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Epigenetic Control of Prolyl and Asparaginyl Hydroxylases in Prostate Cancer

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In many solid tumors, including prostate cancer, hypoxia-inducible factors (HIF) are up-regulated compared to their normal tissue counterparts. These HIF molecules are transcription factors, and up-regulate metabolic and angiogenic proteins in the cancer. Recently, a set of proteins known as prolyl and asparaginyl hydroxylases (PHD and AHD respectively) have been shown to be essential in the regulation of HIF, and in some cancers have been transcriptionally and translationally silenced. We therefore proposed a study that focuses on the epigenetic control of these crucial enzymes. In this report, we present data demonstrating our first findings of PHD expression in prostate cancer cell lines as well as expanding our studies to relevant human samples. Furthermore, we begin to identify specific epigenetic mechanisms that may play a major role in the transcriptional and translational control of these enzymes. Last, we will explain our future direction of the project after the award period has expired.

Hypoxia Inducible Factor, HIF, Prolyl Hydroxylase, PHD, Prostate Cancer
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Introduction

In many solid tumors, including prostate cancer, hypoxia-inducible factors (HIF) are up-regulated compared to their normal tissue counterparts. These HIF molecules are transcription factors, and up-regulate metabolic and angiogenic proteins in the cancer. Recently, a set of proteins known as prolyl and asparaginyl hydroxylases (PHD and AHD respectively) have been shown to be essential in the regulation of HIF, and in some cancers have been transcriptionally and translationally silenced. We therefore proposed a study that focuses on the epigenetic control of these crucial enzymes. In this report, we present data demonstrating our first findings of PHD expression in prostate cancer cell lines as well as expanding our studies to relevant human samples. Furthermore, we begin to identify specific epigenetic mechanisms that may play a major role in the transcriptional and translational control of these enzymes. Last, we will explain our future direction of the project after the award period has expired.

Body

In our previous reports, we had begun the initial characterization of the PHD’s and AHD in an array of prostate and breast cancer cell lines. In contrast with our initial hypothesis, the expression levels of PHD1, PHD2, and AHD did not significantly differ amongst the cancer cell lines compared to normal controls. On the contrary, the mRNA and protein level of PHD3 was quite variable amongst the cell lines assayed. Due to this observation, our initial statement of work was changed to focus on this specific isozyme. We attempted to examine the role of epigenetic silencing of PHD3 in these cell lines, and found that DNA methylation appeared to be a key regulator of the expression level of this protein. Furthermore, it was found that PHD3 could be reactivated if given a potent DNA de-methylating agent (5-aza-dC) demonstrating a causal relationship to the methylation of the PHD3 promoter and gene silencing. Since the time of our last report, we have confirmed and extended these results in numerous cancer cell lines of prostate, breast, and melanoma origins (Figure 1). The epigenetic silencing of PHD3 was not limited to DNA methylation, as it was observed that global chromatin structure was also altered around the PHD3 promoter (data shown in previous report). While these results are novel and exciting, using only carcinoma cell lines leaves questions about clinical relevance of our findings. To
address this issue, we have finalized studies upon prostate tumor samples extracted from human patients. It was discovered that on average the clinical tumors demonstrate significantly lower levels of PHD3 mRNA than normal prostate tissue (Figure 2). Unfortunately, the limited quantity of tumor per sample we were able to acquire prevented our ability to confirm these results at the protein level. In contrast, DNA methylation analysis was performed on these tissues and demonstrated little to no methylation at the PHD3 promoter regardless of expression level of PHD3 (data not shown). Two possibilities exist to explain this finding: 1) The region within the PHD3 promoter that was analyzed is methylated in cancer cell lines but not clinical tumor samples, or 2) other epigenetic processes may be involved in the silencing of PHD3 in vivo. Both of these hypotheses provide enticing endeavors for the future of this project.

With the understanding that DNA specific epigenetic marks may not explain the entire story of control of PHD3, we attempted to examine other avenues of gene expression regulation. When examining the specific prostate cancer cell line DU145, it was observed that these cells had attenuated PHD3 expression that could be reversed by hypoxia but exhibited no methylation at the PHD3 promoter (see attached manuscript for full data). These data suggest another level of mRNA regulation beyond DNA methylation. Using the miRNA prediction software TargetScan, we were able to discover 4-5 potential miRNA binding sites in the PHD3 3’ untranslated region (UTR) (Figure 3). These miRNA sites may play a major role in the regulation of PHD3 mRNA levels post-transcriptionally, and could be a novel regulatory element not yet described for this gene. To examine if these miRNA binding sites do play a role in the control of PHD3, we cloned the 3’ UTR of PHD3 into a vector downstream of a luciferase gene. By doing this, we allowed the stability of luciferase to be controlled by the same regulator elements that may impact the stability of PHD3. Interestingly, when this vector was transfected into DU145 prostate cell lines it was observed that the stability of luciferase was decreased almost 50%, and hypoxia demonstrated no significant change in this reduction (Figure 4). While these data are preliminary, they are highly suggestive of expression regulatory regions within the 3’ UTR of PHD3. Future work on this project will attempt to elucidate the exact mechanism by which PHD3 is controlled post-transcriptionally; a new aspect of this project not fully appreciated at the start.
Finally, a very recent discovery has shed new light on the functional role of PHD3. Luo et al. (Cell. 2011 May 27;145(5):732-44.) have shown that PHD3 may interact with and disrupt the function of pyruvate kinase in a hydroxylase independent manner. This finding implies that PHD3 may have a direct impact on cellular metabolism, as opposed to an indirect effect through the hypoxia inducible factors (HIFs). To query if this indeed was true, we examined if the presence of PHD3 could change the amount of oxidative stress (an indirect measure of metabolism) within cancer cells. Using an adenovirus to over-express PHD3, we infected the pancreatic cell line Mia-PaCa2 (this cell line was used due to its high infectivity with adenovirus, as well as undetectable expression of endogenous PHD3). It was observed that the over-expression of PHD3 in these cell lines decreased dihydroethidium oxidation by approximately half, suggesting a 50% reduction in cellular reactive oxygen species (Figure 5). These data support a direct metabolic alteration by the presence of PHD3, and warrants further investigation into the exact mechanism of how this may affect tumor cell growth and proliferation.

In summary, we have thoroughly characterized the epigenetic processes of DNA methylation and chromatin structure at the PHD3 promoter as initially outlined in our statement of work. While certain avenues of this project did not produce the anticipated results, the project produced significant findings in the regulation of PHD3. It is our goal to continue elucidating novel mechanisms of control of this enzyme in the hopes of potential new therapies for patients with prostate cancer.

**Key Research Accomplishments**

- Completion of PHD3 expression analysis on clinical tumor samples
- Completion of PHD3 promoter methylation analysis on clinical tumor samples
- Identification of putative microRNA binding sites, and preliminary regulatory studies using the 3’UTR of PHD3 on a luciferase reporter
- Oxidative stress analysis on prostate carcinoma cell line in the presence and absence of PHD3, with the potential for understanding metabolic changes due to PHD3
Reportable Outcomes

Peer-reviewed publications:


Abstracts:


Awards/Degrees:

- Case AJ. - PhD in Free radical and radiation biology – May, 2011

- Case AJ. – Travel Award – Society for Free Radical Biology and Medicine annual conference – November 2010
Conclusion

In summary, this project has surpassed its initial goals as outlined prior to the funding period. Initially, our study was limited to basic expression and DNA methylation analysis, but we have expanded our abilities to include genetic manipulations, in-depth oxygen tension studies, miRNA studies, as well as metabolic examinations. This work has produced a peer-reviewed journal article in a highly regarded journal, and others are in preparation. We hope to continue our findings on PHD3 in the future, and anticipate new students to take on the challenges of elucidating even more novel findings with this enzyme.

The PI of this work has achieved numerous other manuscripts, research awards, and has presented his work at a variety of national conferences around the United States. Furthermore, as of May, 2011 and the end of the award period, the PI successfully completed his PhD studies and is pursuing a 1-year transition post-doctoral fellowship with Dr. Domann prior to pursuing his career in medicine/research.
Figure 1. Cell lines representative of PHD3 expressers and PHD3 non-expressers were either not treated (C), treated with hypoxia (H), treated with 5-Aza-dC (Aza), or treated with 5-Aza-dC and then subjected to 1% oxygen (Aza + H). Real time quantitative PCR was performed to determine the mRNA expression of PHD3. The data were normalized to GAPDH and expressed relative to PHD3 mRNA in the corresponding untreated controls. Error bars = SEM. n = 3.
Figure 2. PHD3 mRNA expression is down-regulated in multiple primary human prostate cancer specimens. A) Total mRNA was isolated from frozen sections of primary human prostate cancer specimens with Gleason scores ranging from 7–9. Quantitative real-time PCR was performed using PHD3 specific TaqMan primer-probe. Relative PHD3 mRNA expression for each tumor sample is represented as (average dCt of n = 3 benign prostate tissue samples) – (dCt tumor sample). Samples were normalized to GAPDH. Dotted lines represent +/2 1 SD for benign tissue PHD3 mRNA expression. B) Box plot depicting PHD3 mRNA expression from samples shown in A. p value is based on ANOVA between 3 benign samples and 10 malignant samples.
Figure 3. TargetScan analysis of 3'UTR of PHD3. When analyzing the untranslated region of PHD3 four major putative microRNA binding sites are elucidated.
Figure 4. The 3' UTR of PHD3 destabilizes the reporter gene luciferase. After successfully cloning the 3' UTR of PHD3, the segment of DNA was placed downstream of a luciferase construct. When transfected into DU145 cells, it was discovered that the placing luciferase under the regulatory control of the 3' UTR demonstrated an approximate 50% decrease in luciferase stability.
Figure 5. The over-expression of PHD3 reduces cellular reactive oxygen species. Mia-PaCa2 cells were transfected with a mammalian expression vector of PHD3, and then exposed to dihydroethidium as a direct measure of cellular reactive oxygen species. The presence of PHD3 reduced dihydroethidium staining by approximately 50%.
Aberrant Promoter CpG Methylation Is a Mechanism for Impaired PHD3 Expression in a Diverse Set of Malignant Cells

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Abstract

Background: The prolyl-hydroxylase domain family of enzymes (PHD1-3) plays an important role in the cellular response to hypoxia by negatively regulating HIF-α proteins. Disruption of this process can lead to up-regulation of factors that promote tumorigenesis. We observed decreased basal expression of PHD3 in prostate cancer tissue and tumor cell lines representing diverse tissues of origin. Furthermore, some cancer lines displayed a failure of PHD3 mRNA induction when introduced to a hypoxic environment. This study explores the mechanism by which malignancies neither basally express PHD3 nor induce PHD3 under hypoxic conditions.

