Proton export upregulates aerobic glycolysis

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Abstract

Introduction: Aggressive cancers commonly ferment glucose to lactic acid at high rates, even in the presence of oxygen. This is known as aerobic glycolysis, or the “Warburg Effect.” It is widely assumed that this is a consequence of the upregulation of glycolytic enzymes. Oncogenic drivers can increase the expression of most proteins in the glycolytic pathway, including the terminal step of exporting H+ equivalents from the cytoplasm. Proton exporters maintain an alkaline cytoplasmic pH, which can enhance all glycolytic enzyme activities, even in the absence of oncogene-related expression changes. Based on this observation, we hypothesized that increased uptake and fermentative metabolism of glucose could be driven by the expulsion of H+ equivalents from the cell.

Results: To test this hypothesis, we stably transfected lowly glycolytic MCF-7, U2-OS, and glycolytic HEK293 cells to express proton-exporting systems: either PMA1 (plasma membrane ATPase 1, a yeast H+-ATPase) or CA-IX (carbonic anhydrase 9). The expression of either exporter in vitro enhanced aerobic glycolysis as measured by glucose consumption, lactate production, and extracellular acidification rate. This resulted in an increased intracellular pH, and metabolomic analyses indicated that this was associated with an increased flux of all glycolytic enzymes upstream of pyruvate kinase. These cells also demonstrated increased migratory and invasive phenotypes in vitro, and these were recapitulated in vivo by more aggressive behavior, whereby the acid-producing cells formed higher-grade tumors with higher rates of metastases. Neutralizing tumor acidity with oral buffers reduced the metastatic burden.

Conclusions: Therefore, cancer cells which increase export of H+ equivalents subsequently increase intracellular alkalization, even without oncogenic driver mutations, and this is sufficient to alter cancer metabolism towards an upregulation of aerobic glycolysis, a Warburg phenotype. Overall, we have shown that the traditional understanding of cancer cells favoring glycolysis and the subsequent extracellular acidification is not always linear. Cells which can, independent of metabolism, acidify through proton exporter activity can sufficiently drive their metabolism towards glycolysis providing an important fitness advantage for survival.

Keywords: Cancer, Glycolysis, Proton, pH, Warburg, PMA1, CA-IX, Metastasis

Background

In 1924, Otto Warburg and colleagues demonstrated that cancer, even in the presence of oxygen, ferments glucose to lactic acid at high rates [73], and this was contemporaneously confirmed by the Coris [10]. Aerobic glycolysis, commonly termed the “Warburg Effect” in cancer [14, 71], is undeniably a hallmark of primary tumors and aggressive invasive disease [25]. This preference of tumors for increased glucose uptake is exploited in diagnostic PET imaging of fluorodeoxyglucose (18F-FDG)
uptake [36]. It is commonly believed that this increased fermentative glycolysis is driven by activity of oncogenic pathways, such as RAS, MYC, HIF, and AKT [34, 77] and that this augmented flux out-competes the ability of mitochondria to oxidize pyruvate, leading to the net production and export of lactic acid and reversion of NADH to NAD\(^+\) to maintain redox balance. Hence, the Warburg Effect could solely be an epiphenomenon of oncogene activation, which is consistent with the observation that fermentation under aerobic conditions is energetically unfavorable and does not confer any obvious evolutionary benefits. It must be acknowledged that glycolytic ATP generation is more rapid than oxidative phosphorylation and that this may confer some selective advantage for transporters [16, 70]. Nonetheless, it is clear that under conditions of similar proliferation and motility, more aggressive cancer cells ferment glucose at much higher rates.

We thus propose an alternative to this canonical view, building on principles of evolutionary dynamics [21]. Aerobic glycolysis is such a commonly observed phenotype of aggressive cancers [57, 79], and we argue that it **MUST** confer some selective advantage for tumor growth [13]. An inherent consequence of glycolysis is lactic acid production, and we propose that acid secretion per se renders cells more competitive, despite the energetic cost [20, 22]. Indeed, oncogenic drivers upregulate expression of proton-exporting systems, e.g., sodium hydrogen anti-porters, NHEs [6, 8], carbonic anhydrase 9, CA-IX [35, 42, 68], and sodium bicarbonate co-transporters, NBCs [3]. Such activities will exacerbate the intra- to extracellular pH gradient, raising the intracellular pH (pHi) and acidifying the extracellular pH (pHe). These transporters have been associated with breast cancer aggressiveness [1]. Additionally, glycolytic enzymes all exhibit significant pH dependence [54], and thus maintenance of an alkaline pH would directly promote increased glycolytic flux.

Simultaneously, lowering pHe provides invading cancer cells competitive benefits that can enhance colonization, invasion and metastasis. These benefits include extracellular matrix remodeling via release and activation of proteases to increase invasion [33, 64, 75], inhibition of immune surveillance [4], promotion of an epithelial-to-mesenchymal transition (EMT) [47, 51, 55, 56], and anchorage-independent growth [12, 30, 49, 52]. This is further supported by the observation that neutralization of pHe acidity can inhibit invasion and metastases [17, 58].

The current study investigates if and how aerobic glycolysis can be driven by proton export and further investigates the impact of this on cancer aggressiveness. We demonstrate that over-expression of proton exporters is sufficient to increase aerobic glycolysis, through enhanced glucose uptake and lactate production. We further observed that proton export increased intracellular pH and increased metabolic flux upstream of pyruvate kinase, PK. Finally, we observed in vivo that these proton-exporting cell lines were more aggressive, generating higher-grade tumors and increased metastases. There is a known association between acid production, aerobic glycolysis, and metastatic potential. Further, experimental metastases can be inhibited with acid-neutralizing buffers. The current work adds to this literature by demonstrating that acid production per se can be sufficient to drive the Warburg Effect and promote metastasis.

**Results**

**Over-expression of CA-IX in cancer cells increases glycolytic metabolism**

CA-IX hydrates extracellular CO\(_2\) to H\(^+\) + HCO\(_3^-\). This facilitates CO\(_2\) diffusion away from the cell reducing pH gradients across tissues. The bicarbonate generated from CO\(_2\) hydration, a reaction that occurs either spontaneously or is sped up by carbonic anhydrases, can then reenter the cell via Na\(^+\) + HCO\(_3^-\) co-transporter [3, 65].

Numerous studies in, e.g., breast, ovarian [9], and astrocytoma [48] cancers, have shown that CA-IX expression correlates with poor prognosis and reduced survival. Figure 1A [24] shows the overall survival of luminal B, ER+ breast cancer patients with low and high (autoselect cutoff) CA-IX expression was 108 and 57.53 months, respectively, \(p=0.0433\). Although the sample size was smaller, a similar pattern was seen for metastasis-free survival in patients with low and high CA-IX expression, 130 vs. 50 months, respectively, \(p=0.0012\). Metastasis-free survival is also reduced in other cancers with high CA-IX expression, including cervical and colorectal [69]. CA-IX’s role in outcome makes it a clinically significant target warranting further investigation.

