Identification of Novel Hypoxia Response Genes in Human Glioma Cell Line A172

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Abstract

Objectives(s): Hypoxia is a serious challenge for treatment of solid tumors. This condition has been manifested to exert significant therapeutic effects on glioblastoma multiform or (WHO) astrocytoma grade IV. Hypoxia contributes numerous changes in cellular mechanisms such as angiogenesis, metastasis and apoptosis evasion. Furthermore, in molecular level, hypoxia can cause induction of DNA breaks in tumor cells. Identification of mechanisms responsible for these effects can lead to designing more efficient therapeutic strategies against tumor progression which results in improvement of patient prognosis.

Materials and Methods: In order to identify more hypoxia regulated genes which may have a role in glioblastoma progression, cDNA-AFLP was optimized as a Differential display method which is able to identify and isolate transcripts with no prior sequence knowledge. Using this method, the current study identified 120 Transcription Derived Fragments (TDFs) which were completely differentially regulated in response to hypoxia. By sequence homology searching, the current study could detect 22 completely differentially regulated known genes and two unknown sequence matching with some Expressed Sequence Tags (ESTs).

Results: By sequence homology searching, the current study could detect 22 completely differentially regulated known genes and two unknown sequence matching with some Expressed Sequence Tags (ESTs).

Conclusion: Further characterizing of these genes may help to achieve better understanding of hypoxia mediated phenotype change in tumor cells.

Introduction

Tumor hypoxia is a common feature of most solid tumors, and serves as a critical factor in tumor progression (1-5).

Several studies have shown that impaired balance of oxygen supplies, results from structurally and functionally irregular and disrupted diffusion conditions of tumor vascular system. These conditions along with unlimited fast proliferation of tumor cells are the main causes of hypoxic stress in tumor cells (6-8). This stress condition exerts some adverse effects on neoplastic cells, in genome, transcriptome and metabolome homeostasis levels. These changes act as a natural selection force which affects tumor cells and leads to either tumor cells death or adaptation of these cells to hypoxia. Adapted cells represent a more aggressive behavior (4, 9-11). Hypoxia causes clonal expansion of tumor cells which are resistant to apoptosis, more invasive, metastatic and refractory to treatment (12-14).

As a clinical result, tumor hypoxia has direct correlation with poor prognosis in majority of solid tumors.

According to these findings, good understanding of molecular mechanisms of hypoxia mediated changes in tumor cells may help to find more effective therapeutic targets (8, 10, 12).

Glioblastoma multiform (GBM) or grade IV astrocytoma (based on World Health Organization classification) originating from glial tissue of brain, is the most common and aggressive form of intracranial primary brain tumors (8).

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Fast growing, deep infiltration of tumor cells into surrounding healthy brain parenchyma, existence of hypoxic and necrotic regions and also poor prognosis are some inherent features of these lethal tumors (7, 8).

The median survival time for patients with glioblastoma is only 12–15 months, and the 3% 5-year survival rate is significantly lower than the 60% survival rate note for other brain tumors such as oligodendroglia and medulloblastoma (10).

Usual presence of excessive hypoxic and necrotic regions indicates that hypoxic stress plays a vital role in growth and invasiveness of tumor cells.

Since hypoxia is a potent controller of gene expression, characterization of hypoxia-regulated genes is a means to study the molecular response to hypoxic stress.

The current study aimed to elaborate the current knowledge of altered gene expression in high-grade astrocytoma and to identify additional oncogenes or tumor suppressor genes involved in evolution of glioblastoma multiform tumors.

The present experiment used CDNA-AFLP method to assess A172 cell line gene expression profile in response to hypoxia condition.

Materials and Methods

Cell lines

Human Glioma cell line A172 was obtained from National Cell Bank of Iran, Pasteur Institute of Iran (Tehran, Iran). 5×10^5 cells were cultured in RPMI-1640 medium supplemented with 10% fetal bovine serum, 2 mM l-glutamine, 100 U/ml penicillin and 100 μg/ml streptomycin to 60% confluence. Cells were synchronized by overnight serum deprivation before hypoxia treatment.

