Choline phosphorylation and regulation of transcription of choline kinase α in hypoxia

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Running footnote: Regulation of choline kinase in hypoxia

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ABSTRACT

Choline kinase catalyzes the phosphorylation of choline, the first step of phospholipid synthesis. Increased phosphorylation of choline is a hallmark characteristic of the malignant phenotype in a variety of neoplasms. However, in hypoxic cancer cells, choline phosphorylation is decreased. To understand the mechanism behind this altered metabolic state we examined the expression and regulation of major choline kinase isoform, ChKα in hypoxic PC-3 human prostate cancer cells.

Hypoxia decreased choline phosphorylation, choline kinase activity and ChKα mRNA and protein levels. Promoter analysis studies revealed a region upstream of the ChKα gene bearing a conserved DNA consensus binding motif, hypoxia response element-7 (HRE7) at position -222 relative to +1 translation start site, for binding the hypoxia dependent master regulator transcription factor, hypoxia-inducible factor 1α (HIF-1α). Electrophoretic mobility shift competition/supershift assay and chromatin immunoprecipitation assay confirmed binding of HIF-1α to HRE7. A putative promoter of ChKα was isolated from PC-3 genomic DNA and cloned into a luciferase (Luc) based reporter vector system. In PC-3 cells, hypoxia decreased the expression of Luc under the control of the ChKα promoter. Mutation of HRE7 abrogated this hypoxia effect, further demonstrating the involvement of HRE7 in hypoxia-sensitive regulation of ChKα. The results strongly suggest that transcriptional control of choline phosphorylation is largely mediated via HIF-1α binding to the newly identified HRE7.

KEY WORDS

Phospholipids/Phosphatidylcholine; Gene Expression; Lipid Kinases; Phospholipids/Metabolism; cancer; choline, hypoxia, choline kinase, HIF
In recent years, choline phospholipid metabolism has been widely studied in cancer research. Choline is an important precursor of phospholipids. Its phosphorylation by choline kinase to form phosphocholine is the first step of multistep Kennedy pathway (1) that synthesizes the major membrane phospholipid, phosphatidylcholine. Increased choline phosphorylation (primarily by choline kinase alpha, ChKα) in tumor as compared to normal tissue has been reported in lung, breast, colorectal, and prostate cancers (2-5). This has motivated evaluation of cancer therapies involving choline kinase inhibition (6) and use of cancer imaging techniques with choline phosphorylation as a diagnostic metabolic step (7,8).

Metabolism of choline in cancer cells is known to be sensitive to its microenvironment. Tracer studies in the mouse atrial cardiomyocyte tumor lineage, AT-1 (9) and 9L glioma allografts (10) showed that radiolabeled choline phosphorylation and accumulation is significantly diminished in hypoxia. Our previous tracer studies with two human prostate cancer cell lines, PC-3 and LNCaP showed 15% and 28% decreases in choline accumulation, respectively, after 4h of anoxia (0% O2) (11). After 24h of anoxia, choline accumulation continued to decrease in both cell lines without loss of cellular viability. Since low oxygen exposure is a common factor in the microenvironment of tumors, it is important to understand the mechanisms of this effect and the implications for therapeutic and diagnostic applications involving choline metabolism.

The goals of the present study were to utilize PC-3 prostate cancer cells to assess the effects of hypoxia on 1) steady-state levels of choline metabolites, 2) equilibrium status of choline phosphorylation, 3) radiolabeled choline uptake and phosphorylation, 4) ChKα mRNA and protein levels, and 5) choline kinase activity. In this work, we demonstrate that choline phosphorylation is not at equilibrium in hypoxia. It is further shown that modulation of ChKα expression and choline kinase activity is a HIF-1α dependent response. Characterization of the ChKα promoter for regulation of ChKα expression in hypoxia revealed a novel hypoxia response element, HRE7, which is strongly implicated in this study to mediate the effects of hypoxia on ChKα expression.

**MATERIAL AND METHODS**

**Cell culture, hypoxic conditions, and cell viability assay**

The human prostate cancer cell line, PC-3 (ATCC, Manassas, VA, USA) was cultured in complete RPMI-1640 medium under normoxic (21% O2) or hypoxic (1% O2) conditions for 24h as described in supplemental information available online. It has been well reported that in a cell exposure system, the medium equilibrates with chamber atmosphere within 0.5-3h (12). Cells cultured in medium were maintained under normoxic and hypoxic conditions for 24h in closed chamber which allowed adequate exposure and full equilibration of medium with the chamber environment. Cell viability assays were performed using trypan blue staining (10). In some assays hypoxic conditions were mimicked by exposing cells to 24h of 2mM dimethyloxalylglycine (DMOG). DMOG stabilizes hypoxia dependent master regulator, hypoxia-inducible factor 1α (HIF-1α) thereby acting as a hypoxia mimetic.
Cell counting and calculation of population doubling time

Cells were trypsinized and suspended in medium. Cells were counted and population doubling time was calculated as previously described (10).