To test the hypothesis that proton export can drive aerobic glycolysis, we established models to over-express proton exporters. We transfected MCF-7 cells with a Ca9 expression vector and isolated two individual clones (M1 and M6) and confirmed CA-IX protein expression (Fig. 1B). MCF-7 cells do not express CA-IX under normal conditions, but do express other carbonic anhydrases, CA-II and CA-XII (Fig. 1B). CA-IX is distinct among exofacial CA’s (CA-IV, CA-XII), as it contains a proteoglycan domain, which enables it to maintain enzymatic activity at lower pHe [38]. In tissues, CA-IX can function as a “pH–stat,” which tumors hijack to maintain an acidic pHe [37]. We also transfected MCF-7 cells with an empty pcmv6 vector hereafter referred to as MOCK-2. Additionally, we confirmed by immunocytochemistry (ICC) that CA-IX, an exofacial membrane-bound protein, was expressed on the plasma membrane in both CA-IX clones (Fig. 1C).
To test our hypothesis that proton export can drive aerobic glycolysis, we interrogated the metabolism of our CA-IX expressing clones using a Seahorse XFe96 Extracellular Flux (XF) Analyzer, enzymatic, and radiochemical assays to assess both glycolytic and mitochondrial metabolism. Specifically, the Seahorse glycolytic stress tests (GST) showed that both CA-IX clones exhibited higher proton production rates (PPR) upon glucose stimulation compared to MOCK-2 or parental clones (Fig. 1D). Using glucose and [3H]-2-deoxyglucose (2DG) uptake assays (Fig. 1E & Additional file 1: Fig. S1), and lactate production rate assays (Fig. 1F & Additional file 1: Fig. S2), we further confirmed CA-IX expressing clones had increased glycolysis in normoxic conditions. Mitochondrial metabolism in these clones, as measured by Seahorse mitochondrial stress test (MST), exhibited a decreased reliance on oxidative phosphorylation. In the CA-IX clones, both basal oxygen consumption rates (Fig. 1G) and ATP-linked oxygen consumption rates (Additional file 1: Fig. S3) were decreased compared to MOCK-2 or parental cells. CA-IX expressing clones also had hyperpolarized mitochondria (Additional file 1: Fig. S4). Therefore, CA-IX expression did not globally upregulate all ATP turnover, but upregulated aerobic glycolysis and limited reliance on oxidative phosphorylation.

To investigate whether these metabolic alterations were specific to the MCF-7 cells, we tested other cell lines. CA-IX was over-expressed in U2-OS osteosarcoma human cells and HEK 293 human embryonic kidney cells. CA-IX expression upregulated glycolysis in both cell lines, as seen by increased aerobic lactate production (Additional file 1: Fig. S5 & S6). Even in HEK 293 cells, which have higher basal glycolysis than the other cell lines tested, over-expression of the proton-exporting CA-IX still enhanced their rate of aerobic glycolysis. Therefore, CA-IX expression did not globally upregulate all ATP turnover, but upregulated aerobic glycolysis and limited reliance on oxidative phosphorylation.

To delineate which steps in glycolysis were being impacted by acid export, we analyzed the intracellular metabolites using mass spectrometry and a library
of known metabolites (see “Methods”). Principal component analysis (PCA) showed a statistically significant separation of MOCK-2 and parental MCF-7 from the CA-IX clones (Fig. 2A). Although the parental and MOCK-2 cells had distinct metabolic profiles, neither overlapped with the CA-IX clones, which were analogous. Consistent with this, heatmap visualization of the top 50 most significantly altered metabolites showed that the two CA-IX clones exhibited similar metabolic profiles (Fig. 2B) but were considerably different from the

Fig. 2 Unlabeled metabolic profiling and analysis of intracellular metabolites of CA-IX expressing MCF-7 cells. A Principal component analysis of intracellular metabolites in CA-IX expressing, MOCK-2, and parental MCF-7 cells. B Heatmap (hierarchical clustering) of the fifty most significant fold changes of intracellular metabolites between CA-IX expressing clones, MOCK-2, and parental cells. Numerous glycolytic intermediates were significantly higher in CA-IX expressing cells compared to MOCK-2 and parental. C–J Average peak intensity of each glycolytic intermediate in CA-IX expressing, MOCK-2, and parental MCF-7 cells. N = 5–6 biological replicates per group, statistical analysis using ordinary one-way ANOVA. *p < 0.05, **p < 0.01, ***p < 0.001, ****p < 0.0001
MOCK-2 and parental clones. MOCK-2 cells grow more rapidly in vitro compared to the parental MCF-7 or the CA-IX clones which is likely why they exhibit a different metabolic profile from the parental line (Additional file 1: Fig. S7). However, this difference in growth rate was not maintained in vivo when grown as primary tumors. Out of all metabolites assessed, the glycolytic intermediates were consistently altered in the CA-IX clones compared to parental or MOCK-2.

The CA-IX clones exhibited increased levels of all glycolytic intermediates upstream of pyruvate kinase, PK (Fig. 2C–J), which catalyzes the penultimate step of glycolysis: the conversion of phosphoenolpyruvate (PEP) + ADP → pyruvate + ATP. Thus, it appears that the activities of the upstream enzymes have increased, leading PK to now become rate-limiting for glycolytic flux in the CA-IX expressing cells (Fig. 2). As the CA-IX cells have higher glycolytic flux (Fig. 1D–F), the most straightforward interpretation is that CA-IX expression de-inhibited all of the enzymatic steps upstream of PK. Overall, CA-IX expression enhances glycolytic intermediates’ flux, resulting in enhanced lactate and acid production.

**CA-IX over-expression increases pH**

Our metabolomics studies suggest multiple glycolytic enzymes were impacted. We therefore hypothesized that CA-IX expression might raise the intracellular pH of cells, which could pleiotropically increase glycolytic enzyme rates. Most glycolytic enzymes have ionizable residues that can alter their enzyme activity. Recently, these residues have been characterized for all glycolytic enzymes through homology modelling, which predicted glycolytic enzyme activities generally increase with pH above neutral [54]. We tested the PPR in MCF-7 parental cells after altering pH in a range of pH media 6.6–7.4, expecting that the PPR rate would increase with increasing pH. Cells were incubated with either a chloride-containing solution or an iso-osmotic low-chloride formulation which replaced chloride salts with gluconate equivalents. Low-chloride solutions alter the driving force for Cl⁻/HCO₃⁻ exchangers, effectively loading cells with HCO₃⁻ ions and raising pH at the same pH [61, 78]. At all values of pH from 6.6 to 7.2, cells in gluconate media exhibited significantly higher glycolytic rates compared to those in chloride-containing media (Fig. 3A & Additional file 1: Fig. S8), suggesting that the increased glycolytic rate is strongly dependent on pH.

CA-IX over-expression has been shown to increase pH in other systems [46], leading us to probe pH in our CA-IX expressing cells. We used a pH reporter dye, cSNARF1 to measure the effects of CA-IX expression on intracellular pH (pHi). An in situ cSNARF1 calibration curve was generated using nigericin/K⁺ buffers (Additional file 1: Fig. S9). We then loaded MCF7 parental or the CA-IX expressing clones with cSNARF1 and used fluorescence imaging to measure pH. MOCK-2 cells were unsuccessfully loaded with cSNARF1, possibly due to reduction or loss of esterase activity from the integration of the MOCK-2 plasmid into the MCF-7 DNA; therefore, they are not included in these analyses. We equilibrated the cells in media at pH 6.6, 6.8, 7.1, and 7.4 and co-loaded them with cSNARF1 and nuclear dye Hoechst 33,342 (Fig. 3B–E). The inclusion of a nuclear dye allowed post-processing to mask the nucleus, ensuring measurement of cytoplasmic pH only. At the intermediate pH values of 6.8 and 7.1, the CA-IX expressing clones had significantly higher pH compared to parental (Fig. 3C, D & Additional file 1: Table S1). The enzymatic activity of CA-IX is optimal at pH 6.8 and drops precipitously as the pH is lowered to 6.0 [38, 44]. Consistent with this, at an acidic pH of 6.4, we observed that the MCF7 parental cells had a higher intracellular pH compared to CA-IX clones (Fig. 3B & Additional file 1: Table S1), suggesting CA-IX enzymatic function was reduced resulting in reduced CO₂ venting. There were no significant differences in pHi at pH 7.4 (Fig. 3E & Additional file 1: Table S1). These data show that CA-IX expression can raise the pH at intermediate pH values and thus increasing pH likely results in increased glycolytic flux by globally enhancing glycolytic enzyme activity.

