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Abstract

Mechanism for enhanced 5-aminolevulinic acid fluorescence in isocitrate dehydrogenase 1 mutant malignant gliomas

Ja Eun Kim

Department of Medicine and Major in Neuroscience
The Graduate School
Seoul National University

Fluorescence-guided surgery using 5-aminolevulinic acid (5-ALA) has become main modality in the treatment of malignant gliomas. However unlike glioblastomas, there are inconsistent result about fluorescence status in WHO grade III gliomas. Here, we have shown that mutational status of IDH1 is bind to 5-ALA fluorescence, and investigated its mechanism. Using genetically engineered malignant glioma cells harboring wild type (U87MG-IDH1\textsuperscript{WT}, U87MG-IDH2\textsuperscript{WT}) or mutant (U87MG-IDH1\textsuperscript{R132H}, U87MG-IDH2\textsuperscript{R132K}) versions of IDH1 and IDH2, we confirmed a lag in 5-ALA metabolism and a temporary accumulation of protoporphyrin IX (PpIX) in U87MG-IDH1\textsuperscript{R132H} and U87MG-IDH2\textsuperscript{R132K} cells. To investigate the metabolic aspects of the mechanism responsible, we used liquid chromatography–mass spectrometry (LC-MS) to
screen for tricarboxylic acid (TCA) cycle-related metabolite changes resulting from 5-ALA exposure. We observed low baseline levels of NADPH, an essential cofactor for the rate-limiting step of heme degradation, in U87MG-IDH1<sup>R132H</sup> cells. Abundant levels of NADPH are required to metabolize excessive 5-ALA, giving a plausible reason for the temporary enhanced 5-ALA fluorescence in mutant IDH1 cells. This hypothesis was supported by the results of metabolic screening in human malignant glioma samples. In conclusion, we have discovered a relationship between enhanced 5-ALA fluorescence and IDH1 mutations in WHO grade III gliomas. Low baseline levels of NADPH in tumors with mutated IDH1 is responsible for the enhanced fluorescence from the metabolic aspects of the mechanism.

Keywords : 5-ALA, fluorescence, IDH1, glioma, NADPH

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List of Abbreviations and symbols

5-ALA: 5-aminolevulinic acid

LC-MS: liquid chromatography–mass spectrometry

TCA: tricarboxylic acid

PpIX: protoporphyrinogen IX

IDH1/2: isocitrate dehydrogenase 1/2

α-KG: α-ketoglutarate

2-HG: R-2- hydroxyglutarate

MRMs: multiple reaction monitorings

BCA: bicinchoninic acid

RFUs: relative fluorescence units

HO: heme oxygenase

ABCB6: ATP-binding cassette sub-family B member 6

AA: anaplastic astrocytoma

AO: anaplastic oligodendroglioma

PKM2: pyruvate kinase isoform M2

ROS: Reactive oxygen species

MRS: magnetic resonance spectroscopy

FACS: fluorescence activated cell sort
Introduction

Among recent advances in malignant glioma surgery, fluorescence-guided surgery is becoming popular due to its ability to improve the extent of resection while preserving patient functional status [1, 2]. 5-aminolevulinic acid (5-ALA) highlights otherwise indistinguishable malignant areas with a temporarily red fluorescence under an appropriate blue light source with a specific wavelength [3]. 5-ALA itself is a non-fluorescent endogenous compound that can be metabolized intracellularly into protoporphyrinogen IX (PpIX), a fluorescent intermediate product of the heme synthesis pathway [4]. Although the exact mechanism responsible has yet to be verified, exogenous administration of 5-ALA leads to selective accumulation of PpIX in malignant gliomas but not in low-grade gliomas or normal brain tissue [5]. The 5-ALA-induced fluorescence rate is lower in WHO grade III gliomas (83% in anaplastic oligodendroglioma/oligoastrocytomas, and 22% in anaplastic astrocytomas) than grade IV gliomas (96% in glioblastomas) [6]. The reason why grade III gliomas have a lower 5-ALA fluorescence rate than grade IV gliomas is unknown.