Choline accumulation and metabolism study

Normoxic and hypoxic PC-3 cells were cultured in 6 well format and incubated for different durations with ~ 74KBq $[^3]$Hcholine (GE Healthcare, Piscataway, NJ, USA) per well for cellular uptake measurement and with ~ 444KBq $[^3]$Hcholine per well for metabolite analysis. After incubation with the radiotracers, cellular accumulation measurements and radiolabeled choline metabolite estimations were performed as previously described (10).

Pulse-chase experiment

To investigate the efflux rates of radioactivity from PC-3 cells after initial uptake, pulse chase experiments with $[^3]$Hcholine were performed as previously described (10). Efflux measurements were made every 15min for 2h.

Choline kinase activity assay

Choline kinase activity assays from normoxic and hypoxic cell extracts were performed as previously described (10).

ATP, choline, phosphocholine and mass action ratio (MAR) measurement

ATP concentrations in cultured cells were determined as previously described (10). Choline was measured in the perchloric extracts of cells by spectrophotometric assay (13). Phosphocholine was estimated in cell extracts by dephosphorylation of phosphocholine by phosphatase and measurement of the subsequent increase in choline (14). MAR for choline phosphorylation reaction was calculated by using formula, $\text{MAR} = \frac{[\text{phosphocholine}] \times [\text{ADP}]}{[\text{choline}] \times [\text{ATP}]}$ with measured choline, phosphocholine and ATP. Intracellular concentration of [ADP] is assumed to be 1.1mM (15). For a reaction to be at equilibrium MAR should be equal to $K_{eq}$. Equilibrium status was established by comparing calculated MAR with known equilibrium constant, $K_{eq}(1.24 \times 10^4)$ for choline phosphorylation (16).

Western Blotting

Protein was extracted from PC-3 cells using RIPA lysis buffer (SIGMA, St. Louis, MO, USA) with Complete protease inhibitor cocktail (Roche Diagnostics GmbH, Germany). Western blotting was performed as described in supplemental information available online. Monoclonal anti-HIF-$1\alpha$ (BD Biosciences, San Jose, CA, USA) and polyclonal anti-ChK$\alpha$ (SCBT, Santa Cruz, CA, USA) were used for evaluating HIF-$1\alpha$ and ChK$\alpha$ protein levels and anti–$\beta$-actin (SCBT, Santa Cruz, CA, USA) was used as a loading control.

RT-PCR

Total cellular RNA was isolated from cells using RNeasy Plus mini Kit (Qiagen, Valencia, CA, USA). RT-PCR and data analysis was performed with validated ChK$\alpha$ and VEGF specific TaqMan gene expression assays (Applied Biosystems, Foster City, CA, USA). RT-PCR analysis was performed using 5 independent RT-PCR estimations.
HIF-1α silencing

A lentiviral approach was used to silence HIF-1α in PC-3 cells. The lentiviral expression vector, pLKO.1 (control and HIF-1α targeting-shRNA construct) were kindly provided by Dr. Andrew L. Kung (Dana Farber Cancer Institute) and packaging plasmids (pCMV-dR8.91 and VSV-G/pMD2G) were kindly provided by Dr. William C. Hahn (Dana Farber Cancer Institute). Recombinant lentiviruses were produced by cotransfecting HEK 293T cells with the lentivirus expression vector and packaging plasmids using Lipofectamine 2000 (Invitrogen, Carlsbad, CA, USA) as a transfection reagent. Infectious lentiviruses were collected at 24 and 48h after transfection and the pooled supernatants centrifuged to remove cell debris and filtered through a 0.45 μm filtration unit. PC-3 cells were infected with these lentiviruses and stable transfectants were selected in puromycin for 14 days.

Chkα silencing

ON-TARGET plus SMARTpools siRNA target sequences reagents (Thermo Fisher Scientific, Dharmacon Products, Lafayette, CO, USA) were used to silence Chkα in PC-3 cells according to manufacturer’s protocol. ON-TARGET plus Non-Targeting Pool (Thermo Fisher Scientific, Dharmacon Products, Lafayette, CO, USA) was used as a control.

