A H₂S-Nampt Dependent Energetic Circuit Is Critical to Survival and Cytoprotection from Damage in Cancer Cells

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Abstract

We recently demonstrated that cancer cells that recover from damage exhibit increased aerobic glycolysis, however, the molecular mechanism by which cancer cells survive the damage and show increased aerobic glycolysis remains unknown. Here, we demonstrate that diverse cancer cells that survive hypoxic or oxidative damage show rapid cell proliferation, and develop tolerance to damage associated with increased production of hydrogen sulfide (H₂S) which drives up-regulation of nicotinamide phosphoribosyltransferase (Nampt). Consistent with existence of a H₂S-Nampt energetic circuit, in damage recovered cancer cells, H₂S, Nampt and ATP production exhibit a significant correlation. Moreover, the treatment of cancer cells with H₂S donor, NaHS, coordinate increases Nampt and ATP levels, and protects cells from drug induced damage. Inhibition of cystathionine beta synthase (CBS) or cystathionase (CTH), enzymes which drive generation of H₂S, decreases Nampt production while suppression of Nampt pathway by FK866, decreases H₂S and ATP levels. Damage recovered cells isolated from tumors grown subcutaneously in athymic mice also show increased production of H₂S, Nampt and ATP levels, associated with increased glycolysis and rapid proliferation. Together, these data show that upon recovery from potential lethal damage, H₂S-Nampt directs energy expenditure and aerobic glycolysis in cancer cells, leads to their exponential growth, and causes a high degree of tolerance to damage. Identification of H₂S-Nampt as a pathway responsible for induction of damage tolerance in cancer cells may underlie resistance to therapy and offers the opportunity to target this pathway as a means in treatment of cancer.

Introduction

In recent years, it has been shown that gasotransmitters, nitric oxide (NO), carbon monoxide (CO) and hydrogen sulfide (H₂S), play a critical role in diverse physiological functions, including vascular tone, host defense against pathogens, neuromodulation, apoptosis, and energy metabolism in mammalian cells [1]. Among these gaseous molecules, H₂S is produced during amino acid metabolism by the trans-sulfuration and cysteine desulfuration pathways [2]. Endogenous H₂S production is catalyzed by three enzymes, cystathionine beta synthase (CBS), cystathionase (CTH) also known as cystathionine gamma-lyase, and 3-mercaptopyruvate sulfurtransferase (MST) [3–5].

H₂S functions as a stimulator of cellular bioenergetics, contributes to the increased reliance of cancer cells on the glycolytic pathway for ATP production and promotes angiogenesis and cytoprotection [6–8]. H₂S protects cells from oxidative stress [9], it can affect cellular responses to injury [10,11] and was shown to exhibit both pro-apoptotic and anti-apoptotic effects [12–14]. Although some suggested that H₂S has anti-cancer effects [15], others reported that it promotes proliferation of HCT116 and SW480 colonic cancer cells [16]. Fu et al. showed that Ca²⁺ stimulation causes increased CTH expression and increases H₂S and ATP production [17]. It was shown that endogenous H₂S production driven by 3-MST complements and balances the cellular bioenergetics and maintains electron flow in mitochondria [18]. Colonic cancer cells have been shown to exhibit up-regulated expression of CBS and increased formation of H₂S which directs cell proliferation and angiogenesis in colon cancer [6]. It is known that tumor cells can recover from potential lethal damage induced by hypoxia, acidosis, or by radiation and drug treatment [19–22]. We recently reported that cancer cells that recover from damages induced by hypoxia, acidosis and glucose deprivation show mitochondrial remodeling, increased aerobic glycolysis, and exhibit a high rate of ATP production [23]. In this study, we explore the role of H₂S in the process of recovery of cancer cells from damage. Damaged cancer cells exhaust their
energy supply due to repair mechanisms. Both ATP and NAD⁺ (Nicotinamide adenine dinucleotide) are the main energy sources. Nicotinamide phosphoribosyltransferase (Nampt), an enzyme required for NAD synthetic salvage pathway [24], is vital to the maintenance of cellular energy supply. Therefore, we examined the role of Nampt in conjunction with H₂S in cancer cells that recover from damage. We demonstrate that H₂S controls the recovery of cancer cells from damage by regulating Nampt directed change in energy expenditure, which drives adoption of aerobic glycolysis and increase in ATP and NAD⁺ synthesis. The interaction of H₂S and Nampt confers the cancer cells a high proliferation rate and a high degree of tolerance to damage.