The identification of isocitrate dehydrogenase 1 (IDH1) mutations in glioma is one of the major discovery mediating metabolomics and oncogenesis [7]. The majority of IDH1 mutations occur at the active site arginine 132, the most common substitution being histidine (R132H) [7-10]. The role of IDH1 is to catalyze the oxidative decarboxylation of isocitrate into α-ketoglutarate (α-KG), simultaneously converting NADP(+) to NADPH in the cytoplasm and
peroxisomes [11, 12]. IDH1 mutations in glioma cells diminish the canonical enzymatic activity of IDH1 while conferring a neomorphic enzymatic activity: the production of R-2-hydroxyglutarate (2-HG) via the consumption of NADPH [13-16]. The IDH1 mutation is observed in 6% of WHO grade IV glioblastomas and 55% of grade III gliomas [17].

Endogenously, 5-ALA is produced from succinyl-CoA, which is converted from α-KG [18]. Based on the idea that IDH1 mutations can alter the 5-ALA metabolic pathway and stimulate exogenous 5-ALA fluorescence, we confirmed an association between the R132H IDH1 mutation and enhanced 5-ALA fluorescence in WHO grade III gliomas, then investigated metabolic aspects of the underlying mechanism. Although the IDH1 mutation is not the core mechanism of 5-ALA fluorescence considering the discrepancy between 5-ALA fluorescence and IDH1 mutation prevalence in grade II or IV gliomas, we showed that the IDH1 mutation can act as a sensitizer when the cell is under the condition of altered 5-ALA metabolism.
Materials and Methods

1. Acquisition of patient and sample data

We used a cohort of 49 patients with histologically-proven WHO grade III gliomas to acquire clinical data and samples, based on a program approved by the Institutional Review Board. For fluorescence-guided surgery, 20 mg/kg of 5-ALA (Gliolan®; Medac, Wedel, Germany) was administered orally 3–4 hours before the induction of anesthesia. The main procedure was carried out at 5–7 hours after 5-ALA intake. 5-ALA positive fluorescence was confirmed if there were any red fluorescent spots in the tumor under blue-violet light. The fluorescence status of each area was documented as fluorescence positive or negative by the performing neurosurgeon. The tumor was defined fluorescence positive at least a focal area was identified as positive fluorescence. Positive areas were collected separately for histological diagnosis and study. All tumor samples used in this study were snap-frozen in liquid nitrogen as soon as possible during the surgery and stored at -80°C. Fluorescence-guided surgeries were done with a Leica M720 OH5 microscope (Leica, Wetzlar, Germany) equipped with an FL400 Fluorescence module or a Zeiss Pentero equipped with a fluorescent 400 nm UV light and filters (Zeiss, Oberkochen, Germany).

Histological diagnosis was performed using the criteria described in the 2007 WHO Classification of Tumours of the Central Nervous System [39]. IDH1 mutations in patient samples were detected via targeted sequencing, as described previously.[40]
2. Construction and establishment of wild type and mutant IDH1 cell lines

Human IDH1 (GenBank accession number NM_005896) and IDH2 (GenBank accession number NM_002168) cDNA Clone were obtained from Origene (Rockville, MD, U.S.A), and amplified using the polymerase chain reaction. To generate the IDH1\textsuperscript{R132H} construct, the G395A base pair change was introduced in the IDH1 open reading frame, and to generate the IDH2\textsuperscript{172K} construct, the G515A base pair change was introduced in the IDH2 open reading frame. Amplified wild type, mutant IDH1 and mutant IDH2 constructs were subcloned into lentiviral vector CD526A-1 (System Biosciences, Mountain View, CA, U.S.A) via a proprietary Eco cloning method. This vector contains a green fluorescent protein (GFP)-Puromycin fusion dual marker under a Rous sarcoma virus (RSV) promoter, and the cloned insert is expressed under an enhanced, constitutive cytomegalovirus (CMV) promoter (Fig. 1A). The primers designed for IDH1/2 cDNA cloning were

\[
\text{IDH1-F}, \; 5'\text{-TCCAGCCTCCGGACTCTAGA-3'}, \\
\text{IDH1-R}, \; 5'\text{-AGCAGATACTGGCTTAACTA-3'} \\
\text{IDH2-F}, \; 5'\text{-GCTGCAGTGGGACCACATT-3'}, \\
\text{IDH2-R}, \; 5'\text{-TGTGGCCTTGTACTGCAGAG-3'}. \\
\]

The sequence of the cloned cDNA was confirmed by direct DNA sequencing. A control sequence was cloned in the same vector. The control vector could produce a mock virus (not expressing any target, but with the same marker expressed), which was used for negative controls. The recombinant lentivirus...
was produced by GenTarget Inc (http://www.gentarget.com/, San Diego, CA, USA). Expression lentiviral particles were produced in 293T cells in DMEM medium, collected, and filtered through a 0.45 μm membrane filter (Millipore, Billerica, MA, U.S.A.) before being immediately stored at −70°C. Virus titers were measured via GFP-positive cell counting after transduction into TH1080 cells; the concentration titers were 1.21×10⁷ IFU/mL (mock), 1.09×10⁷ IFU/mL (IDH1<sup>WT</sup>), and 1.17×10⁷ IFU/mL (IDH1<sup>R132H</sup>).