**CA-IX expression increases metastasis**

We then investigated the effect of CA-IX expression on migration, invasion, and metastasis, as there are reported correlations between increased glycolytic flux, reduced extracellular pH, and metastasis [2, 17, 26, 32]. Additionally, CA-IX has been associated with increased invasion and metastasis in several systems [66, 74].

In vitro, we utilized scratch assays and gel escape to measure migration and invasion in the CA-IX clones. Scratch assays showed that clone M1 had increased migratory ability (Additional file 1: Fig. S10), closing the wound significantly more rapidly than controls. However, this was not observed in the M6 CA-IX clone. In the gel escape assay, expansion out of the gel is due to a combination of proliferation, invasion, and migration. Both CA-IX clones invaded the area surrounding the gel drop substantially quicker than the parental MCF-7 clone (Additional file 1: Fig. S11). Compared to the MOCK-2 cells, however, only the M6 clone was significantly more invasive. The apparent increased invasion rate by MOCK-2 compared to parental is likely due to their increased proliferation rate as correcting for proliferation eliminates the difference between MOCK-2 and parental (Additional file 1: Fig. S7). It is important to note
that these assays were carried out at neutral pH and that the invasive behavior might be further enhanced by low pH, as we have shown previously for melanoma cells [45]. Together, these studies suggest that CA-IX expression can enhance cell motility, increasing migration and local invasion.

Furthermore, aggressive cancers with stem-like properties can resist anoikis, which can be measured in vitro by cells’ ability to form spheroids independent of attachment to a basement membrane. Utilizing the hanging droplet technique, we observed that CA-IX expression enabled robust, compact spheroid formation, compared to the MOCK-2 and parental MCF-7 clones, which could not (Additional file 1: Fig. S12). This spheroid forming ability suggests CA-IX not only enhances cell:cell adhesion but also suggests that increased proton export can contribute to anoikis resistance when detached from the basement membrane.

Although the phenotype of our proton-exporting CA-IX clones appeared to be more aggressive, this could be an in vitro only phenomenon. We thus investigated the clones in vivo. We studied the effect of CA-IX expression on primary tumor growth, as well as the ability of these clones to form both spontaneous (from the mammary fat pad) and experimental (tail vein injected) metastasis. For primary and spontaneous metastasis models, MOCK-2, M1 or M6 CA-IX MCF-7 cells were implanted in the mammary fat pads of mice, and growth was monitored by caliper measurement. Although in vitro, mock cells proliferated faster (Additional file 1: Fig. S7), this was not observed in vivo, as the tumors from CA-IX expressing clones grew significantly faster. At all of the time points measured, primary tumor volume was significantly increased in the CA-IX expressing clones, compared to controls (Fig. 4A & Additional file 1: Fig. S13). Regression-based analysis determined the growth rate of tumors was significantly different between the groups, \( p < 0.0001 \). Survival surgery was performed and the primary tumors were resected and subsequently sectioned, stained with H & E, and blindly graded by a board-certified pathologist (M.B.), who identified increased stromal invasion in the CA-IX expressing tumors (Fig. 4B).
The mice were monitored for an additional 12 weeks post-primary tumor removal, at which time the animals were sacrificed, and lungs stained for evidence of micro and macro metastases, scored blindly by a board-certified pathologist (M.B.). As shown in Table 1, 6/21 mice developed spontaneous metastasis from the M6 clone, whereas no spontaneous metastases were formed from the M1 clone and 1/10 were formed from the MOCK-2 clone.

Because formation of spontaneous metastases is a multi-step, time-consuming, and complex process, we also investigated the ability of these clones to form experimental metastases following tail vein injection, which only involves the final steps of extravasation and

| Mouse Model         | Experimental metastasis # of mice with lung mets | p-value (**p<0.01)** | Spontaneous metastasis # of mice with lung mets | p-value  |
|---------------------|-----------------------------------------------|----------------------|-----------------------------------------------|----------|
| MCF7/parental       | 0/5 (0%)                                     |                      |                                               |          |
| MCF7/MOCK-2         | 0/6 (0%)                                     |                      |                                               |          |
| MCF7/CA-IX M1       | 4/7 (57%)                                    | 0.07 (n.s.)          | 1/10 (10%)                                    |          |
| MCF7/CA-IX M6       | 6/6 (100%)                                   | 0.0022**             | 6/21 (29%)                                    | 0.37 (n.s.)|

**p<0.01
colonization. Metastases to the lung were scored blindly by a board-certified pathologist (M.B.), who observed that neither of the control groups developed metastases and that both CA-IX clones had significant macrometastases (Table 1). Consistent with the spontaneous model, the M1 formed fewer metastases compared to the M6; however, both CA-IX clones exhibited gross metastasis to the lungs (Fig. 4C), which were confirmed histologically (Fig. 4D). The resulting macrometastases significantly reduced overall survival of the mice (Fig. 4E). Additional immunohistochemistry (IHC) staining confirmed the lung metastases expressed both CA-IX and human estrogen receptor, which we used as a marker for MCF7 cells (Fig. 4F).

Prior studies have shown that neutralization of acidity using oral buffers inhibits metastasis [27, 28]. As we hypothesized that our M6 CA-IX clones were metastatic by virtue of increased acid production (Fig. 1D, E), we asked whether buffer therapy would reduce the metastatic burden. Using the experimental metastasis model, we compared untreated to buffer-treated M6 or MOCK-2 mice. As in the first experimental metastasis study, all mice injected with M6 developed macrometastases, and these were significantly reduced by bicarbonate (Fig. 4G, H). Two out of ten MOCK-2 mice developed very small micrometastases, and no metastases were found in the buffer therapy MOCK-2 group (Fig. 4G).

Over-expression of yeast proton pump PMA1 increases motility and metabolism
While we hypothesize that CA-IX is acting as a proton equivalent exporting system, there are many other activities of this protein, including non-enzymatic activities that could be activating glycolysis and promoting metastasis. To test whether the observed effects on glycolysis could be due to increased proton export, we utilized another model, PMA1, which electrogenically pumps H+ out of cells at the expense of ATP [19]. Prior work has shown ectopic expression of PMA1 in murine 3T3 fibroblasts led to tumorigenesis [53] and to increased aerobic glycolysis with elevated intracellular pH [23, 43]. We engineered MCF-7 cells to express PMA1 and selected two clones following zeocin selection (PMA1-C1 and PMA1-C5) as well as an empty vector transfected control (MOCK-1). PMA1 over-expression in C1 and C5 was confirmed by qRT-PCR (Additional file 1: Fig. S14) and western blot (Fig. 5A). Moreover, using immunocytochemistry of non-permeabilized cells (Fig. 5B & Additional file 1: Fig. S15) or permeabilized cells (Additional file 1: Fig. S16), we verified PMA1 protein expression on the plasma membrane and in the cytoplasm of both PMA1 clones.

