On activation of PDH, however, pyruvate can be decarboxylated to acetyl-coenzyme A, enter the Krebs cycle, and complete glucose oxidation in the mitochondrial matrix, generating up to 36 moles of ATP per glucose molecule in the presence of oxygen. Glucose oxidation does not take place when pyruvate does not enter the mitochondria (for example, in diseased mitochondria or if PDH is inhibited) or in the absence of oxygen.

Warburg (6) first showed that the metabolism of cancer cells, even under normoxia, is characterized by an increase in the ratio of cytoplasmic glycolysis to mitochondrial glucose oxidation. Although the mechanism of this “Warburg effect” is unknown, and whether it is etiologically related to carcinogenesis remains unproven (7), there is increasing interest in metabolism as a target for cancer therapies (8–11). The energetic switch from mitochondrial glucose oxidation to cytoplasmic glycolysis may offer a proliferative advantage to cancer cells (11). For example, most glycolytic enzymes also have direct anti-apoptotic actions (12); lactic acid promotes angiogenesis and interstitial matrix breakdown, facilitating metastasis (11); and decreased mitochondrial function is associated with inhibition of mitochondria-dependent apoptosis (3). GBM has a strong glycolytic phenotype, and a number of the molecular abnormalities that occur in GBM are known to suppress mitochondrial glucose oxidation and promote cytoplasmic glycolysis (1), including the activation of the phosphatidylinositol 3-kinase–Akt or myc pathways or suppression of the p53 pathway (9, 10).

The mitochondria of cancer cells are hyperpolarized, with respect to those of noncancer cells (3, 13), a condition associated with suppressed mitochondrial function. Although controversial [reviewed in (14)], the efflux of prosapoptotic mediators through the mitochondrial transition pore (MTP) depends in part on mitochondrial membrane potential (ΔΨm), and thus, mitochondrial hyperpolarization may mark an apoptosis resistance state (3, 15). We have shown that this state can be reversed in cancer cells by DCA, which by inhibiting PDK promotes
pyruvate entry into the mitochondria, reversing the increase in glycolysis to glucose oxidation ratio, improving mitochondrial function, and reversing mitochondrial hyperpolarization (3). DCA therefore decreases tumor growth in vitro and in vivo, without affecting noncancer mitochondria and tissues (3, 16–20). The increase in mitochondrial respiration is associated with an increase in production of mitochondrial reactive oxygen species (mROS), predominantly superoxide. Superoxide can be dismutated to H$_2$O$_2$, a relatively stable mROS that can reach other cellular structures beyond the mitochondria. For example, H$_2$O$_2$ can activate redox-sensitive voltage-dependent potassium channels in the plasma membrane and, at least in some tissues, promote a decrease in intracellular calcium (3, 4). Other redox-sensitive targets may include p53, which is activated when oxidized (21, 22). The p53 axis is inhibited in GBM, contributing to the increased proliferative state of GBM cells (1). p53 also represses hypoxia-inducible factor–1α (HIF-1α)–stimulated transcription because p53 and HIF-1α compete for the same cotranscription factor (23, 24). HIF-1α increases the expression of glucose transporters and several glycolytic enzymes as well as PDK, thus sustaining the glycolytic phenotype (25, 26). In addition, HIF-1α increases the expression of vascular endothelial growth factor (VEGF), enhancing angiogenesis. Angiogenesis may also be enhanced by normoxic HIF-1α activation. Because mitochondria are important oxygen sensors (27), inhibited mitochondria may transmit pseudohypoxic redox signals and activate HIF-1α even during normoxia (28–30). In addition, a decrease in α-ketoglutarate, a direct product of the Krebs’ cycle, may also promote HIF activation because it is a cofactor for the prolyl hydroxylation reaction that degrades HIF-1α (30).

We hypothesized that orally administered DCA, which crosses the blood-brain barrier, would decrease GBM growth in vivo. We further suggested that this could occur by (i) reversing the glycolytic phenotype and normalizing ΔΨm, which would promote mitochondria-dependent apoptosis; (ii) increasing mROS and promoting p53 activation; and (iii) increasing α-ketoglutarate concentrations. The last two effects would lead to inhibition of HIF-1α, a decrease in VEGF, and inhibition of angiogenesis.