Human U87MG cells were purchased from the American Type Culture Collection (ATCC, Rockville, MD, USA) and maintained in Roswell Park Memorial Institute 1640 medium (RPMI; WELGENE) containing 10% fetal bovine serum (FBS; GIBCO) in a humidified incubator at 37°C and 5% CO₂. To express IDH1/2<sup>WT</sup>, IDH1<sup>R132H</sup> or IDH2<sup>172K</sup> in cells, U87MG cells were transduced with lentivirus for 24 hours in the presence of 4–8 µg/mL polybrene. Transduced cells were selected using puromycin (Cat. No. A7793, Sigma-Aldrich, USA), and transduction efficacy was analyzed via fluorescence microscopy using green filters (Leica, Wetzlar, Germany) and by fluorescence activated cell sorting (FACS) analysis (Fig. 1B and 1C). The transduction efficiency of U87MG cells expressing GFP abundantly was over 90% using the lentiviral vector. GFP fluorescence in transduced cells was analyzed using a FACS Calibur cell sorter (BD Bioscience, San Jose, CA, USA) equipped with a 530-nm filter (bandwidth, 15nm), a 585-nm filter (bandwidth, 21 nm), and Cell-Quest software (BD Bioscience). Sorted cells were used for further studies.

Direct sequencing of DNA from each cell line showed the expected sequences for IDH1/2<sup>WT</sup>, IDH1<sup>R132H</sup> or IDH2<sup>172K</sup> (Fig. 1D). We confirmed the presence of
IDH1/2\textsuperscript{WT}, IDH1\textsuperscript{R132H} or IDH2\textsuperscript{172K} expression via Western blot (Fig. 1E). Cells were lysed in ice-cold lysis buffer [20 mM Tris-HCl (pH 7.5), 150 mM NaCl, 1 mM Na\textsubscript{2}EDTA, 1 mM EGTA, 1% Triton, 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, 1 mM Na\textsubscript{3}VO\textsubscript{4}, 1μg/mL leupeptin, and protease inhibitor cocktail (Sigma)], and the concentration of lysate protein was evaluated using the BCA method (Cat. No. 23227, Pierce\textsuperscript{TM} BCA Protein Assay Kit, Thermo scientific, Rockford, IL, USA). Approximately 30 μg protein was loaded in each lane of a polyacrylamide denaturing gel for electrophoresis. After electrophoresis, protein was transferred to nitrocellulose membranes for blotting. We used a rat monoclonal anti-IDH1 antibody (Cat No. DIA-W09, Dianova, Hamburg, Germany), a mouse monoclonal anti-IDH1 R132H antibody (Cat No. DIA-H09, Dianova, Hamburg, Germany), a rabbit polyclonal anti-IDH2 antibody (Cat No. 15932-1-AP, Proteintech, USA), a mouse monoclonal anti-IDH2 R172K antibody (Cat No. 26163, Neweastbio, PA, USA) and a rabbit polyclonal antibody to β-actin (Abcam, Cambridge, England). Primary antibodies were detected using horseradish peroxidase-conjugated antibodies (Santa Cruz Biotechnology, CA, U.S.A.).

Since aberrant overproduction of 2-HG is a hallmark of malignant gliomas with IDH1 mutations [13], we used LC-MS to confirm elevated 2-HG levels in U87MG- IDH1\textsuperscript{R132H} cell lines vs. U87MG- IDH1\textsuperscript{WT} or U87MG-mock cell lines (Fig 1F).

3. In vitro 5-ALA treatment

We purchased 5-ALA from Sigma-Aldrich (Cat. No. A7793, USA) and
dissolved it in phosphate buffered saline (PBS, pH 7.4) to prepare a 500 mM stock solution. It was sterilized by filtration using a 0.2 um pore filter, and wrapped in aluminum foil to avoid light exposure and stored at −20°C. Next, 5×10⁴ cells were seeded into 24-well plates and incubated in 500 μl media overnight. Cells were incubated with 5-ALA (final concentration 1 mM) for 1 hour, rinsed twice with PBS, then incubated in complete medium until analysis. Special care was taken to avoid exposure to light during the experiments.