Electrophoretic mobility shift assays (EMSA)

Nuclear proteins were prepared using the NE-PER nuclear and cytoplasmic extraction kit (PierceNet, Rockford, IL, USA). Complementary PAGE purified biotinylated or unbiotinylated oligonucleotides (Eurofins MWG Operon, Huntsville, AL, USA) were annealed to make double-stranded probes. The labeled probes were incubated with hypoxic PC-3 cell nuclear extracts according to manufacturer’s protocol in LightShift Chemiluminescent EMSA Kit (PierceNet). The DNA-protein complexes was resolved in 5% nondenaturing TBE polyacrylamide gel (Biorad, Hercules, CA, USA), followed by wet transfer to Biodyne B Nylon membrane (PierceNet). Complex was detected using Chemiluminescent Nucleic acid Detection Module (PierceNet). For competition assays, 4 pmol (200 times excess) unlabeled probes and 0.5 μg monoclonal anti-HIF-1α (BD Biosciences) was used. For supershift assay, 0.5 μg polyclonal anti-HIF-1α (Novus Biologicals, Littleton, CO, USA) was used. Incubation without nuclear extract was used as a negative control. Details of probes are described in Supplemental Table S1.

Chromatin Immunoprecipitation (ChIP) assay

A standard ChIP assay protocol was followed (17,18). Chromatin was sonicated to 500- to 1,000-bp fragments and immunoprecipitation was carried out with polyclonal anti-HIF-1α (Novus Biologicals). Chromatin (sonicated or immunoprecipitated) was purified by a standard phenol-chloroform procedure followed by column purification using minElute PCR purification kit (Qiagen). DNA was quantified by Quant-IT Picogreen quantification kit (Invitrogen, USA). Equal amount of unprocessed (Input) and immunoprecipitated (ChIP) chromatin was used for PCR. Traditional PCR was performed to standardize conditions for quantitative PCR. Q-solution (Qiagen) additive was included in amplification of the promoter region spanning the HRE7 site due to high GC content of the amplicon. Fold enrichment of a promoter region was assessed by performing quantitative PCR using QuantiTect SYBR Green PCR Kit (Qiagen) with chromatin samples taken
before (input) and after immunoprecipitation (ChIP). Details of the primers and calculations for fold enrichment are described in Supplemental Table S2.

**Promoter isolation, mutation, cloning and luciferase assay**

The genomic DNA was extracted from PC-3 cells using DNeasy Blood & Tissue Kit (Qiagen) and quantified. The putative promoter region upstream of ChKα was amplified from genomic DNA using Phusion High-Fidelity PCR Master Mix with GC Buffer (NEB, Ipswich, MA, USA). The identity of the promoter was confirmed using nested PCR amplifications and sequencing. Details of primers used are described in Supplemental Table S3. The amplified full length promoter had restriction sites, Acc651 and HindIII at the 5’ and 3’ end, respectively, for directed cloning into the pGL4.10 [luc2] vector (Promega, Madison, WI, USA). The sequence alignment of the putative promoter of ChKα from rat, mouse, human and chimpanzee was analyzed using on-line EBI Clustal W program (http://www.ebi.ac.uk/Tools/clustalw2/index.html). Mutation at the conserved HRE site (5’-TCGTGC-3’) to (5’-AGCATT-3’) was performed using QuikChange Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA, USA). Promoter assays were performed with pGL4.74[hRluc/TK] as transfection control using Dual-Glo® Luciferase Assay System (Promega).

**Statistical Analysis**

Results are presented as means ± standard deviation. Student’s t-test was applied for statistical evaluation and a p-value <0.05 was considered significant.

**Supplemental material**

Supplemental detailed information (SI) on materials and methods are available online (editor to add URL).

**RESULTS**

**Effect of hypoxia on cell population doubling time and viability**

The cell population of hypoxic PC-3 cells doubled at a 75% slower rate than their normoxic counterparts (Figure 1a), while cellular viability was preserved at nearly 100% over 24h.

**Effect of hypoxia on choline accumulation and metabolism**

In normoxic PC-3 cells incubated with radiolabeled choline, the choline tracers were taken up avidly, and rapidly phosphorylated. The accumulation of choline increased in a linear fashion with increasing incubation time in normoxic PC-3 cells (Figure 1b). The accumulated radioactivity was nearly completely (>95%) in the form of phosphocholine at incubation periods as short as 5min. The nature of accumulated choline was very different in hypoxic PC-3 cells. Following exposure to chronic hypoxia, at early incubation times (<60min), there was a linear increase in choline accumulation, but at later time points (>60 min) choline accumulation saturated with no change in tracer accumulation (Figure 1b). In hypoxic PC-3 cells, potent inhibition of choline phosphorylation was seen with <10% of the radioactivity contributed by phosphorylated choline at all incubation time points. Choline accumulation rate in normoxic cells was slightly slower in early incubation times.
(< 20min) as compared to hypoxic cells but significantly higher at later incubation times (>60min).