Methodology/Principal Findings: Using bisulfite sequencing and methylated DNA enrichment procedures, we identified human PHD3 promoter hypermethylation in prostate, breast, melanoma and renal carcinoma cell lines. In contrast, non-transformed human prostate and breast epithelial cell lines contained PHD3 CpG islands that were unmethylated and responded normally to hypoxia by upregulating PHD3 mRNA. Only treatment of cell lines containing PHD3 promoter hypermethylation with the demethylating drug 5-aza-2-deoxycytidine significantly increased the expression of PHD3.

Conclusions/Significance: We conclude that expression of PHD3 is silenced by aberrant CpG methylation of the PHD3 promoter in a subset of human carcinoma cell lines of diverse origin and that this aberrant cytosine methylation status is the mechanism by which these cancer cell lines fail to upregulate PHD3 mRNA. We further show that a loss of PHD3 expression does not correlate with an increase in HIF-1α protein levels or an increase in the transcriptional activity of HIF, suggesting that loss of PHD3 may convey a selective advantage in some cancers by affecting pathway(s) other than HIF.

Introduction

The cellular response to reduced oxygen availability (hypoxia) is controlled by a class of proteins called hypoxia-inducible factors (HIF-α). There are 3 known isoforms of HIF-1α, HIF-2α and HIF-3α. HIF-1α and HIF-2α are transcription factors. HIF-3α appears to lack transcriptional activity and may play a role in negative regulation of the HIF pathway [1]. Thus, from here on, when referring to HIF-α, we are referring to only HIF1 and HIF2. Transcriptionally active HIF1 and 2 are heterodimers composed of the HIF-α subunit and aryl hydrocarbon nuclear translocator receptor [ARNT/HIF-β];HIF-1α activates the transcription of EPO, VEGF, heme oxygenase-1 and several other critical intracellular responses to hypoxia including enzymes of the glycolytic pathway [2,3]. While less is known about HIF-2α transcriptional targets, HIF-2α appears to play a lesser role in the glycolytic response with more emphasis on EPO and VEGF transcription [4].

HIF-α mRNA levels are generally stable in cells. It is not until after translation that HIF-α subunits are kept at low levels by constant proteolytic degradation. First, a hydroxylation reaction is catalyzed by a family of prolyl hydroxylase domain-containing proteins (PHD/EGLN/HPH) which utilize iron, oxygen and 2-oxoglutarate as co-factors to enzymatically catalyze hydroxylation on the oxygen-dependent
degradation domain (ODD) of the HIFα-subunit [5]. Hydroxylated proline residues on HIF-α are recognized by Von Hippel-Lindau (VHL) protein, an E3 ubiquitin ligase that ubiquitinates the HIF-α subunit, targeting it to the proteosome [6]. Under hypoxic conditions, HIF prolyl hydroxylase activity is decreased and HIF-1α protein accumulates. HIF-α subunits translocate to the nucleus and dimerize with the constitutively expressed ARNT subunit [7,8]. This hetrodimer acts to turn on transcription of genes involved in oxygen homeostasis and glucose metabolism [2].

Three main isoforms of HIF prolyl-hydroxylase domain containing proteins, PHD1-3, have been identified [9]. These isoforms have been reported to have different specificities for HIF-1α and HIF-2α [10], and also differ in their subcellular localization. It has been shown that PHD1 is exclusively present in cytoplasm, PHD2 is mainly located in the nucleus and PHD3 is evenly distributed in both cytoplasm and nucleus [11]. PHD2 and PHD3, however, are considered to be the major isoforms that contribute to HIF-1 and -2α degradation in cells [12,13]. In normoxia, PHD2 is the primary enzyme that hydroxylates HIF-1α [14], whereas PHD3 has been reported to play an important role in HIF-2α hydroxylation and also in retaining cellular hydroxylase capacity in a hypoxic environment [10,15].

In normal cells, PHD3 mRNA and protein are expressed at low levels during normoxia, but are significantly induced upon exposure to hypoxia. In contrast, PHD3 expression in a significant number of cancer cell types has been shown to be low or absent not only during normoxia, but also under hypoxic conditions [10,16]. To date, no mechanism has been discovered to explain this defect in hypoxic inducibility. Interestingly, Hatzimichael et al. have recently demonstrated that the promoter of PHD3 is methylated in certain primary B-cell dyscrasias [17]. We had observed a decrease in PHD3 mRNA expression in human breast and prostate carcinoma cell lines, with an absence of PHD3 upregulation in response to hypoxia. Therefore, we were interested to determine whether PHD3 promoter methylation was responsible for this aberrant expression pattern. In this study, we show that the promoter region of PHD3 is methylated in representative human prostate carcinoma, melanoma, renal carcinoma and breast cancer cell lines. Furthermore, we show that neither HIF-1α protein levels nor hypoxic response through an HRE-luciferase reporter vector are compromised in PHD3 methylated compared to non-methylated cell lines. These results indicate that PHD3 promoter methylation is utilized by malignancies derived from diverse human cell types. Furthermore, these data suggest that loss of PHD3 expression may not affect the transcriptional response through the HIF pathway, leaving open the possibility that PHD3 silencing in tumors is selected through the loss of specific interactions with other cellular pathways.

Methods

Cell culture

Normal human prostate epithelial cells (NPrEC) were purchased from Clonetics, Lonza Inc. (Walkersville, MD) and were grown on the recommended PrEGM media supplied by Clonetics, Lonza Inc. The hTERT-HME1 cells were cultured in mammary epithelial basal medium MEGM (Lonza Inc.) at 37°C and 5% CO₂ according to the manufacturer’s instructions (Lonza Inc.). The DU 145, 22RV.1, PC-3, MDA-MB-435 (MB-435), and MCF7 cell lines were obtained from ATCC (Manassas, VA). MCF7, DU145 and MDA-MB-435 cells were cultured in Eagle’s Minimum Essential Medium (MEM) supplemented with 10% fetal bovine serum (FBS); PC-3 cells were cultured in F12 medium supplemented with 10% FBS, 2 mM L-glutamine, 1 mM Na Pyruvate, and supplemented with 100 U/ml pen/strep. All cell lines were routinely maintained at 37°C in a humidified atmosphere with 5% CO₂. Fresh media was replaced every three days while routine subculture was performed by washing with 1X PBS and detaching cells with TrypLE Express.

Semi-quantitative RT-PCR

Total RNA was extracted from individual cell lines using RNaseasy Mini Kit (Qiagen, Valencia, CA) and quantified using a NanoDrop 1000. To assess PHD3 and GAPDH expression, 500 ng of total RNA was used for reverse transcription using a OneStep RT-PCR Kit (Qiagen). The PHD3 forward primer is 5’-GGGCAAAATAC-TAGCTCAAGGAG-3’ and the reverse primer is 5’-AGTCTT-CAGTGAGGCCAGATTC-3’. GAPDH expression was assessed using GAPDH-specific primers. PCR conditions for PHD3 and GAPDH were the same except that 28 cycles of PCR were performed for PHD3 analysis and 25 cycles were performed for GAPDH. The parameters used were: 95°C for 5 minutes followed by the stated number of cycles of 94°C for 1 minute; 56°C for 1 minute, and 72°C for 1 minute, ending with a final extension at 72°C for 7 minutes. The amplified products were electrophoresed on a 1% agarose gel and stained with ethidium bromide to visualize the bands.

Quantitative real time RT-PCR

Total RNA was isolated from cells using Trizol, followed by DNase treatment and NaOAc precipitation. The reverse transcription was carried out with High-Capacity cDNA Archive Kit (Applied Biosystems, Foster City, CA). PHD3 TaqMan primer-probe was utilized from Applied Biosystems (Hs00229966_m1). The quantitative real-time PCR was set up as follows: 10 ng of RNA was used as template for each real-time PCR reaction (10 μg reaction volume); primer pairs at 0.3 μM for GAPDH with Syber Green Master Mix (Applied Biosystems). For PHD3, TaqMan universal master mix was used. The DNA polymerase was activated by heat at 95°C for 10 min followed by 40 cycles, denaturing at 95°C for 15 s, annealing and elongating at 60°C for 1 min. Data were collected with ABI PRISM 7000 sequence detection system. Data were analyzed using the ΔΔCt method.

Western Blot analysis

Cells were immediately washed with ice cold phosphate-buffered saline (pH 7.4). Cells were lysed on the plate in 200 μl RIPA cell-lysis buffer (50 mM Tris pH 8.0, 150 mM NaCl, 0.1% SDS, 0.5% Na Deoxycholate, 1% TX-100) plus 1 mM NaF, 10 mM NaVO₄, 10 mM PMSF, and 1/100 protease inhibitor cocktail (Sigma), immediately boiled for 2 minutes and then sonicated. SDS-polyacrylamide gels (7%, PHD3; 15% HIF1, HIF2) were used for protein electrophoresis. Proteins were electrotransferred onto nitrocellulose membranes and treated with anti HIF-1α (Abcam, Cambridge, MA) 1:500 overnight at 4°C. Anti PHD3, NB100-139 and anti HIF-2α antibodies (Novus Biologicals, Littleton, CO) were used at 1:500 and 1:200 respectively overnight at 4°C. Equal protein loading was confirmed on all immunoblots using human actin antibody (Sigma, St. Louis, MO) at a dilution 1:2000. Goat anti-rabbit IgG (BD Transduction Laboratories, San Diego, CA) was used as a secondary antibody against all primary antibodies. Bands were visualized by chemiluminescence with ECL plus reagent (Pierce, Rockford, IL) on a Typhoon FLA 7000.

Sodium bisulfite sequencing

Genomic DNA was extracted with the use of the DNeasy Tissue Kit (Qiagen, Valencia, CA), and sodium bisulfite conversion was...
performed with the use of the EZ DNA Methylation Kit (Zymo Research Corporation, Orange, CA). A pair of primers was designed to amplify the PHD3 promoter of both bisulfite modified methylated and unmethylated DNA but not unmethylated DNA. Nested PCR amplification on converted DNA used the following primers: outside forward: 5′-GTGTGGGATTTAGGGTTTTTTAG-3′ (SB1); outside reverse: 5′-CCAATCTCAAGCTCAATATATAC-3′ (SB2); and nested inner primers (SB3) and (SB4) whose sequences and locations are described in detail below.

The resulting PCR products were gel-extracted with the use of the Qiagen Gel Extraction Kit, or gel digestion with β-agarase followed by EtOH precipitation, and cloned with the TOPO TA Cloning Kit (Invitrogen). Plasmid DNA was extracted with the use of the Qiagen Plasmid Miniprep Kit (Qiagen). Sequencing was performed by the sequencing core facility maintained by the University of Iowa and results were tabulated for methylation status of each of the 58 CpGs contained in the amplicons from each cell line.