Materials and Methods

Materials

H₂O₂, NaHS, bleomycin, O-(carboxymethyl) hydroxylamine hemihydrochloride (CHH) DL-propargylglycine (PAG) and FK866 were purchased from Sigma-Aldrich (St. Louis, MO). Antibodies against CBS (A-2), CTH (G-1) and β-Actin (G-2) were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Antibodies against Nampt (Visfatin, PBEF) and Bax were purchased from Abcam (Cambridge, MA). Antibody against γ-H2AX was purchased from Millipore (Billerica, MA). Secondary antibodies conjugated to horseradish peroxidase were purchased from Jackson ImmunoResearch Laboratories (Baltimore, PA).

Cell culture

HepG2, MDA-MB-231 and MDA-MB-435S were obtained from ATCC (Manassas, VA) cultured in DMEM with 2 mM glutamine, 25 mM glucose and 10% fetal bovine serum, in 37°C incubator with 5% CO₂. Damage was induced in cancer cells by hypoxia, glucose deprivation and hydrogen peroxide. For induction of hypoxia (DRH), cells were incubated for 18 hr in 1% O₂. For glucose deprivation (DRG), cells were cultured for 18 hr in a medium without glucose in presence of 5% CO₂. For induction of damage by hydrogen peroxide (DRH₂O₃), cells were treated with 800 μM H₂O₂ for 3 hr.

Animal experiments

Animal care and all procedures were carried out following approval of the Institutional Animal Care Committees of University of California, Irvine (protocol #2012-3042). Eight-week-old athymic nude (Nu/Nu) male mice (n=10) were purchased from Charles River Laboratories (San Diego, CA). Mice were anesthetized by inhalation of Isoflurane before injection of cells. Mice were euthanized by carbon dioxide inhalation when tumors grew to the size of 10–15 mm in diameter. Tumors were harvested and snap frozen in liquid nitrogen, then stored at −80°C for frozen western blot analysis.

Measurement of H₂S production in extra and intra-cells

Measurement of extracellular H₂S level was performed using Free Radical Analyzer (TBR4100 and ISO-H2S-2, World Precision Instruments, Sarasota, FL) following manufacturer’s instruction. Briefly, cell number was adjusted to 1×10⁵ viable cells in PBS and the cell suspensions were incubated at 37°C for 1 hr. Cells were then centrifuged and the supernatants were subjected to measurements. Prior to each measurement, the sensor was polarized and calibrated by adding four aliquots of the Na₂S stock solution at the final concentrations of 0.25, 0.5, 1.0 and 2.0 μM. Detection of intracellular H₂S was performed by H₂S fluorescent probe HSNI2 (a kind gift from Professor Michael D. Pluth, University of Oregon, Department of Chemistry, Eugene, Oregon).

Whole cell protein extraction and Western blotting

Proteins from cells were extracted in lysis buffer [50 mM Tris-HCl, pH 7.5, 150 mM NaCl, 1% NP-40, 2 mM EDTA, 1 mM PMSF, 1 mM Na3VO4, 50 mM NaF, and protease inhibitor cocktail]. Protein measurements were carried out by Bio-Rad protein assay based on Bradford dye-binding method (Bio-Rad Lab, Hercules, CA). Blotting bands were detected by ECL enhanced chemiluminescence (Amersham ECL Plus Western Blotting Detection Reagents GE Healthcare Life Sciences, Pittsburgh, PA) using C-Digit Digital Imager (LI-COR, Lincoln, NE) and densitometric analysis was performed using myImage Analysis software (Thermo Scientific). β-actin served as a loading control.

Cell viability measurement

Relative cell number was measured by XTT assay (Sigma-Aldrich, St. Louis, MO). Cells were incubated with XTT and phenazine methosulfate (PMS) at 37°C for 2 hr and absorbance was read at 450 and 650 nm as a reference.