To further investigate the fate of radiolabeled choline metabolites, a pulse-chase experiment was performed. After administering a pulse of radiolabeled choline tracers for 2h in normoxic conditions, moderate levels of radioactivity were washed out of the cells in successive 15min incubations. The washout rate was 0.05 ± 0.02 %/min (Fig. 2). In contrast in hypoxic PC-3 cells, the washout rate was 0.6 ± 0.05 %/min, a factor of 12 times higher than for normoxic cells. Nearly all the radioactivity was washed out of the hypoxic cells in successive incubations with fresh buffer (Figure 2).

In normoxic conditions, PC-3 cells contained 0.031±0.012 μmol choline /200mg protein and 2.1 ± 0.2 μmol phosphocholine/200mg protein (Table 1). Marked changes in steady state concentrations of these metabolites were seen in hypoxia. Hypoxia caused a 666% increase in concentration of choline and 14% decrease of phosphocholine (Table 1). Consistent with the decreased choline phosphorylation, a 30% decrease in choline kinase activity was observed in hypoxic cells (1.7 ± 0.1 nmol/min/mg protein) as compared to normoxic cells (2.43 ± 0.02 nmol/min/mg protein). Hypoxia also decreased the energy state of PC-3 cells as evidenced by ~19% reduction in concentration of ATP (Table 1). The phosphocholine/choline ratio was decreased by 10 fold in hypoxic cells (phosphocholine/choline = 7.8) as compared to normoxic cells (phosphocholine/choline = 70) (Table 1).

Equilibrium status of choline phosphorylation step
Choline phosphorylation was not at equilibrium in normoxic and hypoxic cancer cells based on comparison of the estimated MAR and $K_{eq}$ constant. MAR for choline phosphorylation was 23.3 ± 9.4 for normoxic cells and 3.3 ± 0.9 for hypoxic cells (Table 1) which was far from the value of $K_{eq}$ constant 1.24 X 10$^4$ (15). Estimation of MAR was made after assuming ADP concentration to be 1.1mM (15). This is a reasonable assumption based on physiological range of ADP (15) that should not severely affect the MAR estimations. The measured choline, phosphocholine, ATP and MAR estimates further supports assumption of ADP concentration and non-equilibrium status of choline phosphorylation in PC-3 cells because it would require non-physiological ADP concentrations of approximately 0.6M or 4M to achieve equilibrium status (MAR=$K_{eq}$) in normoxic or hypoxic PC-3 cells, respectively.

ChKα gene expression in hypoxia
Hypoxic incubations resulted in significant elevation of levels of hypoxia inducible factor (HIF-1α) protein in PC-3 cells (Figure 3). Existence of a HIF-1α dependent response in hypoxic PC-3 cells was confirmed by increased mRNA levels of the hypoxia marker VEGF gene under chronic hypoxia using RT-PCR analysis. VEGF levels in hypoxic cells were 2.44 ± 0.4 fold increased relative to normoxic cells for PC-3 cells, respectively. Hypoxic exposure to PC-3 cells resulted in ~30% decreases in choline kinase activity. Likewise, RT-PCR and Western blot assay (Figure 3) showed ~ 26% decrease in ChKα mRNA (hypoxia: normoxia = 0.74 ± 0.12) and ~20% protein levels in hypoxic PC-3 cells. For the Western blot assay, highly specific commercially available polyclonal antibody against ChKα protein was
used and identity of the ChKα protein band (doublet) in the blot was validated using siRNA based ChKα silencing (Figure 4).

As shown earlier, hypoxia decreased choline kinase expression in PC-3 cells but it was not clear if this reduction was mediated by HIF-1α. In order to investigate a potential mediatory role of HIF-1α, stable PC-3 cell lines were generated with suppressed HIF-1α expression. The HIF-1α expression status of these cell lines were confirmed by Western blot assay (Figure 3). ChKα expression was decreased in hypoxic control PC-3 cells as compared to normoxic control PC-3 cells whereas in PC-3 cells with suppressed HIF-1α expression, hypoxia did not decrease ChKα expression (Figure 3) suggesting a possible HIF-1α involvement.

**HIF-1α binding sites and promoter alignment**

HIF-1α regulates gene expression by binding to hypoxia response core elements (HREs, 5'- CGTG - 3') in the proximal promoter region of the hypoxia responsive genes. In the promoter region upstream of human choline kinase α (hChKα), 6 previously identified HRE sites relative to +1 translation start site (19) were confirmed as HRE1 (-1723); HRE2 (-1460); HRE3 (-1027); HRE4 (-880); HRE5 (-851); and HRE6 (-825) (Figure 5). In addition to these, two putative HRE sites were newly identified as HRE2B (-1422) and HRE7 (-222). DNA sequence alignment of promoter region upstream of the hChKα (19), chimp (from NCBI), rat (20) and mouse (from NCBI) showed that out of these 8 putative HRE sites in the proximal promoter region only HRE7 positioned at -222 nucleotide was conserved across all species (Figure 5).