4. Quantification of PpIX fluorescence

To quantify intracellular PpIX fluorescence, we followed a previously described method with modifications [41]. Specifically, after 5-ALA treatment, cells were lysed with 60 μl 0.2% (vol/vol) Triton X-100 on ice for 5 minutes and centrifuged at 13,000 rpm for 5 minutes at 4°C. Next, 5 μl cell supernatant was transferred for the BCA Protein Assay to normalize detected relative fluorescence units (RFUs) of PpIX. To detect PpIX fluorescence, 100 μl methanolic perchloric acid, consisting of 5.6% (vol/vol) perchloric acid in 50% (vol/vol) methanol, was added to the sample and incubated at 37°C in darkness for 15 minutes, then centrifuged at 13,000 rpm for 5 minutes at 4°C. Then, 100 μl supernatant was transferred into 96-well black plates, and fluorescence intensity was measured using the Infinite M200 PRO (TECAN, Switzerland) and Magellan™ software at a fluorescence excitation wavelength of 400±9 nm and an emission wavelength of 645±20nm.

To visualize PpIX fluorescence, microscope coverslips (12 mmΦ, Cat No. 01
115 20, Labnet, Germany) were coated with 0.1% gelatin in a 37°C incubator for 1 hour and washed with PBS. Next, $2 \times 10^4$ cells were plated on the coverslips in 24-well plates. A 5-ALA solution (final concentration 1 mM) was added, the 24-well plates were wrapped in aluminum foil to avoid light exposure for 1 hour, and were then rinsed with PBS. After incubation for 2 hours, cells on the coverslip were washed with PBS twice and fixed with 4% paraformaldehyde in PBS at room temperature for 10 minutes, washed again with PBS twice, and mounted using Slow Fade Gold (Cat No. S36936, Invitrogen, USA). Cells were analyzed under 400-fold magnification with a confocal laser scanning microscope (LSM 510 META, Carl Zeiss, Germany). PpIX fluorescence was excited at 405 nm, and images were collected in the red channel using a 560 nm long-pass filter.

5. Metabolite screening via LC-MS

Thawed samples were centrifuged at 21,000 g for 20 minutes to remove insoluble materials, then supernatants were transferred into glass vials with micro-inserts for small volume injections. Analysis was carried out using an Agilent 1100 Series liquid chromatography system equipped with a binary LC pump, a degasser, and an auto-sampler set at 4°C (Agilent, CO, USA). Chromatographic separation was performed by injecting 2 μl sample on a ZIC-pHILIC Polymeric Beads Peek column (150×2.1 mm, 5 μm, Merck, Germany) at 35°C and a 0.15 mL/min flow rate, using 10 mM ammonium carbonate (pH=9.0) in distilled water as mobile phase A and acetonitrile (ACN) as mobile
phase B. The linear gradient was as follows: 80% B at 0 minutes, 35% B at 10 minutes, 5% B at 12 minutes, 5% B at 25 minutes, 80% B at 25.1 minutes, and 80% B at 35 minutes.

During the LC-MS experiments, mass spectra were acquired in the negative ion mode using an API 2000 Mass Spectrometer (AB/SCIEX, Framingham, MA, USA) equipped with an electrospray ionization (ESI) source. Parameters of the ESI source operation were as follows: -4.5 kV of ion spray voltage; curtain gas (nitrogen), ion source gas 1 (nitrogen), and ion source gas 2 (nitrogen) pressures at 30, 40, and 80, respectively; temperature of the heater (turbo) gas at 350°C.

MRMs were performed and controlled with Analyst 1.6 Software. Chromatographic peaks were identified by comparing retention times, m/z values and MS2 fragmentation patterns with those in the HMDB (www.hmdb.ca), METLIN (metlin.scripps.edu) and Massbank (www.massbank.jp) databases and those of standards.

