Metabolic Plasticity of Metastatic Breast Cancer Cells: Adaptation to Changes in the Microenvironment

Abstract

Cancer cells adapt their metabolism during tumorigenesis. We studied two isogenic breast cancer cell lines (highly metastatic 4T1; nonmetastatic 67NR) to identify differences in their glucose and glutamine metabolism in response to metabolic and environmental stress. Dynamic magnetic resonance spectroscopy of 13C-isotopomers showed that 4T1 cells have higher glycolytic and tricarboxylic acid (TCA) cycle flux than 67NR cells and readily switch between glycolysis and oxidative phosphorylation (OXPHOS) in response to different extracellular environments. OXPHOS activity increased with metastatic potential in isogenic cell lines derived from the same primary breast cancer: 4T1 > 4T07 and 168FARN (local micrometastasis only) > 67NR. We observed a restricted TCA cycle flux at the succinate dehydrogenase step in 67NR cells (but not in 4T1 cells), leading to succinate accumulation and hindering OXPHOS. In the four isogenic cell lines, environmental stresses modulated succinate dehydrogenase subunit A expression according to metastatic potential. Moreover, glucose-derived lactate production was more glutamine dependent in cell lines with higher metastatic potential. These studies show clear differences in TCA cycle metabolism between 4T1 and 67NR breast cancer cells. They indicate that metastases-forming 4T1 cells are more adept at adjusting their metabolism in response to environmental stress than isogenic, nonmetastatic 67NR cells.
suggest that the metabolic plasticity and adaptability are more important to the metastatic breast cancer phenotype than rapid cell proliferation alone, which could 1) provide a new biomarker for early detection of this phenotype, possibly at the time of diagnosis, and 2) lead to new treatment strategies of metastatic breast cancer by targeting mitochondrial metabolism.

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Introduction

Breast cancer is the most prevalent type of cancer among women in the United States [1], and mortality is primarily caused by metastatic disease. The complex mechanisms of breast cancer invasion and metastasis [2] are intrinsically related to the malignant cell type [3], their interaction with stromal cells [4,5], and changes in the tumor microenvironment, related to poor perfusion, intermittent hypoxia, transient nutrient deprivation, and acidity [6,7]. Cancer cells adapt to dynamic stresses and proliferate by reprogramming their metabolism to support synthesis of an expanding biomass [8,9]. Due to oncogene-driven upregulation of key glycolytic enzymes [10], most cancer cells exhibit aerobic glycolysis known as the Warburg effect [11]. This metabolic phenotype has been studied by noninvasive techniques, such as 18F-fluorodeoxyglucose positron emission tomography and magnetic resonance spectroscopy (MRS of 13C-labeled substrates) [12,13]. Due to enhanced glycolysis, tumor cells synthesize high levels of lactate and export H+, resulting in acidification of the microenvironment, which in turn promotes invasion and dissemination [14,15]. Recent studies with two isogenic murine breast cancer cell lines derived from the same spontaneous breast tumor, 4T1 and 67NR [16], have shown differences in lactate dehydrogenase (LDH) A expression during normoxia and hypoxia [17]. However, other studies have highlighted the importance of oxidative phosphorylation (OXPHOS) in tumorigenesis and progression [18–20].

Because it is well recognized that tumor cells are often hypoxic and nutritionally deprived in vivo [7], we have monitored in real time the metabolic changes in live 4T1 and 67NR cells under conditions that reflect these common, often transient, physiologic stresses. Our hypothesis is that the metabolism of metastases-forming breast cancer cells (4T1) is more adaptable to changes in the microenvironment than the metabolism of isogenic, nonmetastatic cells (67NR), thus providing 4T1 cells with a distinct advantage to grow, invade, and proliferate under different conditions. Previous metabolic studies using cell extracts have demonstrated marked differences in basal glucose consumption, lactate production, and oxygen consumption between these two cell lines grown in standard bidimensional tissue culture conditions [17,21]. However, these studies have not investigated the — potentially reversible — metabolic adaptations of these cancer cells to in vivo tumor conditions, which include changing microenvironmental stresses during tridimensional growth. We used a magnetic resonance (MR)–compatible cell perfusion system and time-course MRS of 13C isotopomers to investigate how living cancer cells adapt their metabolism and growth to selective supply/deprivation of glucose and glutamine under both aerobic and hypoxic conditions. In contrast to standard 2D tissue culture studies, the MR-compatible cell perfusion system allows high-density 3D cancer cell growth and exposing cells dynamically and reversibly to various tissue growth environments in a single sample, more similar to the cellular microenvironment of small (<100 mm3), well-perfused tumors. Moreover, compared with indirect metabolic measurements based on dynamic extracellular pH and O2 changes (e.g. Seahorse XF analyzer studies), the cell perfusion system allows measuring real-time changes in intra- and extracellular metabolite levels and cellular energetic profiles by sequential multinuclear (13C, 31P) MRS. Our assessment of the dynamic interplay between various environmental stresses and tumor cell metabolic response clearly demonstrates that 4T1 cells are more capable of adapting their metabolic responses to changes in the microenvironment than 67NR cells. This is largely accomplished in 4T1 cells by their greater plasticity and ability to more effectively metabolize glucose through either glycolysis or OXPHOS than 67NR cells, providing greater adaptability to a changing tumor and metastatic microenvironment.