Reverse transcription-Polymerase chain reaction (PCR) and Quantitative PCR (qPCR)

Total RNA was isolated using GenElute Mammalian Total RNA Miniprep Kit (Sigma-Aldrich, St. Louis, MO). Reverse transcription was performed using High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems, Carlsbad, CA). RT-PCR was carried out using the primers specific for the human CBS (forward: GAACCGAGCGGAGACGACA; reverse: GTGGCTCAGGAACCTGTGA), for the human CTH (forward: AAAAGACGGCTTCTCGACAGG; reverse: AGGGCACTCTTGATGGGATTC) and for the human MTS (forward: CGCCGGTGGATCTGTTGATGT; reverse: CAGGTTCAATGGCGCTCTCTG). Gene expression was assessed by PCR using Taq 5× Master Mix (New England Biolabs, Ipswich, MA) with an initial denaturation step 94°C for 5 min, followed by 30 cycles with each at 94°C for 30 sec, 55°C for 30 sec, and 68°C for 1 min.

Quantitative evaluation was performed by using myImage Analysis software (Thermo Scientific, New Hampshire). For normalization of data, β-actin was amplified with specific primers (forward: AAGCCACCCACCTTCCTCTCT; reverse: GAGACAAAAAGCGCTGATACTCT). qPCR was performed using Quanti Tect SYBR Green PCR Kit, qPCR was carried out using the primers specific for the human NAMPT (forward: ATCACTGTGTTATGCTCTTG; reverse: TGCGCTGTCACTACGTTG). Gene expression was analyzed using the primers specific for human MTS (forward: ATACTGTGTTATGCTCTTG; reverse: TGCGCTGTCACTACGTTG).

XF (extracellular flux) bioenergetic analysis

XF24 Extracellular Flux Analyzer from Seahorse Bioscience (N. Billerica, MA) was utilized for extracellular fluid bioenergetic analysis [25]. Under typical in vitro cell culture conditions, extracellular acidification rate (ECAR) is contributed by lactic acid production generated by glycolysis. To examine the aerobic glycolysis, five biological replicate cultures of control (3×10⁴) and five independently generated DR cells (3×10⁴) were plated in each well of the XF 24 culture plate and cells were incubated overnight at 37°C in presence of 5% CO₂. All cultures were examined in

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**H₂S-Nampt Circuit in Cancer Survival**
The potential to recover from damage, we transferred floating cells to new culture vessels to allow cells that recover from damage to bind to the culture substrates. Serial weekly passages of floating cells to new culture vessels allowed isolation of damage recovered cells with different recovery time. Throughout this paper, we refer to these cells as damage-recovered (DR) cells with indicated recovery time of one (DRW1), two (DRW2) or three weeks (DRW3) from damage, while we refer to the parental control cells as Pc cells (Figure 2A). This method enabled us to separate three populations of cells that reflect the length of time required for recovery. In order to assess whether isolated DR cells recovered from damage, we examined the expression of pro-apoptotic molecule, Bax. DR cells showed a decrease in Bax expression in a recovery time dependent manner as an indication of repair, while as predicted, Pc cells exposed to H2O2 for 3 hr (acute damage) expressed a high level of Bax (Figure 2B). Furthermore, cancer cells recovered from damage also exhibited an increased production of H2S as compared to the undamaged Pc cells in a recovery time dependent manner (Figure 2C, D). Since endogenous hydrogen sulfide is generated by three enzymes, CBS, CTH and MST, we examined mRNA and protein levels of these enzymes in Pc and DR cells. As shown in Figure 2E, mRNA levels of CBS and CTH increased in DR cells also in a recovery time dependent manner. However, MST was absent in Pc or DR cells. As compared to parental undamaged Pc cells, cancer cells recovered from damage had increased CBS and CTH proteins in direct correlation with the recovery period. Among the DR cells, those cells with a longer recovery time from damage show the highest up-regulation of CBS and CTH (Figure 2F). CBS was up-regulated up to 2 fold after recovery from damage induced by glucose deprivation, hypoxia and hydrogen peroxide (Figure 2G). As compared to Pc cells, DR cells generating higher levels of H2S had a higher proliferation rate and showed a higher level of tolerance to damage induced by bleomycin (Figure 2H, I). These findings show that in cells recovered from damage, H2S production might play a role in their increased tolerance to damage.

DR cells exhibit increased glycolysis and enhanced cellular bioenergetics

Because rapidly dividing cells adopt aerobic glycolysis to promote an increased biomass followed by cell division, we examined the level of key glycolytic intermediates and enzymes in cells recovered from damage. DR cells showed an elevated extracellular acidification rate (ECAR), which is an indicator of level of glycolysis (Figure 3A). DR cells generating high levels of H2S had a better bioenergetic profile as evidenced by increased level of cellular ATP (Figure 3B).

