Acetate Recapturing by Nuclear Acetyl-CoA Synthetase 2 Prevents Loss of Histone Acetylation during Oxygen and Serum Limitation

Highlights

- Acetyl-CoA synthetase 2 (ACSS2) expression controls acetate uptake and utilization
- Exogenous acetate is used for fatty acid synthesis rather than histone acetylation
- The main function of nuclear ACSS2 is to retain acetate released from histones
- Nuclear and cytosolic acetyl-CoA pools are largely compartmentalized

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In Brief

Acetyl-CoA synthetase 2 (ACSS2) generates acetyl-CoA from acetate and is important for tumor growth. Bulusu et al. show that, in hypoxic tumor regions, ACSS2 is prominently expressed in the nuclei of tumor cells. Here it maintains histone acetylation by recapturing acetate released by histone deacetylases so it can be re-used for acetylation.
Acetate Recapturing by Nuclear Acetyl-CoA Synthetase 2 Prevents Loss of Histone Acetylation during Oxygen and Serum Limitation

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SUMMARY

Acetyl-CoA is a key metabolic intermediate with an important role in transcriptional regulation. The nuclear-cytosolic acetyl-CoA synthetase 2 (ACSS2) was found to sustain the growth of hypoxic tumor cells. It generates acetyl-CoA from acetate, but exactly which pathways it supports is not fully understood. Here, quantitative analysis of acetate metabolism reveals that ACSS2 fulfills distinct functions depending on its cellular location. Exogenous acetate uptake is controlled by expression of both ACSS2 and the mitochondrial ACSS1, and ACSS2 supports lipogenesis. The mitochondrial and lipogenic demand for two-carbon acetyl units considerably exceeds the uptake of exogenous acetate, leaving it to only sparingly contribute to histone acetylation. Surprisingly, oxygen and serum limitation increase nuclear localization of ACSS2. We find that nuclear ACSS2 recaptures acetate released from histone deacetylation for recycling by histone acetyltransferases. Our work provides evidence for limited equilibration between nuclear and cytosolic acetyl-CoA and demonstrates that ACSS2 retains acetate to maintain histone acetylation.

INTRODUCTION

Cancer is a disease of unrestrained cell proliferation. This necessitates the continued production of cellular components, including lipid membranes, for the generation of daughter cells (DeBerardinis et al., 2008). The bulk elements of membranes are fatty acids, and their de novo synthesis by cancer cells places a high demand on the precursor acetyl-coenzyme A (AcCoA) (Menendez and Lupu, 2007; Currie et al., 2013). In addition to its role in biomass production, the central position of AcCoA in both anabolic and catabolic pathways makes it a rheostat of the metabolic state of the cell (Pietrocola et al., 2015; Shi and Tu, 2015; Fan et al., 2015). AcCoA abundance directly affects the metabolic activity of many enzymes, and acetylation events on histones and other proteins control cellular functions at the transcriptional and post-translational levels. Disrupting acetylation homeostasis by inhibiting deacetylases (i.e., histone deacetylases [HDACs] and sirtuins) was shown to influence cancer cell survival (Falkenberg and Johnstone, 2014). It is therefore crucial for cells to maintain AcCoA homeostasis.

In solid tumors, cells frequently experience reduced oxygen availability because of aberrant vascularization (Brown and Wilson, 2004). Although, in oxygenated cells, most AcCoA is made from glucose, during hypoxia, stabilization of HIF1 causes cancer cells to shunt most glucose-derived carbon toward lactate (Kim et al., 2006; Papandreou et al., 2006). This diminishes AcCoA production from glucose, and how cancer cells adapt has been a topic of interest. Tracing experiments revealed that glutamine, through reductive carboxylation, is a precursor for AcCoA that is used for fatty acid biosynthesis, also known as lipogenic AcCoA (Wise et al., 2011; Metallo et al., 2011; Mullen et al., 2011). It has later been argued that a reduction in citrate levels in hypoxic cells causes increased reversibility with α-ketoglutarate. Because of this, at least some fatty acid labeling from glutamine could be explained as isotope exchange (i.e., mixing of label) rather than a net reductive flux (Fan et al., 2013).

More recently, it was found that 13C-acetate significantly labels the lipogenic AcCoA pool of hypoxic cells (Kamphorst et al., 2014) and, consequently, lipids (Schug et al., 2015), suggesting that acetate can act as an alternative carbon source for biomass production under these conditions. In a separate study, glioblastomas and brain metastases were shown to oxidize acetate for energy production, indicating that the metabolic fate may depend on tumor type (Mashimo et al., 2014). Importantly, silencing the nuclear-cytosolic isoform of AcCoA synthetase (ACSS2), which converts acetate to AcCoA, reduced tumor growth in a xenograft model of breast cancer (Schug et al., 2015). Furthermore, ACSS2 deletion diminished tumor burden in a genetically engineered mouse model of hepatocellular carcinoma (Comerford et al., 2014).
The extensive fatty acid labeling from 13C-acetate in hypoxic cancer cells may indicate that the increased ACSS2 expression supports biomass production. It is important to note, however, that the actual carbon contribution to fatty acid synthesis remains to be determined. As with glutamine, the fractional labeling of AcCoA from acetate may not reflect net synthesis because of isotope exchange (Fan et al., 2013). A rapid equilibration between the acetate and AcCoA pools could occur as a consequence of a fast protein acetylation-deacetylation cycle, with the cellular acetate pool also exchanging with labeled medium acetate. This would result in pronounced labeling of AcCoA and, subsequently, fatty acids from 13C-acetate without a net carbon transfer. Therefore, a more extensive evaluation to quantify the contribution of acetate to biomass production is needed. Furthermore, it has been reported that a significant proportion of ACSS2 localizes to the nuclei of tumor cells (Comerford et al., 2014), and how much exogenous acetate can contribute to nuclear processes such as histone acetylation remains unknown.