Hypoxia treatment

After synchronization (as performed in triplicate), the medium were replaced by 10% FBS containing RPMI 1640 medium and cells were either placed overnight in an CO2/O2 incubator with 5% O2 and 5% CO2 at 37°C as hypoxic condition, and 21%O2, 5%CO2 at 37°C as normoxic condition.

Total RNA isolation

Total RNA from cells was extracted by using TriPure reagent (Roche) according to manufacturer’s instruction.

In summary, after removal of medium, 1 ml of TriPure reagent was added to each flask and cell lysates were homogenized by pipetting. RNA containing phase was isolated by adding 200 μl of chloroform and centrifugation. Total RNA was precipitated by adding 600 μl isopropanol and centrifugation. Finally RNA pellet was washed with 75% ethanol and after air drying; RNA was resolved in 40 μl of distilled water (DW).

DNase treatment

To eliminate genomic DNA contamination, RNA samples were treated by RNase free DNasel (promega) according to manufactures instruction. Finally, integrity and concentration of RNA samples were evaluated by agarose gel electrophoresis and spectrophotometry with Nanodrop 2000.

cDNA synthesis

1 μg of RNA was applied to cDNA synthesis by MuMLV reverse transcriptase (Fermentas) according to manufacturer’s instruction.

20 μl of single strand cDNA was used to synthesize the second strand of DNA by using Klenow fragment of E. Coli DNA polymerase I. Reaction mixture was containing 1X DNA Pol I buffer, 0.5 mM from each dNTP, 20 μl of CDNA and 80 unit of Klenow fragment. Finally, reaction mixture was adjusted to 150 μl by adding distilled water.

Reaction was performed by incubation at 16°C for 1 hr. After purification by ethanol precipitation, concentration of resulted dsDNA was evaluated by Nanodrop 2000.

Enzymatic digestion and linker ligation

dsDNA was partially digested by Mbol and Msel enzymes to obtain a heterogeneous dsDNA mixture with 100-1000 bp in length. Reaction mixture was containing ~ 1 μg of dsDNA, 3 units of each restriction enzymes and 1x of reaction buffer. Reaction was performed in 37°C for 1hr.

Digested DNA samples were purified using silica gel and applied in ligation reaction by adding AFLP Adaptors to the end of DNA fragment. Reaction mixture was containing all of double digested DNA, 0.4 μM of each adaptor, 1X ligation buffer and 5 unit of T4 ligase. Reaction volume was adjusted to 20 μl by adding distilled water. Reaction was performed in 4°C overnight.

Ligation product was purified by ethanol precipitation and used as PCR reaction template.

Preamplification and selective amplification

Amplification of DNA fragment was performed in three steps by three sets of primers (Table 1). In each step DNA product of previous step was diluted with the ratio of 1:10 in DW (distilled water). Then, 1 μl of diluted DNA was applied as template in 25 μl reaction buffer containing 1x PCR buffer, 100 μM of each dNTP, 2 mM MgCl₂, 0.2 μM of each primer and 1 unit of Taq Hot Start DNA polymerase.

PCR reaction was performed in 2720 ABI thermal cycler which is programmed for 30 cycles of denaturation at 94°C for 40 sec, 64°C for 40 sec and 72°C for 2 min; an additional step of denaturation at 95°C for 10min preceded the first cycle and another step of extension at 72°C for 7 min was extended at the end of reaction.

PCR products of hypoxia and normoxic samples from selective PCR reaction were subjected to electrophoresis on 10% polyacrylamide gel in parallel.
**Isolation, re-amplification and sequencing**

After silver staining of gels, transcription-derived fragments (TDFs) were compared in hypoxia and normoxic sample lanes. TDFs which were absent in one of the lanes, were scored as completely differentially expressed. Desired TDFs were excised from the gel and after DNA extraction, were subjected for re-amplification with a temperature profile as mentioned above.

Subsequently PCR products were electrophoresed through 2% agarose gel and then reamplified products were extracted from agarose gel by DNA extraction kit (bioneer) as recommended by the manufacturer. Finally, these fragments were sequenced and were analyzed by homology searching databases such as BLAST software.