5-Aza-dC treatment

Cells were counted, and seeded (day 0) at approximately 750,000 cells/100 mm dish. Fresh 5-Aza-dC (5 mM) was added to the dish on days 1, 3, and 5 while a control flask was left untreated. On day 3, 5-Aza-dC treated cells were split into two 60 mm dishes in media supplemented with 5 mM 5-Aza-dC. On day 6, one of each of the 60 mm dishes was placed in a hypoxia chamber and placed under 1%O2, 94%N2 and 5% CO2 at 37°C. On day 7, all wells were harvested with 500 μl Trizol for RNA extraction.

Chromatin Accessibility

Chromatin accessibility experiments were conducted as previously described by Rose et al. [18]. The primers CA1 and CA2 were located within the region of the PHD3 gene queried for DNA methylation (primer sequences and locations described in detail below). After nuclei extraction and a 3 minute DNase I digestion the DNA was extracted and real-time PCR was conducted on an ABI 7000 Sequence Detection System. The accessibility index for each amplicon was then determined by the following formula (accessibility index = 2^n (A cleaved treated) – (A cleaved untreated)). GAPDH chromatin accessibility was also determined as a positive control for a constitutively expressed gene to control for equivalent DNase digestion between the cell lines examined.

HRE-Luciferase assay

Cell lines ~85% confluent in 60 mm dishes were transfected with an HRE-luciferase reporter vector [19] (2.5 μg) and Renilla luciferase (1.5 μg) according to Lipofectamine 2000 transfection reagent protocol. Transfection media was removed after 6 hours and replaced with fresh medium. Cells were then placed under 94% N2, 5% CO2, 1% O2 gas mixture in a hypoxia chamber, or normoxia for 24 hours and then lysed according to the Dual luciferase reporter assay system (Promega, Madison, WI) protocol. Luminescence was measured 3 times per sample using a Tecan SpectraFluor Plus luminometer.

Adenoviral Transduction

The adenoviral PHD3 construct was a generous gift from Dr. Robert Freeman from Rochester University (unpublished). Briefly, the human PHD3 coding sequence was engineered into the pDC315 vector and contains an N-terminal FLAG tag. PC3 cells were grown to ~85% confluency and then transduced with 20, 40 or 60 MOI of Ad-PHD3. Approximately 36 hours following transduction, cells were lysed with RIPA buffer and western blotted according to the procedures outlined above.

Clinical Samples

Clinical prostate tumor samples were received as frozen blocks in OCT. Sections were cut and ground with mortal and pestle. DNA and RNA were extracted in with Qiagen DNeasy Tissue kit and Trizol respectively.

Methylated DNA Enrichment

Genomic DNA was harvested from cells and tissue using Qiagen DNeasy Tissue kit. 2 μg of DNA in 120 μl of 10 mM Tris pH 8.0 was sonicated into fragments of approximately 150 bp using a Covaris S2. Fragmentation was done according to the Covaris protocol. 1 μg (60 μl) of sheared input gDNA was used according to the protocol supplied by the MethylMiner kit (Invitrogen). Binding reactions between beads containing methyl-CpG binding domains and sheared genomic DNA were performed at 4°C overnight. Bound DNA was eluted using progressively increasing NaCl concentrations. Eluates were precipitated using NaOAc and ETOH precipitation and resuspended in 60 μl 10 mM Tris pH 8.0. Real time PCR was performed using 1 μl resuspended DNA, SYBR Green master mix and 100 mM Fwd Primer: 5′-GAGGCTCCACACCCGTTTCT-3′ and Rev Primer: 5′-GCAGTGGTGGTGGCTTCCAT-3′ in a 10 μl reaction volume. The kit was validated using samples from human tumor cell lines with known PHD3 CpG island methylation status as determined by bisulfite sequencing (Figure S1).

Statistical Analysis

Significant differences between groups of data were determined using a t-test for all bar graphs and ANOVA for box plot powered by SigmaPlot 11.0 software, n = 3 was used for each data set unless otherwise noted.

Results

PHD3 mRNA is aberrantly silenced in human melanoma, prostate and breast carcinoma cell lines

A panel of human carcinoma cell lines was screened for PHD3 mRNA expression. This panel consisted of three prostate cancer cell lines (DU 145, 22RV.1 and PC-3), two breast cancer cell lines (MCF7, HS578T), one melanoma (MDA-MB-435), a non-transformed prostate epithelial cell line (NPESC) and a non-transformed immortalized breast cell line (hTERT-HME1). We found that PHD3 mRNA was expressed at different levels varying from abundant to almost undetectable levels as determined by conventional RT-PCR (Figure 1a). The prostate cancer cell lines showed decreased PHD3 mRNA expression compared to the normal prostate epithelial cells. A comparison of three prostate cancer cell lines showed that PHD3 is expressed in DU 145 and 22 RV.1, whereas in PC-3, PHD3 mRNA is nearly undetectable. The melanoma cell line, MDA-MB-435 was also found to have very low PHD3 mRNA expression. Among the mammary cell lines, HS578T had much lower PHD3 mRNA than the HME1 mammary epithelial cells and MCF7 cells expressed far more PHD3 mRNA than its normal HME1 mammary epithelial cell counterpart. To confirm and extend the results shown in Figure 1a, quantitative real time RT-PCR analysis of PHD3 mRNA expression were conducted and the results are shown in Figure 1b. Similarities in expression were found with both the methods used, and PHD3 mRNA expression was nearly undetectable in PC-3, MDA-MB-435 and HS578T cell lines.
Re-expression of PHD3 with DNA methyltransferase inhibitor 5-Aza-dC

The near absence of PHD3 mRNA expression in a subset of cell lines suggested an epigenetic mechanism might be responsible for their silencing. Unlike genetic mutations that accumulate in cancer, epigenetic modifications are reversible [20]. We hypothesized that if DNA methylation of the PHD3 gene was responsible for its reduced expression, a DNA methyltransferase inhibitor such as 5-Aza-2'-deoxycytidine (5-Aza-dC) should induce its expression. This is in line with 5-Aza-dC’s purported ability to re-activate genes previously silenced by DNA methylation in cancer cells [21].

We chose 2 cell lines displaying the most marked decrease in PHD3 expression, MB-435 and PC-3, and 2 cell lines displaying moderate to high basal PHD3 expression, MCF7 and DU 145 for treatment with 5-Aza-dC. When MB-435 and PC-3 cells were treated with 5-Aza-dC, there were significant increases in the PHD3 mRNA expression compared to their respective untreated controls [Figure 2]. Furthermore, MB-435 cells became responsive to PHD3 mRNA upregulation by hypoxia to a significant degree following hypoxic exposure. Cells that already expressed PHD3 at moderate levels did not respond to 5-Aza-dC by significantly upregulating PHD3. Although this finding suggests that CpG methylation is involved in silencing, further direct queries of epigenetic alterations at this locus were necessary to more deeply address this question.

The PHD3 CpG island is aberrantly CpG methylated and displays decreased chromatin accessibility in human carcinoma cells

To determine whether PHD3 gene methylation is present at the CpG island in cell lines that have reduced PHD3 mRNA expression and respond to 5-Aza-dC by upregulating PHD3 mRNA, we utilized sodium bisulfite sequencing to identify methylated CpG sites. Figure 3A, illustrates the CpG island in the 5’-end of the PHD3 gene. The 58 CpG sites in the region analyzed are represented by vertical lines. A putative hypoxia response element (HRE) in the promoter region of the gene is also indicated. We found that these CpGs were highly methylated in
the PHD3 negative cell lines PC-3, MB-435, and HS578T (Figure 3B). These represent examples of human melanoma, prostate, and mammary carcinoma cells respectively. We also noted that many cell lines appeared to be heterogeneous with respect to PHD3 promoter methylation status. Within certain cell lines, some clones display high levels of methylation, whereas other have very few to no methylated CpGs. This small population of unmethylated or hemimethylated cells within a cell line may explain our ability to detect very low levels of PHD3 mRNA in cell lines displaying largely methylated PHD3 CpG islands. Furthermore, areas of CpG methylation in some methylation positive cell lines overlap with a putative HRE in the PHD3 promoter region, which could hinder the ability of PHD3 to be induced upon hypoxic stimuli. In contrast to the positive methylation status in PHD3 negative cells, the CpGs in the PHD3 CpG island were largely unmethylated in the PHD3 positive cells, NPrEC, DU 145, 22RV.1, HME1, and MCF7.

DNA methylation is typically associated with other alterations to chromatin structure that participate in cell-type specific gene expression patterns. Aberrant cytosine methylation in the 5’-regulatory regions of genes is typically associated with deacetylated histones, and thus a state of DNA that is generally inaccessible to transcription factors and other enzymes that act on DNA, such as polymerase II. This is a mechanism of gene silencing often exploited by cancer cells [22]. In a chromatin accessibility assay, we found the promoter region of PHD3 in PC-3 cells was resistant to cutting by DNase I when compared to the MCF7 PHD3 promoter, whereas there was little change in GAPDH promoter accessibility between the two cell lines (Figure S2). This evidence further supports the hypothesis that PHD3 promoter methylation and heterochromatin formation are part of the mechanism for reduced expression of PHD3 in these human breast and prostate cancer cell lines.

PHD3 expression is not induced upon exposure to hypoxia in cell lines containing PHD3 promoter methylation

Unlike PHD1, both PHD2 and PHD3 genes contain hypoxia response elements, and can be induced by hypoxia by the HIF-1 and HIF-2 transcription factor complex. In the case of PHD3, mRNA and protein expression can be relatively low during normoxic conditions, with marked increases upon hypoxic insult [10]. Therefore, we tested PC-3, DU 145, MB-435 and MCF7 cells for their ability to upregulate PHD3 following 24 hours of hypoxia (1% O2) (Figure 4A). We found cell lines that contained PHD3 promoter methylation (PC-3, MB-435) failed to appreciably upregulate PHD3 mRNA under these conditions. However, we did note a very small upregulation of PHD3 mRNA in PC-3 cells. This can likely be attributed to the heterogeneity of PHD3 promoter methylation between specific clones in this cell line (see figure 3B). In contrast, PHD3 mRNA was much more prone to upregulation in the unmethylated cell line MCF7. Upregulation of PHD3 in DU 145 cells varied by experiment, and averaged as a non-significant trend toward hypoxic upregulation.