Materials and Methods

Cell Lines

The 67NR, 168FARN, 4T07, and 4T1 cell lines were initially derived from a spontaneous breast tumor growing in a BALB/c mouse [16]. These cell lines were kindly provided by Dr. Fred Miller (Karmanos Cancer Institute, Detroit, MI) and grown in Dulbecco’s modified Eagle’s media

| Study | Label (°C) | Stage | Experiment Time (h) | Medium (mM) | Oxygenation |
|-------|------------|-------|---------------------|-------------|-------------|
|       |            |       |                     | Glucose     | Glutamine   |
| A     | –          | A-1   | 0-1                 | 25          | 6           | Ox         |
|       |            | A-2   | 1-6                 | 25          | 2           | Ox         |
|       |            | A-3   | 6-32                | 25          | 6           | Ox         |
| B     | –          | B-1   | 0-1                 | 25          | 6           | Ox         |
|       |            | B-2   | 1-6                 | 25          | 6           | Ox         |
|       |            | B-3   | 6-11                | 25          | 6           | H          |
|       |            | B-4   | 11-16               | 0           | 6           | Ox         |
|       |            | B-5   | 16-21               | 0           | 6           | H          |
|       | –          | B-6   | 21-22               | 25          | 6           | Ox         |
| C     | –          | C-1   | 0-1                 | 25          | 6           | Ox         |
|       |            | C-2   | 1-6                 | 25          | 0           | Ox         |
|       |            | C-3   | 6-11                | 25          | 0           | H          |
|       |            | C-4   | 11-16               | 25          | 2           | H          |
|       |            | C-5   | 16-21               | 25          | 2           | Ox         |
|       | –          | C-6   | 21-22               | 25          | 6           | Ox         |

Three different studies were carried out: A, B, and C. The first experimental hour of each experiment was reserved for loading the sample into the MR spectrometer, MR probe tuning, and matching, followed by sample shimming. Study A consisted of well-oxygenated cells perfused continuously for 31 hours with 99% 13C-glucose-containing culture medium, initially with 2 mM glutamine (1-6 hours) and then with 6 mM glutamine (6-32 hours). In separate experiments, cells were studied during four sequential 5-hour stress conditions in the presence of either 99% 31P-glutamine (study B) or 99% 13C-glucose (study C) in the culture/perfusion medium. In study B, the experiment with 3-13C-Gln–labeled medium consisted of the four conditions/stages, B2 to B5, evaluating the effects of the presence or absence of 25 mM glucose under well-oxygenated or hypoxic conditions on glutamine metabolism. In study C, the experiments with 99% 13C-glucose–labeled culture medium were carried out in four stages/environmental conditions: C-2 to C-5. The perfusion media for stages C-4 and C-5 contained in addition to 2 mM Glu also a basal level of pyruvate (1 mM). Abbreviations: Glc C1: 13C-glucose, Glu C3: 31C-glutamine, Ox: well-oxygenated conditions, H: hypoxia.
containing 25 mM glucose (Glc), 6 mM glutamine (Gln), 100 U/ml of penicillin, 100 μg/ml of streptomycin, and 10% fetal calf serum, referred to as DMEcompl. Cells were cultured in 5% CO2 / 95% air at 37 °C in a humidified chamber, split every 2 to 3 days, and used up to passage 10.

Cell Perfusion Studies

For MR cell perfusion studies, 4T1 and 67NR cells were grown to >70% confluence on microcarrier beads. The cell perfusion studies were carried out on a 500-MHz AVANCE III Bruker MR system equipped with a 10-mm BBO probe for 1H and 31P/13C acquisition (Bruker BioSpin, Billerica, MA). Three different experimental studies were performed for 4T1 and 67NR cell lines, as detailed in Table 1, each consisting of three independent cell perfusion experiments per cell line. During the experiments, the cells were continuously perfused with media containing either 98% to 99% 1-13C-glucose (CLM-420-0; Cambridge Isotope Labs, Billerica, MA) or 99% 3-13C-glutamine (604941-SPEC; Sigma-Aldrich, St. Louis, MO), whereas 13C MR spectra and 31P MR spectra were acquired repeatedly to assess metabolism of the 13C-labeled nutrients and cellular energy status, respectively. Tumor cell–conditioned medium (TCM) samples were also collected at the end of the cell perfusion studies A (Table 1) and stored at −20°C until high-resolution 1H-decoupled 13C MRS studies were performed. Additional details about the cell perfusion system, MR data acquisition, processing, and isotopomer analysis are provided in the Supplementary Experimental Procedures.

Cell Culture Studies

The four isogenic cell lines (67NR, 168FARN, 4T07, and 4T1) were seeded separately in the same 96-well plate at 1.0 × 10⁴ and 2.0 × 10⁴ cells/well, respectively. After attachment and growth for 18 hours, the DMEcompl medium in each well was changed to XF assay medium (102365-100; Seahorse Bioscience, Billerica, MA) with the same glutamine and glucose concentration as in DMEcompl and incubated for 1 hour in a CO2-free atmosphere. The cells were then studied in an XF96 Analyzer (Seahorse Bioscience, Billerica, MA), measuring the extracellular acidification rate (ECAR; studied during the first 27 minutes) and the oxygen consumption rate (OCR; 2-hour studies) in each well. During the OCR studies, the specific respiratory chain inhibitors oligomycin (inhibition of ATP synthase), Carbonyl cyanide 4-(trifluoromethoxy) phenylhydrazone (FCCP; H⁺ ionophore, uncoupler), and antimycin A plus rotenone (inhibition of complexes I and III, respectively, disabling the electron transport completely in the respiratory chain) (Sigma-Aldrich, St. Louis, MO) were sequentially injected into each well at 30, 60, and 90 minutes, respectively [22]. The time course of OCR values collected during data acquisition were normalized to the total protein content in each well. The latter were determined at the end of the experiment from the cell lysates (RIPA buffer; Thermo Scientific, Rockford, IL) using the bicinchoninic acid protein assay (BCA kit; Thermo Scientific Pierce, Rockford, IL). Measurements were summarized by taking the geometric mean across wells for each cell line, at each time point, and in each replicate of the experiment. The geometric mean was used because of the skewed distribution of data across wells within each experimental condition. In the specific case of ECAR, the logarithm of ratios between paired 4T1 and 67NR results was calculated based on data skewness and because a ratio of 1 translates into a log-ratio of 0 when applying a one-sample t test across experimental replicates. The same analysis was applied to the log-ratio of the geometric means across the four time points and to the linear contrast, estimating the slope of log-ratios versus time points. Finally, an additional experiment was carried out to compare OCR measurements between all four isogenic cell lines according to the protocol previously described. Results were analyzed as above using the geometric means of the dynamic readings in each microplate well (4T1, n = 21; 4T07, n = 22; 168FARN, n = 23; 67NR, n = 22).

