Pyruvate Carboxylase Is Critical for Non-Small-Cell Lung Cancer Proliferation

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Pyruvate carboxylase is critical for non–small-cell lung cancer proliferation

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Introduction

Accelerated glycolysis under aerobic conditions (the “Warburg effect”) has been a hallmark of cancer for many decades (1). It is now recognized that cancer cells must undergo many other metabolic reprogrammings (2) to meet the increased anabolic and energetic demands of proliferation (3, 4). It is also becoming clear that different cancer types may utilize a variety of metabolic adaptations that are context dependent (5–11), commensurate with the notion that altered metabolism is a hallmark of cancer (12). Enhanced glucose uptake and aerobic glycolysis generates both energy (i.e., ATP) and molecular precursors for the biosynthesis of complex carbohydrates, sugar nucleotides, lipids, proteins, and nucleic acids (3, 13, 14). However, increased glycolysis alone is insufficient to meet the total metabolic demands of proliferating cancer cells. The Krebs cycle is also a source of energy via the oxidation of pyruvate, fatty acids (15), and amino acids such as glutamine (10, 16, 17). Moreover, several Krebs cycle intermediates are essential for anabolic and glutathione metabolism, including citrate, oxaloacetate, and α-ketoglutarate (Figure 1A).

Anabolic biosynthesis requires precursors supplied by the Krebs cycle, which in turn requires anaplerosis to replenish precursor intermediates. The major anaplerotic sources are pyruvate and glutamine, which require the activity of pyruvate carboxylase (PC) and glutaminase 1 (GLS1), respectively. Due to their rapid proliferation, cancer cells have increased anabolic and energy demands; however, different cancer cell types exhibit differential requirements for PC- and GLS-mediated pathways for anaplerosis and cell proliferation. Here, we infused patients with early-stage non–small-cell lung cancer (NSCLC) with uniformly 13C-labeled glucose before tissue resection and determined that the cancerous tissues in these patients had enhanced PC activity. Freshly resected paired lung tissue slices cultured in 13C6-glucose or 13C5,15N2-glutamine tracers confirmed selective activation of PC over GLS in NSCLC. Compared with noncancerous tissues, PC expression was greatly enhanced in cancerous tissues, whereas GLS1 expression showed no trend. Moreover, immunohistochemical analysis of paired lung tissues showed PC overexpression in cancer cells rather than in stromal cells of tumor tissues. PC knockdown induced multinucleation, decreased cell proliferation and colony formation in human NSCLC cells, and reduced tumor growth in a mouse xenograft model. Growth inhibition was accompanied by perturbed Krebs cycle activity, inhibition of lipid and nucleotide biosynthesis, and altered glutathione homeostasis. These findings indicate that PC-mediated anaplerosis in early-stage NSCLC is required for tumor survival and proliferation.

Continued functioning of the Krebs cycle requires the replenishment of intermediates that are diverted for anabolic uses or glutathione synthesis. This replenishment process, or anaplerosis, is accomplished via 2 major pathways: glutaminolysis (deamination of glutamine via glutaminase [GLS] plus transamination of glutamate to α-ketoglutarate) (18, 19) and carboxylation of pyruvate to oxaloacetate via ATP-dependent pyruvate carboxylase (PC) (EC 6.4.1.1) (refs. 3, 20, 21, and Figure 1A). The relative importance of these pathways is likely to depend on the nature of the cancer and its specific metabolic adaptations, including those to the microenvironment (20, 22).

For example, glutaminolysis was shown to be activated in the glioma cell line SF188, while PC activity was absent (17), despite the high PC activity present in normal astrocytes (23, 24). However, SF188 cells use PC to compensate for GLS1 suppression or glutamine restriction (20), and PC, rather than GLS1, was shown to be the major anaplerotic input to the Krebs cycle in primary glioma xenografts in mice (22). It is also unclear as to the relative importance of PC and GLS1 in other cancer cell types or, most relevantly, in human tumor tissues in situ. Our preliminary evidence from 5 non–small-cell lung cancer (NSCLC) patients indicated that PC expression and activity are upregulated in cancerous (CA) compared with paired noncancerous (NC) lung tissues (21), although it was unclear whether PC activation applies to a larger NSCLC cohort or whether PC expression was associated with the cancer and/or stromal cells. Nor has the relationship between PC activation and glutaminase status been explored in human patient...
studies. Moreover, the role of PC in cell survival and proliferation and whether glutaminolysis can compensate for this role under PC suppression in lung cancer cells is unknown.

Here, we have greatly extended our previous findings (21) in a larger cohort (n = 86) by assessing glutaminase 1 (GLS1) status and analyzing in detail the biochemical and phenotypic consequences of PC suppression in NSCLC. We found PC activity and protein expression levels to be, on average, respectively, 100% and 5- to 10-fold higher in cancerous (CA) lung tissues than in paired NC lung tissues resected from NSCLC patients, whereas GLS1 expression showed no significant trend. We have also applied stable isotope–resolved metabolomic (SIRM) analysis to paired freshly resected CA and NC lung tissue slices in culture (analogous to the Warburg slices; ref. 25) using either [U-13C] glucose or [U-13C, 15N] glutamine as tracers. This novel method provided information about tumor metabolic pathways and dynamics without the complication of whole-body metabolism in vivo. We used immunohistochemical analysis to verify the specific localization of PC in cancer cells within the tumor tissue. We further determined the functional role of PC in NSCLC cell lines using shRNA, which showed that attenuation of PC activity inhibited cell proliferation, colony formation, and tumor growth in mouse xenografts, while compromising the cells’ ability to quench oxygen radicals. These phenotypic effects can be linked to several important metabolic outcomes of PC suppression in the lung adenocarcinoma A549 cell line cultured in vitro and in vivo, including reduced lipid and nucleotide biosynthesis and altered glutathione homeostasis.