**RESULTS**

Effects of DCA on mitochondria from 49 freshly isolated GBM tumors

To determine whether human GBM could be a target for metabolic therapy with DCA, we studied 49 freshly excised consecutive primary GBMs (60% male, 48 ± 11 years). In addition to the clinical and neuropathology reports, we confirmed GBM identity with Fig. 1.
immunohistochemistry, which showed expression of glial fibrillary acid protein (GFAP) but no βIII-tubulin or oligodendrocyte markers (fig. S1). ΔPm was increased in the freshly isolated GBMs compared to noncancer brain tissues obtained in epilepsy surgery (n = 3) (Fig. 1A). DCA, but not vehicle (normal saline), caused mitochondrial depolarization in GBM but not in normal brain tissue. DCA also increased GBM mROS (Fig. 1A). This suggested that the metabolic and mitochondrial remodeling in GBM is partially reversible and that this remodeling is at least in part regulated by PDK. The response to DCA is consistent with a higher concentration of PDKII [the most ubiquitously expressed isoform and the one with the lowest $K_i$ for DCA (31)] in GBM than in noncancer brain tissue, as seen with immunohistochemistry and immunoblots (Fig. 1, B and C). Cells exhibiting the highest PDKII concentrations also contained proliferating cell nuclear antigen (PCNA), suggesting that these cells were proliferating (Fig. 1B). These data, collected over a 2-year period, strengthened the rationale for subsequently administering DCA to patients with GBM (4).

**Clinical effects of DCA on five patients with GBM**

We then treated with DCA five consecutive patients with primary GBM, referred from our brain cancer program and from whom tissue was available from the last debulking surgery. Three patients (patients 1 to 3) had recurrent GBM with disease progression after several chemotherapies (in addition to the standard treatment with surgery, RT, and TMZ) and were considered appropriate for palliative therapy. Two additional patients (patients 4 and 5) were newly diagnosed, and after the initial debulking surgery, DCA was administered in addition to the standard treatment of RT and TMZ. In patient 4, a 3-month pretreatment with DCA was followed by the addition of RT and TMZ, whereas in patient 5 DCA was initiated simultaneously with RT and TMZ, after debulking surgery. If the patients required reoperation or autopsy, tissue from the last debulking surgery (before DCA administration) was compared to the post-DCA treatment tissue. Their clinical information is summarized in table S1. DCA has been administered to patients for >30 years, mainly in the treatment of inborn errors of mitochondrial metabolism, and pharmacokinetic and pharmacodynamic data are available (5, 32–34). We treated patients with a starting dose of 12.5 mg/kg orally twice a day for 1 month, at which point the dose was increased to 25 mg/kg orally twice a day. We then followed a dose de-escalation protocol, decreasing the dose by 50% when dose-limiting toxicity occurred. The patients were followed clinically for up to 15 months. None of the patients had hematologic, hepatic, renal, or cardiac toxicity (table S1). Peripheral neuropathy was the only apparent toxicity. Patients had variable dose-dependent degrees of peripheral

![Fig. 1.](image1.png)

**Fig. 2.** In vivo effects of DCA in patients with GBM. (A) T1 gadolinium–enhanced axial MRI images (left, midventricular level; right, supraventricular level) and merged positron emission tomography (PET)–MRI images taken before and after treatment with DCA for patients 1 (left) and 2 (right). After 15 months of treatment with oral DCA as the only therapeutic agent, apparent resolution of the tumor is seen in patient 2. In patient 1, the metastatic paraventricular tumor mass regressed after 9 months of therapy with DCA. Baseline, month 3 of DCA therapy; +9 months, month 12 of DCA therapy. In patient 1, the primary tumor site (which is not seen at the level of these images but can be seen in fig. S2) remained unchanged in this interval. (B) Representative micrographs from tissue taken from patient 3 (see text for clinical details) and summary data (patients 2 to 4), quantifying tumor proliferation (% PCNA-positive cells) and apoptosis (% TUNEL-positive cells) before and after chronic DCA therapy. There is a decrease in the number of cells [shown by the number of nuclei, in blue (DAPI)], a significant decrease in the expression of PCNA, and a significant increase in apoptosis after treatment with DCA. Percent of PCNA- or TUNEL-positive cells were measured blindly in eight random fields per slide; a minimum of three slides per experiment was used (n = ~350 to 400 cells per patient). *P < 0.01. (C) PDH activity is significantly increased in GBM tissues from patients treated with DCA, compared to the baseline tissues from the same patients, taken before DCA treatment. This suggests effective inhibition of PDK within the tumor tissue in vivo (n = 3 patients). *P < 0.001.
neuropathy, which was reversible, confirming previous studies (35–37). When the dose was decreased to 6.25 mg/kg orally twice a day, none of the patients had clinically significant peripheral neuropathy (table S1). Initially, the half-life of DCA is <1 hour. DCA inhibits its own metabolism and serum concentrations increase, eventually reaching a plateau (34). The plasma trough concentrations of DCA in our patients remained undetectable for the first 2 to 3 months but thereafter reached therapeutic concentrations. At a dose of 6.25 mg/kg orally twice a day for at least 3 months, trough DCA concentrations were 0.44 ± 0.16 mM (mean ± SD; n = 4) (table S1). These values are similar to those seen in chronic DCA treatment of adults with mitochondrial defects (34) and are in the same range as the $K_i$ of DCA for PDKII (0.2 mM) (31). Patients 1, 4, and 5 showed some evidence of radiologic regression on magnetic resonance imaging (MRI) (Fig. 2A and figs. S2 to S4). Patient 3 had a very large tumor with brain edema at baseline (fig. S5), despite being on high steroid doses, and a low Karnofsky score and continued to deteriorate. He died from brain edema complications 3 months after initiation of DCA therapy. Patient 2 required drainage of a cyst and debulking in month 11 of DCA therapy. Patient 4 showed radiologic progression on month 3 of DCA therapy, at which point further debulking was performed and RT plus TMZ was given in addition to DCA. All, except patient 3, were clinically stable at month 15 of DCA therapy and alive at month 18 (telephone follow-up). Further clinical details are described in the Supplementary Material.