To characterize the metabolic activity of PMA1 transfectants, we again utilized the glycolysis and mitochondrial stress tests (GST and MST, respectively) of the Seahorse (XFe) Analyzer. The glucose-induced PPR (Fig. 5C) and the glycolytic reserve (Additional file 1: Fig. S17) were significantly higher in the PMA1 clones compared to empty vector MOCK-1 or parental clones, suggesting functional activities of the transfected pump. In contrast to the CA-IX transfectants, there were no significant differences in oxygen consumption rates (OCR) between PMA1 clones and controls (Fig. 5D). This could be due to increased energy demand from the ATPase proton pump. We further confirmed these metabolic alterations by measuring glycolytic flux. PMA1 clones had significantly higher glucose consumption rates (Fig. 5E) and lactate production rates (Fig. 5F) compared to MOCK-1 or parental MCF-7 clones. These data, together with the CA-IX results, indicate that acid export can drive cells to exhibit a Warburg phenotype.

As with the CA-IX transfectants, we also measured invasion and migration using gel escape and circular wound healing assays, respectively. Compared to the MOCK-1 cells, both PMA1 clones expanded significantly more out of the gel drop (Fig. 5G & Additional file 1: Fig. S18). In the circular “wound healing” assay, we monitored the migration of cells into a cell-free area. Again, compared to MOCK-1, both PMA1 clones had increased migration rates (Fig. 5H & Additional file 1: Fig. S19). Together, these results indicate that cellular invasion and migration were significantly enhanced by PMA1 expression and acid production.

PMA1 induces a metastatic phenotype in vivo and alters the expression of proteins involved in metabolism and pH regulation
To investigate if proton export enhanced aggressiveness, as seen in the CA-IX model, we measured the PMA1 cells’ metastatic ability in vivo in both spontaneous and experimental metastasis models. In our in vitro studies, the proliferation rates of PMA1-C1 and the empty vector MOCK-1 clones were similar, whereas the growth rate of PMA1-C5 was significantly slower (Additional file 1: Fig. S20). Hence, we omitted PMA1-C5 in our in vivo studies to reduce the possibility of proliferation being a confounding variable. In both the spontaneous and tail vein metastases models, only 1 of 10 mice in each MOCK-1 group developed lung metastases. In contrast, 7 of 12 PMA1-C1 mice developed lung metastases following tail vein injection and 4 of 9 formed spontaneous metastases (Additional file 1: Table S2).

Lung metastases of PMA1 cells, visualized by H&E staining (Fig. 6A), were further validated by IHC of PMA1 (Fig. 6A) and RNA analysis of FFPE lung tissue for
PMA1 gene expression (Additional file 1: Fig. S1). Notably, primary tumors revealed no significant growth differences between PMA1 and MOCK-1 tumors (Fig. 6B) or final tumor volume (Fig. 6C). However, blind grading (1 to 4+) of H&E-stained tumor sections by a board-certified pathologist (A.L.) indicated that PMA1 primary tumors were of significantly higher grade compared to MOCK-1 controls ($p = 0.016$). The average grade was 2.7 ± 0.52 for MOCK-1 compared with 3.4 ± 0.67 for the PMA1 primary tumors (Fig. 6D, E). Maintenance of
PMA1 expression in vivo was confirmed with quantitative IHC of the resected primary tumors, demonstrating a significant difference in PMA1 protein expression between the MOCK-1 (19% ± 3.0, n = 10) and PMA1-C1 (90% ± 2.5, n = 9) tumors (Fig. 6F, G).

Additional IHC of PMA1 tumors showed they had higher levels of the monocarboxylate (lactate) transporter, MCT1 (Fig. 6H,I), which has been associated with increased aggressiveness in breast cancer [15]. Conversely, PMA1 tumors had significantly lower CA-IX expression than MOCK-1 (Fig. 6 J,K). As CA-IX plays a vital role in regulating tumor pH [15, 37, 67], we postulate that its activity may have been made redundant by PMA1. Notably, other proteins, such as glucose transporter 1, GLUT1 (Additional file 1: Fig. S22), and the sodium hydrogen exchanger 1, NHE1 (Additional file 1: Fig. S23) showed no differences between the PMA1 and MOCK-1 groups. These data suggest that the increased glycolytic flux, which requires glucose uptake by GLUT1, can be accommodated by native protein levels of GLUT1 (i.e., it is not rate-limiting).

**Discussion**

The primary goal of this study was to determine if acid export per se could drive aerobic glycolysis, the “Warburg Effect,” in cancer. Aerobic glycolysis, a cancer hallmark, is often associated with more aggressive tumors. Numerous studies have attempted to determine why tumors favor fermentative glycolysis, even in the presence of sufficient oxygen [21, 70]. Aerobic glycolysis is bioenergetically inefficient, producing only 2 ATP per glucose, compared to 36 ATP upon complete oxidation. Thus, it is not obvious why this should be such a prevalent phenotype. It is axiomatic that common phenotypes confer a selective advantage, yet the evolutionary drivers of aerobic glycolysis are not clear. Theories proposed include glucose addiction [5], dysfunctional mitochondria [63], which is not the case in most cancers, and to meet the rapid energetic requirements of membrane transporters [16]. Another theory is that glycolysis enhances tumor cell proliferation by generating the anabolic building blocks for macromolecules. However, most of the glucose-derived carbons are exported out of the cell as lactate [41]. Moreover, TCA cycle intermediates derived from glutamine consumption are considered more important for lipid, amino acid, and nucleotide synthesis, compared to glycolytic intermediates [7, 76].

However, none of these theories clearly demonstrate why so many cancers favor aerobic glycolysis. We hypothesize that acid export per se and extracellular acidification provides a distinct selective advantage and that it is enabled by increased glucose fermentation. This theory was first proposed as the “acid-mediated invasion hypothesis” [20] and has been subsequently elaborated [22].

To test the hypothesis that acid export increases glycolysis in cancer, we over-expressed a proton exporter, CA-IX, in an OXPHOS dominant cell line, MCF-7. In MCF-7 cells, CA-IX is not expressed under normoxic conditions, and over-expression resulted in cells that more rapidly exported acid and upregulated glycolysis. Both glucose consumption and lactate production rates increased. Other studies have shown that CA-IX over-expression can increase lactate production, albeit under hypoxic conditions [29]. As glycolysis was specifically increased, we investigated the mechanisms whereby acid export could be causing this shift in metabolism. We broadly interrogated cellular metabolism using a targeted metabolomics panel and found that the most significantly altered metabolites in the CA-IX expressing clones were glycolytic intermediates. Specifically, all glycolytic intermediates upstream of pyruvate kinase were increased. Enzyme activity can be affected by pH, including those in the glycolytic pathway, and are most active at the pH of their subcellular compartment from acidic lysosomes to mitochondrial [54]. For glycolytic enzymes, pH optima are slightly on the alkaline side of neutral (7.2–7.4), meaning that raising pH above neutrality will globally increase activities of glycolytic enzymes. Because...
the data suggested pleiotropic increases in enzyme activity, we measured intracellular pH using fluorescence ratio imaging in our CA-IX expressing clones. These clones had a higher pHi in biologically relevant extracellular pH conditions, compared to the parental. Specifically, at pH 6.8 and 7.2, at which CA-IX can function as a proton equivalent exporter, the CA-IX clones had increased pHi compared to parental. One caveat of this experiment...
was that the MOCK cells were unquantifiable as they did not accumulate SNARF-1, possibly due to reduced esterase activity or increased activity of multidrug resistance transporters. In addition, the experiments in low-chloride directly implicate increased pH in regulating aerobic glycolytic flux. These data indicate loading cells with HCO$_3^-$ ions raises the intracellular pH sufficiently to enhance enzyme activity and result in increased glycolytic flux.