Here we apply innovative stable isotope tracing and mass spectrometry approaches to quantify acetate consumption and utilization by downstream pathways in a panel of cancer cell lines with varying levels of ACSS2 expression. We find that the combined expression of ACSS1 and ACSS2 determines the net acetate uptake rate. Exogenous acetate is used extensively by the mitochondria and for lipogenesis, and the demand for acetate substantially exceeds its uptake. As a consequence, exogenous acetate only modestly labels histone-bound acetate. However, nuclear localization of ACSS2 increases during oxygen and serum limitation, and nuclear ACSS2 is prominent in poorly perfused, hypoxic tumor regions of a mouse model of breast cancer. We find that the primary function of nuclear ACSS2 is to retain endogenous acetate released by deacetylases to maintain histone acetylation and propose that this is especially relevant in hypoxic and nutrient-limited areas of the tumor.

RESULTS

ACSS2 Expression Dictates Lipogenic AcCoA Labeling from U-13C-Acetate in Hypoxic Cancer Cells

A substantial fraction of the AcCoA used for fatty acid biosynthesis (i.e., lipogenic AcCoA) is produced from glucose and glutamine (Figure 1A). Alternatively, lipogenic AcCoA can be produced from acetate by ACSS2, and this pathway has been reported to be induced in hypoxic tumor cells (Schug et al., 2015; Comerford et al., 2014). Importantly, we determine the contribution of these different precursors to lipogenic AcCoA from fatty acid labeling without the need to analyze AcCoA directly, which is a mixture of all pools in the cell (Kamphorst et al., 2014; Tumanov et al., 2015).

We previously reported a substantial contribution of acetate to lipogenic AcCoA in hypoxic cancer cells, including MDA-MB-468 breast cancer cells (Kamphorst et al., 2014). Because the concentration of acetate in human plasma is 50–250 μM (Schug et al., 2015), we sought to confirm the propensity of these cells to use acetate in the lower end of this physiological range. Normoxic and hypoxic (1% O2) MDA-MB-468 cells were exposed...
to 90 μM U-13C-acetate. Although a relatively small fraction of lipogenic AcCoA labeled from U-13C-acetate in normoxic cells, this increased to approximately 30% in hypoxia (Figure 1B). Next, cells were cultured in the presence of U-13C-glucose, U-13C-glutamine, and U-13C-acetate to confirm that, combined, these are the major substrates for lipogenic AcCoA production (Figure 1C). As expected, the contribution from glucose dropped considerably in hypoxia, and labeling from both U-13C-glutamine and U-13C-acetate increased. Under both conditions, the majority of lipogenic AcCoA was labeled from all substrates combined. The somewhat lower labeling under hypoxic conditions could, in part, be explained by the presence of residual unlabeled acetate (≈15 μM) in the medium. Therefore, when provided in a physiologically relevant concentration, acetate is an important substrate for lipogenic AcCoA.

ACSS2 is reportedly responsible for acetate utilization for lipogenesis, but an accurate quantitation of its contribution is lacking. When silencing ACSS2 expression in MDA-MB-468 cells using small interfering RNA (siRNA) (Figure 1D), we observed a pronounced reduction in labeling of lipogenic AcCoA from U-13C-acetate (Figure 1E). Similar results were obtained for BT-474 cells (Figures S1A and S1B). Under these conditions, labeling from U-13C-glucose and U-13C-glutamine increased to ≈80% (Figure 1F). We found that unlabeled acetate in the medium suppressed labeling of lipogenic AcCoA from U-13C-glucose and U-13C-glutamine, indicating that acetate is preferentially used for lipogenesis under these conditions (Figure S1C). The increased labeling from U-13C-glucose and U-13C-glutamine upon ACSS2 knockdown may not indicate increased AcCoA biosynthesis from these substrates; net production may remain equal while fractional labeling increases because of the drop in the contribution from acetate. This supports earlier findings that acetate, glucose, and glutamine are the main substrates for lipogenic AcCoA (Kamphorst et al., 2014).