**6. Statistical Analyses**

Significance of deviance was indicated by Pearson residuals ≥2 or ≤-2, calculated from a chi-square test and expressed by a mosaic plot. All statistical analyses were performed using the free statistical software package R (version 3.0.2; [http://www.r-project.org/](http://www.r-project.org/)), and IBM SPSS statistics software (version 21; SPSS Inc). Statistical significance was accepted at a level of p<0.05 (*), p<0.01 (**) or p<0.001 (**). Error bars in histograms represent 95% confidence
intervals. Quantitative heat maps visualizing the relative expression of metabolites were produced using Gene Pattern software modules (www.broadinstitute.org/cancer/software/genepattern; Comparative Marker Selection version 10, Hierarchical Clustering version 6, and Heat Map Viewer version 13) [42].
Results

Clinical evidence of an association between IDH1 mutations and 5-ALA fluorescence

Among 35 patients with WHO grade III gliomas who were operated upon using 5-ALA fluorescence-guided surgery, IDH1 mutations were observed in 24 and intraoperative fluorescence was observed in tumor tissues of 19 patients (Table 1). There was a significant association between IDH1 mutations and 5-ALA fluorescence in tumor tissue (p=0.03). The distribution of patient data and associations between variables are shown in a mosaic plot (Fig. 2).

5-ALA metabolism lags in IDH1/2 mutant malignant glioma cell lines

To verify the relationship between IDH1 mutations and 5-ALA fluorescence, we established human U87 MG glioma cells that stably expressed R132H mutant (IDH1\textsuperscript{R132H}), wild type (IDH1\textsuperscript{WT}) IDH1, R172H mutant (IDH2\textsuperscript{R172K}) or wild type (IDH2\textsuperscript{WT}) IDH2, using lentiviral vectors (Fig. 1). We used U87MG cells transduced only with vector as controls (U87MG-mock). We also confirmed that U87MG-IDH1\textsuperscript{R132H} cells produce 2-HG at significantly higher levels than U87MG-IDH1\textsuperscript{WT} cells (Fig. 1).
Figure 1: Establishment of wild type and mutant IDH1/2 cell lines using U87MG cells and a lentiviral vector. A. Schematic drawing of the lentiviral IDH1/2 gene construct. B. Transduction efficacy of vector containing the IDH1/2 gene construct, evaluated by GFP. C. FACS results showing high purity of GFP-positive cells. D. Direct sequencing analysis confirming expected sequences in U87MG-IDH1WT, U87MG-IDH1R132H, U87MG-IDH2WT, and U87MG-IDH2R172K cells. E. Western blot results showing expression of IDH1WT, IDH1R132H, IDH2WT, and IDH2R172K from U87MG-IDH1WT, U87MG-IDH1 R132H, U87MG-IDH2WT, and U87MG-IDH2R172K cells, respectively. F. 2-HG levels in U87MG-IDH1R132H cell extracts were approximately 18-fold the levels in U87MG-IDH1WT or U87MG-mock cells.
Table 1: Association between intraoperative 5-ALA fluorescence and IDH1 mutations in WHO grade III glioma patients.

<table>
<thead>
<tr>
<th></th>
<th>Intraoperative fluorescence</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Mutated IDH1</td>
<td>16</td>
<td>8</td>
</tr>
<tr>
<td>Wild type IDH1</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td>19</td>
<td>16</td>
</tr>
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P=0.03