**Effects of DCA on GBM tumors in vivo, primary GBM cell lines, and putative GBM-SC derived from the DCA-treated patients**

We conducted experiments on tissues derived from these five patients and were able to make comparisons in tissues before and after DCA treatment in patients 2 to 4; we only had “before” tissues in patients 1 and 5. Compared to pre-DCA tissue, post-DCA GBM tissue in all three patients showed decreased number of cells per unit volume, decreased proliferation, and increased apoptosis (Fig. 2B), as well as increased tissue enzymatic activity of PDH, suggesting effective inhibition of PDK in vivo (Fig. 2C). Putative GBM cancer stem cells (GBM-SCs) may be responsible for posttreatment resistance and recurrence of GBM (38–43). These cells are characterized as CD133+/nestin+ GBM-SC and form niches around capillaries (41). In such vascular GBM-SC units, GBM-SC can induce angiogenesis, whereas their molecular stem cell phenotype is maintained by their accessibility to circulating growth factors (44). GBM-SC proliferation is associated with particularly poor clinical outcome (42). CD133+/nestin+ GBM-SC expressed PCNA in vivo in all pre-DCA tumors, indicating that they are dividing, but the percentage of CD133+/nestin+ cells that expressed PCNA was significantly decreased after DCA therapy in patients 2 to 4 (Fig. 3A). Simultaneous staining with a CD133 antibody and tetramethyl rhodamine methyl ester (TMRM) showed that CD133+ cells had the highest $\Delta\Psi_m$ compared to neighboring non–GBM-SC in vivo (fig. S6). In tumor-derived primary cell lines, ~10% of cells expressed both CD133 and nestin, whereas >90% of the cells expressed the mature marker GFAP (but not βIII-tubulin or oligodendrocyte) (fig. S7), similar to the histopathology of GBM (fig. S1). We isolated putative GBM-SC from GBM tumors and cultured them with the appropriate growth factors (human fibroblast growth factor, 20 ng/ml; human epidermal growth factor, 20 ng/ml). These cells had a very high expression of both CD133 and nestin, had very low expression of mature glial markers (fig. S7), and formed characteristic neurospheres (Fig. 4 and fig. S7), an independent predictor of poor clinical outcome (43). We measured $\Delta\Psi_m$ in freshly excised tumors, in primary cell lines, and in GBM-SC isolated from those tumors as well as in differentiated cells derived from GBM-SC (Fig. 3B). The highest potential was found in
the putative GBM-SC. Both the primary and the GBM-SC–derived secondary GBM cells (15-day differentiation) had mitochondrial potentials similar to that of the parent tumors. DCA (0.5 mM for 24 hours) decreased the potential in all groups of cells. Although the cause of the increased ΔΨm in cancer (3, 13) remains to be fully defined, it has been proposed to be caused in part by a translocation of hexokinase II (HXKII), a key glycolytic enzyme, from the cytoplasm to the outer mitochondrial membrane (45, 46). There, HXKII may bind to and inhibit the voltage-dependent anion channel (a component of the MTP), increasing the ΔΨm and the apoptotic threshold. Inhibition of this translocation decreases cancer ΔΨm and reverses the resistance to apoptosis (45, 46). Our primary cell lines generated from pre-DCA tumors showed a sustained mitochondrial translocation of HXKII, potentially explaining the increase in ΔΨm. HXKII translocation was not present in primary cell lines from tumors after DCA treatment (fig. S8), compatible with the notion that DCA induced suppression of glycolysis and decreased ΔΨm. As in the tumors, PKDII was present at high concentrations in the GBM cell lines generated from patients 2 to 4, although the other known isoenzymes were also expressed (fig. S9A). When GBM-SCs were allowed to differentiate into secondary GBM cell lines, the proportion of cells with GBM-SC markers decreased to a value similar to that of the primary cell lines (~10%). When allowed to differentiate in the presence of DCA (0.5 mM), however, the proportion of cells with GBM-SC markers was decreased even further to ~5% (fig. S7). Indeed, DCA induced apoptosis in GBM-SC in vitro (Fig. 4 and fig. S9B) as well as in GBM primary cell lines (fig. S9C). Apoptosis was further increased in GBM-SCs by the combination of DCA plus TMZ (Fig. 4 and fig. S9B), providing a rationale for combination therapy. GBM-SC apoptosis also took place in vivo in the post-DCA treatment tumors, shown by the colocalization of nestin, CD133, and terminal deoxynucleotidyl transferase–mediated deoxyuridine triphosphate nick end labeling (TUNEL) staining (Fig. 4 and fig. S9D).