**Electrophoretic mobility shift assay**

An electrophoretic mobility shift assay (EMSA) was performed to assess the binding ability of HIF-1α to HRE7 in *in vitro* conditions (Figure 6). Lane 1 and 5 showed only a single band of unbound or free biotinylated probe (containing HRE7 site). With the addition of the nuclear protein extract of hypoxic PC-3 cells in the binding reaction, an extra band of decreased mobility was observed (Figure 6; Lane 2 and 6). This extra band represented the binary HIF-1 - biotinylated probe complex. The binary complex was not visible when the binding assay was performed with unlabelled probe (with HRE7 site) added 200 times in excess of the biotinylated probe (Figure 6; Lane 3), demonstrating saturability of binding. A significant decrease in intensity of the extra band was also seen when an unlabelled probe with a known functional HRE site upstream of the VEGF gene was added in 200 times in excess of the biotinylated probe (Figure 6; Lane 4). Incubation with a specific monoclonal antibody against HIF-1α prevented binding of HIF-1α to the biotinylated probe (Figure 6; Lane 7). Incubation with a specific polyclonal antibody against HIF-1α did not disrupt the HIF-1-biotinylated probe complex but formed a tertiary complex of anti-HIF-1α, HIF-1α, and biotinylated probe. This tertiary complex was seen as an extra band on the acrylamide gel that was above the band represented by the HIF-1α-biotinylated oligo binary complex (Figure 6; Lane 8). Thus, a supershift of biotinylated probe was caused due to a significant decrease in mobility of the tertiary complex.
Chromatin Immunoprecipitation (ChIP) assay

A ChIP assay was performed to determine whether HIF-1α binds to the putative HRE7 site in normoxic and hypoxic PC-3 cells. For this assay, hypoxia was mimicked by incubation with 2mM DMOG. Using traditional PCR we observed significant enrichment of promoter regions with HRE7 in the DMOG treated cells (Figure 7a). The enrichment in hypoxia and normoxia was quantified from threshold Ct values obtained by quantitative PCR (Figure 7b). Based on the quantitative PCR results, following ChIP, HRE7 showed 7.7-fold enrichment in hypoxia as compared to 2.3-fold enrichment in normoxia, suggesting increased binding of HIF-1α to HRE7 in hypoxic conditions.

Promoter isolation, cloning, mutation and luciferase assay

Further evidence for a primary role of the newly identified HRE7 site was obtained by investigating the effects of specific mutation at this site. The proximal promoter (~2.2 Kb) upstream of hChKa was successfully isolated from PC-3 genomic DNA and cloned into a luciferase vector construct. This was confirmed with nucleotide sequencing and amplification with nested primers. Mutation at the conserved HRE7 site (5'-TCGTGC-3' to 5'-AGCATT-3') was successful and confirmed by nucleotide sequencing. In normoxic conditions (21% O2 or 0mM DMOG), there was no difference in promoter mediated luciferase signal from PC-3 cell lines expressing promoter with wild type HRE7 and mutated HRE7 (Figure 8). In contrast, in hypoxic conditions (1% O2 or 2mM DMOG), a significant decrease in luciferase signal was observed in PC-3 cells expressing wild type HRE7 as compared to PC-3 cells expressing mutated HRE7(Figure 8). Thus, the HRE7 site represents a predominant role in response to hypoxia. Taken together with the previous results on the critical interaction of HIF-1α at the HRE7 site, it is strongly implied that the wild-type HRE7 but not the mutated HRE7 site is responsive to increased levels of HIF-1α in hypoxic and DMOG treated cells.

DISCUSSION

The goal of the present study was to utilize prostate cancer cells to understand the mechanisms behind low oxygen mediated reduction in rate of choline phosphorylation in cancer cells (9-11). It should be noted that in our previous study (11) we focused on the effect of anoxia (0% O2) on uptake of glucose, acetate and choline in cancer cells. The longest exposure to anoxia studied was 4h. To understand the effect of low oxygen environment on regulation of choline kinase expression (mRNA and protein), a chronic exposure is required. We found that PC-3 cells cannot be exposed to 0% O2 for more than 24h due to significant decrease in viable cell population in chronic anoxia. Chronic hypoxic exposure (1% O2) avoids this problem. A significant reduction in choline uptake was also observed in hypoxic (1% O2) cells with no loss in cell viability in PC-3 cells in the present study. Therefore, hypoxia (1% O2) was used in order to study the effect of chronic low oxygen environment on regulation of choline kinase expression (mRNA and protein) and choline metabolism. In addition, the results of our study can be compared to other literature studies as hypoxia (~1% O2) is commonly utilized (21-23). The oxygen concentration in solid tumors varies considerably (0.1 – 2% O2). Low concentration of oxygen has been reported in HGL-21 tumor (1% O2) (21), SCC-21 tumor (1% O2) (21), FaDu tumor (1% O2) (21), fibrosarcoma (0.3-1% O2) (22), 9L tumor (0.1-0.9% O2) (22), L8174T tumor (2% O2) (21), HP-555 tumor (0.7% O2) (21)
and rat mammary tumor, R3230Ac (0.5% O₂) (23). Accordingly, we chose 1% O₂ as oxygen concentration for the hypoxic condition in the present study.