To further investigate the role of ACSS2 expression in acetate utilization for lipogenic AcCoA, we performed U-13C-acetate tracing in a set of six cancer cell lines originating from cervical (HeLa), lung (A549), pancreatic (AsPC-1), and breast tumors (HeLa, BT-474). This set covers a wide range of ACSS2 expression and facilitates assessment of the relation between ACSS2 expression and lipogenic AcCoA labeling in cells with the same or different tissue backgrounds (Figures 1G and 1H). Across cell lines, a clear correlation was found between ACSS2 expression and lipogenic AcCoA labeling from U-13C-acetate. ACSS1 expression generally appeared to be low in most cell lines, except for BT-474 cells (Figure S1D). In this cell line, silencing ACSS1 had no effect on ACSS2 expression but did cause a small increase in lipogenic AcCoA labeling from U-13C-acetate (Figures S1E and S1F), perhaps because of some degree of competition between ACSS1 and ACSS2 for the available acetate. Together, these results show a strong correlation between ACSS2 expression and lipogenic AcCoA labeling from U-13C-acetate.

Cancer Cells Exchange Acetate with the Medium, and Net Exchange Is Determined by ACSS2 Expression

As stated earlier, fatty acid labeling from U-13C-acetate is not sufficient to determine the carbon contribution from acetate because of possible cycling and exchange (Fan et al., 2013; Kamphorst et al., 2014). To measure acetate directly and quantify its net uptake by cancer cells, we developed a method based on alkylation of acetate to its propyl ester derivative using a methyl chloroformate (MCF)-catalyzed derivatization reaction and subsequent gas chromatography-mass spectrometry (GC-MS) analysis (Experimental Procedures; Figure 2A; Tumanov et al., 2016). For our analyses, we used [2H3]-acetate as an internal standard, enabling within-run quantification of both 13C-acetate and U-13C-acetate. We applied this method to analyze uptake of 90 μM U-13C-acetate spiked into the medium by both normoxic and hypoxic MDA-MB-468 cells. The hypoxic cells consumed almost all U-13C-acetate (Figure 2A). Interestingly, although, in normoxic cells, lipogenic AcCoA labeled only marginally from acetate (Figure 1B), U-13C-acetate uptake was comparable with hypoxic cells.

To our surprise, in addition to a reduction in U-13C-acetate because of consumption, unlabeled acetate (12C-acetate) accumulated in the cell-conditioned medium (Figure 2B). We concluded that this acetate was being released by cells because the initial medium 12C-acetate and U-13C-acetate concentrations remained stable when incubated in the absence of cells (Figure S2B). Furthermore, this acetate labeled from U-13C-glucose and U-13C-glutamine (Figure S2C). Acetate release was approximately equal to acetate consumption, resulting in an unchanged total medium acetate concentration (Figure 2C). This was also observed with physiological concentrations of glucose (5.5 mM) and glutamine (0.65 mM) (Figure S2D). Under the conditions described here, lipogenic AcCoA and, hence, fatty acid labeling from U-13C-acetate was prominent (Figure 1B). However, because no net uptake of acetate could be observed, it is caused by cycling between acetate and lipogenic AcCoA and exchange between cellular and medium acetate.

To expand our analysis of acetate uptake (u), release (r), and net exchange (e) flux independently of cell number and growth, we cultured cells in medium containing various concentrations of U-13C-acetate. The net exchange rate was calculated as the change in total medium acetate divided by the area under the curve growth curve. To deconvolute the uptake and release components of the exchange, we used the balance equations for the exchange of total acetate (e = r − u) and 12C-acetate (xe = ar − bu), where x is the concentration change in medium 12C-acetate relative to the concentration change of total acetate, a is the average 12C-acetate fraction inside cell, and b is the average 12C-acetate fraction in the medium. The parameters e, x, and b can be estimated from the measurement of medium 12C- and U-13C-acetate at different time points (see Supplemental Experimental Procedures for more information). The intracellular free acetate concentration was too low to quantitively accurately. We therefore considered all biologically relevant values of the intracellular acetate fractions (Supplemental Experimental Procedures). Under all conditions tested, both uptake and release of acetate were ≈3–4 mmol/hr/L cell volume, with little variation because of extracellular acetate concentration or oxygenation (Figure 2D). In contrast, the net exchange rate remained close to zero, with a tendency toward net acetate release.

We expanded our net acetate exchange experiments to the panel of six cancer cell lines (Figure 1G) cultured in hypoxia.
and high (10%) or low (1%) serum (Figure 2E). The cell lines with low ACSS2 expression (HeLa and A549) showed a net release of acetate that was reduced in low serum. Generally, culturing cells in low serum and hypoxia caused induction of ACSS2 expression (Figure S2E; Schug et al., 2015), and this led to a net acetate uptake for MDA-MB-231 and MDA-MB-468 cells. For the latter, net exchange was comparable with AsPC-1 cells, which have similar levels of ACSS2. BT-474 cells express high levels of ACSS2 because of gene amplification (Schug et al., 2015). They were found to most avidly take up acetate, especially in low serum (≈8 mmol/hr/L cell volume). Together, these results demonstrate that cells exchange acetate with the medium and that high ACSS2 expression promotes net acetate uptake.