Figure 2: Mosaic plot for 5-ALA fluorescence, IDH1 mutations, and histological diagnosis (n = 35). The colored cells show the magnitude of the Pearson residuals obtained from an independence model. (AA, anaplastic astrocytoma; AO, anaplastic oligodendroglioma; AOA, anaplastic oligoastrocytoma.)
From our clinical experience with 5-ALA fluorescence-guided brain tumor surgery, we hypothesized that the difference in fluorescence between malignant glioma and normal brain tissue is a temporary event best observed between 3 and 9 hours after 5-ALA administration [1]. Thus, we performed an in vitro time-course analysis measuring 5-ALA-derived PpIX concentrations in a glioma cell line after exposure to 5-ALA. After a 1-hour incubation with 5-ALA, the fluorescence intensity of intracellular PpIX was measured at various time points between 0 to 4 hours with a fluorescence plate reader and normalized to total cell protein concentrations measured using the bicinchoninic acid (BCA) protein assay. Intracellular PpIX fluorescence increased rapidly up to 1 hour after incubation, then diminished in U87MG-IDH1/2\(^{WT}\) cells (Fig. 3A, 4A), whereas there was a delay in 5-ALA metabolism in U87MG-IDH1\(^{R132H}\), U87MG-IDH2\(^{R732K}\) cells; PpIX fluorescence peaked at 2 hours after incubation, at which time there was a significant difference in PpIX concentration between U87MG-IDH1\(^{R132H}\) and U87MG-IDH1\(^{WT}\) cells (Fig. 3B) or between U87MG-IDH1\(^{R172K}\) and U87MG-IDH2\(^{WT}\) cells (Fig. 4B). This difference was associated with differences in fluorescence visualization as well (Fig. 3C, 4C). These results indicate that the IDH1/2 mutation is associated with a lag in 5-ALA metabolism, resulting in temporary differences in the fluorescence activity of malignant glioma cells.
Figure 3. In vitro evidence for an association between IDH1 mutations and 5-ALA fluorescence. A. PpIX concentrations after 5-ALA treatment, expressed as relative fluorescence units (RFUs) normalized against total cell protein levels. B. Accumulation of intracellular PpIX 2 hours after 5-ALA treatment. C. Confocal laser scanning microscope images taken 2 hours after 5-ALA treatment. PpIX is visible as red fluorescence, located mainly in the cytoplasm. (λex=405nm, λem>500nm)
Figure 4. *In vitro* evidence for an association between *IDH2* mutations and 5-ALA fluorescence. A. PpIX concentrations after 5-ALA treatment, expressed as relative fluorescence units (RFUs) normalized against total cell protein levels. B. Accumulation of intracellular PpIX 2 hours after 5-ALA treatment. C. Confocal laser scanning microscope images taken 2 hours after 5-ALA treatment. PpIX is visible as red fluorescence, located mainly in the cytoplasm. (λex=405nm, λem>500nm)
Supplementary Figure 1. Confocal laser scanning microscope images after 5-ALA treatment in U87MG-IDH2 cell lines. A–F: Accumulation of intracellular PpIX 0–4 hr hours after 5-ALA treatment or absence of 5-ALA.
Citrate, α-KG, and 2-HG levels are selectively altered in IDH1 mutant malignant glioma cells upon exposure to 5-ALA

To screen for metabolic changes that could explain the difference in 5-ALA metabolism between U87MG-IDH1\textsuperscript{R132H} and U87MG-IDH1\textsuperscript{WT} cells, we profiled tricarboxylic acid (TCA) cycle intermediates in the lysates of both cell lines, with or without 5-ALA exposure, using liquid chromatography–mass spectrometry (LC-MS). A total of 17 multiple reaction monitorings (MRMs) were performed, and values of detected metabolites were normalized to total protein levels in the cell, as measured by the BCA method. To select candidate metabolites related to enhanced 5-ALA fluorescence in IDH1 mutant malignant glioma cells, we identified three metabolites (citrate, α-KG, and 2-HG) of which the levels were significantly altered in U87MG-IDH1\textsuperscript{R132H} cells after 5-ALA treatment (Fig. 5). After 5-ALA treatment, citrate increased (3.5 fold; p<0.001) and α-KG decreased (0.6 fold; p<0.001) in U87MG-IDH1\textsuperscript{R132H} cells but not in U87MG-IDH1\textsuperscript{WT} cells. Non-treated IDH1 mutant glioma cells produce an excess of 2-HG relative to wild type glioma cells (7.8 fold; p<0.001); 2-HG levels increased further after 5-ALA treatment (1.7 fold; p<0.005). Levels of other observed TCA intermediates were similar in U87MG-IDH1\textsuperscript{WT} and U87MG-IDH1\textsuperscript{R132H} cells.
Figure 5. Heat map illustrating the relative ratio of the metabolites expressed in U87MG cells with vector only, the IDH1 wild type construct, and the IDH1 mutant construct. Metabolites whose levels were significantly different in IDH1 mutant cells but not wild type cells after 5-ALA treatment are shown in separate histograms.
Hypothesis for the mechanism of enhanced 5-ALA fluorescence in IDH1 mutant malignant glioma cells