Our discovery of PHD3 promoter methylation in melanoma, breast and prostate cancer cell lines prompted us to ask whether cell lines from other malignant tissues contained methylation at the PHD3 locus. Therefore, we further performed real-time PCR and bisulfite sequencing on a panel of 3 human renal cell carcinoma cell lines (A-498, ACHN and 769-P), and on cDNA prepared from normal kidney tissue (Figure 4B). We found that ACHN and 769-P cells express nearly undetectable levels of PHD3 mRNA, whereas A-498 expresses levels comparable to normal tissue. Of these cell lines, the PHD3 positive A-498 displayed an unmethylated PHD3 promoter whereas 769-P cells displayed an aberrantly methylated PHD3 promoter. Interestingly though, we did not detect any CpG methylation at the promoter of PHD3 negative
ACHN cells, suggesting an alternative mechanism for silencing in this cell line.

PHD3 promoter methylation status does not correlate with hypoxia induced HIF-1α protein accumulation or HIF transcriptional activity

The presence of PHD3 promoter methylation in such a broad range of epithelial malignancies suggests that it may be a selective advantage for tumor survival. One hypothesis is that PHD3 silencing by promoter methylation may allow for an increased HIF transcriptional response during hypoxic conditions. In order to determine whether PHD3 promoter methylation specifically affects the hypoxia response pathway, we performed western blots on cell lysates from MCF7, PC-3, MB-435 and DU 145 cell lines to compare the HIF protein levels and the HIF transcriptional response to hypoxia (Figure 5a). Following 24 hours of hypoxia, HIF-1α protein was upregulated in all the cell lines regardless of PHD3 expression status. We also observed that DU 145 cells appear not to express detectable levels of PHD3 protein. We are unsure whether this is due to limits of detection by our PHD3 antibody, or if DU 145 cells downregulate PHD3 expression by a posttranslational mechanism. Our antibody appears to be specific to PHD3 as transduction of an adenoviral-PHD3 expression vector into PC3 cells produces a band at an identical molecular weight as the band seen in MCF7 cells (Figure S3). Interestingly though, MCF7 cells, which do express basal levels of PHD3 mRNA and protein, displayed the largest induction of HIF-1α protein. Thus, HIF-1α protein levels, in general, did not show any correlation with presence or absence of PHD3. We also found HIF-2α to be expressed under normoxic conditions in MCF7. Moreover, MB-435 cells, which express the lowest levels of PHD3 mRNA out of all the cell lines tested appear not to express HIF-2α at an appreciable level. Thus, loss of PHD3 does not appear to be significantly correlated with an accumulation of HIF-1α or HIF-2α levels in these cell lines.

To further investigate the effect of PHD3 promoter methylation on the transcriptional response of the hypoxia response pathway,
we transfected MCF7, PC-3, MB-435 and DU 145 cancer cell lines with an HRE-luciferase reporter construct [19]. Following 24 hours of hypoxia, luciferase activity was measured and plotted relative to luciferase activity in normoxic cells (Figure 5B). There was no correlation between PHD3 promoter methylation status and hypoxic induction of luciferase. The results of this experiment resembled the pattern of HIF-1α protein accumulation seen in Figure 5A. MCF7 cells showed the largest induction luciferase activity, whereas the other cell lines, which do not express detectable levels of PHD3 protein, were comparable to one another. Although not absolute, these data are highly suggestive that HIF protein stabilization and transcriptional activity is largely independent of PHD3 expression.

**Figure 4. The methylated PHD3 gene in melanoma, prostate, breast and renal carcinoma cell lines is refractory to induction by hypoxia.** A) Melanoma, prostate and breast carcinoma cell lines were treated with hypoxia (1% O2) or normoxia (21% O2) for 24 hours. Total RNA was extracted and converted to cDNA by reverse transcription. Quantitative real-time reverse transcription-PCR analysis of PHD3 was performed with normalization to GAPDH gene expression. Relative quantitation was determined by the DDCt method. ND = not detectable. Error bars = SEM. n = 3. B) Renal clear cell carcinoma cell lines were either untreated or treated with hypoxia as in A). The right panel depicts relative PHD3 mRNA levels compared to mRNA extracted from normal renal tissue. The left panel depicts the methylation status at each of 58 CpG dinucleotides present in the PHD3 CpG island of the representative renal carcinoma cell lines. Error bars = SEM. n = 3. doi:10.1371/journal.pone.0014617.g004

**PHD3 promoter methylation is absent in primary human prostate adenocarcinomas**

Recent data published by Hatzimichael et al. suggested that acquisition of PHD3 promoter methylation may be a relatively common event in certain plasma cell neoplasias [17]. Therefore, we asked whether primary human prostate neoplasias contained methylation at the PHD3 CpG island. We extracted DNA and RNA from frozen sections of 10 prostate cancer specimens containing a minimum of 70% malignant tissue with a Gleason score ranging from 7–9 as well as 3 benign prostate specimens. Real-time PCR of extracted RNA showed that all 10 tumors contained decreased PHD3 mRNA expression compared to 3 benign tissue specimens (Figure 6A and B).
Probing for methylated CpGs in select tumor specimens using the MethylMiner kit indicated the possible presence of methylated CpGs at the PHD3 CpG island in tumor sample 1 (Figure S4), which also contained the lowest PHD3 mRNA levels. However, further bisulfitene sequencing of 8 clones from tumor sample 1 and tumor sample 4 did not detect any methylated CpGs (data not shown). This could be due to better sensitivity to population average methylation states as compared to the single molecule at a time approach of bisulfite sequencing.
Perturbations in the cellular responses to hypoxia are well known to play a role in the malignant process. Familial mutations in \textit{VHL}, a negative regulator of the HIF-\(\alpha\) proteins, results in vascular tumors of the brain, spinal cord and retina, as well as appearance of renal clear-cell carcinomas [23]. PHD proteins play a role upstream of VHL regulation; they hydroxylate HIF-\(\alpha\) proteins, creating a binding site for VHL [24,25,26]. Thus, it is feasible that deregulation of PHD activity or expression could also contribute to the malignant process. In fact, an absence of PHD3 upregulation following hypoxia has been observed in multiple human cell lines from tumors of the breast, prostate and brain [10,16]. A recent clinical study of breast tumors containing \textit{BRCA} mutations supports the hypothesis that PHD3 plays an important role in malignancy. This study found a positive correlation between decreased PHD3 expression and a basal phenotype, which is considered a higher grade and more aggressive tumor [27].
Here, we report aberrantly silenced basal mRNA expression of PHD3 in breast, prostate, melanoma and renal cell carcinoma cell lines, and the absence of PHD3 mRNA induction upon hypoxic stimulus. PHD3 expression could be recapitulated in some PHD3 negative cell lines after treatment with 5-aza-dC, a DNA methyltransferase inhibitor, implicating DNA methylation as a mechanism for the decreased expression of PHD3 mRNA in these cell lines. PHD3 promoter methylation was verified by sequence analysis of PCR products cloned from bisulfite-treated genomic DNA. We found that among the human cancer cell lines investigated, PC-3, MB-435, HS578T, and 769-P cell lines have hypermethylated PHD3 CpG islands. Furthermore, the PHD3 promoter region was more resistant to DNase I in PC-3 cells (hypermethylated PHD3 promoter) compared to MCF7 cells (hypomethylated PHD3 promoter). The methylation of the PHD3 promoter in these carcinoma cell lines appears to be aberrant since insignificant DNA methylation was found in the non-transformed cell counterparts of prostate and mammary epithelial cell lines NPrEC and HME1 respectively. The apparently aberrant PHD3 promoter methylation status in these cell lines is the likely mechanism, at least in part, for PHD3 transcriptional suppression because DNA methylation is typically associated with a condensed heterochromatin state, and is known to inhibit transcription factor binding to promoter regions of genes [28].

Although all members of the PHD family have the ability to hydroxylate both HIF-1 and HIF-2α, the specificities appear to differ slightly. PHD2 has been reported to play a more pronounced role in the regulation of HIF-1α, whereas PHD3 more strongly affects HIF-2α stability [10]. Therefore, it would seem likely that loss of PHD3 expression by promoter methylation would convey a cellular advantage mediated through increased HIF-1α and/or HIF-2α stability during hypoxia. This could lead to increased expression of VEGF and erythropoietin with subsequent vascular recruitment. Tumors of the breast, skin, kidney and prostate, being solid tumors, would certainly benefit from an increase in vascular supply to hypoxic areas. In fact, there is evidence that cell lines from other solid tumors downregulate PHD3 as well. Henze et al. have shown that several glioma cell lines display little to no PHD3 protein expression during normoxia, with no induction upon hypoxia when compared to other glioma cell lines studied [16]. Our results would predict that a subset of those cell lines have aberrant methylation of the PHD3 CpG island. Interestingly, HIF-2α protein levels after 18 hours of hypoxia appeared lower in PHD3 non-expressing cells than those in cells expressing relatively high levels of PHD3 [16]. This is the opposite of what we had expected, and may demonstrate the ability of other PHD family members to substitute for the loss of PHD3 expression in regulating HIF-1α and HIF-2α stability. This hypothesis is supported by data from Appelhoffer et al., who measured relative protein amounts of PHD isoforms in multiple other glioma cell lines studied [16]. Our results would predict that a subset of those cell lines have aberrant methylation of the PHD3 CpG island. Interestingly, HIF-2α protein levels after 18 hours of hypoxia appeared lower in PHD3 non-expressing cells than those in cells expressing relatively high levels of PHD3 [16]. This is the opposite of what we had expected, and may demonstrate the ability of other PHD family members to substitute for the loss of PHD3 expression in regulating HIF-1α and HIF-2α stability. This hypothesis is supported by data from Appelhoffer et al., who measured relative protein amounts of PHD isoforms in multiple other glioma cell lines studied [16].

In fact, the results of our study support a mechanism whereby PHD3 silencing by PHD3 CpG island methylation affects pathways outside of the conventional hypoxic response pathway. When HIF transcriptional activity was measured through a hypoxia responsive HRE-luciferase reporter, we observed nearly equal transcriptional responses to hypoxia in 3 out of 4 cell lines, which included both PHD3 silenced as well as PHD3 expressing cell lines. In MCF7 cells, which express PHD3 at the protein level and do not contain PHD3 promoter methylation, we observed nearly a 15-fold increase in luciferase induction upon hypoxic treatment. These results suggest that PHD3 expression status does not significantly affect HIF protein stabilization or HIF transcriptional activity through an HRE containing promoter upon exposure to 1% oxygen. Therefore, modulation of an alternative cellular pathway(s) remains an open candidate for mediating the effects of PHD3 loss in malignancies.