In order to address whether suppression of glycolysis attenuates H2S production, we measured H2S level in DR cells treated with HK1 inhibitor, 100 μM bromopyruvic acid, or LDH-A inhibitor, 1 mM sodium oxamate, for 15 hr. As shown in Figure S2, there was no statistically significant difference in the level of H2S after treatment with either bromopyruvic acid or sodium oxamate, whereas, as expected, glycolytic activity was substantially diminished by both inhibitors. These data indicate that glycolytic enzymes do not regulate H2S-Nampt circuit.

Cells recovered from damage heavily relied on glycolysis that increased demand for NADH/NAD+. Therefore, as a measure of the bioenergetic state, we evaluated the levels of NAD+ and NADH. Although acute damage led to a decrease in the level of NAD+, NAD+ was significantly increased in cancer cells that recovered from damage induced by H2O2 and hypoxia (Figure 3G, D; Figure S3). Reduced form of NAD+, NADH, was significantly increased in DR cells (Figure S4).
Figure 1. Endogenous hydrogen sulfide increases in response to acute damage in cancer cells. (A) Amount of H$_2$S released by 293 cells, fibroblasts (Fibro.), HepG2, MDA-MB-231 and MDA-MB-435S cells. (B) The levels of H$_2$S in HepG2, MDA-MB-231 and MDA-MB-435S Pc cells subjected to hypoxia (0.5% O$_2$, 18 hr), glucose deprivation (glucose free medium, 18 hr) or treatment with bleomycin (35 nM, 18 hr), H$_2$O$_2$ (800 mM, 3 hr). Data are expressed as percent of H$_2$S released from untreated cells. (C) Intracellular CBS and Bax (left panel) assessed by western blot analysis in MDA-MB-435S Pc cells, and Pc cells treated with 400 or 800 mM H$_2$O$_2$ for 3 hr. (D) The level of CBS and γH2AX with or without treatment with H$_2$O$_2$ (800 mM, 3 hr) in MDA-MB-435S cells. Protein density was normalized using β-Actin. (E) Amount of H$_2$S and cell viability after treatment with a range of concentration of H$_2$O$_2$. *: p<0.05; **: p<0.005; ***: p<0.0005.

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Nampt, which is required for NAD$^+$ synthetic salvage pathway [24], was significantly increased upon recovery in DR cells and this increase coincided with up-regulation of H$_2$S generating CBS (Figure 3E). The DR cells with longer recovery time from damage showed the highest increase in Nampt (Figure 3F). Moreover, we found that Nampt levels correlated with both the level of H$_2$S released from cells ($R^2 = 0.9$, $p<0.0005$ in ANOVA statistical analysis) and the intracellular level of ATP ($R^2 = 0.94$, $p<0.0005$) (Figure 3G).

Together, these findings show that cancer cells recovered from damage generate a high level of H$_2$S and Nampt, and that they are heavily dependent on the production of H$_2$S and aerobic glycolysis for their efficient metabolism.

H$_2$S, in a dose-dependent manner, up-regulates Nampt, increases aerobic glycolysis and affords cytoprotection

To further assess the importance of H$_2$S in acquisition of new metabolic characteristics, Pc cells were treated with H$_2$S donor, NaHS. Similar to our observations in DR cells (Figure 3A), treatment with NaHS enhanced ECAR in a dose dependent manner, increased ATP and NAD$^+$ output and led to significant rise in the level of Nampt (Figure 3E). Consistent with previous reports showing that H$_2$S provides cytoprotection [8,26], Pc cells pre-treated with NaHS exhibited a higher resistance to damage induced by either hydrogen peroxide or bleomycin (Figure 4G).