Results

PC expression and activity, but not glutaminase expression, are significantly enhanced in early stages of malignant NSCLC tumors. PC protein expression was significantly higher in primary NSCLC tumors than in paired adjacent NC lung tissues (n = 86, P < 0.0001, Wilcoxon test) (Figure 1, B and C). The median PC expression was 7-fold higher in the tumor, and the most probable (moderal) overexpression in the tumor was approximately 3-fold higher (see Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI72873DS1). We found that PC expression was also higher in the tumor tissue compared with that detected in the NC tissue in 82 of 86 patients. In contrast, GLS1 expression was not significantly different between the tumor and NC tissues (P = 0.213, Wilcoxon test) (Figure 1C and Supplemental Table 1).

To estimate in vivo PC activity, the production of 13C3-Asp from 13C6-glucose (Figure 1A) infused into NSCLC patients was determined by gas chromatography–mass spectrometry (GC-MS). A bolus injection of 10 g 13C6-glucose in 50 ml saline led to an average of 44% 13C enrichment in the plasma glucose immediately after infusion (Supplemental Table 2). Because the labeled glucose was absorbed by various tissues over the approximately 2.5 hours between infusion and tumor resection, plasma glucose enrichment dropped to 17% (Supplemental Table 2). The labeled glucose in both CA and NC lung tissues was metabolized to labeled lactate, but this occurred to a much greater extent in the CA tissues (Supplemental Figure 1A), which indicates accelerated glycolysis in these tissues.

The enrichment of 13C3-Asp (a PC marker, Figure 1A) was on average 117% higher (n = 34, P < 0.005) in NSCLC tumors than in NC lung tissue (Figure 1D). 13C3-Asp can also be produced after the second turn of the Krebs cycle via 13C4-citrate, which is the condensation product of 13C2-acetyl CoA and 13C5-oxaloacetate (OAA). Since the 13C enrichment of the 13C3-citrate precursor was lower than that of the 13C3-Asp product (Supplemental Figure 1, B and C, and Figure 1D), it argues against a significant contribution of this route to 13C3-Asp enrichment. We also measured PC activity by the fractional 13C enrichment of other markers, 13C4-citrate, 13C6-citrate, and 13C7-malate, which were, respectively, 27%, 88%, and 113% higher in CA lung tissues than in NC lung tissues (Figure 1D). We also observed that 13C7-citrate was 5- to 6-fold higher in fractional enrichment than was 13C6-citrate for both CA and NC lungs. This could be attributed to the expected low 13C enrichment in pyruvate and thus acetyl CoA due to a large dilution of the 13C-glucose tracer in the human body, leading to a lower probability of condensation of 13C5-OAA with 13C4-acetyl CoA than with unlabeled acetyl CoA to produce less 13C7-citrate than 13C6-citrate. Moreover, the fractional enhancement of 13C6-citrate in CA over NC lungs was less than that of 13C7-citrate (Figure 1D). This could be a result of increased flux through the pyruvate dehydrogenase–initiated (PDH-initiated) Krebs cycle reactions in CA lung compared with that in NC lung, as evidenced by the elevated fraction of 13C5-isotopologs of Krebs cycle metabolites in CA versus NC lung (Supplemental Figure 1B). This could in turn lead to a relatively higher enrichment of 13C4-acetyl CoA and thus its product (13C5-citrate) of condensation with 13C5-OAA in CA lung compared with that in NC lung. Altogether, the significant increase in the fractional enrichment of these PC markers reflects increased PC activity (which we refer to herein) in the cancer tissue.

In addition to enhanced production of labeled PC markers, we observed a higher percentage of enrichment in the 13C5-isotopologs of Krebs cycle metabolites (citrate+2, glutamate+2, succinate+2, fumarate+2, malate+2, and aspartate+2 in Supplemental Figure 1B) in NSCLC tumors than in NC lung tissue, which is indicative of enhanced glucose oxidation in the Krebs cycle via PDH.

Fresh tissue (Warburg) slices confirm enhanced PC and Krebs cycle activity in NSCLC. To further assess PC activity relative to GLS1 activity in human lung tissues, thin (<1 mm thick) slices of paired CA and NC lung tissues freshly resected from 13 human NSCLC patients were cultured in 13C4-glucose or 13C9,15N2-glutamine for 24 hours. These tissues maintain biochemical activity and histological integrity for at least 24 hours under culture conditions (Figure 2A, Supplemental Figure 2, A and B, and ref. 26). When the tissues were incubated with 13C5-glucose, CA slices showed a significantly greater percentage of enrichment in glycolytic 13C1-lactate (3 in Figure 2B) than did the NC slices, indicative of the Warburg effect. In addition, the CA tissues had significantly higher fractions of 13C1, 13C7, 13C7, and 13C5-citrate (4, 5, and 6 of citrate, respectively, in Figure 2B) than did the NC tissues. These isotopologs require the combined action of PDH, PC, and multiple turns of the Krebs cycle (Figure 2C). Consistent with the labeled citrate data, the increase in the percentage of enrichment of 13C1, 13C7, 13C7, and 13C5-glutamate (3, 4, and 5 of glutamate, respectively, in Figure 2B) in the CA tissues indicates enhanced Krebs cycle and PC activity.