We tested our hypothesis using two more models and another acid exporting protein to minimize cell line and protein-specific effects. Similar to the MCF-7 results, over-expression of CA-IX in U2-OS and HEK293 cells increased glycolysis compared to controls. Although parental HEK293 cells are more glycolytic than the other cell lines chosen, over-expression of CA-IX still enhanced glycolysis. We tested another acid exporting protein, PMA-1, which has an unequivocal activity of exporting protons at the expense of ATP. PMA-1 over-expression in MCF-7 cells similarly resulted in increased glycolysis, as measured by increased glucose uptake and lactate production. These findings, together with our CA-IX results, suggest that expression of proton-exporting activity can upregulate aerobic glycolysis, likely through a global increase of intracellular pH. Notably, while the CA-IX transfecants had reduced oxygen consumption, this was not observed in the PMA1 cells. The cause of this difference is not known and may reflect differences in the bioenergetic requirements for the two transporting systems.

We hypothesize that acid export-driven glycolysis would make them more aggressive, as measured in vitro with motility and invasion assays, and in vivo with experimental and spontaneous metastases studies. Glycolysis and acidity have been correlated with poor prognosis and metastasis [72, 75]. Our focus on CA-IX as an acid exporter was due to its clinical relevance in many cancer types, such as breast, ovarian, and astrocytoma, where CA-IX over-expression correlates with poor prognosis, reduced survival, and reduced metastasis-free survival. This suggests CA-IX specifically and perhaps acid export generally enhances cancer aggressiveness and subsequently metastasis. In our models, acid export driven by CA-IX or PMA1 was linked to enhanced migration and invasion in vitro, which is consistent with prior studies [11, 62, 65]. In vivo, cells at the invasive edge tumor periphery are known to be more acidic and express CA-IX [39, 59]. CA-IX has also been shown to enhance matrix metalloproteinase activity, in particular MMP-14 which is active at an acidic pH, resulting in stromal degradation that aids cancer migration into the periphery [66].

Our encouraging in vitro results led us to take these models, CA-IX and PMA-1, in vivo. We found that aggression and metastasis were higher in both PMA-1 and CA-IX transfectants. Primary tumor growth was enhanced in the CA-IX model compared to controls. In the PMA-1 model, a pathologist blindly scored the primary tumors a higher grade compared to control tumors. However, spontaneous metastasis after primary tumor resection was not significantly increased in PMA-1 or CA-IX transfectants compared to controls (Table 1 & Additional file 1: Table S2). Although not statistically significant in individual experiments, it is notable that combined data showed that in spontaneous models 10/45 mice with PMA-1 or CA-IX clones had metastases, compared to 2/20 control mice with parental or MOCK-transfections. In contrast, our experimental metastasis model, which skips the intravasation step, showed enhanced experimental metastasis in both models, with 17/25 mice in PMA-1 or CA-IX transfectants developing metastasis, compared to 1/21 metastases in control mice (Table 1 & Additional file 1: Table S2). We did not quantify the number or size of metastatic lesions, because we contend that the important metric is binary: i.e., whether or not these clones were able to metastasize at all. A related study in 4T1 breast cancer showed that inhibiting CA-IX reduced tumor growth and experimental metastasis [40]. However, inhibition is different than induction, and 4T1 are highly glycolytic and acidic to begin with. Further, this study does indicate the importance of acid export and its role in enhancing tumor growth. Due to the robust enhanced metastasis formation in the experimental metastasis studies, it indicates that acid export can facilitate tumor cell extravasation out of the blood vessels and colonization of metastatic sites.

In our CA-IX model, buffer therapy significantly reduced tumor burden in the lungs compared to their untreated counterparts. Although this did not completely prevent metastasis, combinations of buffer therapy with specific acid exporter inhibitors may be necessary. CA-IX is minimally expressed in normal tissue and could be a viable therapeutic target [50] and other proton pump inhibitors are currently in clinical trials [18].

Conclusions

Many studies have hinted at the importance of acidity, and many have proposed reasons as to why cancer cells favor aerobic glycolysis, but few have proposed that acidity is the driver. This study represents the first to test and show that acid export can increase aerobic glycolysis and enhance cancer aggressiveness, rather than acid merely being an epiphenomenon of glycolytic metabolism.
Methods

Construction of stable cell lines

Plasmids

Yeast plasma membrane ATPase 1 (PMA1) cDNA construct was designed based on the sequence (Accession Number: NM_001180873; Saccharomyces cerevisiae S288c PMA1). The codons were optimized for the suitable expression in mammalian cells, and restriction enzyme sequences Hind III and Xho I were inserted at the 5’ and 3’ ends of the full-length sequence, respectively. The fully designed DNA sequence was commercially synthesized (Blue Heron Biotechnology, Bothell, WA, 98,021). This was cloned into pcDNA3.1/Zeo (+) vector in which PMA1 gene expression was driven under the CMV promoter. The sequence of the pcDNA/PMA1 construct and the identity of the parental cell line were confirmed by the molecular genomics core facility (Moffitt Research Institute, Tampa, FL). Carbonic anhydrase 9 (CA9) construct was designed by Origene based on the sequence (Accession Number: NM_001216; Homo sapiens) and cloned into a pCMV6 vector (PS10001, Origene, MD) to form pCMV6/CA-IX vectors (CQ10630, RC204839 subclone, Origene, MD) in which CA9 gene expression was driven under the CMV promoter.

Cell lines

The MCF-7 cells and HEK 293 cells as transfection host cell lines were acquired from American Type Culture Collection (ATCC HTB-22, Manassas, VA) and maintained in RPMI media 1640 (Life Technologies Gibco®, 11,875–093) supplemented with 10% FBS (Hyclone Laboratories, UT) under standard cell culture conditions. The U2-OS cell line was a gift from Jillaina Menth, Moffitt Cancer Centre Translational Research Core, and maintained in RPMI media 1640 (Life Technologies Gibco®, 11,875–093) supplemented with 10% FBS (Hyclone Laboratories, UT) under standard cell culture conditions. The MCF-7 cells were transfected with empty pcDNA, pcDNA/PMA1, pCMV6 (PS10001, Origene, MD), and pCMV6/CA-IX vectors (CQ10630, RC204839 subclone, Origene, MD) respectively, in MCF-7/MOCK-1 cells, MCF-7/PMA1, MCF-7/MOCK-2, and MCF-7/CA-IX cell lines by standard clonogenic stable cell construction procedures using Fugene HD (Promega, E 2311). The U2-OS and HEK-293 cells transfected with empty pCMV6 (PS10001, Origene, MD) and pCMV6/CA-IX vectors (CQ10630, RC204839 subclone, Origene, MD) respectively, resulted in U2-OS/MOCK-2, HEK/MOCK-2, and U2-OS/CA-IX and HEK/CA-IX clones. Briefly, a number of individual single clones were selected in the media containing 300 µg/ml zeocin (Invitrogen, 450,430, Carlsbad, CA), or 300 µg/ml G418 and stable expression in individual clones were verified using western blotting. Cell lines were tested for mycoplasma using MycoAlert assay (Lonza).