Expression and Activity of Succinate Dehydrogenase (SDH)

In both 4T1 and 67NR cells, SDH protein expression was assessed by Western blot of the SDH subunit A (SDH-A) using a monoclonal antibody and a methodology analogous to our previous studies [17]. We also used a polyclonal antibody to assess SDH-A expression by Western blot in the four isogenic cell lines: Cells were exposed for 5 hours to media with different concentrations of glucose (25 and 0 mM) and glutamine (6, 2, and 0 mM) under normoxia and hypoxia, paralleling the media conditions in MR cell perfusion studies as described in Table 1. The Western blot protocols are provided in the Supplementary Experimental Procedures. All results are averaged over three independent experiments. The total SDH activity of 4T1 cells and 67NR was assessed using the Complex II Enzyme Activity Microplate Assay Kit (ab109908; Abcam, Cambridge, MA). The activity of SDH was measured in parallel for fresh lysates of 4T1 and 67NR cells, as described by the manufacturer.

Statistical Analyses

Statistical analyses were carried out with the t test (unless indicated otherwise): unpaired when comparing 4T1 and 67NR data acquired under the same experimental conditions ( indicates statistical significance: P < .05) and paired when comparing the same cell line in consecutive stages during the same study, as specified for each experiment ( indicates statistical significance: P < .05). Results are displayed as mean ± standard error (SEM). In graphs, error bars that are smaller than the symbol size are not visible.

Results

Glycolytic Activity of 4T1 and 67NR Cells

We used an MR-compatible cell perfusion system to investigate the differences in glucose metabolism between 4T1 and 67NR cells in the presence of 6 mM and 2 mM glutamine (DMEcompl versus more physiologic glutamine concentrations, respectively) (Study A, Table 1). The bioenergetic profile and phospholipid metabolism were measured dynamically during all cell perfusion studies by 31P
Figure 1. Basal glucose metabolism of live 4T1 and 67NR cells. MR cell perfusion studies \( (n = 3 \text{ for each cell line}) \). (A) Representative nuclear Overhauser effect–enhanced, 1D \( ^1H \)-decoupled \( ^{13}C \) MR spectra acquired from 4T1 (top) and 67NR (bottom) cells after 24-hour perfusion with culture medium containing \( ^{1-13}C \)-glucose (Glc), as detailed in Study A, Table 1. The regions from 80 to 50 ppm and 40 to 25 ppm are shown enlarged above the full spectra. All spectra are displayed with 10-Hz exponential line broadening. Signal assignments of metabolites, with C1-6 depicting the position of the incorporated \( ^{13}C \) label: alanine (Ala), dihydroxyacetone phosphate (DHAP), \( \alpha \)- and \( \beta \)-glucose (\( \alpha \)-Glc and \( \beta \)-Glc, respectively), glutamate (Glu), glutamine (Gln), glutathione (GSH-Glu), glycerol (Gly), glycerol-3-phosphate (G3P), lactate (Lac), methionine (Met), myo-inositol (myo-Ino), pyruvate (Pyr), succinate (Succ). (B) Average time-course changes for the major peak areas, normalized to the initial \( \beta \)-NTP values (\( \beta \)-NTP\(_{\text{init}} \) determined from \( ^{31}P \) MRS; Figure S1). (C) Average \( ^{13}C \)-labeling rates of metabolites at different time intervals for the data shown in B.
MRS and were characteristic for each cell line (Figure S1). The metabolism of 1-13C-glucose (Figure 1) was measured by 13C MRS. Multiple 13C-labeled metabolites synthesized de novo by the tumor cells from 1-13C-glucose were detected (Figure 1A). Their peak areas were normalized to the initial number of cells (Figure 1, B and C) by referencing to the corresponding β-NTP_int levels (Figure S1). Because glucose was labeled only in the C1 position, 50% of the metabolic pool downstream of fructose 1,6-diphosphate incorporated the 13C-label and was detected.

As shown in Figure 1, 4T1 cells demonstrated significantly higher glycolytic activity than 67NR cells, independent of glutamine concentrations in the perfusion medium (2 mM, 1-6 hours, P < .05; and 6 mM, 6-32 hours, P < .05). During the initial 10 hours of perfusion with 13C-labeled medium, the glucose consumption rate was slightly higher in 4T1 cells than in 67NR cells and increased significantly in 4T1 cells when changing the medium from 2 mM to 6 mM glutamine (after 6 hours) (Figure 1C). Under both experimental conditions, the glycolytic-lactate synthesis rates were also approximately 1.5-fold higher in 4T1 than in 67NR cells (1-6 hours, P < .05; 6-11 hours, P = .06), closely matching the difference in doubling times between the two cell lines. In standard tissue culture medium (25 mM glucose and 6 mM glutamine), 4T1 cells have a doubling time of 13.6 ± 1.5 hours (Figure 2D). Their peak areas were slightly higher in 4T1 cells than in 67NR cells and increased significantly in 4T1 cells when changing the medium from 2 mM to 6 mM glutamine (after 6 hours) (Figure 1C).

The 13C-labeled pools of glycerol-3-phosphate [3-13C-G3P, obtained by reduction of dihydroxyacetone phosphate (1-13C-DHAP) and essential for membrane phospholipid synthesis] and myo-inositol [4(6)-13C-myo-Ino, precursor of phosphatidyl inositol phosphates] were also observed in 4T1 and 67NR cells (Figure 1, B and C). Compared with 67NR cells, the higher levels of 3-13C-G3P and 4(6)-13C-myo-Ino in 4T1 cells also indicate their higher glycolytic turnover and membrane biosynthesis requirements, in both 2 mM and 6 mM glutamine medium, and relate to their invasive/malignant phenotypes, in agreement with Smirnova et al. [26].