When tissue slices were incubated with 13C5,15N2-glutamine, 13C9,15N2-glutamate (6 of glutamate in Supplemental Figure 2C) was produced in both CA and NC tissues, indicating active gluta-
minase (see below). In addition, the \(^{14}\text{C}_4\) isotopologs of succinate, fumarate, malate (data not shown), and citrate (Supplemental Figure 2C) were detected, which confirms glutaminase action. However, the enrichment patterns of all but 1 isotopolog of the Krebs cycle intermediates derived from \(^{13}\text{C}_5,^{15}\text{N}_2\)-glutamine were comparable between CA and NC tissues (Supplemental Figure 2C). These data, along with the GLS1 expression data (Figure 1C), indicate that GLS1 is active and likely a significant player in the anaplerosis of the Krebs cycle in both types of tissues. But unlike PC, GLS1 was not significantly activated in NSCLC tumors relative to that seen in NC lung tissues.

**PC is localized in the cancer cells within NSCLC tumors.** NSCLC tumors are very heterogeneous and comprise various cell types including cancerous epithelial cells, pneumocytes, fibroblasts, endothelial cells, and immune cells such as macrophages. To determine whether PC is expressed specifically in cancer cells, we performed immunohistochemical analysis of paired CA and NC tissues from 9 patient samples. Figure 3 and Supplemental Figure 3 illustrate elevated PC levels in 4 NSCLC tumors compared with levels in the paired NC lung tissue, which was also evident in 5 other pairs of samples (data not shown) and is consistent with the Western blot data (Figure 1C). Moreover, PC was localized in the perinuclear region of the cancerous epithelial cells but not in stromal cells in the tumor tissue (Figure 3, bottom left panel, and Supplemental Figure 3D). Such a perinuclear staining pattern is consistent with the localization of PC in the mitochondria, which is supported by the perinuclear costaining of PC and the mitochondrial marker in NSCLC A549 cells (Supplemental Figure 3, C and D).

In contrast, in the NC lung tissue, only macrophages were enriched in PC, while PC in other cell types stained weakly or was absent (Figure 3 and Supplemental Figure 3). Thus, PC is preferentially localized in the cancer cells of heterogeneous lung tumor tissues.

**PC suppression by shRNA in lung cancer cells leads to growth arrest and decreased tumor burden in vivo.** In order to assess the role of PC in the survival and growth of lung cancer cells, we used PC-specific shRNA to suppress PC mRNA expression in human A549 adenocarcinoma cells. As shown in Supplemental Figure 4A, vector shPC54 reduced PC protein levels to 20% of the control levels, whereas PC expression was barely detected in cells transduced with shPC55. To verify the inhibition of PC activity in cells, we incubated empty vector– (shEV), shPC54–, and shPC55–transduced A549 cells in \(^{13}\text{C}_6\)-glucose for 24 hours and measured the markers for PC activity by GC-MS (Figure 1A). The levels of \(^{13}\text{C}_3\)-citrate, \(^{13}\text{C}_3\)-aspartate, and \(^{13}\text{C}_3\)-malate were reduced by both constructs; however, shPC55 was
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**Figure 2.** Ex vivo CA lung tissue slices have enhanced oxidation of glucose through glycolysis and the Krebs cycle with and without PC input compared with that of paired NC lung slices. Thin slices of CA and NC lung tissues freshly resected from 13 human NSCLC patients were incubated with 13C6-glucose for 24 hours as described in the Methods. The percentage of enrichment of lactate, citrate, glutamate, and aspartate was determined by GC-MS. (A) 1H13C HSQC NMR showed an increase in labeled lactate, glutamate, and aspartate. In addition, CA tissues had elevated 13C abundance in the ribose moiety of the adenine-containing nucleotides (1′-APX), indicating that the tissues were viable and had enhanced capacity for nucleotide synthesis. (B) CA tissue slices (n = 13) showed increased glucose metabolism through glycolysis based on the increased percentage of enrichment of 13C6-lactate (3′), and through the Krebs cycle based on the increased percentage of enrichment of 13C6–4–6-citrate (4–6′) and 13C6–3–5-glutamate (3–5′) (see 13C fate tracing in CA). *P < 0.05 and **P < 0.01 by paired Student’s t test. Error bars represent the SEM. (C) An atom-resolved map illustrates how PC, PDH, and 2 turns of the Krebs cycle activity produced the 13C isotope of citrate and glutamate in B, whose enrichment were significantly enhanced in CA tissue slices.

more effective than shPC54 (Supplemental Figure 4B). It is interesting to note that increased PC activity may occur in other unrelated transformed cell lines (3, 27). PC knockdown slowed cell growth, again with shPC55 being more effective than shPC54 (Supplemental Figure 4C). This corroborates the greater extent of PC knockdown by shPC55 (Supplemental Figure 4A). Furthermore, PC knockdown greatly decreased the number of colonies formed in soft agar (Supplemental Figure 4D), reduced cell viability (Supplemental Figure 4E), and altered morphology (Supplemental Figure 4F).

Since shPC55 was the more effective construct in suppressing both PC expression and activity, we used it to probe the effects of PC suppression on the additional NSCLC adenocarcinoma cell lines H1299, H2030, HCC827, and PC9. PC knockdown inhibited effective in vivo as in cell culture. We found that PC expression in the knockdown tumors after 36 days had increased to 30% to 60% of that in tumors with control empty vector (Supplemental Figure 7A). Interestingly, the growth rate of individual tumors correlated significantly (P = 0.006) with their level of PC expression (Figure 4D). Therefore, residual PC expression is likely to account for the limited ability of the shPC55 tumors to grow. Altogether, these data indicate that PC expression is important for in vivo tumor growth.