Spheroid formation

Spheroids were formed as previously described [60]. Briefly, cells were suspended in Perfecta 3D hanging drop plates (HDP1096, 3D Biomatrix, MI) at 25,000 cells/40 µl droplet. Spheroids were allowed to form for 5 days and then centrifuged at 450 rpm, with no brake, into media-containing Costar Ultra low attachment U-bottom 96-well plates (CLS3474, Corning, NY). Spheroids were imaged in the Celigo Imaging Cytometer (Nexcelom Bioscience, MA) using bright-field imaging single colony verification analysis.

Western blotting

Chemiluminescence

The cell membrane protein samples were collected using Mem-PER eukaryotic membrane protein extraction reagent kit (Thermo Scientific, 89,826, MA) according to the protocol instruction, and the protein samples were further purified and concentrated by Pierce SDS-PAGE sample prep kit (Thermo Scientific, 89,888). Thirty micrograms of protein per sample was separated on polyacrylamide-SDS gels and electrophoretically transferred to nitrocellulose membranes. Membranes were incubated with primary antibody against PMA1 (1:1000, Abcam, ab4645) and GAPDH (1:1000, Santa Cruz Biotechnology, TX, sc-25778). For visualization, horseradish peroxidase (HRP)-conjugated secondary antibodies: goat anti-rabbit IgG HRP and goat anti-mouse IgG HRP, followed by ECL kit (Thermo Scientific, 32,209) were used.

Fluorescence

Fifteen micrograms of protein per sample was separated on a Bio-Rad Mini-protein 4–15% precast 12-well 20 µl gels (4,561,085, Bio-Rad, CA) and electrophoretically transferred to Odyssey Nitrocellulose membrane (P/N 926–31,092, LI-COR, NE). Membranes were blocked with Odyssey TBS Blocking Buffer (P/N 927–50,000, LI-COR, NE) and incubated with primary antibody against CA-IX (1:1000, M75 mouse monoclonal CA-IX, Biocscience Slovakia), ER alpha (1:2000, Rabbit polyclonal ab 75,635, Abcam), CA-XII (1:10,000, rabbit Anti-CA12 antibody [EPR14861]—C-terminal, ab195233), CA-II (1:1000, rabbit Anti-Carbonic Anhydrase II antibody
Western blots are available in Supplementary Studio Version 2.1 (LI-COR). Uncropped versions of LIDYCOR Odyssey Blot Imager and quantified using Image anti-rabbit IgG (H+L), IRDye 800RD goat anti-mouse IgG (H+L), and IRDye 800CW donkey anti-rabbit IgG (H+L). Membranes were imaged on LIDYCOR Odyssey Blot Imager and quantified using Image Studio Version 2.1 (LI-COR). Uncropped versions of western blots are available in Supplementary.

**Immunocytochemistry**

Cells were grown on glass coverslips and fixed in 4% paraformaldehyde (Sigma-Aldrich) for 10 min at room temperature. Cells were blocked in 5% BSA for 1 h at room temperature. Cells were stained with the PMA1 antibody (1:100; SC-33735, Santa Cruz Biotechnology) or CA-IX antibody for 2 h (1:500, ab184630, Abcam) and washed in PBS. Cells were further incubated for 1 h with secondary anti-rabbit-Alexa Fluor 594 antibody (1:2000; A11072, Invitrogen) or anti-mouse-Alexa Fluor 594 (1:2000; A1105, Invitrogen) and additionally incubated in WGA, cell membrane marker (W6748, Invitrogen) for 10 min on ice. The cells were mounted for fluorescence with DAPI (H-1200, Vector). The slides were viewed by Leica inverted SP5 AOBS confocal microscope, and micrographs were taken, and images were subsequently acquired in the Moffitt Analytic Microscopy Core Facility by using dual photomultiplier tube detectors and LAS AF software (Leica Microsystems). For detection of intracellular PMA1, cells were fixed and permeabilized with 1:1 mixture of methanol and acetone, and immunostained with PMA1 antibody for 1 h, followed by 1 h of incubation with the secondary anti-rabbit Alexa488 antibody (Molecular Probes, Invitrogen). The cells were mounted and viewed by fluorescence microscopy.

**Proliferation rate assay**

Cells were cultured in a 24-well plate under standard cell culture conditions for 24, 48, 72, and 96 h, and the cell number and viability were determined with a trypan blue dye by using the Countess automated cell counter (Invitrogen).

**Glucose consumption and lactate production assays**

In Fig. 1E and F, cells were seeded in a 96-well plate in the growth media containing 10% FBS. Once cells reached 90% confluence, the growth media were removed, and the cells were washed twice in PBS and media was replaced for 24 h. The media were collected from 24 h incubation for both glucose consumption and lactate production assays. The cells were then trypsinized and the cell densities were determined. Glucose quantification was conducted using glucose bioluminescent assay kit (Glucose Glo Assay, Promega) as described per manufacturer instruction. The lactate assay kit (Lactate Assay kit, Sigma-Aldrich) was used to measure L (+)-Lactate in the culture media according to the manufacturer's instructions. Data were normalized by protein concentration of cells.

In Fig. 5E and F, cells were seeded in a 6-well plate in growth media containing 10% FBS. Once cells reached 90% confluence, the growth media were removed, and the cells were washed twice in PBS and incubated in serum-free and phenol-red free media for 24 h. The media were collected from 24 h incubation for both glucose consumption and lactate production assays. The cells were trypsinized and the cell densities were determined. Glucose quantification was conducted
using glucose colorimetric/fluorometric assay kit (BioVision, K606-100) as described per manufacturer instruction. The lactate assay kit II (BioVision, K627-100) was used to measure L (+)-Lactate in the culture media according to the manufacturer’s instructions. Data were normalized by cell density per well and were reported as lactate production and glucose consumption as pmol per cell.

Glucose uptake radioactive assay

Cells were seeded in 24-well plates to 80% confluence. Cells were incubated for 1 h with 1 μCi of deoxy-D-glucose, 2-[1,2-3H(N)] (NET549250UC, Perkin Elmer, MA) at 37°C. Cells were washed 2× in DPBS and lysed with 300 μl of NaOH, and cell extract was added to a vial with 6 ml of HPLC grade water relative to protein concentration. Data were analyzed using Metaboanalyst online software, there was no data filtering, and data were normalized to protein concentration.

YSI 2950D biochemical analysis of lactate

Cells were seeded in a 96-well plate in growth media containing 10% FBS. Once cells reached 90% confluence, the growth media were removed, and the cells were washed twice in PBS and media was replaced for 24 or 48 h. The media were collected from 24 and 48 h incubation for lactate production measurement. The cell densities per well were determined by Celligo imaging cytometer bright-field cell count application. Lactate quantification was measured by the YSI. Data were normalized by relative cell number and were reported as lactate production in mmol/L/10 K cells (Additional file 1: Fig. S5&S6) and mg/L/10 K cells (Additional file 1: Fig. S2).

Untargeted metabolomics

Samples were prepared according to Beth Israel Deaconess Medical Centre Mass Spectrometry Core and run on a Thermo QEExactive Plus/HS Orbitrap LC–MS/MS. Briefly, cells were grown to 80% confluence in 10-cm2 dishes. Cells were changed into fresh media 2 h prior to collection, media was aspirated, cells were washed in ice-cold PBS, and 1 ml of 80% methanol (−80°C) added to plate on dry ice, then transferred to −80°C freezer for 15 min. The cell plate was scraped on dry ice and contents collected. Sample was spun in cold centrifuge at max speed for 20 min and supernatant removed. Supernatant was dried in speed vac for 5 h then stored at −80°C. Before mass spectrometry analysis, samples were resuspended in HPLC grade water relative to protein concentration. Data were analyzed using Metaboanalyst online software, there was no data filtering, and data were normalized by sum of all metabolites per sample, log transformed and underwent Pareto scaling.