Mitochondrial and Lipogenic Utilization of Exogenous Acetate Limits Its Use for Histone Acetylation

Net acetate uptake is highest in low serum, and this may be more reflective of the tumor microenvironment. We therefore used low serum for subsequent analyses. After being activated to AcCoA, exogenous acetate can be directed toward mitochondria for oxidation (Mashimo et al., 2014), used for biomass production (particularly lipogenesis), or used for acetylation. To determine whether acetate feeds into mitochondria, we analyzed tricarboxylic acid (TCA) cycle intermediate labeling from U-13C-acetate. Substantial (15%–25%) labeling occurred in BT-474 cells and, to a lesser extent, in MDA-MB-468 cells despite the hypoxia and low serum (Figures 3A and 3B). ACSS1 is responsible for mitochondrial utilization of acetate, and its knockdown substantially, although not completely, reduced labeling of TCA cycle intermediates (Figures S3A and S3D). Conversely, overexpressing ACSS1 in MDA-MB-468 cells led to a 4-fold increase in labeling of TCA cycle intermediates (Figures S3B and S3C). In BT-474 cells, ACSS1 silencing caused a concomitant reduction in exogenous acetate uptake by approximately 30% (Figures 3C and 3D). Because ACSS1-generated AcCoA does not appear to feed into fatty acid biosynthesis, it provides an estimate of mitochondrial acetate utilization.

A major AcCoA-consuming pathway in the cytosol is lipogenesis. To determine the AcCoA demand of this pathway,
we performed kinetic flux profiling by analyzing the incorporation of 13C from U-13C-glucose, U-13C-glutamine, and U-13C-acetate into the palmitate pool over time (Supplemental Experimental Procedures; Figures 3E and 3F; Tumanov et al., 2015). The time-based label incorporation reflects the synthesis rate, which can be deduced by fitting a differential equation (Yuan et al., 2008). In agreement with previous reports, hypoxia caused cells to reduce palmitate synthesis rates in the presence of high serum, particularly in MDA-MB-468 cells, because of the presence of exogenous lipids (Kamphorst et al., 2013; Young et al., 2013). Exposing cells to low serum led to a potent induction of the palmitate synthesis rate, increasing the demand for lipogenic AcCoA. Strikingly, in low oxygen and serum, the two-carbon acetyl demand for palmitate synthesis alone outstripped the exogenous acetate uptake rate 2- to 3-fold.

The exact flux of acetate into lipogenesis could not be determined because of the exchange. However, labeling of lipogenic AcCoA from U-13C-acetate is profound (Figure 1H), and exogenous acetate suppresses labeling from U-13C-glucose and U-13C-glutamine (Figure S1C). It is therefore likely that a
large proportion of the exogenous acetate is consumed for lipogenesis.

Last, we determined histone-bound acetate labeling from U-\textsuperscript{13}C-glucose, U-\textsuperscript{13}C-glutamine, or U-\textsuperscript{13}C-acetate. We did this by hydrolyzing acetate from purified histones rather than measuring individual acetylated peptides (Experimental Procedures). Therefore, the measured labeling reflects the aggregate of the entire histone-bound acetate pool. Not surprisingly, glucose was the major carbon donor under normoxic conditions (Figures 3G and 3H). In hypoxia and low serum, the contribution from glucose dropped, and the percent labeling from U-\textsuperscript{13}C-glutamine increased \(\sim 2\)-fold, indicating that reductive carboxylation also occurs for histone acetylation (Figures 3G and 3H). Acetate was the smallest contributor in both cell lines (<10%). Although hypoxia and low serum caused this to increase to \(\sim 15\%\), it was still much less than either glucose and glutamine, and multiple-fold less than for lipogenic AcCoA (Figure 1H; Figure S3F).

ACSS1 overexpression in MDA-MB-468 cells did not affect this (Figure S3E). Thus, exogenous acetate is inefficiently used for histone acetylation.

We hypothesized that, upon cell entry, exogenous acetate would be rapidly activated to AcCoA before being able to reach the nucleus and that this AcCoA would locally feed into the carbon-demanding pathways of mitochondrial metabolism and lipogenesis rather than being used for histone acetylation. We therefore supplied the free fatty acids palmitate and oleate to cells to reduce lipogenic demand for AcCoA. Free fatty acids are avidly consumed by cells (Kamphorst et al., 2013), and this led to an \(\sim 50\%\) reduction in de novo lipogenesis (Figure 3I). This was accompanied by a significant increase in labeling of histone-bound acetate from the exogenously supplied U-\textsuperscript{13}C-acetate (Figure 5J). Thus, the high lipogenic consumption of exogenous acetate at least partly explains its limited use for histone acetylation.

**Oxygen and Serum Limitation Promote Nuclear Localization of ACSS2, and ACSS2 Is Prominently Nuclear in Hypoxic Tumor Regions**

Although exogenous acetate is only sparingly used for histone acetylation, ACSS2 has been reported to be substantially nuclear (Comerford et al., 2014). To quantify this, we performed an analysis of ACSS2 localization in MDA-MB-468 cells using immunofluorescence (Figures 4A and 4B). Under normoxic and high serum conditions, \(\sim 15\%\) of the ACSS2 pool was localized to the nucleus. However, exposing the cells to low serum and oxygen caused a significant, \(\sim 3\)-fold increase in the fraction of nuclear ACSS2 (Figure 4B). A similar trend was observed for BT-474 cells (Figure S4). Further assessment of the individual factors indicated that both serum and oxygen limitation contribute to the increased nuclear localization of ACSS2, although the effect of serum limitation appeared to be strongest (Figure 4B). Low glucose did increase the localization of ACSS2 to the nucleus in normoxic cells but not in hypoxic cells.