**Results**

To identify new induced genes expressed under the hypoxic stress, a cDNA-AFLP screen was carried out.

**General and selective amplification of transcriptome**

After partial enzymatic digestion of dsDNA and ligation of adaptors into the end of DNA fragments, PCR amplification was done by three rounds of PCR.

Final PCR products were separated by polyacrylamide gel electrophoresis and visualized by silver nitrate staining (Figure 1). After evaluation of approximately 5000 TDFs resulting from different primer combinations, about 120 bands were scored as completely differentially expressed. Seventy three out of 120 TDFs were hypoxia depressed and the remaining 47 TDFs were hypoxia induced. Desired TDFs underwent DNA extraction and reamplification respectively.

By DNA sequence analysis of differentially expressed TDFs 22 known genes could be detected (Table 2). Furthermore, two TDFs were detected which matched with chromosome contig on chromosome 9 and 10. In addition, four TDFs were matched to Expressed Sequence Tags (ESTs).

![Figure 1. Expression of Hypoxia and Normoxia transcripts displayed by cDNA-AFLP. An example showing selective amplification with three different primer combination (Ms5 as forward primer and MsB5-7 as reverse primers in A1-A3 respectively) of A172 Glioma cell line exposed to Hypoxia stress. Arrows indicate transcript-derived fragments that show differential expression in either in hypoxia (h) or normoxia condition (n). M represents 100bp molecular weight marker.](image)

**Table 1. Sequence of Adaptors and PCR Primers used in cDNA-AFLP analysis**

| Name            | Forward primer | Reverse primer |
|-----------------|----------------|----------------|
| Adaptor sequences |                |                |
| Adap            | GTCGGCCTGCTGGCCGAGCCCTAG | Adap MboI      |
| MseI            | GTCGGCCTGCTGGCCGAGCCCTAG | MseI           |
| Preamplification primers | PreAmp primer |                |
| Ms1             | CCGCTGAGCGCCATTg | Mb1 GTCGGCCTGCTGGCCCTAGg |
| Ms2             | CCGCTGAGCGCCATTc | Mb2 GTCGGCCTGCTGGCCCTAGc |
| Ms3             | CCGCTGAGCGCCATTa | Mb3 CCGCTGAGCGCCCTAGa |
| Ms4             | CCGCTGAGCGCCATTt | Mb4 CCGCTGAGCGCCCTAGt |
| Ms5             | CCGCTGAGCGCCATTg | Mb5 CCGCTGAGCGCCCTAGg |
| Ms6             | CCGCTGAGCGCCATTgc | Mb6 CCGCTGAGCGCCCTAGgc |
| Ms7             | CCGCTGAGCGCCATTg | Mb7 CCGCTGAGCGCCCTAGg |
| Ms8             | CCGCTGAGCGCCATTgt | Mb8 CCGCTGAGCGCCCTAGgt |
| Ms9             | CCGCTGAGCGCCATTg | Mb9 CCGCTGAGCGCCCTAga |
| Ms10            | CCGCTGAGCGCCATTc | Mb10 CCGCTGAGCGCCCTAgt |
| Ms11            | CCGCTGAGCGCCATTca | Mb11 CCGCTGAGCGCCCTAgt |
| Ms12            | CCGCTGAGCGCCATTct | Mb12 CCGCTGAGCGCCCTAgt |
| Ms13            | CCGCTGAGCGCCATTg | Mb13 CCGCTGAGCGCCCTAgt |
| Ms14            | CCGCTGAGCGCCATTac | Mb14 CCGCTGAGCGCCCTAgt |
| Ms15            | CCGCTGAGCGCCATTaa | Mb15 CCGCTGAGCGCCCTAgt |
| Ms16            | CCGCTGAGCGCCATTat | Mb16 CCGCTGAGCGCCCTAgt |
| Ms17            | CCGCTGAGCGCCATTg | Mb17 CCGCTGAGCGCCCTAgt |
| Ms18            | CCGCTGAGCGCCATTt | Mb18 CCGCTGAGCGCCCTAgt |
| Ms19            | CCGCTGAGCGCCATTa | Mb19 CCGCTGAGCGCCCTAgt |
| Ms20            | CCGCTGAGCGCCATTt | Mb20 CCGCTGAGCGCCCTAgt |
Table 2. Summary of cDNA-AFLP analysis results