**PC knockdown perturbs Krebs cycle activity, the anabolic capacity of the cell, and glutathione homeostasis.** To investigate the metabolic consequences of PC knockdown, A549 cells were transduced with shPC55, grown in 13C6-glucose or 13C6,15N2-glutamine, and incorporation of the label was analyzed by NMR and MS. In addition, the...
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The mouse engrafted with shPC55- or shEV-transduced A549 cells were injected with $^{13}$C$_6$-glucose. With the glucose tracer, we found no evidence that PC affects aerobic glycolysis, as PC-knockdown cells had similar glucose consumption and production of total and labeled lactate in the media (Supplemental Figure 7B). In addition, the lactate released into the media had the same percentage of enrichment as evidenced by the reduced incorporation of labeled glucose with PC knockdown were able to recover PC expression to some extent in vivo (Supplemental Figure 8). Consistently, we observed less glutamine carbon into glutamate, aspartate, and citrate (Supplemental Figure 9B). It should be noted that amidotransferase activity can also contribute to the production of the labeled products shown in Figure 5B and Supplemental Figure 9B. However, changes in amidotransferase activity are unlikely to account for the observed blockage of Gln metabolism induced by PC knockdown. This is because PC knockdown in A549 cells or in mouse xenografts led to suppressed GLS1 expression (Supplemental Figure 10A). In addition, A549 cells exhibited high GLS1 expression and GLS activity in cell lysates, suggesting that GLS1 expression is important for A549 cell growth (see below).

To further probe the importance of glutamine utilization in PC-suppressed cells, A549 cells were incubated with either 0.5 or 0.2 mM glutamine. We found that glutamine restriction significantly reduced the growth rate of control cells by approximately 50%, but had no influence on PC-knockdown cells (Supplemental Figure 10B and C). Thus, glutaminolysis does not compensate for the loss of PC as an anaplerotic supplier of precursors to the Krebs cycle intermediates in NSCLC cells.

Beyond supplying reducing equivalents and energy (ATP and GTP), the Krebs cycle also supplies various precursors for anabolism. For instance, citrate is exported from the mitochondria to supply acetyl CoA for fatty acid synthesis; OAA is transaminated to aspartate for pyrimidine biosynthesis; α-ketoglutarate is aminated to glutamate for de novo glutathione synthesis; and malate is decarboxylated to pyruvate, providing NADPH for lipid biosynthesis and glutathione reduction (Figure 1A). To determine how perturbation of the Krebs cycle by PC suppression affects these anabolic pathways, we used SIRM to follow the biosynthesis of nucleotides, lipids, and glutathione in A549 cells grown in $^{13}$C$_6$-glucose or $^{13}$C$_5$-glutamine tracers.

PC knockdown reduced the levels of uracil and adenine nucleotides both in cell culture and in vivo (Supplemental Figure 11, A and B). These changes were at least in part attributed to attenuated biosynthesis of these nucleotides by PC knockdown, as evidenced by the reduced incorporation of labeled glucose into the C1′ ribose carbon observed in HSQC NMR analysis of $^{13}$C($^{1}$H) (Supplemental Figure 9A and Supplemental Figure 11A). This can be accounted for by the reduced production of the uracil and cytosine ring via decreased synthesis of their precursor aspartate from glucose and glutamine both in cells (Figure 5) and in vivo (Supplemental Figure 8). Consistently, we observed less incorporation of the glucose and glutamine carbon into the ring system of the nucleotides in PC-knockdown cells.

Since glutaminolysis is another major anaplerotic pathway, we asked whether this pathway could compensate for PC deficiency in NSCLC cells. shPC55- or shEV-transduced A549 cells were incubated with $^{13}$C$_5$N$_2$-glutamine to track glutaminolysis. As shown in Figure 5B, GLS1 activity can produce $^{13}$C$_5$N$_2$-glutamate (6) from $^{13}$C$_5$N$_1$-glutamate (7), which subsequently enters the Krebs cycle to generate $^{13}$C$_5$-succinate (4), $^{13}$C$_4$- (4) or $^{13}$C$_4$N$_1$-aspartate (5), and $^{13}$C$_4$-citrate (4). A549 cells readily took up and metabolized $^{13}$C$_5$N$_2$-glutamate into these labeled metabolites. However, the levels of all these labeled isotopologs were significantly reduced in the PC-knockdown cells (highlighted by red circles in Figure 5B), which is consistent with reduced GLS activity. Decreased GLS activity was also evident in the $^{1}$H-$^{13}$C HSQC spectra, which showed that PC-suppressed cells incorporated less glutamine carbon into glutamate, aspartate, and citrate (Supplemental Figure 9B).
glycerol backbone (3 13C atoms), fatty acyl chains (even 13C), and phosphate (α)

Pearson’s correlation coefficient. Downloaded from http://www.jci.org on January 22, 2015. http://dx.doi.org/10.1172/JCI72873

less attenuation of PC expression (30%–60% of control) and growth inhibition. In addition, PC knockdown in the mouse xenografts (Figure 6A). FT-ICR-MS analysis (Figure 6A).

growth effect is not simply related to an inability to regenerate GSH from GSSG. Altogether, these results show that PC suppression compromises anaplerotic input into the Krebs cycle, which in turn reduces the activity of the Krebs cycle, while limiting the ability of A549 cells to synthesize nucleotides, lipids, and glutathione. These downstream effects of PC knockdown were also evident when comparing the metabolism of shPC55-transduced A549 cells against that of A549 cells transduced with a scrambled vector (shScr) (Supplemental Figure 13), which suggests that they are on-target effects of PC knockdown.

Glutaminolysis is also a functionally significant source of anaplerosis in NSCLC. To determine whether GLS1 is functionally important for NSCLC cells, we suppressed GLS1 expression by shRNA in both A549 and H1299, which resulted in reduced growth rates in both cells (Supplemental Figure 14, A and B). The growth of A549, H1299, and PC9 cells also decreased in response to BPTES (a known GLS inhibitor) treatment (Supplemental Figure 14C).