To determine the in vivo expression and localization of ACSS2 as a function of oxygenation, serial sections were prepared from established tumors of the mouse mammary tumor virus polyoma middle T antigen (MMTV-PyMT) genetically engineered mouse model of breast cancer (Guy et al., 1992). These were immunostained for ACSS2, the hypoxic marker carbonic anhydrase IX (CAIX), and the blood vessel marker CD31, in addition to H&E staining (Figure 4C). Overall, ACSS2 staining was particularly intense in hypoxic (high CAIX, low CD31) regions of the tumor and exhibited prominent nuclear localization (Figure 4C). Indeed, scoring of nuclear intensity of ACSS2 staining revealed that nuclear ACSS2 levels are substantially higher in hypoxic regions than in perfused, normoxic regions (Figure 4D). These data show that cultured cells in low oxygen and serum more strongly resemble the conditions of the tumor with respect to ACSS2 localization. Additionally, it points to an important role for ACSS2 in the nuclei of hypoxic tumor cells.

**ACSS2 Recaptures Endogenously Produced Acetate**

We considered whether the primary function of nuclear ACSS2 might be to recapture acetate released by histone deacetylation. In support of an acetate-recapturing function of ACSS2, we noticed that cells that avidly consumed U-\textsuperscript{13}C-acetate released relatively little \(^{13}\text{C}\)-acetate (Figure S5A), arguing that ACSS2 captures exogenous acetate but also prevents the release of endogenously produced acetate by recapturing it. To quantify endogenous acetate release independently of cell number, we plotted the ratio of \(^{13}\text{C}\)-acetate released versus U-\textsuperscript{13}C-acetate consumed in relation to ACSS2 expression for the six cell lines in our panel. We found a strong inverse correlation between this ratio and ACSS2 expression (Figure 5A), indicating that high ACSS2 expression not only facilitates uptake of exogenous acetate but also efficient recapturing of endogenously produced acetate.

To further investigate the role of ACSS2 in recapturing endogenously produced acetate, we silenced its expression in MDA-MB-468 cells by siRNA. In agreement with ACSS2 mediating uptake of exogenous acetate, this largely abrogated U-\textsuperscript{13}C-acetate consumption (Figure 5B). Consistent with the recapturing hypothesis, \(^{13}\text{C}\)-acetate release markedly increased upon ACSS2 knockdown (Figure 5C). Under this serum- and oxygen-limited condition, the net result was a stagnant total acetate level (Figure 5D). Further demonstrating that ACSS2 determines \(^{13}\text{C}\)-acetate release/U-\textsuperscript{13}C-acetate uptake, this ratio increased upon ACSS2 knockdown in MDA-MB-468 and BT-474 cells (Figures 5E and 5F). A similar trend was observed in hypoxia and 10% serum (Figures S5B and S5C). To determine whether ACSS1 recaptures endogenous acetate, we silenced its expression in BT-474 cells. This also led to increased \(^{12}\text{C}\)-acetate release-to-U-\textsuperscript{13}C-acetate uptake ratio (Figure S5D). This demonstrates that, in addition to taking up exogenous acetate, ACSS2 and ACSS1 function to recapture endogenously produced acetate. Net acetate uptake results from a combined intake of exogenous acetate and retention of intracellular acetate.

**ACSS2 Maintains Histone Acetylation during Oxygen and Serum Limitation**

The acetate re-capturing function of ACSS2 and its prominent nuclear localization during oxygen and serum limitation prompted us to test whether ACSS2 maintains histone acetylation. This could be important because histones are among the most abundant proteins in mammalian cells and are heavily acetylated. With a residence half-life of only minutes, turnover of most histone acetylation marks is quite rapid (Evertts et al., 2013). Under oxygen- and serum-replete conditions, silencing
ACSS2 did not result in a decrease but, rather, a modest increase in acetylation of histones H3 (AcH3) and H4 (AcH4) (Figures 6A and 6B), which may be due to HDAC inhibition by nuclear acetate accumulation. In contrast, when silencing ACSS2 in low oxygen and serum, a significant decrease was observed in the acetylation of both H3 and H4. No acetate was added to the medium in this experiment, further reinforcing the notion that ACSS2 maintains histone acetylation by acetate recapturing.

We next asked whether any of the acetate released into the medium originated from histone deacetylation. If so, then inhibition of HDACs should lead to a drop in acetate release. We evaluated this with a short incubation protocol (6–8 hr depending on the cell line). Incubation of MDA-MB-468 cells with the pan-HDAC inhibitor panobinostat led to a significant reduction in acetate release into the medium of the control (scramble [SCR]) MDA-MB-468 and BT-474 cells (Figures 6C and 6D; Figure S6A). Comparable results were obtained with the HDAC inhibitor sodium butyrate (Figure S6B). As expected, silencing of ACSS2 resulted in increased release of acetate. Importantly, treatment with panobinostat in ACSS2-silenced cells caused a greater...

