR activation is dispensable for the metabolic remodeling of the preneoplastic thyroid.

AMPK has been recently shown to suppress the Warburg effect in Myc-driven lymphomas, and its genetic inactivation induces a glycolytic shift in MEFs via normoxia stabilization of HIF-1α (45). Conversely, we do not detect changes in HIF-1α protein levels in the thyroid of Pten−/− mice, nor upregulation of Ldh1 or Aldoa. More importantly, acute AMPK ablation did not affect OCR in both lymphomas and MEFs, suggesting the absence of any detrimental effects on the TCA cycle. These differences with our model might be related to the complete loss of AMPK versus its decreased phosphorylation, or reflect tissue-specific wiring of the metabolic pathways.

ERRs and PGC-1 are major regulators of the expression of TCA cycle/OXPHOS genes, and their loss or overexpression have been shown to lead to repression or induction of these metabolic targets, respectively (24, 46, 47).

Inactivation of AMPK by AKT results in its inability to transactivate PGC-1α (25), which in turn leads to lower levels of ERRα (26). Less clear, instead, is how AMPK controls the expression of PGC-1β, which lacks the two residues phosphorylated by AMPK in PGC-1α, and that of ERRγ. Although ERRγ downregulation might be directly responsible for the reduced levels of PGC-1β (24), its link to PI3K and AMPK is still unclear. It has been recently reported that ERRβ2 upregulates (likely through PI3K) microRNA (miR)-378 expression in breast cancer cells (48). In turn, miR-378* targets ERRγ, leading to downregulation of TCA cycle genes, reduced respiration rate, and increased glycolysis. Future studies will evaluate whether P3K activation in thyroid cancer cells targets ERRγ through miR-378*, and whether AMPK plays a role in this pathway.

The fact that AICAR treatment rescued only partially the expression of mtDNA-encoded genes and the altered mitochondria morphology in Pten−/− mice might be a consequence of the extensive damage sustained by these organelles. Alternatively, additional PI3K-driven pathways might contribute to this aspect of the phenotype. An additional still open question is whether PKA affects AMPK activity in thyroid cancer cell lines, but does not seem to do so in vivo, in mouse thyrocytes. Further work will be needed to clarify these issues.

One important aspect of our findings is that the metabolic remodeling seems to contribute directly to the hyperproliferative phenotype, as AICAR-mediated restoration of normal levels of TCA cycle enzymes and reduction of glycolytic rate in vivo drastically reduces thyrocyte proliferation. However, we cannot completely exclude that the reduced proliferation in AICAR-treated mice is also the consequence of an energy-deprived status due to persistent mitochondrial damage and dysfunction in conditions (i.e., AICAR treatment) that decrease the metabolic flux through glycolysis.

It is tempting to propose that the noncanonical glycolytic switch that takes place in preneoplastic Pten−/− cells might contribute to the neoplastic transformation of thyroid, breast, and colon epithelial cells in patients with Cowden Disease, carrying germline Pten mutations. Similar to SDHB or SDHD mutation carriers, patients with Pten-mutant Cowden Disease have recently been found to have elevated plasma succinate (49). The downregulation of succinate dehydrogenases expression observed in the thyroid of Pten−/− mice also results in increased succinate levels (Antico and colleagues, manuscript in preparation). A similar protransformation contribution might also take place in endometrial epithelial cells in sporadic type I (endometrioid) endometrial cancer, where loss of PTEN is the earliest recognized genetic alteration (50).

Because the analysis of early hyperplastic lesions with PI3K activation in human tissues is not easily feasible, and relevant expression datasets are not publicly available, to extend our
model to additional human tissues, we used the Oncomine database, and found several datasets in which PDIM down-regulation coexists with global repression of TCA cycle genes. Interestingly, one of these datasets is derived from early-stage colorectal cancer (51), supporting our hypothesis that these metabolic changes may take place well before full neoplastic transformation. Finally, a recent proteomic and metabolomic study of gastric cancer, a tumor type often associated with PI3K activation, found extensive repression of TCA cycle genes (52).

In conclusion, our characterization of an in vivo model of PI3K activation in thyroid epithelial cells has led to the discovery of a novel pathway, in which the glycolytic switch is not achieved through direct upregulation of glycolytic enzyme expression and activity, but rather through the active inhibition of AMPK and the consequent repression of the expression of TCA cycle and OXPHOS genes, leading to the impairment of mitochondrial metabolism.

Disclosure of Potential Conflicts of Interest
No potential conflicts of interest were disclosed.

References


Inhibition of AMPK and Krebs Cycle Gene Expression Drives Metabolic Remodeling of *Pten*-Deficient Preneoplastic Thyroid Cells


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