Intracellular pH (pHi)

Solutions and media

(i) Solutions for Seahorse experiments were as follows: 2 mM HEPES, 2 mM MES, 5.3 mM KCl, 5.6 mM NaPhosphate, 11 mM glucose, 133 mM NaCl, 0.4 mM MgCl₂, 0.42 mM CaCl₂, titrated to given pH with NaOH. For reduced Cl⁻ experiments, 133 mM NaCl was replaced with 133 mM NaGluconate and MgCl₂ and CaCl₂ were raised to 0.74 mM and 1.46 mM, respectively, to account for gluconate-divalent binding. Calibration solutions for nigericin were as follows: 145 mM KCl, 1 mM MgCl₂, 0.5 mM EGTA, 10 mM HEPES, 10 mM MES and adjusted with NaOH to required pH. pH e media to measure pH were as follows: Solution A: 125 mM NaCl, 4.5 mM KCl, 1 mM CaCl₂, 1 mM MgCl₂, 11 mM glucose base media. Split solution A were divided into two parts—Solution B: 22 mM HCO₃⁻ and Solution C: 22 mM NaCl. Mix B&C were as follows: 40 ml B and 0 ml C = 22 mM HCO₃⁻ (pH 7.41), 20 ml B and 20 ml C = 11 mM HCO₃⁻ (pH 7.11), 10 ml B and 30 ml C = 5.5 mM HCO₃⁻ (pH 6.81), and 4 ml B and 36 ml C = 2.2 mM (pH 6.41).

Fluorescent labeling and image analysis

Cells were loaded with 10 μM cSNARF1 and 2.7 μM Hoechst 33,342 for 10 min at 37°C. Cells were washed in neutral pH media and imaged in various pH media (6.4–7.4) on Leica SP5 Confocal Microscopy × 40 objective. Cytoplasmic pH was measured by gating pixels according to a threshold level of Hoechst signal within cSNARF1-positive pixels. Fluorescence at 580 and 640 nm was averaged, background offset, and ratioed for each particle representing a cell.

Cell invasion and migration assay in vitro

In vitro cell motility and invasiveness were measured by methods as previously reported with some modifications [31]. The motility change was measured by the circular wound healing assay using OrisTM Cell Migration Assay Kit (Platypus, CMAU101). Cells were plated on a 96-well plate at 1 × 10⁶ cells/ml while a cell seeding stopper masked the circular area at the center of each well. The cell seeding stoppers were removed 24 h after the plating and cells were cultured a further 30 h to monitor the closing of the cell-free area (wound area). The area covered by live cells was measured by labeling cells with Calcein-AM (Life Technology, C3099) and analyzing fluorescent microscopic images (×2.5) by ImageJ (NIH). The smaller wound size represents the higher motility.

Wound healing scratch assay was used to measure motility into CA-IX clones. Ninety-six-well plates were seeded with 1 × 10⁶ cells/ml per well in 10% FBS and 1% PenStrep RPMI-1640 and incubated overnight. The plate was uniformly scratched with Essen Bioscience 96
Woundmaker, media was removed, and the plate washed with DPBS and then 200 µl of media was added to each well. The wound was imaged on Celigo Imaging Cytometer, and the number of cells that migrated into the wound was quantified by direct bright-field cell counting at 0, 24, and 48 h.

Cell invasiveness was measured by monitoring cells escaping from Matrigel (Becton Dickenson, 356,231). Cells were suspended in 50% Matrigel in serum-free RPMI-1640 at 1 × 10⁷ cells/ml. A Matrigel droplet (volume = 5 µl) was placed on each well of a 24-well plate. The Matrigel was solidified by incubating at 37 °C overnight and then FBS-containing normal growth media was added. The cells escaped from the Matrigel droplet were monitored in real-time by using the IncuCyte ZOOM system (Essen BioScience) or Celigo Imaging Cytometer (Nexcelom Bioscience, MA). After 7 days of culture, the cell expansion from the droplets was quantified by Celigo single colony verification algorithm or ImageJ after fixing cells in 3.7% formaldehyde and staining in crystal violet solution. The larger area occupied by cells represents the higher invasion potential.

**Normalization of invasion and migration assays**

The results of invasion and migration assays were normalized by the proliferation rates of the cells. Proliferation rates were calculated by a linear fit of cell growth, 48 h after seeding the cells (Additional file 1: Fig. S20) and weighted by the standards error. The normalization was carried out in order to eliminate inherent differences between clones making it possible to compare them. In the case of the invasion assay, final growth was divided by the proliferation rate of each of the cells. For the migration assay (i.e., wound healing assay), the relative healed area was divided by its corresponding growth rate and then multiplied by the growth rate of the MOCK-1 cells. This allowed comparison between the normalized healed areas of each clone, to one of the MOCK-1 cells. We used the Matlab R2012a curve fitting toolbox (The MathWorks, MA). Matlab code is available in the Supplementary materials.

**SCID mice**

Six-week-old female SCID Beige mice were purchased from Charles River Laboratories. Mice were given a week to acclimate to the animal facility before they were studied. To minimize the risk of any exogenous infection, the SCID mice were maintained and cared for in a sterile, static micro-isolation cage. Mice received irradiated food (Harlan Laboratories) and sterile water ad libitum. All animal experiments were performed under a protocol approved by the University of South Florida Institutional Animal Care and Use Committee.

**Metastasis assays in vivo**

Since MCF-7 cells are estrogen-dependent for tumor formation, estrogen pellets, 17 β estradiol, 0.72 mg/pellet, 60-day release (Innovative Research of America, SE-121) were subcutaneously implanted in the shoulder region of the mice 2 days prior to tumor inoculation. For the primary tumor growth study with MCF-7/MOCK-2 or MCF-7/CA-IX M6 cells, mice were given 200 nM 17 β estradiol in drinking water to try and prevent adverse side effects of pellet use, including balder stones and urinary tract infections.

For the experimental metastasis, SCID mice were injected through tail veins with 1 × 10⁶ cells in 200 µl of PBS solution (either MCF-7/MOCK-1, MCF-7/PMA1-C1, MCF-7/MOCK-2, MCF-7/CA-IX M1, or MCF-7/CA-IX M6 cells). Three months after injection of the cancer cells, the mice were euthanized, and the lung tissues were surgically excised, fixed, and stained with hematoxylin and eosin (HE). Lung sections (at least three histologic sections for each lung specimen) were examined for metastatic nodules under a light microscope by a breast cancer pathologist (A.L. or M.M.B) who was blinded to identifiers.

For the spontaneous metastasis studies, approximately 10 × 10⁶ cells (either MCF-7/MOCK-1 and MCF-7/PMA1-C1) in 100 µl of PBS + 100 µl of Matrigel were injected into the mammary fat pads of mice. Once tumors reached approximately 400 mm³, or 6 weeks post-cell injection, the tumors were resected, fixed, and stained with H&E, or PMA1 antibody. Three months after resection, the mice were sacrificed, and lung sections were examined for lung metastases. The tumors were measured twice every week throughout the study with a digital caliper and volume values were calculated with the formula $V = \frac{(\text{Length} \times \text{Width}^2)}{2}$. The body weights were monitored twice a week throughout the study.