Figure 2. Damage-Recovered (DR) cells show increase in H$_2$S and proliferation rate and exhibit tolerance to damage. (A) A scheme for isolation of Damage-Recovered (DR) cells. (B) Bax expression in H$_2$O$_2$ treated Pc, DR$_{H2O2w1}$, DR$_{H2O2w2}$ and DR$_{H2O2w3}$ HepG2 cells. Significance between Pc and three DR cells was $p<0.0005$ in ANOVA statistical analysis. (D) H$_2$S staining of Pc, DR$_{H2O2w1}$, DR$_{H2O2w2}$ and DR$_{H2O2w3}$ HepG2 cells with 5 μM H$_2$S fluorescent probe, HSN2. Scale bars, 50 μm. (E) PCR analysis of CBS, CTH and MTS genes in Pc, DR$_{H2O2w1}$ and DR$_{H2O2w3}$ HepG2 cells. (F) Western blot analysis of CBS and CTH in Pc, DR$_{H2O2w1}$, DR$_{H2O2w2}$ and DR$_{H2O2w3}$ HepG2 cells. (G) Western blot analysis of CBS in Pc and DR$_{H2O2w1}$, DR$_{H2O2w2}$ and DR$_{H2O2w3}$ HepG2 cells. (H) Proliferation of HepG2 recovered from H$_2$O$_2$, DR$_{H2O2w1}$, DR$_{H2O2w2}$ and DR$_{H2O2w3}$ cells as a percentage of that in Pc cells. (I) Viability of Pc, DR$_{H2O2w1}$ and DR$_{H2O2w2}$ HepG2 cells with and without treatment with bleomycin. *; $p<0.05$, **; $p<0.005$, ***; $p<0.0005$. doi:10.1371/journal.pone.0108537.g002
Figure 3. Changes in glycolysis and bioenergetics in DR cells. (A) ECAR in Pc HepG2 cells treated with or without 800 μM of H₂O₂ and DRHV2 HepG2 cells. (B) ATP levels in Pc, DRHVW1, DRHVW2 and DRHVW3 HepG2 cells. Significance between Pc and three DR cells was p<0.005 in ANOVA statistical analysis. (C) NAD⁺ levels in Pc HepG2 (with or without H₂O₂), DRHVW1, DRHVW2 and DRHVW3 HepG2 cells. NAD⁺ was normalized to the level of total protein. Significance between Pc and three DR cells was p<0.005 in ANOVA statistical analysis. (D) NAD⁺ levels in Pc and DRHV1 HepG2 cells. (E) Western blot analysis of intracellular Nampt and CBS in Pc and DRHV MDA-MB-231 cells. (F) Nampt levels assessed by ELISA in Pc, DRHVW1, DRHVW2 and DRHVW3 HepG2 cells. Significance between Pc and three DR cells was p<0.05 in ANOVA statistical analysis. (G) Correlation between Nampt expression and production of H₂S and level of ATP in Pc, DRHVW1, DRHVW2 and DRHVW3 HepG2 cells. Significance of H₂S and Nampt was p<0.05, and significance of ATP and Nampt was p<0.05 in ANOVA statistical analysis. *, p<0.05; **, p<0.005; ***, p<0.0005.
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Together, our findings show that up-regulation of H2S and Nampt leading to glycolysis and bioenergetic changes are important mechanisms in the development of drug resistance and survival of cancer cells.

**Figure 4. H2S increases ECAR, ATP, NAD$^+$ and Nampt in a dose-dependent manner in cancer cells.** (A) ECAR in Pc HepG2 cells treated for 48 hr with 0, 1, and 100 μM of NaHS and DR$^{WT}$ HepG2 cells. (B) Comparison of ATP levels in Pc HepG2 cells treated for 48 hr with 0, 10 and 100 μM of NaHS, DR$^{H2O2}$ HepG2 cells and Pc MDA-MB-231 cells treated for 48 hr with 0, 10 and 100 μM of NaHS. Data are expressed as a percent of level of ATP in untreated cells. (C) Levels of NAD$^+$ in HepG2 and MDA-MB-231 Pc cells treated with 0, 10, and 100 μM of NaHS for 48 hr. (D) Western blot analysis of intracellular Nampt in MDA-MB-231 Pc cells treated for 48 hr with 0 and 100 μM of NaHS. (E) qPCR analysis of NAMPT expression in Pc HepG2 cells treated for 48 hr with 0 and 100 μM of NaHS. (F) Quantitation of intracellular Nampt by ELISA in Pc HepG2 cells treated for 48 hr with 0, 10 and 100 μM of NaHS. (G) Viability in HepG2 cells pre-treated for 24 hr with 0, 1, and 10 mM of NaHS and then subjected to H2O2 (800 μM) or bleomycin (35 nM, 18 hr). Viability was assessed by Trypan blue exclusion. *; p<0.05; **; p<0.005; ***; p<0.0005.