The up-regulation of choline phosphorylation in normoxic tumors is well recognized but the reasons for decreased choline phosphorylation in hypoxic (~1% O₂) tumors are poorly understood. Hypoxia-mediated inhibition of choline phosphorylation has been reported in cancer cell lines derived from mouse (9), rat (10) and humans (11). In these previous studies, hypoxic treatment of the cells was followed by sample processing in normoxic laboratory conditions. This brief exposure to normoxia could mask the overall changes in cells due to hypoxia. Therefore, to rule out any artifact from re-oxygenation, we used a hypoxia chamber in our study that maintained uninterrupted hypoxia for the entire experiment including sample processing. The present results on choline tracer phosphorylation and steady-state choline metabolites in PC-3 cells confirm that hypoxia significantly reduces choline phosphorylation.

The MAR estimates from our study suggest that the choline phosphorylation step catalyzed by choline kinase was far from equilibrium in normoxic and hypoxic cancer cells. This implies that the rate of choline phosphorylation should be sensitive to changes in choline kinase expression and choline kinase activity. Our finding of intracellular ATP levels (~3mM) that are close to the Kₘ of choline kinase for ATP (~1.5 mM) (24) suggests that decreases in ATP levels in hypoxic cancer cells could also contribute to reduction in choline phosphorylation, an effect that could compound with changes in choline kinase expression and activity. In cells under moderate hypoxic stress, there are mechanisms that maintain ATP levels in a range that keeps the cell viable (25-27). This is usually done by coordinating various energy-supplying and energy-consuming processes. In our study, the level of hypoxic stress resulted in moderate decrease in cellular ATP levels, while significant increases in levels of hypoxia markers HIF-1 alpha and VEGF were observed.

In present study, we have shown that hypoxia decreases expression of ChKα in human prostate cancer cells. The decreased Chkα mRNA and protein levels correlated with decreased in vitro choline kinase activity observed in hypoxic cell extracts. These results suggest a hypoxia-induced transcriptional control of the Chkα gene. It was also shown that HIF-1α silencing in hypoxic prostate cancer cell line, PC-3 is sufficient to abolish the negative effect of hypoxia on the expression of Chkα. Our observations corroborate a HIF-1α silencing microarray study performed in a breast cancer cell line, MCF-7 that showed possible involvement of HIF-1α in down-regulation of many genes including choline kinase (28). Thus, there is compelling evidence of HIF-1α mediated negative control of choline kinase transcription in hypoxic cancer cells.

Transcriptional control of hypoxia responsive genes by HIF-1α is mediated by binding of HIF-1α to HRE sites in the regulatory promoter region of a target gene (29). In our analysis of the promoter region of the hChkα we confirmed the 6 HRE sites previously reported with the sequence 5’-G/CCGTG-3’ (19). In addition, we identified 2 novel putative HRE sites (HRE2B and HRE7) with a nucleotide sequence 5’-TCGTG-3’. This nucleotide sequence has been previously reported to bind HIF-1α in promoter region of a different hypoxia responsive gene, phosphoglycerate kinase -
To find conserved putative HRE sites that could potentially be mediating the hypoxic response across species, we aligned the nucleotide sequence for the promoter of ChKα gene from rat, mouse, chimp and human. All putative HRE core sites are not conserved across species as we found only one well conserved HRE site, HRE7 at the -222 nucleotide position. An interaction of HRE7 with HIF-1α was confirmed by in vitro and in vivo assays. In vitro binding of HIF-1α to HRE7 was confirmed by electrophoretic mobility shift assays (EMSA). Following hypoxia or DMOG exposure, significant enrichment of chromatin fragments with HRE7 in the in vivo ChIP assay, confirmed in vivo binding of HIF-1α to HRE7 in hypoxia.