Besides the HIF family proteins, other interacting partners of PHD3 have already been discovered. PHD3 is a known player in both neuronal apoptosis and in myoblast differentiation [30,31]. PHD3 also appears to interact with Bcl-2 to induce apoptosis in H9c2 cells in response to doxorubicin [32]. Furthermore, PHD3 has also been reported to destabilize ATF-4 through a novel oxygen-dependant domain on ATF-4 [33]. ATF-4 is involved in the regulation of angiogenesis and metabolism [34]. Thus, upregulation of ATF-4 by a loss of PHD3 could promote cell survival. In addition, PHD3 has also been recently reported to inhibit IKKβ. An increase in IKKβ activity in the absence of PHD3 could confer a growth advantage to cells through an increase NFkB signaling [35]. Constitutive NFkB activity is an important and common event in T- and B-cell derived malignancies [36,37] and may explain the PHD3 promoter methylation recently reported in plasma and B-cell neoplasia by Hatzimichael et al. [17]. These interesting prospects related to identification of the downstream targets of PHD3 signaling will undoubtedly become a focus of future investigations.

Our inability to detect PHD3 promoter DNA methylation in primary human prostate tumors was surprising, however our results are supported by a recent study by Huang et al. [38], who screened 168 invasive breast carcinomas and did not find evidence of PHD3 DNA methylation using melting curve analysis of bisulfite converted DNA. We cannot rule out the possibility that a low level of semi-methylated PHD3 CpG islands are present in their samples as the “intermediately methylated” controls were not truly intermediately methylated, but rather were mixtures of 100% methylated DNA with 0% methylated DNA. Nonetheless, it seems clear that a large proportion of primary epithelial tumors does not contain a high degree of PHD3 promoter methylation, and may not be the ideal specimen for detection of methylation at this locus. Given PHD3’s purported ability to negatively regulate the NFkB pathway [30,35], and the widely reported involvement of NFkB in cell migration and metastasis [39,40], future studies on PHD3 CpG island methylation in clinical samples of metastatic disease as opposed to primary tumors may yield more positive results.

Here, we are the first to report DNA methylation of the PHD3 CpG island in solid tumor cell lines derived from diverse cell types. PHD3 methylation in carcinoma cells was associated with their inability to appropriately upregulate PHD3 mRNA upon exposure to hypoxia. We are also the first to show evidence that this aberrant expression fails to correlate with an increase in HIF protein accumulation and transcriptional activity upon exposure to hypoxia in the cell lines examined. The presence of PHD3 promoter hypermethylation and PHD3 silencing in such a wide range of cancer types suggests this might be a common event that elicits a selective advantage for tumors. Our data suggest that at least in some cell lines, the nature of this advantage may extend beyond hypoxia resistance. Furthermore, the selective event may occur during or after the process of invasion/metastasis, as we and others have not found evidence of methylation in primary solid tumors [38].
Supporting Information

Figure S1 Validation of methylated DNA enrichment as a tool for detecting methylated CpG regions in human genomic DNA. A) Genomic DNA from human melanoma and breast cancer lines was enriched for methylated CpG dinucleotides at the PHD3 CpG island using the MethylMiner kit, followed by quantitative real-time PCR analysis using PHD3 CpG island-specific PCR primers. S = supernatant, representing unmethylated DNA. E1 - E5 represent elutions with 200, 650, 1100, 1550, 2000 mM NaCl respectively. Input = 1/60th total input DNA. Total amounts of eluted DNA from each fraction are represented as a fraction of input (left). Methylated CpG sequences from bisulfite-converted DNA in the corresponding cell lines are depicted for comparison (right). B) Control 100% methylated and 0% methylated oligos supplied by the MethylMiner kit were subjected to the MethylMiner protocol. S = supernatant, representing unmethylated DNA. E1-E5 represent elutions with 200, 650, 1100, 1550, 2000 mM NaCl respectively. Identical volumes of DNA from each eluate were subjected to quantitative real-time PCR using primers supplied by the MethylMiner kit. DNA content in each fraction is represented as arbitrary units.

Found at: doi:10.1371/journal.pone.0014617.s001 (10.36 MB TIF)

Figure S2 The methylated PHD3 gene in non-expressing cells is maintained in a less accessible state than the non-methylated PHD3 gene in expressing cells. A) Nuclei from PHD3-positive MCF7 and PHD3-negative PC-3 carcinoma cell lines were isolated and enzymatically restricted with DNase I. Primers CA1 and CA2 (see Fig. 3A) were used for quantitative real-time PCR (right panels) to amplify a region also assessed for cytosome methylation. Accessibility indices (left panels) were calculated as follows: AI = 2[(Ct DNase treated) - (Ct Untreated)]. B) GAPDH accessibility indices were simultaneously assessed as a control for a constitutively expressed gene in both cell lines.

Found at: doi:10.1371/journal.pone.0014617.s002 (3.00 MB TIF)

Figure S3 PHD3 antibody specificity. PC3 cells were transduced with an increasing MOI of adenovaliral-PHD3 vector. Western blot using Novus100-139 antibody co-incubated with anti-actin with an increasing MOI of adenoviral-PHD3 vector. Western blot

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Figure S4 Methylated DNA enrichment of genomic DNA isolated from primary human prostate cancer. A) Total genomic DNA was isolated from frozen sections of 7 malignant prostate cancers and 3 benign prostate samples and subjected to the MethylMiner protocol. Tumor sample number corresponds to samples shown in figure 6A. S = supernatant, representing unmethylated DNA. E1-E4 represent elutions with 300, 550, 800, and 2000 mM NaCl. PHD3 CpG island DNA content in each fraction is represented as a fraction of total PHD3 CpG island DNA present in input.

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Author Contributions

Conceived and designed the experiments: PEP. Performed the experiments: TLP MF SUV MLT AJC FED. Contributed reagents/materials/analysis tools: PEP AJC FED. Wrote the paper: PEP. Assisted with writing and editing: TLP MF SUV. Provided funding resources for the work: SUV MJT MLT AJC FED. Isolated proteins from human tumor samples and all cell lines under normoxia and hypoxia; performed transfection studies for promoter reporter experiments; performed bisulfite sequencing experiments and provided key data for figures 6a (right panel) and 6b. TLP. Extracted, quantified, and analyzed RNA and DNA from all cell lines; designed primers and performed bisulfite sequencing to determine methylation patterns. Provided key data for figures 1, 3, and 4: MP. Extracted RNA, DNA and protein from breast cancer cell lines treated with 5-aza-dC for initial RT-PCR and western blotting experiments; provided key data for figures 1, 2, 5: SV. Extracted RNA, DNA and protein from human cell lines and tumor samples, performed bisulfite and Methylminer experiments on cell lines and tumor samples and assisted with producing real time PCR replicate data for PHD3 expression experiments: SUV. Extracted RNA and protein from prostate and renal cell lines for initial RT-PCR and western blotting experiments; provided key data for parts of figures 1 and 3: AJC. Isolated proteins for western blotting; provided reagents and expertise for western blots and trained the other authors in protein techniques; provided key data for figure 6a (left panel): MLT. Provided overall guidance and direction in all phases of experimental design, implementation, data acquisition and interpretation, and manuscript writing, editing and preparation: FED.

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The term reactive oxygen species (ROS) describes a large group of free radical and non-free radical oxygen-containing compounds (e.g., superoxide, $\text{O}_2^-$; hydrogen peroxide, $\text{H}_2\text{O}_2$; peroxynitrite, ONOO$^-$; hydroxyl radical, 'OH) [1]. It is commonly accepted that ROS are by-products of normal metabolism and as such act to damage cellular components such as nucleic acids, proteins, or lipids [2–4]. Because of this, ROS have been implicated in many different diseases including cancer, atherosclerosis, amyotrophic lateral sclerosis, Alzheimer disease, and many others [5–8]. One specific role of ROS is their ability to enhance the pathogenesis of infections, such as influenza [9,10]. It has been demonstrated that during times of influenza infection ROS may damage lung parenchyma cells, but that this injury may be ameliorated by antioxidant supplementation [11–14]. Current theories propose the mechanism behind this benefit to be attenuation of ROS produced by the innate immune system, but this is not commonly accepted and is still highly debated. More recently it has been shown that cells possess the ability to exploit ROS for signaling and functional purposes. For example, many transcription factor pathways are sensitive to oxidative stress and as such are able to help cells adapt to large deviations in redox status [15–18]. Moreover, ROS are essential in the development of certain organ systems and even whole organisms [19,20]. With this knowledge, the importance of ROS in biology is being elucidated, but many questions about tissue-specific dependence, specific ROS functions, molecular targets of ROS, and ROS mechanisms of action remain unanswered.

One organ system in which ROS have been widely described is the immune system. The biological relevance of ROS in this system was first depicted when it was found that leukocytes generated ROS during the oxidative burst to neutralize pathogens [21]. More recent studies have demonstrated the importance of ROS in the downstream intracellular signaling post-T-cell activation [22–24].
Furthermore, evidence has shown that hydrogen peroxide acts as an important chemotactic agent to direct leukocytes to wound margins at sites of injury, which was pivotal in demonstrating that immune cells are able to respond to exogenous ROS in addition to producing endogenous levels [25]. Recent studies have demonstrated the role of ROS in priming the development of the primitive immune system in Drosophila, illuminating the first described function of ROS in the development of this organ system in an invertebrate species [26]. Finally, it has been demonstrated that an intracellular pro-oxidant shift occurs before normal thymocyte apoptosis during selection [27], and numerous groups have shown increased ROS levels in pathologic T cell systems [28–30]. In contrast, few studies to date have focused on how pro- or antioxidants affect the normal mammalian adaptive immune system, and to our knowledge no published studies have directly addressed the role of excess mitochondrial superoxide in the growth or function of the adaptive immune system. With the understanding that ROS play a major part in intracellular signaling and cellular damage, we hypothesized that mitochondrial superoxide may play a distinct role in normal development and maintenance of the mammalian adaptive immune system, which could be central to intercellular communication between the innate and the adaptive branches of the immune system.