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**H2S-Nampt pathway regulates bioenergetics in DR cells**

To further address the relation between Nampt and H2S production, DR cells were treated with an inhibitor of Nampt, FK866. Treatment of cells with 200 nM of FK866 did not affect
Figure 5. H₂S-Nampt pathway regulates bioenergetics. (A) Amount of H₂S released from DRW₁ HepG2 cells treated with CTH inhibitor, PAG (100 μM, 18 hr), and Nampt inhibitor, FK866 (200 nM, 24 hr). (B) Western blot analysis of CBS in DR cells in the absence and presence of FK866 (200 nM, 24 hr). (C) Western blot analysis of CTH in DR cells in the absence and presence of FK866 (200 nM, 24 hr). (D) ATP levels in DRH₂O₂ W₂ and DRH₂O₂ W₃ HepG2 cells in the absence (−) and presence (+) of PAG (100 μM, 18 hr). (E) ATP levels in Pc, DRH₂O₂ W₂ and DRH₂O₂ W₃ HepG2 cells treated with CBS inhibitor, CHH (500 μM, 18 hr) or CTH inhibitor, PAG (100 μM, 18 hr). *; p<0.05, **; p<0.005, ***; p<0.0005.

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H₂S has been reported to have cytoprotective effects against hydrogen peroxide and doxorubicin induced toxicity and hypoxia [30–33]. We show here that DR cancer cells that recover from damage demonstrate cross tolerance to damaging conditions that induce death in a high number of parental control cancer cells. Moreover, when Pc cells are treated with NaHS, the cells develop resistance to damage. Together, our data suggest that an increase in H₂S can occur in response to damage regardless of cell type or mode of damage. Based on such evidence, we suggest that increased H₂S may be significant in the recovery of cancer cells from damages (e.g., oxidative stress, hypoxia, glucose deprivation, acidosis) that these cells endure in the harsh tumor microenvironments.

H₂S is also essential to an increased proliferation in cancer cells recovered from damage. Consistent with our data, Cai et al. showed that treatment of HCT116 and SW480 colonic cancer cells with NaHS increased cell proliferation in these cancer cells [16]. The increase was dependant on Akt and ERK phosphorylation and blockade of Akt and ERK activation attenuated NaHS-induced cell proliferation. However contrary to such an observation, Jurkoska et al. showed that treatment of human neuroblastoma SH-SY5Y cells with NAC and ribose-cysteine, which results in elevation of hydrogen sulfide, leads to inhibition of cancer cell proliferation [34]. Furthermore, Cao et al. reported that treatment of WiDr colonic epithelial cancer cells with 50 to 200 μM NaHS for 24 hr suppresses viability [35]. One possible explanation for such paradoxical results may be attributed to sensitivity of certain cancer cells towards hydrogen sulfide mediated changes in the redox state. Another possibility is that this might be due to the fact that the effects of H₂S are dose dependent. Thus, we show that H₂S, in a dose-dependent manner, up-regulates Nampt, increases aerobic glycolysis and provides cytoprotection.

We show that in cancer cells that exhibit a high level of H₂S, aerobic glycolysis and level of ATP and NAD⁺ are coordinately increased. We further show that such an increase in level of ATP and NAD⁺ in damage survivors is due to up-regulation of H₂S production since forced increase in intracellular level of H₂S leads to a concomitant rise in cellular pool of ATP and NAD⁺. In line with these observations, H₂S is shown to improve mitochondrial ATP production following hypoxia [17]. Increased generation of H₂S plays an important role in dictating cell survival after severe damage by promoting a reduction-oxidation balance, suppressing oxidative stress in mitochondria and increasing glutathione production [36].

We show that regardless of the type of injury, cancer cells that recover from damage show increased reliance on glycolysis as a main source of energy. Synthesis of ATP depends on the content of NAD⁺. Under glycolytic conditions, cells regenerate NAD⁺ via one of the recycling pathways; by either converting pyruvate into lactate or recycling nicotinamide (NAM) to NAD [37]. As shown here, the cyto-protective effects of H₂S is related to its ability to increase cellular pool of ATP and NAD⁺ through a Nampt mediated response. Cancer cells, which are treated with H₂S donor, NaHS, and cancer cells, which recover from damage, show an increased CBS and CTH driven H₂S synthesis, and increased cell viability indicating the absence of FK866 cytotoxicity at this dose in HepG2 (Figure S5). Suppression of Nampt by its inhibitor, FK866, as well as inhibition of CTH by D, L-proparglyglycine (PAG), led to decreased H₂S production (Figure 5A). Consistent with such a change in H₂S production, the expression of both CBS and CTH was attenuated by treatment of cells with FK866 (Figure 5B, C). The level of ATP was also significantly decreased by treatment of DR cells with PAG and FK866 (Figure 5D, E). This reduction of ATP was consistent with previous study showing that inhibition of Nampt by FK866 suppressed the production of ATP in ovarian cancer cells [27]. Interestingly, treatment of DR cells with inhibitor of CBS (CHH) and inhibitor of CTH (PAG) reduced Nampt (Figure 5F). These data suggest that Nampt may function as a stimulator of H₂S-producing enzymes and thereby production of H₂S, whereas H₂S causes up-regulation of Nampt.