The synthesis rate of glucose-derived alanine (3-13C-Ala, an alternative end-product of glycolysis obtained by transamination of 3-13C-pyruvate with glutamate) was higher in 4T1 than in 67NR cells [Figure 1, B and C]. Consistent with ongoing glycolytic activity and synthesis of 3-13C-glutamate (Figure 1, A and B) via pyruvate dehydrogenase, indicative of mitochondrial TCA cycle activity in both cell lines. However, the synthesis rate of 4-13C-glutamate was two-fold higher in 4T1 cells than in 67NR cells at 6 to 11 hours of perfusion with 1-13C-glucose in the perfusate (Figure 1D). The TCA cycle activity in 4T1 and 67NR cells was also measured for cells in 2D tissue culture by the ECAR (Figure 2A). The ECAR measurements were significantly higher for 4T1 than for 67NR at each successive time point (P < .021, .023, .041, and .05, respectively), and also for the geometric mean across time points (P = .034). The mean of geometric ratios, between paired 4T1 and 67NR results, indicated a 1.4-fold higher ECAR in 4T1 (Figure 2A), consistent with the ratios of the glucose-derived lactate synthesis rates for 4T1 and 67NR obtained with the cell perfusion system within the first 18 hours (1.7, 1.5, and 1.4, respectively (Figure 1C).

**OXPHOS and Metastatic Potential**

To investigate the role of OXPHOS in each cell line, we carried out mitochondrial functional analyses under 2D growth conditions (25 mM Glc, 6 mM Gln) by selective inhibition of different respiratory complexes while measuring the effects on the cellular OCR (Figure 2B). Consistent with the 31P MRS results on live, perfused cells (Figure S1B), 4T1 cells showed a significantly higher basal respiration rate (2.3-fold) and ATP production (2.2-fold) than 67NR cells (Figure 2C). When extending these measurements to two additional cell lines from the same isogenic breast cancer family (168FARN and 4T07), we noticed higher basal respiration and ATP production with increasing metastatic ability: 4T1 (overt metastasis) > 4T07 and 168FARN (micrometastasis) > 67NR (nonmetastatic) (Figure 2, D and E).

**Tricarboxylic Acid (TCA) cycle activity in 4T1 and 67NR cells**

Both 4T1 and 67NR cells incorporated 1-13C-glucose into 4-13C-glutamate (Figure 1, A and B) via pyruvate dehydrogenase, indicative of mitochondrial TCA cycle activity in both cell lines. However, the synthesis rate of 4-13C-glutamate was two-fold higher in 4T1 cells than in 67NR cells at 6 to 11 hours of perfusion with 1-13C-glucose in the perfusate (Figure 1D). To further investigate the TCA cycle activity and metabolite contributions to anaplerosis [29], we considered a dynamic flux analysis based on the time-course changes of the different 13C-glutamate isotopomers (Figure 3, A and B). This model estimated a 2.2 ± 0.6-fold higher TCA cycle flux [F(tca)] in 4T1 cells than in 67NR cells (Figure 3C), which is consistent with the differences in OCR measurements (Figure 2, B and C). The TCA flux analysis model projected identical relative exchange fluxes between α-ketoglutarate and glutamate in both cell lines [F(αk)/F(tca), Figure 3C]. However, the total exchange estimated at the levels of succinyl-CoA, fumarate, and oxaloacetate (OAA), i.e., anaplerotic/catabolitic fluxes [F(OAA)], was 3.2 ± 0.8-fold higher in 67NR cells than in 4T1 cells, although not significantly. Moreover, the estimated pyruvate carboxylase flux F(pC) (OAA-pyruvate exchange) was significantly higher in 67NR cells than in 4T1 cells (Figure 3C). Accordingly, the time-course changes in the 2-13C/4-13C glutamate ratio (<<1, Figure 3D) suggest that pyruvate is channeled to the TCA cycle mostly via pyruvate dehydrogenase [30] in both cell lines, whereas the changes in 2-13C-glutamate to 3-13C-glutamate ratio (Figure 3D) reflect differences in pyruvate carboxylase activity according to glutamine availability. In the presence of 2 mM glutamine (Figure 3D, 1-6 hours), both cell lines displayed similar 2-13C-glutamate to 3-13C-glutamate ratios (~1.5-2), suggesting similar pyruvate carboxylase activity. However, after increasing glutamine to 6 mM (Figure 3D, 6-11 hours), the 2-
C. TCA Cycle Restriction in 67NR Cells

The nonmetastatic 67NR cells showed a strong accumulation of 1-13C-glucose–derived succinate (2-13C-Succ) under full nutrient conditions, whereas glucose-derived succinate was essentially not detected in 4T1 cells (Figure 1, A–C). High-resolution MR analysis of TCM samples collected at the end of each experimental stage (Supplementary Experimental Procedures) also showed that 67NR

\[ ^{13}\text{C}\text{-glutamate to } 3-^{13}\text{C}\text{-glutamate ratio remained essentially unchanged in 67NR cells but significantly decreased in 4T1 cells (−1)}, \text{ suggesting a reduction in pyruvate carboxylase activity, likely due to more glutamine availability to replenish the TCA cycle at the level of OAA. Altogether, our results indicate a rate-limiting step of the TCA cycle downstream of } \alpha\text{-ketoglutarate in 67NR cells. This is summarized in our proposed model (Figure 4, A and B).} \]
cells extruded 5.8-times more succinate than 4T1 cells (data not shown). The accumulation of glucose-derived succinate observed in 67NR cells was further investigated by perfusing each cell line with 3-13C-glutamine during regular growth conditions (1-6 hours), under reversible hypoxia (6-11 and 16-21 hours), and under glucose starvation (11-21 hours) (Table 1, Study B). The time-course changes in phospholipid metabolism, bioenergetic profiles, pH<sub>i</sub>, and pH<sub>e</sub> essentially showed the same trends in both cell lines under glucose deprivation or hypoxia (Figure S4).