Figure 4. ACSS2 Localizes to the Nucleus during Oxygen and Serum Limitation and Is Prominently Nuclear in Tumors
(A) Representative images of DAPI (nuclear) and ACSS2 staining (separately and merged) in MDA-MB-468 cells in normoxia and 10% serum or hypoxia and 1% serum. Scale bar, 20 μm.
(B) Quantification of the nuclear fraction of ACSS2 (percent) in MDA-MB-468 cells cultured under different conditions. Data are mean ± SD of three independent experiments (seven or more images per experiment).
(C) Immunohistochemical staining of serial sections from a representative tumor of the MMTV-PyMT mouse model for carbonic anhydrase 9 (CAIX, a hypoxic marker), CD31 (a marker for blood vessels), ACSS2, and H&E staining.
(D) Scoring of ACSS2 nuclear intensity in normoxic and hypoxic tumor regions. Ten different ROIs were selected for normoxic and hypoxic regions of two tumors (five ROIs each), and the percentages of cells with weak, moderate or strong nuclear ACSS2 staining were scored. Data are mean ± SD (n = 10 ROIs). See also Figure S4.
absolute reduction in acetate release compared with SCR cells, giving further credence to the acetate recapturing function of ACSS2. We additionally tested the involvement of other enzymes in acetate release. Incubation with the sirtuin inhibitors sirtinol (Figure S6C) and nicotinamide (Figure S6D) led to a maximum ~30% reduction in acetate release. In contrast, knockdown of amino acid deacetylases aminoacylase 1 (ACY1) and ACY3, which deacetylate free acetylated amino acids, and aspartoacylase (ASPA), which deacetylates N-acetyl aspartate, did not noticeably affect acetate release (Figure S6E). Thus, acetate release occurs as a consequence of histone deacetylation reactions, and acetate recapturing by ACSS2 maintains histone acetylation during limited oxygen and serum availability.

**DISCUSSION**

ACSS2 was recently found to be important for tumor growth (Schug et al., 2015; Comerford et al., 2014; Mashimo et al., 2014). However, a quantitative understanding of how it controls acetate exchange and supports downstream metabolic processes is still limited. Here we used innovative stable isotope tracing approaches to interrogate acetate metabolism in cancer cells. One of the findings is that cellular acetate freely exchanges with the medium and that net acetate uptake is controlled by ACSS2 expression. ACSS2 appears to act in a manner similar to hexokinase. Just like hexokinase phosphorylates glucose to effectively trap it inside the cell to commit it to downstream metabolism, acetate is captured by ACSS2 and “primed” for metabolic use. In addition to taking up exogenous acetate, our results demonstrate that ACSS2 recaptures acetate released by deacetylation processes to retain it in the cell. It is the combination of these two activities that determines the net acetate uptake rate.

An important aspect of our work is the determination of acetate utilization fluxes. For example, we were able to estimate mitochondrial utilization from the reduction in net acetate uptake upon ACSS1 silencing. This revealed that approximately one-third of the consumed acetate is dedicated for mitochondrial use in BT-474 cells, presumably mostly for oxidation but, perhaps, also for mitochondrial protein acetylation (Baeza et al., 2016). A major consumer of acetate carbon in the cytosol is fatty acid biosynthesis. We find that, in high serum, hypoxia lowers fatty acid biosynthesis, in line with earlier observations (Kamphorst et al., 2013; Young et al., 2013). In hypoxia and low serum, on the other hand, despite reduced growth of both MDA-MB-468 and BT-474 cells, fatty acid biosynthesis is significantly elevated (Figures 3E and 3F). This may explain the increased efficiency of lipogenic inhibitors under these conditions (Schug et al., 2015; Peck et al., 2016). In both cell lines, the demand for lipogenic AcCoA alone exceeds acetate uptake by at least 2-fold. The cycling between acetate and acetyl-CoA
and exchange of intracellular acetate with the medium prohibits accurate quantification of the flux from acetate into fatty acids (Fan et al., 2013), but given the avid and preferential labeling of lipogenic AcCoA from acetate (Figure 1H; Figure S1C), it likely consumes most of the acetate taken up.

To investigate the use of exogenous acetate in the nucleus, we analyzed labeling of histone-bound acetate. This labeling is representative of the aggregate of the histone acetate pool because we measure total (tail and core) histone-bound acetate by subjecting isolated histones to hydrolysis to release bound acetate (Tumanov et al., 2016). This approach enables comparison of the nuclear acetate pool versus other pools in the cells in terms of gross fluxes and pool sizes. As expected, histone-bound acetate labeled substantially from U-13C-glucose and, to a lesser extent, from U-13C-glutamine under normoxic conditions (Evertts et al., 2013; Lee et al., 2014). The relative contribution from U-13C-glutamine increased in hypoxia, showing that reductive carboxylation in addition to fatty acids also affects histone acetylation. In line with our observation that the combined mitochondrial and lipogenic two-carbon demand exceeds uptake of exogenous acetate, we did not observe substantial labeling from U-13C-acetate under either normoxic or hypoxic conditions and certainly less than what is observed for lipogenic AcCoA. This agrees with a recent study demonstrating the inefficient use of acetate-derived AcCoA for histone acetylation (Zhao et al., 2016). Supplementing free fatty acids to reduce lipogenesis increased histone-bound acetate labeling from exogenous U-13C-acetate (Figures 3I and 3J). An explanation for our observations is that the majority of exogenous acetate is activated to AcCoA, either by the mitochondrial ACSS1 or the cytosolic ACSS2, before it can reach the nucleus. Because of the high demand of AcCoA-consuming pathways, especially lipogenesis, much of the produced AcCoA is used before it can find its way into the nucleus.