Figure 4. PC suppression via shRNA inhibits proliferation and tumorigenicity of human NSCLC cell lines in vitro and in vivo. Proliferation and colony-formation assays were initiated 1 week after transduction and selection with puromycin. A549 xenograft in NSG mice was performed 8 days after transduction. *P < 0.01, **P < 0.001, ***P < 0.0001, and ****P < 0.00001 by Student t test, assuming unequal variances. Error bars represent the SEM. (A) NSCLC cells lines were transduced with shPC55 or shEV. Proliferation assays (n = 6) revealed substantial growth inhibition induced by PC knockdown in all 5 cell lines after a relatively long latency period. (B) Colony-formation assays indicated that PC knockdown reduced the capacity of A549 and PC9 cells to form colonies in soft agar (n = 3). (C) Tumor xenografts from shPC55-transduced A549 cells showed a 2-fold slower growth rate than did control shEV tumors (P < 0.001 by the unpaired Welch version of the t test). Tumor size was calculated as ab/4, where a and b are the x,y diameters. Each point represents an average of 6 mice. The solid lines are the nonlinear regression fits to the equation: size = a + b \cdot t^2, as described in the Methods. (D) The extent of PC knockdown in the mouse xenografts (n = 6) was less than that in cell cultures, leading to less attenuation of PC expression (30%–60% of control) and growth inhibition. In addition, PC expression in the excised tumors correlated with the individual growth rates, as determined by Pearson’s correlation coefficient.

of UTP and CTP by Fourier transform ion cyclotron resonance (FT-ICR-MS) analysis (Figure 6A).

Lipid biosynthesis was assessed in the nonpolar fractions of the same cell extracts by FT-ICR-MS, which can resolve a large number of intact lipids and determine their 13C enrichments in the glycerol plus fatty acyl residues (odd >3 13C) in shEV control cells (28). With PC suppression, almost all phosphatidylcholines from both glucose and glutamine tracers (Figure 6B). Using 13C 6-glucose as a tracer, 13C was incorporated extensively into the glycerol backbone (3 13C atoms), fatty acyl chains (even 13C), and glycerol plus fatty acyl residues (odd >3 13C) in shEV control cells (28). With PC suppression, almost all phosphatidylcholine species were either unlabeled (as 0) or labeled at the glycerol backbone only (as 3), while the 13C isotopologs reflecting fatty acyl synthesis from glucose (even and odd >3) were greatly diminished in A549 cells. In addition, the glycerol backbone precursor α-glycerophosphate (αGP) (such as 13C 6α-GP derived from 13C 6-glucose) accumulated in the PC-knockdown cells (Figure 6B). Likewise, fatty acyl synthesis from 13C 6-glucose ("Even" in Figure 6B) via glutaminolysis and the Krebs cycle was greatly attenuated in PC-suppressed cells. Taken together, these results suggest that PC knockdown severely inhibits lipid production by blocking the biosynthesis of fatty acyl components but not the glucose-derived glycerol backbone. This is consistent with decreased Krebs cycle activity (Figure 5), which in turn curtails citrate export from the mitochondria to supply the fatty acid precursor acetyl CoA in the cytoplasm.

Finally, de novo glutathione synthesis was analyzed by 1H[13C] HSQC NMR. Glutathione synthesis from both glucose and glutamine was suppressed by PC knockdown (Supplemental Figure 9, A and B). Reduced de novo synthesis led to a large decrease in the total level of reduced glutathione (GSH; Supplemental Figure 12, A and B). At the same time, PC-knockdown cells accumulated slightly more oxidized GSH (GSSG; Supplemental Figure 12, A and B), leading to a significantly reduced GSH/GSSG ratio both in cell culture and in vivo (Supplemental Figure 12C). To determine whether this perturbation of glutathione homeostasis compromises the ability of PC-suppressed cells to handle oxidative stress, we measured ROS production by DCFDA fluorescence. PC-knockdown cells had over 70% more basal ROS than did control cells (0 mM H2O2; Supplemental Figure 12D). When cells were exposed to increasing concentrations of H2O2, the knockdown cells were less able to quench ROS, as they produced up to 300% more ROS than did control cells (Supplemental Figure 12D). However, N-acetylcysteine (NAC) at 10 mM did not rescue the growth of PC-knockdown cells, suggesting that such a growth effect is not simply related to an inability to regenerate GSH from GSSG. Allogreter, these results show that PC suppression compromises anaplerotic input into the Krebs cycle, which in turn reduces the activity of the Krebs cycle, while limiting the ability of A549 cells to synthesize nucleotides, lipids, and glutathione. These downstream effects of PC knockdown were also evident when comparing the metabolism of shPC55-transduced A549 cells against that of A549 cells transduced with a scrambled vector (shScr) (Supplemental Figure 13), which suggests that they are on-target effects of PC knockdown.
Further, $^{13}$C isotopolog analysis of these 3 cell lines after incubation with $^{13}$C$_5,^{15}$N$_2$-Gln showed that they used glutamine as a significant source of carbon for the Krebs cycle (Supplemental Figure 14D). Therefore, it appears that both GLS1 and PC are important for supporting proliferation and Krebs cycle activity in NSCLC cells. $^{13}$C$_5,^{15}$N$_2$-Gln as a substantial carbon source for the Krebs cycle was also recapitulated in the human tissue slice data (Supplemental Figure 2C). However, unlike PC-based anaplerosis (Figure 2B), GLS-fueled anaplerotic activity was comparable between CA and NC tissue slices (Supplemental Figure 2C), which is consistent with the lack of discernible distinction in GLS1 protein expression between bulk CA and NC tissues (Figure 1C).