3-13C-glutamine–derived succinate levels (2-13C-Succ) were always significantly higher in 67NR cells than in 4T1 cells in the cell perfusion system for all the microenvironment conditions (Study B, Table 1) tested (Figure 5A). 67NR cells synthesized 2-13C-succinate under aerobic conditions, either from 1-13C-glucose (Figure 1, A–C) or from 3-13C-glutamine (Figure 5, A and B; 1-6 and 11-16 hours). The accumulation of 2-13C-succinate from 3-13C-glutamine in 4T1 cells could only be detected during hypoxia with glucose-containing perfusate (Figure 5, A and B; 6-11 hours). This is consistent with a reductive turnover of the TCA cycle, as recently shown in other cancer cell lines growing in hypoxia [31] or with defective mitochondria [32]. In 4T1 cells (and not in 67NR cells), 13C-labeled aspartate (2-13C and 3-13C, in equilibrium with OAA labeled in the same positions) was detected only when oxygen was available (Figure 5A, 1-6 and 11-16 hours) and not under hypoxia (Figure 5A, 6-11 and 16-21 hours).

Full glucose deprivation under well-oxygenated conditions caused a slight (not statistically significant) increase in the levels of 3-13C-glutamine–derived aspartate in 4T1 cells and of succinate in 67NR cells (Figure 5A, 1-6 vs. 11-16 hours), with no corresponding differences between the rates of synthesis (Figure 5B). These findings indicate that the glucose concentration does not have a significant effect on the oxidation of glutamine through the TCA cycle in both cell lines. This backs up a prior observation in a human lymphoma cell line that supports the existence of an alternative carbon source pathway into the TCA cycle driven by glutamine and independent of glucose [33]. Our results, together with the flux model predictions based on glutamate isotopomer analysis (Figure 3), suggest that the lower TCA cycle activity observed in 67NR cells is due to a restriction in TCA cycle flux at the level of SDH (respiratory complex II) (Figure 4).

**SDH-A Protein Expression Pursuant to Metastatic Ability and Cell Line-dependent Response to Environmental Stresses**

We investigated if the lower TCA cycle activity in 67NR cells and the accumulation of succinate could be explained by a deficiency in SDH expression and/or activity. Western blots showed a significant four-fold lower expression of SDH-A (FAD-containing subunit that binds succinate) in 67NR cells than 4T1 cells (Figure 6A). The enzyme activity of SDH was also significantly lower (six-fold) in 67NR cells than in 4T1 cells (Figure 6B). Because an impairment of SDH should directly affect mitochondrial respiration, these results are consistent with the approximately two-fold higher basal respiration rate and ATP production observed in 4T1 cells compared with 67NR cells (Figure 2C). The expression of SDH-A in regular medium conditions (25 mM Glc, 6 mM Gln) was noted to increase with...
metastatic ability across the isogenic cell line panel for oxygenated and hypoxic cells (Table 2). Hypoxia itself affected SDH-A expression depending on the metastatic potential (Table 2, Figure 6C): downregulation in 67NR (nonmetastatic), upregulation in 4T1 (metastatic) \((P = .003)\), and intermediate effects in the other two cell lines characterized by local micrometastases. Relative to 4T1 cells, glutamine deprivation (0 mM and 2 mM Gln) lowered SDH-A protein expression in the three less metastatic cell lines more than glucose deprivation (Table 2). These results suggest a better ability of 4T1 cells to use the TCA cycle, even under stress, than the other three cell lines.

**Metabolic Adaption to Nutrient (Glucose, Glutamine) Supply in 4T1 than in 67NR Cells**

To further understand the metabolic differences between 4T1 and 67NR cells, we investigated the effects of glucose and glutamine deprivation on glucose metabolism. Using standard 2D tissue culture conditions (Figure S5), we found that the cell growth significantly decreased by approximately five-fold in both cell lines when they were fully deprived of either glutamine or glucose. In response to partial glutamine deprivation (2 mM), an approximately two-fold decrease was observed in both cell lines. Therefore, both 4T1 and 67NR cells depend on glucose and glutamine to grow.

For the cell perfusion experiments performed under cellular stress conditions (Table 1, Study C), we selected 2 mM glutamine over 6 mM to more closely simulate the glutamine levels observed \(\text{in vivo} \) (~0.5 mM Gln, \([34,35]\)). Under aerobic conditions, metastatic 4T1 cells perfused with medium containing 2 mM glutamine (Figure 1, B and C; 1-6 hours) responded to full glutamine deprivation by significantly reducing their glycolytic lactate synthesis (2.3-fold) to levels similar to those observed in 67NR cells under the same conditions (Figure 7, A and B; 1-6 hours). In comparison, the 67NR cells did not significantly alter their lactate synthesis rate with glutamine deprivation under comparable conditions. This differential response to glutamine deprivation is likely related to its importance as a major anaplerotic precursor of the TCA cycle together with higher TCA cycle function in 4T1 cells compared with 67NR cells.

Based on our experimental data and our proposed metabolic model (Figure 4), glutamine deprivation would have a higher impact in replenishing the OAA pool in 4T1 cells than in 67NR cells. As a result, glutamine-deprived 4T1 cells would become more dependent on glucose-derived carbon (pyruvate) entering the TCA cycle, even under stress, than the other three cell lines.