To develop a hypothesis regarding the mechanism of enhanced 5-ALA fluorescence in IDH1 mutant malignant glioma cells, we reviewed crosstalk between the TCA cycle and the heme synthesis pathway (Fig. 6A). Succinyl-CoA from the TCA cycle and glycine are combined by ALA synthase to form 5-ALA [19-23]. ALA synthase is also believed to be the rate-limiting synthetic enzyme in the heme synthesis pathway and is negatively regulated by heme, which is usually degraded by heme oxygenase (HO), via a feedback mechanism [19]. In this step, HO degrades heme in concert with NADPH cytochrome P450 reductase, and it has been shown that HO activity can be increased by NADPH or NADH [24]. Interestingly, one production source of NADPH is the conversion of isocitrate to α-KG by IDH1 or IDH2. Moreover, IDH1/2 mutations not only decrease the production capacity of NADPH, but also increase consumption of NADPH during the aberrant production of 2-HG from α-KG, a NADPH-dependent reduction process [13]. We confirmed that baseline NADPH levels were lower in U87MG-IDH1<sup>R132H</sup> cells than U87MG-IDH1<sup>WT</sup> cells (0.3 fold; p<0.05) and that NADPH was almost depleted after 5-ALA treatment in both cell lines (Fig. 6B). Therefore, we may explain the lag in 5-ALA metabolism in IDH1 mutant malignant glioma cell lines by invoking alterations in NADPH metabolism (Fig. 6A). If exogenous 5-ALA is administered, the heme pathway is normally hyperactivated to digest the excess 5-ALA; heme is degraded unrestrictedly by HO, given sufficient
Figure 6. A. Schematic of our hypothesis that the lag in 5-ALA metabolism in IDH1 mutant cells is due to insufficient NADPH. B. Intracellular NADPH levels measured by LC-MS. C. Schematic illustration of the metabolic effects of 5-ALA treatment in IDH1WT and IDH1218H cells.
NADPH, with no accumulation of PpIX. However, if IDH1 is mutated, heme degradation is disturbed as a result of insufficient NADPH for HO activity, resulting in underexpression of ferrochelatase and, in turn, accumulation of PpIX. This hypothesis is supported by our data on alterations in metabolite levels (Fig. 5). In U87MG-IDH1WT cells, no interference exist from the TCA cycle with the heme synthesis pathway, whereas the conversion of isocitrate to α-KG proceeds normally (Fig. 6C). However, ineffective conversion of isocitrate to α-KG, the production of 2-HG from α-KG, and retarded metabolism at the junction of the TCA cycle and heme synthesis pathway due to insufficient NADPH in U87MG-IDH1R132H cells leads to accumulation of citrate and the depletion of α-KG (Fig. 6C).

Validation of metabolic changes in human WHO grade III gliomas

We created an independent validation set of 14 WHO grade III gliomas treated with 5-ALA fluorescence-guided surgery. The 7 samples were with mutated IDH1 and positive fluorescence, and the other 7 samples were with wild-type IDH1 and negative fluorescence. We measured metabolites in frozen tumor samples using LC-MS. Among the 13 available MRMs with normalized values, there was a significantly higher level of citrate in fluorescence positive samples than negative ones (Fig. 7, p<0.001). However, NADPH, α-KG, and succinyl-CoA were not detected in any of the samples, probably because of excessive consumption due to 5-ALA-induced metabolism.
Figure 7. Citrate levels in tumor samples, measured by LC-MS. All samples were taken after 5-ALA administration and grouped according to optical evidence of intraoperative fluorescence (AA, anaplastic astrocytoma; AO, anaplastic oligodendroglioma). Significant differences were observed between wtIDH1 without fluorescence group and mutIDH1 with fluorescence group (mean value: 167639.8 vs 525871.0, p<0.001).
Discussion

Fluorescence-guided surgery using 5-ALA is gaining wide use in malignant glioma treatment. Although researchers have been actively investigating the mechanism at work, the definitive answer is not in our hands yet [25-28]. Most plausible studies explaining the molecular mechanism have focused on expression changes in related enzymes, such as ferrochelatase, or transport molecules of intermediates such as ABCB6 [27, 28]. However, no study has yet tried to investigate the mechanism of 5-ALA fluorescence in malignant gliomas from the perspective of metabolomics. The current study is the first to document the relationship between IDH1 mutations and 5-ALA fluorescence and to suggest a mechanism on the basis of metabolic profiling. It is natural to approach the mechanism of 5-ALA fluorescence in malignant gliomas from the standpoint of view of metabolomics, because the differences of fluorescence between tissues are temporary event indicating differences in metabolism-processing capacity among the tissues. It is a matter of turnover rate of PpIX, rather than accumulation itself, that matters the mechanism of 5-ALA fluorescence. We propose that NADPH is the key mediator in the varying 5-ALA fluorescence observed between wild type and mutant IDH1 malignant glioma cells. In the present study, the heme biosynthesis pathway lags in IDH1 mutant malignant gliomas; 5-ALA overload due to insufficient NADPH can result in a temporary difference in PpIX accumulation relative to wild type IDH1 malignant glioma cells. It has been shown previously that mutant IDH1
is responsible for the inhibition of the mutual conversion of isocitrate and α-KG in a NADP+ or NADPH-dependent manner [11, 13, 29, 30]. Recent research implies that reduction of the NADPH production capacity in IDH1 mutant cells may be a contributing factor in cancer formation [31, 32]. And also, NADPH has been suggested as a promising therapeutic target in glioblastoma cells [33].