The fact that we can measure overall histone-bound acetate labeling and that exogenous acetate is preferentially used for lipogenesis relative to nuclear histone acetylation provides a fundamental insight into metabolic compartmentalization. Histone-bound acetate can be regarded as a proxy for nuclear AcCoA, and lipogenic AcCoA is representative of the cytosolic AcCoA pool. It is thought that metabolites, including AcCoA, in the cytosol and nucleus freely equilibrate. If this occurs on a large scale, this would result in identical labeling patterns of the cytosolic and nuclear AcCoA pools. However, we observe striking differences in their steady-state labeling patterns, indicating that exchange is fairly limited and that the pools are mostly maintained separately. This gives further credence to the notion that local consumption of AcCoA is too rapid to allow extensive exchange between AcCoA in the separate compartments to occur.

An important finding of this work is that the nuclear localization of ACSS2, which increases in low oxygen and serum, prevents loss of histone deacetylation. We find that ACSS2 staining is prominent in tumors of the MMTV-PyMT model of breast cancer and that the intensity of nuclear staining increases in the hypoxic regions of the tumor (Figures 4C and 4D). Given that cells in these regions also likely experience low nutrient availability, it is plausible that retention of acetyl units by ACSS2 to maintain histone acetylation is a relevant mechanism in vivo. Although the depth

Figure 6. ACSS2 Recaptures Acetate Released from Histone Deacetylation

(A) Western blot of AcH3 and AcH4 in MDA-MB-468 cells upon transfection with SCR or ACSS2 siRNAs. Cells were cultured under the indicated conditions for 48 hr after transfection under either condition.

(B) Bar plot of the same data, with AcH3 normalized to histone H3 and AcH4 to histone H4, with normalized values expressed relative to the scrambled RNA.

(C) Acetate release by MDA-MB-468 cells transfected with SCR or ACSS2 siRNAs and with or without 50 μM panobinostat for 6 hr in low oxygen and serum.

(D) The same for BT-474 cells but with 8-hr incubation.

(E) Schematic of acetate metabolism in oxygen- and serum-limited cancer cells.

For (B)–(D), data are mean ± SD (n = 3); *p < 0.05, **p < 0.01. See also Figure S6.
of metabolic labeling will be less in vivo tracing experiments, and comparison of perfused and hypoxic regions is far from trivial, it may be valuable to further explore certain aspects of acetate metabolism. For example, prolonged infusion with $^{13}$C-acetate may help to assess the propensity to incorporate the label as a function of ACSS2 expression. Additionally, to investigate acetate recapturing by ACSS2 in tumors, mice xenografted with either high or low ACSS2-expressing isogenic tumors could be given a bolus of $^{13}$C-acetate. This would lead to higher fractional acetate labeling in high ACSS2 tumors than in low ACSS2 tumors because, in the latter, tumor cells release more unlabelled acetate. Co-administration of an HDAC inhibitor would reduce the unlabeled acetate in the low ACSS2 xenografts and increase fractional labeling.

In conclusion, our work shows that ACSS2 serves a dual function during oxygen and serum limitation. It facilitates consumption of extracellular acetate as an alternative carbon source, but the increased nuclear localization also enables cells to retain much of their endogenously produced acetate. ACSS2, in essence, “recycles” two-carbon units when nutrient availability is low, which is especially relevant in the poorly perfused, hypoxic areas of the tumor. It allows cells to maintain sufficient acetylation of histones, to prevent initiation of apoptosis, and to maintain growth (Liysiotis and Cantley, 2014; Lee et al., 2014). This knowledge may inform the ongoing evaluation of ACSS2 as a therapeutic target.

**EXPERIMENTAL PROCEDURES**

**Cell Culture**
Cell lines were from the ATCC and routinely passaged in DMEM (HyClone) with 25 mM glucose and 2 mM L-glutamine supplemented with 10% (v/v) fetal bovine serum (FBS) (Gibco) and split at 80% confluence. Cells were routinely checked for mycoplasma. Experiments were performed in DMEM with 10 mM glucose and 2 mM glutamine supplemented with 1% or 10% dialyzed FBS (DFBS) (Sigma). Hypoxia was achieved with a hypoxic chamber (Hypoxia station H35, Whitley Scientific) at 1% O2, 3% CO2, and 96% N2 with 80% L water and 300 L hexane, and the hexane fraction was derived from fatty acid labeling by determining the best fit with computed binomial distributions based on a range of AcCoA labeling enrichments (Kamphorst et al., 2014; Tumanov et al., 2015). Metabolite extraction and quantification was done using a macro.