**Figure 4. Metabolic model for 4T1 and 67NR cells.** (A) Extracellular 1,13C-Glc (carbons C1-6 shown as spheres; 13C-labeled carbon in black) in the medium is taken up by the cells via GLUT transporters. Glucose is then metabolized through the glycolytic pathway, labeling 50% of the de novo pools of myo-inositol (C2, C4), DHAP (C1), and glycerol-3-phosphate (C1), which could be detected in the cell perfusion system experiments (Figure 1). Pyruvate (C3 labeling) can be further metabolized in the cytosol to lactate (C3 labeling), regenerating the NAD\(^+\) consumed during glycolysis, or to alanine (C3 labeling) by transamination of glutamate with pyruvate. Alternatively, pyruvate can be channeled to the mitochondria and converted to acetyl-CoA (C2 labeling) via pyruvate dehydrogenase (PDH). Acetyl-CoA enters the TCA cycle and condenses with OAA (initial unlabeled pool) to produce citrate, which then transfers its label to \(\alpha\)-ketoglutarate (C4). The exchange between \(\alpha\)-ketoglutarate (fast turnover) and glutamate (C4 labeling) makes the latter a marker of TCA cycle activity. The C4 label of \(\alpha\)-ketoglutarate is then transferred to succinate C2 (or C3; not distinguishable) and then to OAA C2 (C3). Because succinate C2 (C3) accumulates in 67NR cells, it is likely that their TCA cycle is restricted at the level of SDH, leading to lower flux to OAA (green arrows; higher flux expected in 4T1 cells, displayed by red arrows). After this first turn of the TCA cycle, a significant part of the intermediates has been removed for biosynthetic purposes; the remaining OAA can be used for the second turn — reactions indicated by the dotted gray arrows and the resulting molecules by the gray spheres (13C label, gray; unlabeled, white). In this case, the resulting labeling pattern of glutamate is more complex and generates an additional 1:1 pool of C2 and C3 labeling (highlighted by short gray arrows). (B) Besides PDH, an alternative way for pyruvate to enter the TCA cycle is through pyruvate carboxylase (PC). This anaplerotic pathway becomes more important when glutamine is not sufficiently available to cells or they cannot use it to regenerate OAA. In this case, the first turn of the TCA cycle will generate a 1:1 labeling of glutamate in positions C4 and C2 (indicated by short gray arrows).
cycle and would establish a metabolic phenotype that is closer to the basal metabolic phenotype of 67NR cells. A corresponding reduction of the synthesis rate of lactate would be expected, as we observed (Figure 7, A and B). However, full glutamine deprivation under aerobic conditions reduced TCA cycle flux ~2.3-fold in both cell lines, as evidenced by the significant decrease of glucose-derived glutamate (Figure 7, A and B; 1-6 hours, 0 mM Gln). Our results show that the synthesis rate of glucose-derived lactate is higher in

Figure 5. Cell perfusion stress studies with 3-13C-glutamine. Normalized average time-course changes of metabolite levels (A) and 13C-labeling rates at different time intervals (B) for succinate C2, aspartate C2, and aspartate C3 determined from 13C MR spectra under environmental conditions as detailed in Table 1 (Study B).

Figure 6. SDH expression and activity in breast cancer cells with different metastatic potential. (A) Protein expression of SDH-A was higher in 4T1 than 67NR cells, as shown in a representative Western blot of 4T1 and 67NR cells (left panel) and quantified by averaging the SDH-A/β-actin ratios over three independent experiments (right, bar graph). (B) SDH activity assessed kinetically by absorbance spectroscopy at 600 nm in 4T1 and 67NR cells (dots are average values of at least three independent experiments) and linear fittings for each cell line (slope = SDH activity): 4T1, 1.97 ± 0.28 (R^2 = 0.98); 67NR, 0.32 ± 0.03 (R^2 = 0.99). (C) Ratio of cellular SDH-A/β-actin expression in hypoxia to normoxia for 4T1, 4T07, 168FARN, and 67NR cell lines exposed to culture medium containing 25 mM Glc and 6 mM Gln.
4T1 cells ($V_{\text{max}} = 30.26 \pm 0.12$) than in 67NR cells ($V_{\text{max}} = 18.64 \pm 0.47$) and more dependent on glutamine in the former ($K_{m} = 0.63 \pm 0.01$) than in the latter ($K_{m} = 0.16 \pm 0.03$) cells. Thus, the aerobic synthesis rate of lactate from glucose [$V_{\text{mol}}(4T1/67NR) = 1.6$] requires approximately a four-times higher glutamine concentration in the medium for metastatic 4T1 cells than for nonmetastatic 67NR cells [$K_{m}(4T1/67NR) = 4$] (Figure 7C). This effect is also apparent for the entire panel of isogenic cell lines in 2D cell culture (Figure 8A), as increased glucose-derived lactate synthesis rates were associated with higher metastatic ability and stronger dependency on glutamine levels.

**Interchangeability of Glycolysis and OXPHOS in 4T1 cells**

When 4T1 and 67NR cells are fully deprived of glutamine and subsequently exposed to hypoxia, the glycolytic lactate synthesis rate increased by -1.8-fold in both cell lines (Figure 7, A and B; 6-11 hours), although this increase was significant only in 67NR cells. This observation is expected because 1) hypoxia is a driver of increased glucose uptake and glycolysis [36] and 2) less glucose is being consumed by the TCA cycle under hypoxic conditions, as shown by the low levels of de novo glutamate synthesis (Figure 7, A and B; 6-11 hours). After replenishing glutamine-starved, hypoxic 4T1 cells with glutamine (2 mM), glycolytic synthesis of lactate significantly increased 1.9-fold ($P = .011$, paired t-test), whereas it remained unchanged in 67NR cells exposed to the same nutrient conditions (Figure 7, A and B; 11-16 hours). Although the glutamine-containing medium used in these studies contained pyruvate (1 mM), similar changes in the synthesis rates of lactate and glutamate were obtained for cells in pyruvate-free medium under the same environment stress conditions studied above (results not shown). Furthermore, MR analysis of TCM(2) samples from the four isogenic cell lines grown in hypoxia also indicates a higher lactate synthesis rate in more metastatic cell lines with higher glutamine availability (Figure 8B).