Although the conceptual framework underlying this hypothesis seems simple in nature, the availability of animal models with which to address this hypothesis is limited. Numerous constitutive antioxidant enzyme knockout animals have provided valuable information on the global developmental importance of these proteins [31–36], but these studies have not assessed the role of specific ROS in tissue-specific adaptive immunity. To address these deficits, we used a conditional T cell manganese superoxide dismutase (i.e., SOD2) knockout mouse to examine the role of increased steady-state levels of superoxide during mammalian adaptive immune system development [37]. The superoxide dismutase class of enzymes specifically scavenges superoxide in biological systems [38]. Mammals contain three variants of the enzymes: cytoplasmic Cu/Zn SOD, SOD1; mitochondrial SOD2; and extracellular SOD, SOD3. Because SOD2 has the explicit role of eliminating mitochondrial superoxide, tissue-specific disruption of this activity should provide an excellent in vivo model under conditions where compartmentalized superoxide metabolism is disrupted. A constitutive SOD2 knockout mouse has been created, but because of the postnatal developmental dependence on SOD2 the animal succumbs to numerous organ failures shortly after birth [39,40]. Furthermore, because of the mouse’s limited life span no examination of the immune system was reported. Taken together, our model serves as the first described animal model for studying the effects of perturbing steady-state mitochondrial superoxide levels on the development and function of the mammalian T cell adaptive immune system.

Materials and methods

Mice

Mice homozygous for the floxed SOD2 allele (i.e., B6.Cg-Tg-AbcCreERT2mRFP/J, short hand SOD2fl/fl), in which exon 3 of the SOD2 gene is flanked by two loxP sequences, have been previously described [37]. B6.Cg-Tg-Lck-CreERT2mRFP/J, or Lck-Cre, mice (in which Cre recombinase is exogenously expressed under control of the proximal lymphocyte-specific kinase and Cre recombinase becomes activated in αβ T cells during the CD4+ /CD8− to CD4+/CD8+ stage of development) and B6.Cg-Tg-Vav1-iCreERT2mRFP/J, or Vav-iCre, mice (Cre recombinase is exogenously expressed under control of the Vav promoter, and Cre recombinase becomes activated within the hematopoietic stem cell and affects all lymphoid, myeloid, and erythroid lineages) were generously donated by Dr. Adam Dupuy (The University of Iowa) and have been previously described [41,42]. B6.Cg-Tg-AlbCreERT2mRfp/J mice (Cre recombinase is exogenously expressed under the control of the albumin promoter and as such expression is limited to the liver) were generously donated by Dr. Curt Sigmund (The University of Iowa) and have been previously described [43]. Finally, B6.129-Tg-MMTVCreERT2mRFP/J mice (Cre recombinase is exogenously expressed under the control of the mouse mammary tumor virus promoter, and expression is primarily limited to mammary tissue) were purchased from The Jackson Laboratory and have been previously described [44]. To obtain conditional T cell SOD2 homozygous knockout animals (i.e., SOD2−/−), parent strains of both the floxed SOD2 and the Lck-Cre mice were bred to generate F1 heterozygotes (i.e., SOD2+/−). The F1 generation was then bred back to the parent floxed SOD2 mice to create F2 homozygous knockouts. Lck-Cre was passed only through male parents to limit nonspecific oocyte expression. The mice used were of pure C57BL/6 background, and littermate floxed animals (i.e., SOD2−/−) served as controls. All work was performed under the approval of the Institutional Animal Care and Use Committee at The University of Iowa.

Tissue isolation

For all experiments, fresh tissue was harvested and used from animals at 6 weeks of age unless otherwise noted. For thymocyte, splenocyte, and lymph node preps, organs were freshly harvested upon necropsy and placed in a solution of Hanks’ buffered salt solution (HBSS) containing 10% fetal bovine serum. Tissues were dissociated by physical disruption using ground glass. After this, lymphocytes were isolated by mouse optimized FicoLite-LM gradient (Atlanta Biologicals, Atlanta, GA, USA). Remaining red blood cell contaminants were removed by red blood cell lysis buffer, and the remaining lymphocytes were washed twice with PBS before further analysis. Peripheral blood was isolated by retro-orbital bleeding into heparin-coated capillary tubes.

Real-time PCR

RNA was extracted using the Trizol method and was quantified by the use of a Nanodrop ND-1000. One microgram from each sample was reverse transcribed using the ABI cDNA archive kit. Generated cDNA was then subjected to SYBR green quantitative real-time PCR with primers specific to each individual transcript and the 18S control (supplementary primer sequences) under the following PCR parameters: 95 °C for 10 min, followed by 35 cycles of 95 °C for 15 s and 60 °C for 1 min. A threshold in the linear range of PCR amplification was selected and the cycle threshold (Ct) determined. Levels of transcripts were then normalized to the 18S control and compared relative to the SOD2fl/fl control using the ΔΔCt method.

Western blot analysis

Protein was extracted using standard RIPA buffer and was quantified on the basis of the Bradford assay and a standard curve. Protein was run on an SDS–PAGE gel for separation of different-sized proteins. The products were then transferred from the gel to a nitrocellulose membrane. Identification and quantification of the amount of protein were performed by addition of specific primary antibody and then the addition of a horseradish peroxidase-tagged secondary antibody to the first. Chemiluminescent substrate was added to the blot for exposure of products, and film was exposed and qualitatively quantified for protein amount. Loading errors were controlled for by normalizing to β-actin. Antibodies used for this study were SOD2 (Millipore, Billerica, MA, USA), aconitase (Abcam, Cambridge, MA, USA), SDHB (Abcam), LC3 (Cell Signaling Technology, Beverly, MA, USA), p53 (Abcam), and β-actin (Abcam).
SOD2 activity

To analyze the activity of the superoxide dismutase enzymes, the indirect competitive inhibition assay developed by Spitz and Oberley was used [45]. Briefly, superoxide is generated from xanthine by xanthine oxidase and detected by recording the reduction of nitroblue tetrazolium (NBT). SOD scavenges superoxide and competitively inhibits the reduction of NBT. One unit of SOD activity is defined as the amount of protein required to inhibit 50% of the maximal NBT reduction. To obtain the amount of MnSOD activity, sodium cyanide (5 mM) was added to inhibit the CuZnSOD enzyme activity.

Dihydroethidium/dichlorofluorescin diacetate staining

Superoxide-specific dihydroethidium (DHE) analysis was performed as previously described [46]. Briefly, superoxide production was estimated using the fluorescent dye DHE from Molecular Probes (Eugene, OR, USA). Cells were washed once with PBS and labeled in suspension at 37 °C for 45 min in PBS (containing 5 mM pyruvate) with DHE (10 μmol/L; in 1% dimethyl sulfoxide (DMSO)). Samples were analyzed using an LSR flow cytometer (superoxide-specific excitation 405 nm, emission 585 nm, band-pass filter; nonspecific excitation 488 nm, emission 585 nm, band-pass filter). The mean fluorescence intensity (MFI) of 10,000 cells was analyzed in each sample and corrected for autofluorescence from unlabeled cells. Dihydorodichlorofluorescin diacetate (DCHF-DA) analysis was performed as previously described [47]. Briefly, steady-state levels of peroxides were determined using the peroxide oxidation-sensitive (CDCHF2, 10 μg/ml) and oxidation-insensitive (CDCF, 10 μg/ml) fluorescent dyes (dissolved in 1% DMSO) obtained from Molecular Probes. The cells were washed once with PBS and labeled with the fluorescent dyes for 15 min at 37 °C in PBS. The cells then were resuspended in PBS and analyzed using an LSR flow cytometer (excitation 488 nm, emission 530 nm, band-pass filter). The MFI of 10,000 cells was analyzed in each sample and corrected for autofluorescence from unlabeled cells.

Aconitase and succinate dehydrogenase activities

Aconitase activity assay was adapted from the previously described experiment [48]. Briefly, protein from freshly isolated thymocytes was isolated by sequential freeze/thaws (3×) in a Tris/MnCl2/citrate buffer (50 mM Tris, 600 μM MnCl2, and 5 mM Na citrate). Protein was then quantified by the Bradford assay. After this, 200 μg protein was combined with NADP+ (200 μM) and isocitrate dehydrogenase (10 units). Reaction was monitored at 340 nm for the appearance of NADPH formed every 5 min for 2 h. Rates were determined from slopes determined by regression analysis of data. Succinate dehydrogenase activity assay was adapted from the previously described experiment [49]. Briefly, protein from freshly isolated thymocytes was isolated by sequential freeze/thaws (3×) in a phosphate buffer (20 mM). Protein was then quantified by the Bradford assay. After this, 200 μg protein was combined with succinate, non-complex II inhibitors (antimycin A, rotenone, and cyanide), CoQ, and 2,6-dichloroindophenol (DCIP). Reaction was monitored at 600 nm for the colorimetric change in DCIP accepting electrons every 5 min for 1 h. Rates were determined from slopes determined by regression analysis of data.

Electron microscopy

Isolated thymocytes were fixed overnight with 2.5% glutaraldehyde in 0.1 M cacodylate buffer. Postfixation was carried out for 1 h at room temperature with a buffered 1% osmium tetroxide solution reduced with 1.5% potassium ferrocyanide. Samples were stained en bloc with 2.5% uranyl acetate. Cells were then rinsed and dehydrated using gradually increasing concentrations of acetone to 100%. Infiltration of Spurr’s epoxy resin and acetone was carried out over several days to 100% resin and cured 48 h in a 60 °C oven. Sections of 90 nm thickness were cut using an Ultracut E ultramicrotome (Reichert–Jung). Grids were then counterstained with 5% uranyl acetate for 2 min and Reynolds’s lead citrate for 2 min. Samples were imaged using a JEOL 1230 transmission electron microscope at 120 kV. To perform quantitative morphometry, 100 random images of both SOD2−/− and SOD2+/+ thymocytes were chosen and blindly analyzed using ImageJ (National Institutes of Health) for the following parameters: cellular area, nuclear area, cytoplasmic area, number of mitochondria, mitochondrial perimeter, mitochondrial area, and mitochondrial density.

Flow cytometry

T-cell-specific marker analysis for development or post-influenza infection was performed as previously described [50]. After isolation of fresh cell preparations, cells were suspended in staining buffer (HBSS supplemented with 5% bovine calf serum and 0.1% NaN3) and incubated with conjugated monoclonal antibodies in the presence of normal rat serum (to limit nonspecific antibody binding). After incubation and washing, the cells were suspended in fixative (1% formaldehyde in 1.25× PBS). Stained cells were run on a FACSVantage SE flow cytometer (Becton–Dickinson, Mountain View, CA, USA) with a minimum of 30,000 events collected per sample. Antibodies were semipurified from HB101 serum-free supernatants by 50% ammonium sulfate precipitation and conjugated to specific fluorescent dye by using standard procedures.