Figure 5. PC knockdown perturbs glucose and glutamine flux through the Krebs cycle. $^{13}$C Isotopolog concentrations were determined by GC-MS ($n = 3$). Values represent the averages of triplicates, with standard errors. *$P < 0.05$, **$P < 0.01$, ***$P < 0.001$, and ****$P < 0.0001$ by Student’s t test, assuming unequal variances. The experiments were repeated 3 times. (A) A549 cells were transduced with shPCSS for 10 days before incubation with $^{13}$C$_6$-glucose for 24 hours. As expected, the $^{13}$C isotopologs of Krebs cycle metabolites produced via PC and Krebs cycle activity were depleted in PC-deficient cells (tracked by blue dots in the atom-resolved map and blue circles in the bar graphs; see also Figure 2C). In addition, $^{13}$C$_6$-glucose metabolism via PDH was also perturbed (indicated by red dots and circles). (B) Treatment of PC-knockdown cells with $^{13}$C$_5,^{15}$N$_2$-glutamine revealed that anaplerotic input via GLS did not compensate for the loss of PC activity, since GLS activity was attenuated, as inferred from the activity markers (indicated by red dots and circles). Decarboxylation of glutamine-derived malate by malic enzyme (ME) and reentry of glutamine-derived pyruvate into the Krebs cycle via PC or PDH (shown in blue and green, respectively) were also attenuated. Purple diamonds denote $^{15}$N; black diamonds denote $^{14}$N.
Figure 6. PC suppression hinders Krebs cycle–fueled biosynthesis. (A) 13C atom–resolved pyrimidine biosynthesis from 13C6-glucose and 13C5-glutamine is depicted with a 13C5-ribose moiety (red dots) produced via the pentose phosphate pathway (PPP) and 13C1-3 uracil ring (blue dots) derived from 13C2-4-aspartate produced via the Krebs cycle or the combined action of ME and PC (blue dots). A549 cells transduced with shPC55 or shEV were incubated with 13C6-glucose or 13C5-glutamine for 24 hours. Fractional enrichment of UTP and CTP isotopologs from FT-ICR-MS analysis of polar cell extracts showed reduced enrichment of 13C6-glucose–derived 13C5-ribose (the “5” isotopolog) and 13C6-glucose– or 13C5-glutamine–derived 13C1-3-pyrimidine rings (the “6–8” or “1–3” isotopologs, highlighted by dashed green rectangles; for the “6–8” isotopologs, 5 13Cs arose from ribose and 1–3 13Cs from the ring) (10, 45). These data suggest that PC knockdown inhibits de novo pyrimidine biosynthesis from both glucose and glutamine. (B) Glucose and glutamine carbons enter fatty acids via citrate. FT-ICR-MS analysis of labeled lipids from the nonpolar cell extracts showed that PC knockdown severely inhibited the incorporation of glucose and glutamine carbons into the fatty acyl chains (even) and fatty acyl chains plus glycerol backbone (odd >3) of phosphatidylycerine lipids. However, synthesis of the 13C3-glycerol backbone (the “3” isotopolog) or its precursor 13C3-α-glycerol-3-phosphate (αG3P, m+3) from 13C6-glucose was enhanced rather than inhibited by PC knockdown. These data suggest that PC suppression specifically hinders fatty acid synthesis in A549 cells. Values represent the averages of triplicates (n = 3), with standard errors. *P < 0.05, **P < 0.01, and ***P < 0.001 by Student’s t test, assuming unequal variances.
Discussion

Prior to our earlier report (21), the upregulation of PC in NSCLC was an undocumented phenomenon. Most research on anaplerosis in cancer cells has focused on the upregulation of glutamine metabolism via GLS1 in established cell lines, which cannot recapitulate the influence of the tumor microenvironment (17, 29–32). To gain insights into metabolic reprogramming in human cancer, it is imperative to study human tumors in situ and to compare paired CA and surrounding NC tissues from the same patient to circumvent cell culture artifacts, while eliminating confounding intrinsic genetic and physiological variables. It is also important to implement a fast and consistent liquid N₂-quenching protocol to minimize metabolic artifacts during tissue sampling after surgical resection.

By addressing the two aspects of our study, we found that although PC expression and activity in lung tissues varied substantially from patient to patient, 94% of the NSCLC tumors (82 of 86) overexpressed PC, and 73% (25 of 34) showed elevated PC activity in vivo relative to that in their paired NC lung tissues. This contrasts with the insignificant difference observed in the expression of GLS1 between CA and NC tissues (Figure 1C). PC was highly expressed exclusively in cancer cells of early-stage NSCLC tumor tissues. In contrast, PC was not expressed in the pneumocytes, but its expression was evident in macrophages of paired NC tissue (Figure 3 and Supplemental Figure 3). We surmise that the latter may be related to inflammation, which is common in the lungs of NSCLC patients.

The requirement of PC for NSCLC proliferation and tumorigenesis was demonstrated by PC suppression in several lung cancer cell lines (Figure 4, A and B) and in vivo for A549 cells (Figure 4, C and D) using specific shRNAs. The long latency period between mRNA knockdown and phenotypic effects (Figure 4A) may be related to the long half-life of PC (1–2 d, ref. 33). The intracellular level of PC may need several cell divisions to be attenuated sufficiently to compromise anaplerosis, leading to reduced cell proliferation. We have verified that inhibition of cell proliferation by PC suppression is accompanied by a decrease in anaplerotic input into the Krebs cycle (Figure 5) and the consequent reduction of nucleotide (Figure 6A, Supplemental Figure 11A, and Supplemental Figure 13C) and lipid biosynthesis (Figure 6B and Supplemental Figure 13D). The correlation of the severity of these metabolic effects with the efficiency of the knockdown and the inhibition of cell proliferation both in vitro and in vivo (Figure 4 and Supplemental Figure 4) is consistent with PC knockdown being a significant contributor to the reduced proliferation. Although we have clearly demonstrated the importance of PC for cancer cell proliferation and that enzymic activity is an important aspect of the role of PC, we cannot rule out the possibility that the protein has alternative nonenzymic functions in the cells, as has been observed for several other metabolic enzymes (34).