| Official symbol | Location   | Official full name                                      | Function                                                                 |
|-----------------|------------|--------------------------------------------------------|--------------------------------------------------------------------------|
| DHX8            | 17q21.31   | DEAH (Asp-Glu-Ala-His) box polypeptide 8               | nucleotide, RNA splicing                                                  |
| ODF2L           | 9q34.11    | outer dense fiber of sperm tails 2-like                | maintain the passive elastic structures and elastic recoil of the sperm  |
| TOR1AIP2 (LULL1) | 1q25.2     | torsin A interacting protein 2                         | chaperon like function                                                   |
| CTSC            | 11q13.1    | a lysosomal protease                                   | cysteine proteinase, involved in degradative processes during tumor    |
| HSP90AB2P       | 4p15.33    | heat shock protein 90kDa alpha (cytosolic), class B member 2 (pseudogene) | pseudogene                                                              |
| UBAP1           | 9p13.3     | Ubiquitin associated protein 1                         | ubiquination pathway                                                     |
| ENO2            | 12p13.31   | During neurons development                             | neuro protective                                                         |
| DDHD1           | 14q21      | DDHD domain containing 1                               | cellular trafficking transport                                            |
| CDC144B         | 17p.11.2   | coiled-coil domain containing 144B                    | pseudogene                                                               |
| RPL27A          | 11p15      | ribosomal protein L27a                                 | 60S ribosomal protein L27A                                               |
| TMEM212         | 3q26.31    | transmembrane protein 212                              | unknown                                                                  |
| XRCC2           | 7q36.1     | X-ray repair cross-complementing group 2               | homologous recombination repair (HRR) pathway                           |
| SAT1            | Xp22.1     | spermidine/spermine-N acetyltransferase-1              | Polyamines homeostasis                                                   |
| SMS             | Xp22.1.1   | homo sapiens spermine synthase (SMS), mRNA            | brain development and cognitive function                                 |
| XIAP            | Xq25       | X-linked inhibitor of apoptosis (XIAP)-associated factor 1 | X-linked inhibitor of apoptosis                                          |
| ZNF197          | 3p21       | zinc finger protein 197                                | transcription factor                                                     |
| EIF3K           | 19q13.2    | translation initiation factor 3                       | translation initiation factor 3                                          |
| MTRNR2L1        | 17p11.2    | Humanin(MTRNR2L1)                                      | cell life, antiapoptosis                                                 |
| CFLAR            | 2q33.1     | CASP8 and FADD-like apoptosis regulator                | Apoptosis regulator                                                      |
| PABPC1          | 8q22.3     | poly(A) binding protein, cytoplasmic 1-like           | mRNA translation and stability                                           |
| TOR1A           | 9q34.11    | torsin family 1, member A (torsin A)                   | synaptic functioning                                                     |
| RPLP0           | 12q24.2    | ribosomal protein, large, P0                           | protein, translation/synthesis                                           |
| POP4            | 19q13.11   | processing of precursor 4, ribonuclease P/MRP subunit  | involved in tRNA processing                                              |

Discussion

For a long time, identification of molecular pathways involved in progression and development of solid tumor were the main concern of scientists for cancer therapy. During the past decade, numerous investigations were carried out by focus on amount of oxygenation of solid tumors which result to considerable information about oxygen homeostasis in tumors. Some of these remarkable findings are: lower O₂ pressure of tumor tissue compared to origin, existence of hypoxic regions in tumors regardless of tumor features such as clinical size, grade, histology and position of tumor, as well as severe hypoxic stress in recurrent tumors compared to primary tumors (3).