Apoptosis and cell cycle

For apoptosis, the Annexin V–FITC Apoptosis Detection Kit (Becton–Dickinson) was used to assess annexin V-positive cells. Briefly, fresh cell preparations were incubated with 1× annexin binding buffer and annexin V–FITC (2.5 μg/ml)-conjugated primary antibody for 15 min on ice. After incubation, propidium iodide (PI; 10 μg/ml) was added to the suspension and the cells were analyzed by flow cytometry using an LSR flow cytometer. For cell cycle, analysis was adapted from a previously described report [51]. One million cells were centrifuged and suspended in 0.5 ml of Krishan reagent (0.1% Na citrate, 0.03% NP-40, 0.05 mg/ml PI, 0.02 mg/ml RNase A) before analysis. Analysis was performed by examining propidium iodide-stained nuclei on an LSR flow cytometer.

Influenza and rescue

Mice (weighing approximately 20–25 g) were infected with 1600 TCIU10 of mouse-adapted A/PR/8/34 influenza virus A H1N1 by intranasal administration. For morbidity/mortality experiments, the mice were observed and weighed daily for 14 days. For pulmonary CD8+ T cell analysis, mice were sacrificed at day 8 postinfection and T cells isolated from lung tissue. IAV PA224 (H2D(b)/SSLENFRAYV) and NP366 (H2D(b)/ASNENMETM) tetramers were obtained from the National Institute of Allergy and Infectious Disease MHC Tetramer Core Facility (Atlanta, GA, USA). For rescue experiments, Tempol was obtained from Sigma–Aldrich Co. Mito-CTPO was graciously donated by Dr. Balaraman Kalyanaraman (Medical College of Wisconsin). Both Tempol and Mito-CTPO were dissolved in ethanol vehicle and administered at 25 or 250 nM, respectively. Water bottles were supplemented with sucrose (4 g/100 ml) to offset the taste of the antioxidants. Mice were treated from the time of weaning until the date of experimentation.

Statistics

Data are expressed as means and standard deviation. All experiments were performed on at least three mice. For most experiments,
comparisons between groups were analyzed by the unpaired two-tailed Student t test. For Kaplan–Meier analysis, log-rank analysis was performed. A p value of less than 0.01 was considered significant.

Results

The conditional loss of SOD2 increases superoxide-specific oxidative stress in T cells

Because of the lack of a sufficient model to study ROS stress on in vivo development of the mammalian adaptive immune system, we reasoned that the creation of a T-cell-specific SOD2 knockout mouse would serve as an optimal platform to study the effects of mitochondrial superoxide on immune system development and function. Floxed SOD2 mice were bred with mice expressing Cre recombinase under the control of the proximal promoter of Lck to the F2 generation to create homozygous T cell SOD2 knockouts. PCR analysis of tail sample DNA was used to confirm the proper genotype of animals used for studies (Supplemental Figs. 1A and B). After this, thymocytes were removed from animals to compare the T-cell-specific SOD2 transcript levels were significantly decreased in knockout animals (Fig. 1B). Furthermore, SOD2-specific mRNA extracted from T cells showed the specific excision product in SOD2−/− thymocytes, with no apparent Cre-recombination events in SOD2+/- animals (Fig. 1A). Therefore, SOD2-specific mRNA transcript levels were significantly decreased in knockout animals compared to control animals (Fig. 1B). T-cell-specific SOD2−/− mice were found to have minimal SOD2 protein and activity in thymus-derived T cells (Figs. 1C and D), whereas no significant decreases were noted in any off-target tissues (e.g., heart, liver, brain, pancreas, or muscle) in either SOD2−/− or SOD2+/- mice (data not shown). At every level of analysis, a small amount of residual SOD2 was detectable, suggesting the Cre recombination may have been incompletely penetrant or, alternatively, that our thymocyte preparations may contain non-Lck-expressing cells (i.e., γδ T cells, immature T cell populations, or contaminating stromal cells) and thus do not possess Cre recombinase. In either case, taken together, these data support the T cell specificity and efficiency of the model of SOD2 knockout.

To confirm that SOD2−/− mice did in fact harbor increased oxidative stress compared to their SOD2+/- littermates, we performed DHE stain flow cytometry as a relative measure of superoxide content within the T cell populations of these animals. It was observed that SOD2−/− mice demonstrated significantly increased steady-state levels of superoxide compared to control SOD2+/- mice (Fig. 2A). Importantly, no change was noted in nonspecific DHE oxidation or in the oxidation of a peroxide-sensitive probe, DCFH-DA (Fig. 2B). We confirmed and extended these findings in two separate enzyme activity assays that are highly specific to disruption by superoxide (aconitase and succinate dehydrogenase), as both were shown to be significantly decreased within the SOD2−/− T cells with no significant change in protein levels (Figs. 2C and D). It was previously shown that reactive nitrogen species (RNS) may also inactivate these enzymes [52], but probing cellular proteins for damage by RNS showed nondetectable levels of nitrotyrosine (data not shown). Overall, it appeared that the conditional loss of SOD2 caused a specific and significant increase in steady-state levels of superoxide within the T cell population of SOD2−/− mice; therefore these mice should serve as an effective model system for studying the effects of excess mitochondrial superoxide on the development and function of the mammalian T cell adaptive immune system.

Increased mitochondrial superoxide levels perturb the normal development of murine T cells

To understand the consequences of increased steady-state levels of mitochondrial superoxide during T cell growth and selection,
quantitative morphometry was performed on transmission electron microscopy (TEM) images of developing thymocytes (Fig. 3A). It was observed that cellular cross-sectional area was significantly increased within SOD2−/− T cells, but no change was noted in nuclear area or mitochondrial size (Fig. 3B; Supplemental Figs. 2A and B). Conversely, SOD2−/− thymocytes demonstrated an approximate doubling of the number of mitochondria per cell, but these mitochondria were less electron dense (Fig. 3B), suggesting dysfunctional or degrading organelles, as has been observed previously in oxidatively stressed systems [53]. These three mitochondrial parameters, decreased electron density, increased number per cell, and altered metabolism (Fig. 2C), all further support that the mitochondria within SOD2−/− T cells are defective.

To expand upon these observations, the role of mitochondrial autophagy was explored. It was shown that autophagy was in fact increased in the SOD2−/− T cells as shown by increases in autophagic vesicles and the cleaved form of LC-3 protein (Figs. 3C and D). Overall, the loss of SOD2 seems to have profound effects upon T cell ultrastructure as well as metabolic function.

In addition to the morphological changes, decreases in thymocyte numbers as well as peripheral T cells were noted (Fig. 4A). When total thymocytes were quantified at various ages throughout the mouse immune system development, an approximately twofold decrease was noted in SOD2−/− mice. In contrast to this, mice harboring a SOD2 knockout from an earlier stage in development (i.e., Vav-iCre-mediated recombination) demonstrated no significant decrease.
decrease in thymic cellularity (Fig. 4A, Supplemental Fig. 3A). This finding suggests that various stages in T cell development may be differentially sensitive to the presence of excess superoxide. Moreover, in contrast to the multiorgan failure noted in the aforementioned constitutive SOD2 knockout mouse [40], our preliminary studies have revealed no overt phenotype in unstressed animals harboring either liver-specific or mammary-specific knockouts of SOD2 (Supplemental Figs. 3A, B, and C). Taken together, the identification of a cellular defect in the T cell population due to the loss of SOD2 at a later (Lck-Cre-mediated) rather than an earlier (Vav-iCre-mediated) stage of development suggests that superoxide-mediated effects are tissue and time dependent and not a systemic non-specific phenomenon. Furthermore, the decrease in thymic cellularity of the Lck-Cre SOD2−/− mice was exacerbated by the fact that SOD2−/− mice also displayed reduced percentages of mature CD4+ and CD8+ thymocytes (Fig. 4B), a trend that was also observed in peripheral lymphoid organs as well (Supplemental Fig. 4B). In contrast, the most immature T cell populations in the thymus (i.e., CD4−, CD8−), γδ T cells, as well as nonlymphocyte populations, were shown to be static or increased within SOD2−/− mice (Figs. 4B and C, Supplemental Fig. 4A). This is most likely attributable to the fact that none of these populations activates the Lck promoter and thus they do not express the exogenous Cre recombinase. Conversely, these populations may be exhibiting compensatory up-regulation due to the decrease in mature αβ T cells, a phenomenon that has been described in other models of T cell deficiencies [42,54]. Interestingly, while examining peripheral T cells, we observed that SOD2−/− lymphocytes expressed a greater proportion of Mel14+CD44+ cells, suggesting a more activated state among otherwise naïve T cells (Fig. 4C). This finding was further supported by microarray analysis that demonstrated up-regulation of numerous genes involved in T cell activation in the thymocytes of T cell SOD2−/− mice (Supplemental Figs. 5A and B), and a similar effect on gene regulation by mitochondrial superoxide has been recently reported [55].

Fig. 4. Significant T cell developmental aberrations are observed in SOD2−/− animals. (A) Left: total thymic cellularity counts at various ages of T cell development for SOD2+/+ (blue circles) and SOD2−/− (red diamonds) animals. Open bars show SOD2+/+ averages, and solid bars show SOD2−/− averages. Middle: total cell counts for three peripheral lymphoid organs at 6 weeks of age. Right: total thymic cellularity counts for 6-week-old mice that were created using the Vav-iCre promoter. No significant change in thymic numbers is noted when knocking out SOD2 at a different stage in T cell development. (B) Left: representative flow scatter diagrams of SOD2+/+ and SOD2−/− 6-week-old thymocytes for the developmental markers CD4 and CD8. Right: quantification of CD4+ and CD8+ 6-week-old thymocytes. Colored arrowheads (left) correlate to respective colored quantification (right); solid bars and arrowheads indicate SOD2+/+ and open bars and arrowheads indicate SOD2−/−. (C) Quantification of flow cytometric analysis of γδ+ T cells (left), as well as for markers indicative of T cell activation, MEL14 and CD44, on splenic CD4+ cells (middle) and CD8+ cells (right). In all graphs, solid bars indicate SOD2+/+ and open bars indicate SOD2−/−. At least three mice per experiment were analyzed; data are shown as means and SD. *p < 0.01 by Student’s t-test versus SOD2+/+. 

explain this apparently activated T cell phenotype could be the well-described phenomenon known as homeostatic expansion, a compensatory mechanism to replenish the global loss of T cells throughout the SOD2−/− mouse [56]. Taken together, these data suggest that increased mitochondrial-derived superoxide alters intracellular signaling within the developing thymocytes and may inappropriately mimic the activating conditions of an infection, thus inhibiting proper T cell function during true infection.