In addition, PC knockdown in A549 cells greatly depletes the GSH pool and perturbs glutathione homeostasis (Supplemental Figure 12C) by blocking de novo glutathione synthesis (Supplemental Figure 12B and Supplemental Figure 13B), which presumably leads to greater ROS production (Supplemental Figure 12D) and a compromised ability to survive oxidative stress. Exogenous reductants such as NAC in some circumstances can reduce cellular GSSG to GSH, but NAC cannot overcome the depleted GSH pool in PC-suppressed cells, thus accounting for the failure of NAC to rescue PC-suppressed cells from growth inhibition (Supplemental Figure 12E).

Although active, anaplerotic glutaminolysis could not compensate for lost PC-based anaplerosis in A549 cells (Figure 5B and Supplemental Figure 10), despite the abundant supply of glutamine. This is presumably due to the reduced expression of GLS1 in PC-knockdown cells or in xenograft tumors (Supplemental Figure 10A). Moreover, we noted that in the human NSCLC tumors, in which PC was underexpressed, GLS1 was also underexpressed. These results suggest that PC and GLS1 are not compensatory in their functions in human NSCLC, in contrast to some glioma cells (20).

Finally, the improved variant of Warburg’s tissue slices (25) that we have introduced enables detailed ex vivo analysis of patient-specific tumor metabolism (Figure 2 and Supplemental Figure 2), while maintaining most of the tissue microenvironment and cellular heterogeneity without systemic complications from the blood supply. This ex vivo system also affords versatility of experimental manipulations (e.g., tracer choices or drug treatments) similar to that of the pure cell culture system. In our case, enhanced in vivo PC activity (Figure 1D) in human NSCLC tumor tissues was confirmed by the corresponding ex vivo tracer studies (Figure 2B). Equally important is that the ex vivo system enables unequivocal determination of anaplerotic glutaminolysis using the labeled glutamine tracer (Supplemental Figure 2C), thereby corroborating the lack of in vivo upregulation of GLS1 in human tumor tissues (Figure 1C). This would have been difficult to demonstrate in vivo in human patients due to cost considerations. Thus, the ex vivo tissue culture system can complement existing model systems for translating bench-to-bedside understanding, particularly in an individualized manner (26).

In conclusion, our results in primary human lung tumors and ex vivo tissue slices show that PC protein expression and activity are enhanced in early stages of human NSCLC tumor tissues compared with that seen in NC tissues from the same patient and that the PC protein is localized in the cancer cells within the tumor tissue. Silencing PC expression and activity compromises NSCLC cell growth, tumorigenicity, and antioxidation capacity. These phenotypic effects of PC suppression were related to the vital metabolic functions of PC in NSCLC, including sustaining mitochondrial Krebs cycle activity, nucleotide and lipid biosynthesis, and redox homeostasis.

Methods

Patient accrual. Eighty-six patients with suspected primary lung cancer but without diagnosed diabetes were recruited on the basis of their surgical eligibility. The patients were then randomly grouped into 2 arms. In the first arm, the patients were overnight fasted (>8 hours) before 10 grams of uniformly labeled ¹³C glucose was administered i.v. and again 2.8 ± 0.5 hours prior to video-assisted thoroscopic surgical (VATS) wedge resection, as previously described in detail (35), to enable tracking of metabolic pathways (21, 36–38). Patients in the second arm did not receive a glucose injection. We and others have previously discussed the relative advantages of bolus injections versus slow-infusion protocols (21, 36–41). The bolus injection leads to an initially high blood glucose concentration, which then decays to the baseline level, thus limiting the length of time that effective labeling can be achieved but more closely approximating an initial rate condition. In comparison, a prolonged slow infusion can lead
to a higher degree of isotopic labeling and approach isotopic steady-state conditions, but is a much more involved process, and a true isotopic steady state may be difficult and cost prohibitive to achieve.

The extent of resection was determined by the surgeon according to clinical criteria. The majority of the specimens were obtained from wedge resections to minimize surgical times, while the remaining were acquired in less than 5 minutes after the pulmonary vein was clamped; both practices helped avoid development of significant ischemia in resected tissues. Immediately after resection, the tumor was transsected, and sections of CA tissue and surrounding NC lung tissue at least 2 cm away from the tumor were biochemically quenched by flash-freezing in liquid N$_2$ (36, 38). The margins of the tumor were initially assessed by the surgeon by visual inspection. Parallel tissue samples were sent to on-site pathologists for confirmation of the diagnosis and the cancer-free margins. The remaining specimen was preserved in buffered formalin for detailed pathological examination.

**Sample preparation.** The frozen tissue samples were pulverized to a particle size of less than 10 μm in liquid N$_2$ using a Spex freezer mill (Spex). Polar and nonpolar metabolites were extracted from the ground tissue, which also yielded insoluble protein residues (21, 42). The protein residues were extracted by homogenization in 2% SDS, 62.5 mM Tris, and 1 mM DTT buffer for protein determination using the Pierce BCA method (Thermo Fisher Scientific).

**Electrophoresis, Western blotting, immunohistochemistry, cell proliferation, and colony-formation assays.** Detailed protocols are described in the Supplemental Methods.