The present study provides a novel contribution to the understanding of the regulation of ChKα by its promoter in hypoxia. A recent promoter deletion study by Glunde et al. (19) suggested the existence of two non-overlapping regions in the ChKα promoter that up-regulate or down-regulate ChKα expression in hypoxia. It was shown that ChKα promoter mediated up-regulation occurs in hypoxia only when the highly repressive downstream +1 to -338 nucleotides were deleted from the promoter. Based on our present work we have for the first time shown that this dominant highly repressive downstream ~338bp ChKα promoter region contains the conserved HIF-1α binding HRE7 site at the -222 nucleotide position. The role of HRE7 in transcriptional down-regulation of ChKα was confirmed when the mutation of HRE7 abrogated the transcriptional repressive control of full length ChKα promoter under hypoxia or DMOG exposure in the promoter-luciferase construct assays.

This work provides a mechanistic understanding to the observation that hypoxic regions within tumors exhibit low accumulation of radiotracer choline, for example as seen choline radiotracer studies of tumors (9-11). Choline accumulation is decreased primarily due to decrease in choline phosphorylation. Reduction in choline kinase expression, activity and reduced ATP levels in hypoxic cells contributes to the decrease in choline phosphorylation. Decreases in choline kinase expression in hypoxia are likely mediated by HIF-1α. Therefore, it is suggested that the potential impact of tumor oxygenation on choline phosphorylation in the phospholipid synthesis pathway should be considered when choline based cancer therapy and imaging applications are pursued.

CONCLUSIONS

The present study confirms that hypoxia down-regulates both ChKα expression and choline kinase activity in PC-3 human prostate cancer cells. The repression of ChKα expression is most likely resulting from transcriptional level mediation by HIF-1α at a newly identified HRE site (HRE7) at the -222 position of the ChKα promoter region. Because the choline phosphorylation reaction is far from equilibrium, small changes in ChKα expression, choline kinase activity, or intracellular ATP concentrations can sensitively modulate cellular choline and phosphocholine concentrations.
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REFERENCES


FIGURE LEGENDS

Figure 1 Effect of 24h hypoxia (1% O\textsubscript{2}) on (a) cell population doubling time and (b) [\textsuperscript{3}H]choline uptake time course in PC-3 cells (n=3, each condition). The cell population of hypoxic PC-3 cells doubled at a 75% slower rate than their normoxic counterparts (*p<0.05) and hypoxia blunted the phosphorylation dependent accumulation of [\textsuperscript{3}H]choline at later time points.

Figure 2 Representative plot of efflux of $^3$H-radioactivity from normoxic and hypoxic PC-3 cells upon successive washing with growth medium after incubating the cells for 2h with [\textsuperscript{3}H]choline under normoxia (21% O\textsubscript{2}) and hypoxia(1% O\textsubscript{2}). Hypoxia dramatically enhanced washout of $^3$H-radioactivity from the cells, demonstrating a profound decrease of metabolic sequestration of [\textsuperscript{3}H]choline.

Figure 3 Effect of hypoxia and HIF-1\textalpha silencing on HIF-1\textalpha and ChK\textalpha protein levels in PC-3 prostate cancer cell line. (a) Western blot showing expression of HIF-1\textalpha and ChK\textalpha in PC-3 cells and (b) quantitative densitometry analysis of ChK\textalpha doublet in the Western blot. The expression of ChK\textalpha was decreased in hypoxic control PC-3 cells (shRNA Control) as compared to normoxic control PC-3 cells whereas in PC-3 cells with suppressed HIF-1\textalpha expression (shRNA HIF-1\textalpha), hypoxia did not decrease ChK\textalpha expression (Figure 4) suggesting a possible involvement of HIF-1\textalpha in regulation of ChK\textalpha expression.

Figure 4 Western blot analysis showing ChK\textalpha protein levels in PC-3 prostate cancer cell lysates identified using highly specific commercially available polyclonal antibody against human ChK\textalpha. Inhibition of choline kinase expression by siRNA approach significantly reduced the intensity of ChK\textalpha protein bands as seen in (a) Western blot and (b) quantitative densitometry analysis.
Figure 5 Nucleotide alignments of segments of promoter region upstream of ChKα gene from human, chimp, rat and mouse showing 8 putative HRE sites in the promoter region from human ChKα. The schematic diagram and HRE nomenclature was adapted from Glunde et al. (19).

Figure 6 In vitro binding of HIF-1α to HRE7 within the human ChKα promoter as shown by electrophoretic mobility shift competition and supershift assay using nuclear extract from hypoxic PC-3 cells. HRE7^{CHK} – 200 fold excess of unlabeled probe with HRE7 site; HRE^{VEGF} – 200 fold excess of unlabeled probe with functional HRE upstream of VEGF; mHIF-1Ab – monoclonal anti - HIF-1α and pHIF-1Ab-polyclonal anti-HIF-1α. The experiment was performed in triplicate.