**Effects of DCA on the GBM-SC microvessel unit and angiogenesis in vivo and in vitro**

In the untreated, pre-DCA tissues, niches of GBM-SC were found around microvessel beds [von Willebrand factor (vWF) staining], as reported (41). This GBM-SC microvessel unit (44) was destroyed by DCA treatment because, in addition to GBM-SC, apoptosis was also increased in microvascular endothelial cells (Fig. 4 and fig. S9D), suggesting a potential inhibition of angiogenesis. Indeed, decreased vWF staining in post-DCA treatment tumors suggested that there was decreased vascularity (Fig. 5A). HIF-1α was highly expressed or activated (nuclear localization) in the pre-DCA and inhibited in the post-DCA tumor tissues (Fig. 5B). Post-DCA treatment tumors from patients 2 to 4 showed a significant increase in mROS in vivo (superoxide measured by mitoSOX) compared to the pre-DCA tumors from the same patients (Fig. 5C). In the pre-DCA tumors, acute DCA increased mROS to values seen in the post-DCA treatment tumors. In contrast, in the post-DCA tumors, acute DCA only minimally increased mROS, suggesting an almost maximal effect in vivo. DCA increased mROS in GBM-SC as well (Fig. 5C). Low ROS concentrations in cancer stem cells may reflect resistance to apoptosis, and therapies that increase cancer stem cell ROS are suggested to be more effective (47). Although we studied mROS (mitochondrial superoxide), controversy exists whether HIF-1α activation in cancer is associated with an overall increase or decrease in ROS [reviewed in (28)]. Using a different technique, we also measured whole-cell H2O2. DCA increased GBM cell H2O2 in a dose-dependent manner (Fig. 6). In addition, DCA also increased intracellular α-ketoglutarate concentrations in a dose-dependent manner (Fig. 6). This is compatible with the increase in glucose oxidation that follows PDH activation (3) because α-ketoglutarate is a product of the Krebs’ cycle. The Krebs’ cycle produces the electron donors that feed into the electron transport chain during respiration. Accordingly, DCA increased respiration rates by 44 ± 4% in GBM cells (mean ± SEM; n = 3; P < 0.05), supporting an overall increase in mitochondrial activity.