Taken together, our findings show that up-regulation of H₂S and Nampt and their crosstalk improve bioenergetic efficiency and facilitate recovery from damage.

Accumulation of H₂S leads to higher glycolysis, better bioenergetics and increased proliferation in damage recovered cancer cells isolated from in vivo grown tumors

To identify whether cells similar to in vitro generated DR cells exist or can be generated in tumors, epithelial cancer cells were inoculated into athymic nude mice. We isolated viable cancer cells (T⁴), and damaged recover (T⁸R) cells, as described in our previous study [23]. T⁴R cells isolated from in vivo generated tumors showed the accumulation of H₂S as well as increased expression of CBS and CTH (Figure 6A, B). Similar to in vitro data, the levels of NAD⁺ and Nampt were increased in T⁴R cells as compared to the levels detected in Pc and T⁴ cells (Figure 6C, D); ECAR was increased in T⁴R cells and these cells showed a better bioenergetic profile as evidenced by increased level of ATP and a higher proliferation rate (Figure 6E–G).

Based on these findings, we propose a scheme whereby cancer cells recovered from damage change their metabolic profile via up-regulation of H₂S-Nampt pathway (Figure 6H). Thus, H₂S-Nampt dependent energetic circuit is a critical regulator of stress tolerance, increased glycolysis, improved bioenergetics and increased cell proliferation.

Discussion

The endogenous production of H₂S is highly upregulated in epithelial colorectal and prostate cancer cells, and in tumor-derived endothelial cells [28,29]. In line with this evidence, we show that epithelial cancer cells (liver, breast, skin) innately produce large amounts of hydrogen sulfide independent of their origin. H₂S is increased even further in cancer cells upon acute damage induced by hypoxia, hydrogen peroxide and bleomycin, or following recovery from damage as a result of increased expression of H₂S-producing enzymes, CBS and CTH. These data are consistent with observations reported by others that CTH expression is upregulated by hypoxia leading to increased production of H₂S [17].
Nampt, the rate-limiting factor in NAD$^+$ biosynthesis, and its product NAD$^+$. Such changes should enable cells to survive when supplies of ATP and NAD$^+$ are exhausted. Another possibility to consider is that, some, if not all of the increase in ATP in cancer cells, might be due to the direct utilization of H2S as a substrate in ATP generation by cancer cells. Recent evidence suggests that, in mammalian cells, H2S can serve as an electron donor and an inorganic source of energy. While low concentrations of H2S (0.1–1 μM) elicit a significant increase in mitochondrial function, its higher concentrations (3–30 μM) have an opposite effect on

Figure 6. H2S levels and bioenergetic changes in DR cells isolated from tumors. (A) H2S levels in T$^v$ and T$^{DR}$ cells isolated from HepG2 tumors grown in vivo. (B) Expression of CBS and CTH in T$^v$ and T$^{DR}$ cells. (C) NAD$^+$ levels in T$^v$ and T$^{DR}$ cells isolated from HepG2 tumors grown in vivo. (D) Nampt levels in T$^v$ and T$^{DR}$ cells isolated from HepG2 tumors grown in vivo. (E) ECAR in T$^v$ and T$^{DR}$ cells isolated from HepG2 tumors grown in vivo expressed as the percent of the ECAR level in Pc. (F) ATP level in viable tumor cells (T$^v$) isolated from HepG2 tumors grown in vivo expressed as the percent of the ATP level in Pc. (G) Proliferation of T$^v$ and T$^{DR}$ cells isolated from HepG2 tumors grown in vivo expressed as the percent of the proliferation of Pc. Cells were seeded at a concentration of 2×10$^4$ and the total number of cells was assessed after 24 and 48 hr of culture. (H) A scheme for H2S-Nampt dependent bioenergetic circuit. Cell damage leads to increased H2S and Nampt that coordinate to lead to metabolic changes. These cells exhibit exponential growth and tolerance to damage. *; p<0.05,**; p<0.005, ***; p<0.0005.