An accumulating number of evidence about hypoxia-induced changes from the genomic to the posttranslational levels provide clearer insight into crucial role of oxygen homeostasis in cancer.
promotion, however, more studies are needed to precisely decode hypoxia mediated phenotype changes (15).

For better understanding of glioma cell transcriptome alterations in response to hypoxia, cDNA-AFLP was employed. By 256 primer combinations, about 5000 TDFs could be screened.

Of these bands, 120 bands were scored as completely differentially regulated. By sequencing and homology searching of these bands, 22 genes were known as hypoxia regulated.

These genes can be divided into some functional categories which will be discussed here.

**Genes regulating polyamine homeostasis**

Two of the genes induced by hypoxia were involved in polyamine homeostasis.

Spermine/Spermidine Acetyl Transferase 1 (SAT1) is a key regulator of polyamine homeostasis, which is essential in central cellular processes including proliferation and differentiation (16).

A large number of documents indicate that SAT-1 activity is under precise regulation of several microenvironment factors such as toxic agents, hormones, growth factors and also polyamines (16). Furthermore, recent evidences have shown that cell oxygen pressure directly affects enzyme activity.

According to recent studies, SAT-1 has a contradictory role in neoplastic promotion which is relevant to tumor tissue. Some investigations have shown that high level expression of SAT1, display tumor-suppressive effects while other studies point to neoplastic initiator role of SAT1 overexpression (16, 17).

The second polyamine regulating enzyme which was detected as hypoxia induced by the current experiment is spermine synthase (SMS). This enzyme catalyzes the production of the polyamine spermine from the shorter chain of polyamine spermidine (18). SMS is widely expressed in brain and is responsible for brain development and cognitive function (19). Spermine has neuroprotective effects when brain damage happens (20). Recently Nikhil et al reports that SMS gene is one of the 55 survival finger genes at chromosome 3p21 (32). This regulatory gene families in eukaryotic pre-mRNA splicing machinery and belongs to DEAD box proteins with conserved motif Asp-Glu-Ala-Asp ( DEAD) which are members of helicase super family II. DHX8 encodes a DEAD box protein which is homolog to yeast Prp22. This protein facilitates nuclear transport of mature mRNA from the 5’ snRNP of the spliceosome complex by releasing the RNA. It may have a general regulatory role in cell transcription (30, 31).

One of the most important gene expression regulatory gene families in eukaryotic cell is the zinc finger containing genes. According to the obtained results, ZNF197 gene is a hypoxia regulated gene in glioma cells. ZNF197 is located in a cluster of zinc finger genes at chromosome 3p21 (32). This gene is highly expressed in esophagus, thyroid and reproductive system. ZNF197 is also overexpressed in some thyroid papillary carcinomas. The role of this gene in mechanisms of hypoxia induced response is unknown (32-34).

Torsin A interacting protein 2a or LULL1 is another hypoxia induced gene detected in the current study. Because of sequence similarity between LULL1 protein and chaperons in AAA-domain, it seems that Torsin A interacting protein has chaperon like function in ER lumen and by conformational alteration of substrate such as Torsin A, plays an important role in brain cells (35). Also dysfunction of this transmembrane protein results in an autosomal dominant childhood-onset neurological disease DYT1 Dystonia (35, 36).

**Genes which protect cells from apoptosis**

As mentioned before, one of the most important effects of hypoxia on tumor cells is expansion of cells which are resistant to apoptosis. The current study
found two antiapoptotic genes which are overexpressed in hypoxia condition.

MTNR2L1 or Humanin (HN) is a 24-amino acid peptide with anti-apoptotic features, which was firstly identified as a neuroprotective factor against Alzheimer’s disease (AD) (37).

Several recent in vitro studies have confirmed that the HN peptide protects neurons from apoptosis by mechanisms such as inhibition of OGD-induced neuronal apoptosis, ASK/JNK mediated neuronal cell death, and mitochondrial related Bax apoptosis (38).

More recently, it has been demonstrated that HN involves in physiological mechanisms which are promoting cell survival in stressful conditions, such as neurodegeneration, inflammation or energy crisis (39, 40).