**Western Blotting**
Protein lysates were prepared in radioimmunoprecipitation assay (RIPA) buffer (Pierce) with protease inhibitor cocktail (Sigma), and total protein concentration was determined by protein assay (Bio-Rad). Proteins were separated using precast NuPAGE gels (Invitrogen, Life Technologies) and transferred onto a nitrocellulose membrane. Protein detection and quantification were done with a Li-COR Odyssey infrared flatbed scanner (Li-COR Biosciences). The primary antibodies used were as follows: anti-ACSS1, 1:1,000 (Sigma, SAB1400745); anti-$\beta$-tubulin, 1:5,000 (Sigma, T5201); anti-ACSS2, 1:2,500 (Cell Signaling Technology, 3658S); anti-acetyl-histone H3, 1:1,000 (Millipore, 06-599); anti-acetyl-histone H4, 1:1,000 (Millipore, 06-598); anti-histone H3, 1:5,000 (Active Motif, 39763); and anti-histone H4, 1:5,000 (Active Motif, 61521). Secondary antibodies were IRDye680LT- and IRDye800CW-conjugated (Li-COR Biosciences). Acidic proteins were hydrolyzed overnight (heating at 95 °C overnight in 10 M NaOH) and then neutralized with hydrochloric acid. The resulting free acetate was processed as described.

**Immunofluorescence Analysis and Imaging**
Cells were maintained for 48 hr on glass coverslips in the indicated medium. For fixation, cells were washed three times with PBS and then incubated for 10 min in 4% paraformaldehyde. Cells were washed three times with PBS and permeabilized for 5 min in PBS containing 0.2% Triton X-100. Following additional washing (three times) in PBS, samples were blocked in PBS with 1% Triton X-100 and 5% FBS for 30 min at room temperature (RT). Samples were first incubated with anti-ACSS2 (1:200) antibody (Cell Signaling Technology) and then with anti-rabbit Alexa Fluor 488 secondary antibodies and DAPI, washed, and mounted on a glass coverslip using Dako fluorescence mounting medium. Images were taken on an Olympus FV1000 microscope, and processing was done with ImageJ software. Nuclear-to-cytoplasm staining ratio quantification was done using a macro.
MMTV-PyMT Mice

MMTV-PyMT mice (Guy et al., 1992) were obtained from The Jackson Laboratory. Mammary tumors were measured by caliper twice weekly, and mice were culled humanely when tumors reached the clinical endpoint, according to regulations. Animal experiments were done in accordance with United Kingdom regulations under project license PPL 70/8645, ethical review (University of Glasgow), and EU Directive 2010.

Immunohistochemistry and Image Analysis

Serial sections (4 μm) of paraffin-embedded tumor slices were stained for H&E and immunostained using a Dako autostainer for AceCS1 (ACSS2) (Cell Signaling Technology, 3658S), CD31 (Abcam, ab236364), and CAIX (Novus Biologicals, NB100-417). Slides were analyzed using HALO v2.0 software (Indica Labs). Sections stained for CAIX and ACSS2 were analyzed using Cytonuclear v1.5. Briefly, first hematoxylin staining was set as the nuclear stain and ACSS2 staining at stain 1, and we manually checked for stain separation. ACSS2 staining was specified to localize to the nucleus to get the nuclear staining intensities, and individual cells were assigned to have weak, moderate, or strong nuclear intensities in a given area based on pre-defined threshold intensities. This was done for regions of interest (ROIs) that were manually selected in the CAIX image to represent hypoxic or normoxic areas.

siRNA and Transfections

All siRNAs were from Dharmacon (GE Life Sciences), reconstituted in sterile DNase/RNase-free water to a stock concentration of 20 μM, and transfected using Lipofectamine RNAiMAX reagent (Invitrogen) at a final concentration of 20 nM/well. 48 hr after transfection, cells were seeded for experiments. The following siGENOME siRNAs were used: scrambled RNA control (non-targeting siRNA pool #2, D-001206-14-05), ACSS1 siRNA #1 (D001206-02), ACSS1 siRNA #2 (D001206-03), ACSS1 siRNA #1 (A008549-13), and ACSS1 siRNA #2 (A008549-14).

Statistical Analyses

Two-tailed Student’s t tests and Pearson correlation analysis were done using Graph Pad Prism, version 6. Error bars represent SD of the mean. n is the number of independent wells for each condition and is mentioned in the figure legends for each experiment.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures and six figures and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2016.12.055.

AUTHOR CONTRIBUTIONS

Conceptualization, V.B. and J.J.K.; Methodology, V.B., S.T., Z.T.S., J.V.V., A.V., and E.G.; Investigation, V.B., S.T., E.M., C.N., S.D., K.B., A.V., and E.G.; Funding Acquisition, N.J.V.D.B., G.M., C.N., S.D., K.B., A.V., and E.G.; Investigation, V.B., S.T., Z.T.S., J.V.V., A.V., and E.G.; Supervision, E.G. and J.J.K.

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