**Treatment model**

Female SCID Beige mice received 200 mmol/L of sodium bicarbonate water 2 days prior to tail vein injection for experimental metastasis study. Mice continued receiving bicarbonate water until the end of the experiment. Control mice received regular tap water. Since MCF-7 cells are estrogen-dependent for tumor formation, estrogen pellets, 17 β estradiol, 0.36 mg/pellet, and 90-day release (Innovative Research of America, SE-121) were subcutaneously implanted in the shoulder region of the mice 2 days prior to tumor inoculation. For the experimental metastasis, SCID mice were injected through tail veins with 1 × 10⁶ cells in 200 µl of PBS solution (MCF-7/MOCK-2 and MCF-7/CA-IX M6 cells). Once MCF-7/CA-IX M6 control group had observable lung metastasis
by T2 MRI (~74 days), mice were humanely euthanized, and lungs and kidney collected.

**Histology**

The tissues were harvested, fixed in 10% neutral buffered formalin. Tissue sections (4 µm) were prepared and stained with H&E in the Moffitt Cancer Center Tissue Core. The histological slides of resected primary breast tumor xenografts from MCF-7/MOCK-1 or MCF-7/PMA1-C1 groups, and MCF-7/MOCK-2, MCF-7/CA-IX M1, and MCF-7/CA-IX M6 groups were blindly examined under a light microscope by a pathologist (A.S.L. or M.M.B) for tumor grades using the most common grading system: G1: well differentiated (low grade); G2: moderately differentiated (intermediate grade) G3: poorly differentiated (high grade); G4: undifferentiated (very high grade), as assessed according to histological features of stromal hypercellularity, atypia, stromal mitotic activity, presence of stromal over-growth and mitosis, necrosis, spindle cell differential, and chromatim activity. Tumor burden in the lung was measured by Aperio ImageScope (Leica Biosystems, IL) and calculated as % of total lung tissue and compared between groups. Tumors were drawn around by hand using the Aperio software and confirmed by a pathologist (M.M.B); % area of tumor in lungs was then calculated by comparing the area of total lung tissue to the area of the tumor within lungs. For all lung metastases IHC, three different sections were taken from each lung and analyzed, with 5–6 sections taken between each analyzed slice to ensure entirety of lungs, and metastatic burden was analyzed.

**Immunohistochemical (IHC) staining**

The cross-sections were stained with various antibodies as per normal laboratory protocol in the Moffitt Tissue Core Histology Facility. Positive and negative controls were used for each antibody staining, and staining condition was optimized for each antibody. The antibodies were utilized in this study as follows: rabbit anti-*Saccharomyces cerevisiae* PMA1 (sc-33735, Santa Cruz Biotechnology); rabbit anti-human CA-IX (ab15086, Abcam, Cambridge, MA); rabbit anti-human MCT1 (sc-50324, Santa Cruz, CA); rabbit anti-human GLUT1 (ab15309, Abcam, Cambridge, MA); rabbit anti-human NHE1 (sc-28758, Santa Cruz, CA); rabbit anti-human ER (RM9101, Thermo Fisher Scientific, CA); rabbit anti-human MCF7 cells. Imaged using Celigo, n=1,862 cells, average ± SD, p<0.0001****.

**RNA analyses in formalin-fixed, paraffin-embedded (FFPE) tissue**

FFPE tissue samples were cut in 10-µm-thick sections on a microtome and deparaffinized by deparaffinization solution (Qiagen, 19,093). Total RNA was extracted from deparaffinized FFPE sections with the miRNeasy FFPE kit (Qiagen, 217,504) following the manufacturer’s protocol. Real-time qRT-PCR analyses for PMA1 mRNA were described above in the qRT-PCR section. Experimental Ct values from PMA1 amplification were normalized with GAPDH Ct values and were expressed relative to MCF-7/MOCK-1 control Ct values.

**Statistical analyses**

A two-tailed unpaired Student t test or Welch’s T-test was employed to determine statistical significance. Ordinary one-way ANOVA with Geisser Greenhouse correction and Tukey’s multiple comparison test, with a single pooled variance. A p-value of less than 0.05 was considered statistically significant or otherwise indicated. Kaplan-Meier curve was used to analyze overall survival in mouse models with log-rank test curve comparison.

**Supplementary Information**

The online version contains supplementary material available at https://doi.org/10.1186/s12915-022-01340-0.
Spheroid formation assay to measure ability of cells to grow in 3D, an indicator of metastatic ability in vitro. Imaged using Celigo with single colony verification analysis after 5 days of growth in hanging drop plates. Representative images shown. Figure S13. Effect of CA-IX expression on primary tumor growth of MCF-7 cells. n=12–15 mice per group. Figure S14. qRT PCR of relative PMA1 gene expression in PMA1 transfected clones compared to Mock (empty vector clone). Average ± SD. (Ordinary one-way ANOVA, p<0.05***). Figure S15. Characterization of PMA1-expressing cells by ICC staining (non-permeabilized). Figure S16. ICC of permeabilized cells staining for PMA1. Figure S17. Glycolytic reserve measured by glycolysis stress test with XFe96 Seahorse assay. Reserve is determined by difference in glucose induced proton production rate (PPR) and addition of oligomycin which inhibits mitochondrial energy production. Average ± SD. (Ordinary one-way ANOVA, p<0.0001****). Figure S18. Raw data of gel escape assay and corresponding images, prior to normalization by growth rate. N=4 biological replicates, average ± SD. (Ordinary one-way ANOVA, p<0.001***). Figure S19. Raw data of circular wound healing assay and corresponding images, prior to normalization by growth rate. N=4 biological replicates, average ± SD. (Ordinary one-way ANOVA, p<0.001***). Figure S20. Cellular growth rate was determined by number of cells over time and calculated by a linear fit of cell growth for normalization of invasion and migration assays. Figure S21. Relative PMA1 gene expression in lung micrometastases. Average relative gene expression ± SD, unpaired t-test p<0.001. Figure S22. Quantification of GLUT1 protein staining in IHC samples of resected primary tumors and representative images. N=9–10, Average ± SD, unpaired t-test n.s. Figure S23. Quantification of NHE1 protein staining in IHC samples of resected primary tumors and representative images, n=8–9 Average ± SD, unpaired t-test n.s.

Additional file 2.

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Authors’ contributions

All authors read and approved the final manuscript. SR, LX, YK, DV, MCL, RJG, and JW conceived and designed the study. LX, YK, DV, ASL, JJ, MCL, JW, PS, RJG, and SR developed the methodology. SR, LX, RJG, and PS wrote the manuscript. SR, LX, YK, DA, BO, JJ, TE, ER, PS, DV, and JW performed imaging, in vitro and in vivo experiments and acquisition of data. SR, LX, YK, DA, ASL, ER, BO, MMB, JJ, TE, MCL, PS, DV, RJG, and JW analyzed and interpreted these data. SR, LX, YK, ASL, TE, JJ, JW, MMB, ER, PS, DV, BO, and RJG generated figures and reviewed the manuscript. LX, JW, RJG, and SR supervised the study.

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Availability of data and materials

The authors declare that the data supporting the findings of this study are available within the paper and its Additional files. The raw data are deposited online in the figshare data repository, https://doi.org/10.6084/m9.figsh are19729732v2. Any other data can be requested from the corresponding author.

Declarations

Ethics approval and consent to participate

All animal experiments were performed under a protocol approved by the University of South Florida Institutional Animal Care and Use Committee.

Consent for publication

Not applicable.

Competing interests

RJG is supported by a grant from Helix, Biopharma. The other authors declare that they have no competing interests.

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