In the last stage of the cell perfusion experiments, shown in Figure 7, A and B, reoxygenation (16-21 hours) of the nutrient-replenished 4T1 cells significantly decreases (2.5-fold) the synthesis rate of glucose-derived lactate, whereas no change was observed in 67NR cells. During this period, the synthesis rate of glucose-derived glutamate was 2.4-fold higher in 4T1 cells than in 67 NR cells, consistent with a stronger reactivation of TCA cycle flux in 4T1 cells following the reintroduction of oxygen and glutamine. Thus, less glucose transitioned to lactate in 4T1 cells, and de novo succinate from glucose accumulated in 67NR cells (but not in 4T1 cells) (Figure 7, A and B). These observations agree with the results of the initial cell perfusion experiments performed under standard growth conditions (Figure 1).

No significant differences in Pi, P<sub>i</sub>, or pH<sub>i</sub> were detected between 4T1 and 67NR cells during the periods of full glutamine deprivation and hypoxia (Figure S6, 1-11 hours). Whereas significant differences in extracellular pH were detected during initial glutamine deprivation (Figure S6, 1-6 hours), the time-course changes in both cell lines followed the same trend. During these periods, the nonmetastatic 67NR cells consistently accumulated more PCO<sub>4</sub>/PCho ratios, and had significantly higher levels of β-NTP than 4T1 cells (Figure S6, Table S1). These results suggest that, compared with 4T1 cells, the energy metabolism of 67NR cells is less sensitive to short-term glutamine deprivation and hypoxic stress because they rely less on OXPHOS than 4T1 cells. The slight decrease of β-NTP levels detected in 4T1 cells (compared with initial values; Table S1, Study C) could also reflect an attempt of the metastatic phenotype to maintain a high glycolytic flux by removing ATP-induced inhibition on phosphofructokinase [37].

**Discussion**

We investigated the dynamic flux of glucose through glycolysis and the TCA cycle, and the flux of glutamine through the TCA cycle. Experiments were performed on a panel of isogenic murine cell lines derived from the same spontaneous breast tumor, focusing mostly on the highly metastatic (4T1) and nonmetastatic (67NR) variants [16]. We found significant differences in metabolism between these two cell lines. The more aggressive cell line (4T1) had higher glycolytic, OXPHOS, and TCA cycle activity than 67NR cells to meet their energy and anabolic requirements for rapid proliferation and the formation of metastases. 4T1 cells adapted more readily than 67NR cells to a changing microenvironment. This is accomplished by a greater ability to regulate glucose metabolism through glycolysis and/or oxidative metabolism. This “plasticity” of 4T1 cells is also supported by our previous experiments [17,23]. For instance, 4T1 cells, which have higher basal levels of LDH-A expression than 67NR cells [17], significantly increased their OCR in response to LDH-A knockdown and reduced aerobic glycolysis [23].

Further, we identified a restriction of TCA cycle flux in 67NR cells but not in 4T1 cells. This was found to be due to lower expression of SDH in 67NR in comparison to 4T1 cells. Thus, 67NR cells accumulated significantly more succinate and displayed significantly lower OXPHOS activity than 4T1 cells under standard growth conditions. These data agree well with the study by Lu et al. [21], which used targeted liquid chromatography–mass spectrometry to assess the metabolomic profile of a panel of nonmetastatic and metastatic murine breast cancer cell lines, including 67NR and 4T1. Under standard tissue culture conditions, they also reported higher succinate levels in 67NR than in 4T1 cells (Table S2, [21]). The accumulation of succinate has also been associated with a “pseudohypoxic” cellular microenvironment that stabilizes HIF-1α [38]. Stabilization of HIF-1α is associated with upregulation of angiogenesis-related proteins, e.g., VEGF [39]. This would agree with our previous observations [17] and others [40], demonstrating that 1) 67NR tumors have better vascular perfusion than 4T1 tumors at comparable tumor sizes and 2) 67NR cells are better adapted to growing in hypoxia [17]. Although SDH has been proposed as a tumor suppressor gene [38], no evidence of its epigenetic silencing has been found in clinical breast cancers [41]. Thus, the presence of a fully functional TCA cycle (including SDH and OXPHOS) may be a hallmark of metastatic breast cancer.

Our results suggest lower SDH-A expression in less aggressive cell lines either in replete environmental conditions or under stress conditions.
(glutamine deprivation and hypoxia). This agrees with the study by Kim et al. showing an association between low SDH expression and low-grade histology in a panel of 721 breast cancers [42]. With relatively less efficient mitochondrial OXPHOS activity, less aggressive cell lines could potentially reduce their TCA cycle by channeling glucose into other, more efficient pathways. This would also agree with data from our group indicating upregulation of phosphoenolpyruvate carboxykinase (PCK1) aligning with decreased metastatic ability (67NR > 168FAR > 4T07 >> 4T1, Figure S7). Together with our estimates of relatively lower pyruvate carboxylase flux in 4T1 cells, this would suggest lower gluconeogenic potential in more aggressive breast cancer phenotypes, in agreement with the loss of fructose-1,6-bisphosphatase in basal-like breast cancer [43]. Moreover, higher relative levels of SDH-A in hypoxic 4T1 cells could be a response to the increased levels of glutamine-derived succinate (reductive carboxylation pathway [44]), whereas SDH-A decreases in hypoxic 67NR cells. Therefore, although more adaptable to changes in their microenvironment, more aggressive cell lines may preserve a relatively higher basal potential for aerobic metabolism under stress, enabling faster adaptation to new microenvironments.