Additionally, it has been reported that Humanin increases ATP biosynthesis in human rhabdomyosarcoma TE671 cells cultured under serum-free conditions (41). So, this protein may be involved in mitochondrial related diseases or brain ischemia. Humanin, also through the STAT3 dependent anti apoptotic signal transduction cascade has correlation with oncogenesis (42).

X-linked inhibitor of apoptosis protein (XIAP) is another anti apoptotic gene which is induced by hypoxia in the current experiment. XIAP belongs to inhibitor of apoptosis proteins family of caspase inhibitors. According to recent studies, it seems that overexpression of XIAP protein has apoptosis inhibitory effect on both the initiation and execution phases of the caspase cascade and finally leads to suppression of the programmed cell death (43). Since XIAP overexpression is directly related to many diseases such as cancer progression, autoimmune and neurodegenerative disease, it is an attractive target for novel therapeutic agents for the treatment of malignancy as a new way to counteract cancer and overcome drug resistance and poor clinical outcome (44).

XIAP, by formation of a complex with the TAK1 kinase and TAB1, can result in activation of apoptotic signaling pathways including NF-κB, JNK, MAP kinase, and the ubiquitin proteosome pathways (45). Furthermore, XIAP plays a crucial role in a variety of cellular functions including immune regulation, cell division and differentiation, cell migration, morphogenesis, and heavy metal metabolism (44).

Especially it has been reported that XIAP is overexpressed in glioblastoma (46), and it is supposed to be associated with drug resistance and poor prognosis of these patients.

The current study also identified CFLAR induction by hypoxia. CFLAR (CASP8 and FADD-like apoptosis regulator) gene located on 2q33.1 which encodes a regulatory protein of apoptosis pathway and plays a significant role in the regulation of apoptosis and is structurally similar to caspase-8 which blocks death receptor-mediated apoptosis by inhibiting caspase 8 (47).

Genes involved in DNA repair

One of the well-known effects of hypoxia stress is loss of genomic integrity and DNA breakage. So, hypoxic cells adapt with this environmental pressure by over activating of DNA repair systems.

X-ray repair cross-complementing group-2 (XRCC2) gene is an important member of mammalian RecA/Rad51 family proteins which are related to the homologous recombination repair (HRR) pathway required for correct chromosome segregation and the repair of DSBs (48). Because of its interaction with some tumor suppressor genes such as brcal and brc2, c-abl, p53, it seems that XRCC2 is involved in tumor progression (48, 49).

Several investigations point to the role of XRCC2 mutations and polymorphisms in various cancers such as breast cancer, colorectal cancer, brain cancer, oral and Lynch syndrome and esophageal adenocarcinoma (50).

Overexpression of X-ray repair cross-complementing group 2(XRCC2) is a hallmark of neoplastic cells and especially in glioblastoma tumor cells leads to resistance against anticancer drugs Temozolomide (TMZ) (51).

Cancer/testis specific gene

ODF2 cancer/testis gene was found as a de novo transcriptional target of hypoxia. ODF2 is one of the major components of the cytoskeleton of the sperm tail.

ODF2 by interaction with other proteins, like Plk1, plays a crucial role in spindle formation and the progression into mitosis (52).

ODF2 has a testis restricted expression pattern and does not express in normal tissues. Ectopic expression of this gene has been detected in prostate cancer and basal cell carcinoma.

In addition to these results, the current study detected hypoxic induced expression of 4 EST and also 2 chromosome contig sequences, which indicate the presence of other unknown genes in hypoxic response of cells.

cDNA-AFLP which was first developed by Vos et al. (1995) is a powerful approach that facilitates identification and isolation of transcripts (53). Also, it can discover any variation between two global expression patterns without pervious knowledge of sequence. In addition, it is able to amplify any infrequent transcripts from transcriptome pool.

Conclusion

The present study found significant variations in gene expression among hypoxia and normoxic cultured glioma cells. Some of these results are reported for the first time and when further characterized may give us precise insights to the process of transcriptome re-modeling by hypoxia.
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