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and Nampt implying that H2S-Nampt is the leading cause of damage is accumulation of H2S that enables their recovery via a positive feedback loop between H2S and Nampt [38]. Our preliminary data demonstrated that Sirtinol (Sirt1) that was reported to have impact on the survival of cancer cells mediated inhibition of H2S production might be attributed to reduced biosynthesis of serine and its utilization by CBS, which produces H2S and cystathionine. Cystathionine, in turn, is utilized by CTH to further produce H2S and cysteine. The existence of this positive feedback loop between H2S and Nampt may contribute to glycolytic activity and survival of cancer cells in face of microenvironmental challenges as well as after drug treatment. One of the downstream targets of Nampt is Sirtuin (Sirt1) that was reported to have impact on the survival mechanism [39]. Our preliminary data demonstrated that Sirtinol that acts as an inhibitor of Sirt1, did not affect the level of ATP in H2O2 treated control or DR cells (data not shown).

Consistent with the in vitro data, we show that damage recovered cancer cells derived from tumors generated in vivo exhibit a high level of H2S and Nampt and concomitantly show increased glycolysis, ATP and NADP+ production. According to Oncomine database, H2S-producing enzymes, CBS, CTH and MST as well as Nampt are all overexpressed in cancers of liver and breast and in melanomas (https://www.oncomine.org/) suggesting that this pathway operates in cancer cells of diverse origin. Together, our finding suggests that most cancers rely on H2S and Nampt to survive damages that they endure in their microenvironment.

In summary, a key attribute of cancer cells recovering from damage is accumulation of H2S that enables their recovery via a Nampt mediated metabolic change. H2S-Nampt pathway is a driving force in acquisition of a glycolytic phenotype, exponential growth and tolerance to damage. Therefore, even a small subset of tumor cells that recover from damage can give rise to generation of resistant cells that contribute to the expansion of cancer cell pool and facilitate their adaptation to a new microenvironment.

Supporting Information

Figure S1 Floating cells exhibit higher H2S level compared to bound cells. HepG2 cells were treated with 800 μM H2O2 for 3 hr, then floating or bound cells were collected for H2S measurement. Mean values were compared to untreated control. ***; p<0.0005. (TIF)

Figure S2 Levels of H2S and NAMPT are not affected by glycolytic inhibitor. The DRfK866 W2 HepG2 cells were treated with HK1 inhibitor, 100 μM Bromopyruvic Acid, or LDH-A inhibitor, 1 mM Sodium Oxamate for 15 hr and levels of H2S (A) or Lactic Acid (B) were measured. Lactic Acid was measured by p-phenylphenol based colorimetric assay [40]. *; p<0.05, **; p< 0.005. (C) NAMPT expression using Bromopyruvic Acid or Sodium Oxamate treated DRfK866 W2 HepG2 cells. β-ACTIN was used as a loading control. (TIF)

Figure S3 NAD+ level decreases upon H2O2 damage. HepG2 cells were treated with 800 μM H2O2 for 3 hr. NAD+ assay (A) was performed and Nampt expression (B) was measured by Western blotting. Mean values were normalized to total protein. β-actin served as a loading control. *; p<0.05. (TIF)

Figure S4 Level of NADH is increased in DRH+ cells. Level of NADH was measured in Ptc and DRH+ HepG2 cells by NADH assay kit following manufacturer’s instruction. Level of NADH was normalized to the protein content. (TIF)

Figure S5 Viability of cancer cells treated with FK866. Pc and DRfK866 W2 HepG2 cells were treated with 0, 1, 10, 100 and 200 nM with FK866 for 18 hr and performed XTT assay. Data are expressed as percentage (%) of control. There were no statistically significant changes in cell viability of cancer cells treated with any concentration of FK866. (TIF)

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Author Contributions

Conceived and designed the experiments: RSA EO SA ST. Performed the experiments: RSA EO SA ST. Contributed reagents/materials/analysis tools: SG. Contributed to the writing of the manuscript: SG ST.

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