As summarized in Table S2, recent data support an association between mitochondrial function and metastatic potential in breast cancer, and an association with a short metastasis-free survival [45]. The ability to effectively use OXPHOS in addition to glycolysis has been demonstrated in aggressive breast cancer cell lines [46–48] and proposed as a key adaptation for developing brain metastasis [19]. A recent model of tumor metabolism also suggested that stromal cells fuel breast cancer cell growth and the development of metastasis through mitochondrial metabolism [20]. Lu et al. showed that TCA cycle intermediates increase with metastatic potency in isogenic 67NR, 168FARN, 4T07, and 4T1 cells [21]. In our studies with the same panel of cell lines, we found this to be related to higher OXPHOS, indicating that higher metastatic potential is associated with less reliance on aerobic glycolysis alone. The higher utilization of OXPHOS in primary 4T1 tumors may contribute to the development of necrosis due to outgrow of their vascular oxygen supply [17]. Our previous attempts to inhibit 4T1 cancer growth and metastasis focused on LDH-A knockdown alone [23]. These new observations suggest that mitochondrial metabolism (OXPHOS) is also a potential target for treatment in this breast cancer model.

Figure 7. Cell perfusion stress studies with 1,13C-glucose as detailed in Table 1 (Study C). Normalized average time-course changes (A) and labeling rates at different time intervals (B) of lactate C3, glutamate C4, and succinate C2 as determined from 13C MR spectra. In B, the rates in the gray boxes are from the experiments shown in Figure 1C and displayed again for comparison. (C) The values obtained from experiments C-2, A-2, and A-3 (detailed in Table 1) and displayed in A (1-6 hours, 0 mM Gln) and Figure 1C (1-6 hours, 2 mM Gln; and 6-11 hours, 6 mM Gln) were fitted according to the Michaelis-Menten asymptotic model. Because the cellular washout of glutamine is slow (data not shown), when changing from 6 mM to 0 mM glutamine-containing perfusion media, a 5% residual level of glutamine (0.3 mM) was estimated in the cell microenvironment at the beginning of the study. Error bars indicate the standard error of the means.
Thus, combined metabolic inhibition of both the glycolytic and OXPHOS pathways may prove to be more effective in the treatment of metastatic breast cancer.

Glutamine is an essential nitrogen source for the synthesis of nucleotides, amino acids, and ATP, and a major anaplerotic precursor for the TCA cycle [27]. Glutamine addiction is frequent in cancer cells [49] and has been linked to Myc overexpression [33]. Previous work has shown that Myc expression is 2.3-fold higher in 4T1 than in 67NR cells [50]. Singh et al. observed in a panel of aggressive, glutamine-addicted breast cancer cell lines (including 4T1) a correlation between glutamine addiction and metastatic ability [51]. More recently, glutamine dependency was associated with higher invasion and metabolic “rewiring” in ovarian cancer cell lines; specifically, glutamine downregulated glycolysis (glucose uptake and lactate secretion) in highly invasive cancer cells but not in cancer cells with low invasive potential [52]. Similarly, we observed that lactate production by aerobic glycolysis (Warburg effect) is four times more glutamine dependent in 4T1 cells than in 67NR cells. However, in our study, glutamine stimulated the synthesis of glucose-derived lactate according to metastatic potential. Compared with the studies by Yang et al. [52], this could indicate 1) basal metabolic differences between the two cancer cell types (ovarian and breast) and/or 2) different effects of glutamine during the metabolic reprogramming toward more invasive or metastatic phenotypes. Because lactate is a major driver of invasion and dissemination [14,15], our data suggest that glutamine deprivation might prevent production of lactate and inhibit/ delay breast cancer metastases. Glutamine was found to recycle into OAA only in 4T1 cells (as shown by the labeling of aspartate from 3-13C-glutamine studies), whereas nonmetastatic 67NR cells are more dependent on glucose as an anaplerotic precursor of the TCA cycle via pyruvate carboxylase. Accordingly, the flux analysis of our cell perfusion data (in high-glutamine-containing medium) predicted an exchange at the level of pyruvate carboxylase only in 67NR cells and not in 4T1 cells. This also explains the marked dependency of glycolytic lactate synthesis on glutamine observed in 4T1 cells but not in 67NR cells. To sustain their higher ATP demands by OXPHOS, glutamine-deprived 4T1 cells must rely more on glucose to replenish the OAA pool of the TCA cycle (via pyruvate carboxylase). This is accomplished by increasing the amount (fraction) of glucose used for de novo synthesis of citrate. Both cell lines ultimately respond to full glutamine deprivation by reducing the TCA cycle flux and decreasing their lactate synthesis rates to similar levels. This observation agrees with the results reported by Richardson et al. in a panel of transformed MCF-10 breast cancer cells grown in glutamine-free medium [53], which showed no significant differences in lactate excretion rates with increasing malignancy and metastatic potential. Thus, limiting OXPHOS by targeting glutamine metabolism [28] may provide an additional therapeutic approach for metastases-prone or metastatic breast cancer, although other modulators of cellular adaptation should be considered, such as asparagine [54].

Conclusions

Whereas most previous investigations have focused on identifying gene signatures associated with the pathophysiology of metastases [55], our results suggest that the noninvasive evaluation of a tumor’s metabolic phenotype can provide useful information related to the metastatic propensity of the tumor. We show that 4T1 metastatic breast cancer is associated with metabolic plasticity in response to changing tumor cell microenvironments. Specifically, these cells use OXPHOS efficiently during optimal growth conditions (whereas nonmetastatic 67NR cells do not) and reversibly shut down OXPHOS during transient hypoxia and/or glutamine deprivation. These transitions were dynamically quantified for the first time with tumor cell perfusion experiments that closely mimic the in vivo tumor microenvironment. The observed metabolic differences and the ability to assess adaptability in living cells provide the opportunity to identify new biomarkers for early detection of the metastatic phenotype, possibly at the time of diagnosis. Metabolic therapies directed at inhibiting both the glycolytic and OXPHOS pathways are also suggested by these studies; such combination therapies may be more effective in the treatment of aggressive metastatic breast cancer.

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