Figure 7 In vivo binding of HIF-1α to HRE7 within the hChKα promoter region in PC-3 cells as shown by ChIP assay. Following 24h exposure to normoxia (0mM DMOG) or hypoxia (2mM DMOG), chromatin was cross-linked, sonicated and immunoprecipitated with polyclonal anti-HIF-1α. PCR was performed with reverse-crosslinked chromatin samples taken before (input) and after immunoprecipitation (ChIP). Enrichment of chromatin fragment including the specific HRE in immunoprecipitated chromatin was compared to unprocessed chromatin (input) by (a) semi-quantitative traditional PCR (representative gel) and (b) fold enrichment calculated by quantitative PCR (n=5, each condition). Fold enrichment calculated by quantitative PCR was normalized with fold enrichment in negative control. Negative control used was a region upstream of VEGF gene that contained no HRE site. DMOG caused a significant fold increase in enrichment of HRE7 demonstrating HIF-1α binding to HRE7.

Figure 8 Promoter – reporter construct assay showing ChKα gene promoter dependent luciferase signal from PC-3 cells expressing wild type promoter and mutated promoter in (a) normoxia (21% O2) and hypoxia (1% O2), and in (b) presence of 0mM DMOG and 2mM DMOG. Hypoxia (*p < 0.05 vs normoxia) and DMOG (#p< 0.01 vs 0mM DMOG) reduced luciferase signal with wild type ChKα promoter but had no significant effect with the mutated HRE7. RLU = Relative Luminescence Unit = Luminescence as compared to Luminescence of promoter
construct with wild type HRE7 in normoxia (for Figure 8a) or in presence of 0mM DMOG (for Figure 8b). Means and standard deviations were calculated using 3 independent assays.
**Table 1** Metabolite levels and mass action ratio (MAR) of choline phosphorylation in PC-3 cells after 24h normoxia (21% O$_2$) and hypoxia (1%O$_2$) (n =3, each condition).

Concentration is expressed as $\mu$mol/200mg protein.

<table>
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<tr>
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<th>Choline</th>
<th>Phosphocholine</th>
<th>ATP</th>
<th>MAR</th>
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<tr>
<td>Normoxia</td>
<td>0.031 ± 0.012</td>
<td>2.1 ± 0.2</td>
<td>3.2 ± 0.2</td>
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<tr>
<td>Hypoxia</td>
<td>0.23 ± 0.06*</td>
<td>1.8 ± 0.1*</td>
<td>2.6 ± 0.1*</td>
<td>3.3 ± 0.9</td>
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*P<0.05 Vs values for normoxia, unpaired student t-test, n=3
FIGURES

Figure 1

(a) Cell population doubling time (days) in normoxia and hypoxia. (b) Choline uptake (Dose%/2h/10^6 cells) over incubation time (min) in normoxia and hypoxia.
Figure 2

Fraction of radioactivity in medium transported from cells (%) vs. Incubation time (min)

- **Normoxia**
- **Hypoxia**
Figure 3

(a) Western blot analysis showing the effects of HIF-1α shRNA on Actin, ChKα, and HIF-1α expression.

(b) Bar graph comparing arbitrary scanning units of controls (N) and HIF-1α shRNA treated (H) samples.

- Actin: No significant difference (P=0.05, n=4)
- ChKα: No significant difference (n.s., n=4)
- HIF-1α: Significant difference (P<0.05, n=4)
Figure 4

a

<table>
<thead>
<tr>
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<th>siRNA Control</th>
<th>siRNA ChKα</th>
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<tr>
<td>ChKα</td>
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<tr>
<td>Actin</td>
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</table>

b

P<0.05, n=4

Arbitrary scanning units

siRNA Control | siRNA ChKα
Figure 6

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<tr>
<td><strong>Nuclear extract</strong></td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><strong>Biotinylated HRE7(^{CHK})</strong></td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td><strong>Competitor</strong></td>
<td>−</td>
<td>−</td>
<td>HRE7(^{CHK})</td>
<td>HRE(^{VEGF})</td>
<td>−</td>
<td>−</td>
<td>mHIF1Ab</td>
<td>pHIF1Ab</td>
</tr>
</tbody>
</table>

Free Biotinylated HRE7\(^{CHK}\) →

Supershift →

Shift →
Figure 7

(a) Western blot analysis showing the expression levels of HRE 7 under different DMOG conditions. The blot includes samples for 0mM DMOG Input and ChIP, and 2mM DMOG Input and ChIP. The negative control is also included.

(b) Bar graph depicting the fold enrichment of HRE 7 with 0mM and 2mM DMOG conditions. The graph shows a significant increase in enrichment at 2mM DMOG, indicated by the asterisk.
Figure 8

(a) Normoxia (24h) vs Hypoxia (24h)

(b) 0mM DMOG (24h) vs 2mM DMOG (24h)