The increase in H2O2 and α-ketoglutarate

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**Fig. 4.** DCA induces apoptosis in both putative GBM-SC and microvascular endothelial cells. (Left) Apoptosis, as measured by the TUNEL assay, in response to DCA (0.5 mM for 72 hours) alone and DCA in combination with TMZ (100 μM) is shown in GBM-SC in vitro. (Right) Apoptosis in GBM-SC is shown in tissue, from tumors both before and after therapy with DCA. Representative examples are shown; summary mean data are shown in fig. S9, B and D. The formation of neurospheres in GBM-SC growing in vitro is apparent. Simultaneous quadruple staining in tumor tissue shows that the putative GBM-SCs express high concentrations of PKDII. In addition, they form niches around capillary networks, shown by staining with the endothelial marker vWF. There is no TUNEL staining in the pre-DCA treatment tissues, whereas in the post-DCA tissues there is an increase in TUNEL staining in both GBM-SC and endothelial cells.
can explain the decrease in HIF-1α activity (Fig. 5B), which is also confirmed by the dose-dependent decrease in VEGF production by GBM cells (Fig. 6). These data are consistent with the decrease in the angiogenesis in vivo (Figs. 4 and 5) and suggest that GBM cells signal to endothelial cells in a paracrine manner. To study whether DCA can directly suppress angiogenesis, we used the standard technique of human endothelial cell tube formation in Matrigel. Under physiologic moderate hypoxia, DCA caused a dose-dependent direct inhibition of angiogenesis in vitro (Fig. S10). The post-DCA treatment tumors from patients 2 to 4 showed increased activity of the mROS-sensitive p53 (nuclear translocation), also confirmed by the increased activity and abundance of its downstream target p21 (fig. S11). These effects on p53 or p21 can also explain the decrease in HIF-driven transcription and are consistent with the antiproliferative, in addition to the proapoptotic, effects of DCA in GBM (fig. S12).

**DISCUSSION**

The metabolic modulator DCA exerts anticancer effects in cultured cells and rodents (3, 16–20). We have now shown that DCA can be used in patients suffering from GBM. DCA treatment was associated in some GBM patients with prolonged radiologic stabilization or tumor regression and, in general, displayed an overall good safety profile. This early, first-in-human report provides a rationale for extended studies with this generic small molecule in patients with GBM. Our results indicate that GBM is a good candidate for metabolic intervention. The target of DCA, PDKII, is highly expressed in GBM tumors and cell lines, and DCA can inhibit its activity in vivo. GBM is characterized by mitochondrial hyperpolarization, in keeping with the metabolic remodeling (Warburg effect) and the related apoptosis resistance that characterize GBM and most solid tumors (11). Mutations in the genes for cytoplasmic and mitochondrial isocitrate dehydrogenases have been described in GBMs arising from lower-grade gliomas (secondary GBMs) (48), but the mechanism by which these mutations relate to carcinogenesis remains unclear (49, 50). Our patients had primary GBMs and the mitochondrial remodeling was at least partially reversible with DCA, suggesting that it was not due to irreversible dysfunction. Furthermore, we show that putative GBM-SC may undergo the same metabolic and mitochondrial remodeling, but to an enhanced degree, because GBM-SC had the most hyperpolarized mitochondria both in vivo and in vitro. Reversal of this mitochondrial remodeling by DCA induced apoptosis in GBM-SC both in vitro and in vivo. Although the magnitude of apoptosis induction by DCA is
achieved plasma concentrations at values required for PDK inhibition (31). With the small number of treated patients in our study, no firm conclusions regarding DCA as a therapy for GBM can be made. Our work supports the need for further studies with DCA in GBM, with an emphasis on combination therapy protocols. GBM may also be vulnerable to other drugs in the emerging family of metabolic modulators, pointing to a new approach in the management of this incurable cancer.

**SUPPLEMENTARY MATERIAL**

www.sciencetranslationalmedicine.org/cgi/content/full/2/31/31ra34/DC1

**Materials and Methods**

**Results**

**Discussion**

Fig. S1. Molecular characterization of GBM tumors.

Fig. S2. Evolution of tumor response in patient 1.

Fig. S3. Evolution of tumor response in patient 4.

Fig. S4. Evolution of tumor response in patient 5.

Fig. S5. GBM MRI from patient 3.

Fig. S6. Mitochondrial membrane potential in GBM-SC from freshly excised GBM tissue.

Fig. S7. Characterization of primary GBM cells and GBM-SC.

Fig. S8. HXKII in GBM cells derived from patients before and after chronic DCA treatment.

Fig. S9. Effects of DCA treatment on p53 and p21 activity in vivo.

Fig. S10. Effects of DCA on angiogenesis in vitro.

Fig. S11. Effects of DCA treatment on p53 and p21 activity in vivo.

Fig. S12. A proposed comprehensive mechanism for the anticancer effects of DCA in GBM (see Supplementary Discussion).

Table S1. Laboratory and clinical parameters of five GBM patients before and after DCA treatment.

**REFERENCES AND NOTES**