**shRNA knockdown of PC in lung cancer cells.** NSCLC cells lines and HEK 293T cells obtained from ATCC were grown in DMEM media containing 0.45% glucose and 4 mM glutamine at 37°C, 95% relative humidity, and 5% CO$_2$. HEK 293T cells were transfected with pCMV and pMD2.G vectors carrying viral packaging (gag, pol, rev) and envelope (VSV-G) genes, along with a plasmid containing shRNA against PC (shPC54 or shPC55), shEV, or shScr (Mission shRNA; Sigma-Aldrich). The anti-PC sequences used were: shPC54: CCGGGCCAGTTATTTGGCAGAAATTCGCAGATTCTGACACCATAAACTGGGCTTTTTG; shPC55: CCGGGCCAGAAGGAGAA-CACAAGTGAATGCTCAGATCTACGTTTCTCCTTTGGGCTTTTG.

After 48 hours, medium from the transfected cells was collected, filtered, supplemented with 16 μg/ml polybrene, and added to NSCLC cell lines in an equal volume of DMEM media. Fresh media were added to the 293T cells, and after 24 hours, transduction of the NSCLC cells was repeated. The resulting NSCLC cells bearing the plasmid with the puromycin resistance gene were selected using 1 μg/ml puromycin.

**Tracer studies in cell cultures.** Cells transduced with shEV, shScr, shPC54, or shPC55 were incubated in DMEM in the presence of 0 mM $^{13}$C$_6$-glucose or 2 mM $^{13}$C$_5$, $^{15}$N$_2$-glutamine for 24 hours, quenched in cold acetoniure, and extracted in acetoniure/water/chloroform (v/v 2:1:5:1), as described previously (10).

**Tracer studies in mouse xenographs of PC-knockdown cells.** To assess tumor growth of PC-knockdown A549 cells, 6-week-old female NOD/SCID gamma (NSG) mice (n = 6 mice/group) (The Jackson Laboratory) were injected s.c. with $8 \times 10^6$ cells containing either shEV or shPC55 mixed with Matrigel (BD Biosciences). The tumor diameters were determined 3 times per week by caliper. Tumor sizes were calculated as $\pi r^2/4$, where a and b are the lengths of the maximum and minimum dimensions. After 36 days, the mice were injected via the tail vein with 100 μl 1.075 M [U-$^{13}$C]-glucose 3 times at 15-minute intervals. Blood samples were taken intraorbitally immediately after the first injection and at sacrifice 60 minutes later. The tumors were immediately excised and weighed prior to flash-freezing in liquid N$_2$. Metabolites were extracted as described above (43).

**Tracer studies in ex vivo tissue slices.** These experiments were performed on fresh slices of paired tumor and nontumorous lung tissues resected from individual patients (26). Upon lung resection, thin slices (0.5-1 mm thick) of tissue were excised using a Weck microtome in the operating room. These slices were immediately placed in T-flasks containing DMEM with the appropriate tracer (either 10 mM [U-$^{13}$C]-glucose or 2 mM [U-$^{13}$C,$^{15}$N]-glutamine), and then transferred to a CO$_2$ incubator set at 37°C and 5% CO$_2$. The flasks were continuously rocked for 24 hours for aeration and to maintain constant nutrient supplies at the tissue surface, while avoiding local buildup of waste products such as acids. The slices were then washed in cold PBS, frozen in liquid N$_2$, pulverized as described above, and extracted for polar and nonpolar metabolites and proteins as described above for cell cultures.

**SIRM analysis.** Polarmetabolite fractions from cells, patient tissues, and mouse tumors were analyzed by GC-MS, NMR, and FT-ICR-MS as previously described (10, 42). Nonpolar fractions were analyzed for lipids by FT-ICR-MS as previously described (10, 28). $^{13}$C fractional enrichments were calculated for metabolite isotopomers ($^{13}$C at different atomic positions) and isotopologs (different numbers of $^{13}$C atoms) as the level of a given isotopemer or isotopolog divided by the summed level of all isotopomers or isotopologs. The absolute levels of labeled metabolites were obtained by normalizing the moles of metabolites to the extractable protein content.

**Statistics.** Statistical analyses were conducted using SPSS 19.0 (IBM- Somers) and Kaleidagraph (Synergy Software). Differences between patient cohorts were assessed using the paired Wilcoxon test, the 2-tailed Student t test on the logarithms of the values (i.e., for fractional differences), and by examining the distribution terms of the mean, median, and modal values. The expression ratios (CA/NC) for PC or GLS were highly non-normally distributed, and the mean, median, and modal values were calculated accordingly. The modal values were estimated from the maximum in the frequency-versus-ratio histograms and were thus slightly affected by the bin size.

Mouse tumor sizes were tested using an unpaired Welch’s version of the t test (44). First, the significance was evaluated at each time point after injection. Second, all growth curves were fitted to different models, including exponential, power, and quadratic growth, and best-fit parameters with the associated standard deviations were subjected to Welch’s t test. The simplest form that gave good fitting was a simple quadratic equation: size = $a + b t^2$. In this form, a is the initial tumor size at $t = 0$, and b is the specific growth rate. Third, the tumor weights at necropsy were compared as 2 groups. The significance of differences between shEV and shPC groups in terms of proliferation and metabolites was determined using Welch’s t test. In all analyses, a P value of less than 0.05 was considered significant.

**Study approval.** This study was approved by the IRB of the University of Louisville. Written informed consent was obtained from all subjects prior to inclusion in this study. The murine tumor protocol was performed in compliance with all relevant laws and institutional guidelines and was approved by the IACUC of the University of Louisville.
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