The decrease in SOD2−/− T cell numbers is due to increased apoptosis

During T cell development apoptosis is the major selective mechanism for lymphocyte death [57]. With the understanding that apoptosis may be initiated through cytochrome c release from damaged mitochondria or through the non-caspase-mediated p53 pathway [58], we postulated that increased apoptosis due to dysfunctional mitochondria may be the mechanism for decreased thymocytes in the SOD2−/− mice. When flow cytometry for annexin V/propidium iodide was performed, a twofold increase in the apoptotic fraction of thymocytes was found in the SOD2−/− mice (Fig. 5A). In addition, an approximate twofold increase in thymocyte sensitivity to oxygen toxicity in vitro as well as increased immunoreactive p53 was also noted, further suggesting increased susceptibility of SOD2−/− thymocytes to death by apoptosis (Supplemental Figs. 6B and C). In addition, propidium iodide cell cycle analysis demonstrated a greater cycling population in SOD2−/− mice, which supports a role for compensatory proliferation in more primitive CD3−/CD4−/CD8− cells as previously discussed (Supplemental Fig. 6A). When mice were treated with small-molecule superoxide scavengers (i.e., Tempol [59] or Mito-CTPO [60]) added to the water supply from the time of weaning, the antioxidant supplementation led not only to the complete rescue of the SOD2−/− phenotype, but also to increased thymocyte numbers in control SOD2L/L animals by approximately twofold, while decreasing their apoptotic fraction by 50% (Fig. 5B). There was a strong correlation between apoptotic fractions and total cell counts, suggesting that apoptosis is the primary mechanism for increased T cell death in SOD2−/− animals (Fig. 5B). These data suggest a tight relationship between the regulation of steady-state mitochondrial superoxide levels within the normal developing thymus and immune system aberrations.

Mitochondrial superoxide mediates susceptibility to influenza A H1N1

The lack of proper T cell development observed in the SOD2−/− mice strongly suggested an immunocompromised state. To test this hypothesis, mice were challenged with a sublethal dose of IAV H1N1, a pathogen that causes a T-cell-mediated immune response, and their ability to mount an immune response was observed. Surprisingly, 100% of SOD2−/− animals succumbed to the infection and died, whereas only a small percentage of SOD2L/L animals exhibited mortality, and furthermore, SOD2−/− mice demonstrated increased

![Fig. 5. Apoptosis explains the decrease in SOD2−/− thymocytes, which may be rescued by superoxide scavenger supplementation. (A) Representative annexin V and propidium iodide flow scatter diagrams of 6-week-old thymocytes from vehicle-treated SOD2L/L (left), vehicle-treated SOD2−/− (middle), and Mito-CTPO-treated SOD2−/− (right) mice. (B) Left: total thymic cellularity with and without pharmaceutical antioxidant supplementation. Middle: quantification of annexin V positive and propidium iodide negative thymocytes with and without pharmaceutical antioxidant supplementation. The colors are coordinated to the arrowheads in (A). Right: regression analysis of total thymic cellularity versus apoptotic fraction. At least three mice per experiment were analyzed; data are shown as means and SD. *p<0.01 by Student’s t test versus vehicle-treated SOD2−/−. Ψp<0.01 by Student’s t test versus vehicle-treated SOD2−/−.](image-url)
morbidity (i.e., weight loss) with no recovery compared to SOD2−/− mice (Fig. 6A). Moreover, treatment of SOD2−/− animals with the small-molecule superoxide scavenger Tempol (Supplemental Fig. 7) or Mito-CTPO (Fig. 6A) decreased this weight loss and rescued their ability to survive the influenza infection with delayed or no mortality noted. When examining immunological parameters of the infection, we once again found that SOD2−/− mice manifest their immune defect with decreases in not only total T cells, but also IAV-specific CD8+ IAV-peptide-MHC-tetramer+ cells, interferon-γ (IFN-γ)+ cells, tumor necrosis factor α (TNF-α)+ cells, and dual-expressing IFN-γ+TNF-α+ cells (Fig. 6B; Supplemental Fig. 8A). Interestingly, of the SOD2−/− T cells that were present at the site of infection the quantity of IFN-γ expression per cell was no less than in control SOD2+/+ T cells, and TNF-α expression actually increased within SOD2−/− T-cells (Supplemental Fig. 8B). These findings are consistent with other studies of immunodeficient states and sepsis in which T cells compensate by overproducing proinflammatory markers in what is known as a “cytokine storm” [61] and may in part explain the apparently activated state of the naïve T cells in the SOD2−/− mice. Overall, alterations in mitochondrial superoxide steady-state levels seem to significantly affect the outcomes associated the mammalian adaptive response to viral infection.

**Discussion**

The current findings suggest an excess of mitochondrial superoxide at the stage of Lck activation of development tilts the intracellular redox potential toward apoptosis, resulting in the decreased thymic cellularity observed. A relative increase in the normal steady-state levels of superoxide had severe consequences on the immunocompetence of the mammalian adaptive immune system. This novel finding complements the well-established role of ROS in the innate immune system and adds yet another effect of superoxide and its derived reactive oxygen intermediates on normal tissue growth and development. However, further studies are required to examine specific signaling pathways that are affected by increased mitochondrial superoxide. It would be predicted that redox-sensitive signaling cascades such as NF-κB [17], AP-1 [15], or JAK/STAT [18] may be affected and, as such, may in part explaining the pseudo-activated state and microarray results observed in this study. Furthermore, the role of posttranslational modifications on proteins such as the hypoxia-inducible factor could also provide valuable information on how ROS regulate cellular signaling [62]. Moreover, how ROS affect peripheral lymphocyte mitosis, proliferation, and expansion in antigen-unchallenged and -challenged mice also warrants further examination. It may also be beneficial to understand the role of compartmentalization of superoxide (e.g., cytoplasmic, extracellular), and as such these studies should be followed by similar investigations using SOD1 [63] or SOD3 [64] conditional knockout animals. Because both SOD1 and SOD3 constitutive knockout animals are viable and have no gross pathologic phenotype compared to the constitutive SOD2 knockout mouse [31,39,40,65], it would be postulated that superoxide compartmentalization may explain this phenomenon at least in part. We observed that elevated mitochondrial superoxide levels had severe effects on T cell development, but it may be postulated that elevated superoxide in different cellular and tissue

![Fig. 6](image-url). SOD2−/− mice are immunologically susceptible to influenza virus, but may be rescued by superoxide-scavenging pharmaceuticals. (A) Left: Kaplan–Meier analysis of mice succumbing to IAV H1N1 infection during a 2-week period after viral administration with and without Mito-CTPO administration. Right: relative weight loss of mice over a two-week period after IAV H1N1 infection. At least six mice per experiment were analyzed; data are shown as means and SD. For weights and bar graphs, *p<0.01 by log-rank analysis versus SOD2−/−. For mortality, *p<0.01 by Student’s t test versus SOD2−/−. (B) Left: total lymphocyte counts harvested from lungs of both SOD2+/+ and SOD2−/− animals 8 days post-influenza virus infection. Also, flow cytometric analysis of CD8+ influenza-specific NP366 and PA224 MHC-tetramer-positive cells (middle) and analysis of NP366- and PA224-specific IFN-γ+/TNF-α+ (right) CD8+ T cells isolated from lungs 8 days post-IAV H1N1 infection. At least six mice per experiment were analyzed; data are shown as means and SD. For weights and bar graphs, *p<0.01 by Student’s t test versus SOD2−/−.
compartments may lead to a spectrum of developmental, functional, or disease (e.g., cancer) predispositions. Last, we have examined mitochondrial superoxide and the consequences of its excess on the development of the T cell adaptive immune system. An in-depth analysis of other reactive oxygen species (e.g., hydrogen peroxide, peroxynitrite, hydroxyl radical) may prove highly informative to further understand the part each plays in the development and function of organ systems. These data could serve as a platform for targeting specific immunodeficiencies that are currently not well understood and may be due to alterations in redox status and tailoring antioxidant therapies accordingly.

In this study, it was demonstrated that the loss of immunocompetence in a SOD2−/− background could be rescued using a mitochondria-targeted superoxide scavenger, an example of the aforementioned tailoring of specific therapies to immunodeficiencies. Serendipitously, it was shown that SOD2−/− mice with fully developed and functional adaptive immune systems also responded to the pharmaceutical intervention. These mice were shown to have significantly increased numbers of thymocytes compared to untreated control animals, and this seemed to be due to a decreased apoptotic fraction. At first glance, it seems that this increase in T cells could potentially be a boost to immune system prowess, but SOD2−/− mice treated with superoxide scavengers showed no significant increase/decrease in the ability to recover from influenza A, H1N1, infection (data not shown). This finding is most probably attributable to the hypothesis that the increase in T cells is due to an inhibition of proper apoptotic death of cells destined to die in the thymus (i.e., nonfunctional cells, cells that are self-recognizing, etc.) as opposed to the expansion of pathogen-specific immune cells. Although in the short term this had no observable consequence, it is speculated that the preservation of cells predestined for death could lead to the potential for autoimmunity. Additional studies will be needed to confirm this potential role of superoxide in the development of autoimmune diseases in both mouse and human. In this study, equal numbers of male and female mice were used for all assays, but anecdotally it was observed in many cases that females demonstrated more pronounced outcomes due to the effects of altering steady-state superoxide levels. With the understanding that females are more prone to autoimmune diseases, as well as the recent observation of different antioxidant capacities in females and males [66], this preliminary observation may warrant further examination of this phenomenon in similar models of adaptive immune system antioxidant alterations.

In conclusion, we have used a previously undescribed model of T-cell-specific superoxide scavenging deficiency to examine the development of the adaptive branch of the mammalian immune system and found that excess mitochondrial superoxide late in T cell development is detrimental to normal T cell maturation. In contrast, early developmental loss of SOD2 activity displayed no gross alteration in thymic cellularity, indicating the importance of the temporal nature of the loss of SOD2 function on thymic developmental processes. In addition, we have shown that the loss of SOD2 does not exert a similar damaging effect in development or maturation of other organs, including liver and mammary glands, supporting previous findings by others using SOD2 floxed models [37,67]. At this time the nature of the differences in cell-type specificity and developmental timing of superoxide toxicity are poorly understood, but our findings warrant further investigation into possible mechanisms mediating these differential sensitivities. Using the appropriate Cre-recombinase mouse, this model could be expanded to examining other elements of the immune system (e.g., B cells [68], neutrophils [69], macrophages [69]), different time points in development [41,42,70], or other organ systems altogether [53,71,72]. In addition, because of the simplicity of the breeding scheme the floxed SOD2 mouse may be applied in combination with other protein deletions or additions (e.g., p53 null [73], Myc overexpression [74,75]) to examine the effect of superoxide oxidative stress in alternative immune model systems. Overall, this new model and these observations serve as an initial step in understanding the role of ROS in development and function of mammalian organ systems.

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Appendix A. Supplementary material

Supplementary data to this article can be found online at doi:10.1016/j.freeradbiomed.2010.11.025.

References