*IDH1/2* mutations in glioma are thought to be principal events in early-stage gliomagenesis, and their oncometabolite, 2-HG, is responsible for multiple activities involved in tumor development [34, 35]. In addition, 5-ALA fluorescence is strongly associated with phenotypic tumor aggressiveness in gliomas [5]. Therefore, if there are metabolic changes linking *IDH1* mutations and 5-ALA fluorescence, they may implicate important metabolic steps reflecting malignant transformation in gliomas. One of the key molecules involved in metabolic alterations in cancer is NADPH, which functions as a cofactor in many enzymatic reactions crucial for macromolecular biosynthesis and as a defender against reactive oxygen species (ROS) produced during rapid cancer cell proliferation [36]. There are two sources of NADPH production: the TCA cycle and the pentose phosphate pathway [37]. In the TCA cycle, IDH1/2 and malic enzyme 1 (ME1), of which the levels are altered in many type of cancer cells, contribute to NADPH production [36]. Alternately, overexpression of pyruvate kinase isoform M2 (PKM2) in various type of cancer cells can divert metabolic precursors into the pentose phosphate pathway to produce NADPH [36]. Controlling increased levels of ROS with NADPH is crucial for the survival of cancer cells. Moreover, growing evidences indicate
that oncogenesis of IDH1 mutated tumor involves reduced production of NADPH as well as oncometabolite of 2HG production [38]. The role of NADPH in the present study provides insight into its use as a potential target for diagnostic and therapeutic purposes in the future. In that NADPH is more easily detectable than 2-HG by non-invasive diagnostic modalities such as magnetic resonance spectroscopy (MRS), development of NADPH as a surrogate marker for IDH1 mutation and predictive marker of 5-ALA fluorescence can be a possible example of clinical applications.

We admit that the mechanism suggested in the present study may not be the only one responsible for 5-ALA fluorescence in gliomas. This is because 96% of glioblastomas lack IDH1/2 mutations but show a strong 5-ALA fluorescence [6]. In addition, most low-grade gliomas lack 5-ALA fluorescence and harbor IDH1/2 mutations [6]. Thus, we used the term “enhanced 5-ALA fluorescence” to describe the result of the hypothesized mechanism, cognizant that there must be another way to explain 5-ALA fluorescence in gliomas as a whole, or at least in glioblastomas. However, in light of the growing evidence of aberrant metabolism involving the TCA cycle in gliomas, we believe that crosstalk between the TCA cycle and the heme pathway is the core mechanism for selective 5-ALA fluorescence. There may be diverse ways to trigger this mechanism that have yet to be discovered. Genomic screening in relation to 5-ALA fluorescence will give insight to this question. The other issue to be validated in further study is that the correlation of 5-ALA fluorescence and WHO grade III gliomas with diverse molecular and genetic characteristics. Compared to glioblastomas, both clinical and translational studies on 5-ALA
fluorescence focusing on WHO grade III gliomas are rare.

In conclusion, the present study demonstrate that enhanced 5-ALA fluorescence is associated with \textit{IDH1} mutations in malignant gliomas. Reduced production of NADPH plays a role in the lag of the heme synthesis pathway as observed in mutant \textit{IDH1} cells, resulting in the accumulation of intracellular PpIX and enhanced fluorescence. These discoveries provide new insights into cancer metabolism in malignant gliomas and are expected to contribute in multiple ways to glioma management and research. By using a non-invasive method to detect metabolic characteristics specific to \textit{IDH1} mutants, we may be able to select appropriate candidates for fluorescence-guided surgery. NADPH itself can be developed not only as a biomarker, but as a therapeutic target in \textit{IDH1} mutant gliomas. Further investigations into this aberrant metabolic axis in gliomas will broaden our knowledge of gliomagenesis and the mechanism of malignant transformation.
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초 록

5-아미노레블린산 (5-ALA)을 이용한 형광유도수술은 최근 악성 뇌교종 수술에 많이 이용되고 있다. 그러나 교모세포종 (glioblastoma)과 다르게 WHO grade 3 악성 뇌교종에서는 5-아미노레블린산 (5-ALA)에 의한 특이한 형광발현을 나타내므로 이번 연구에서 IDH1 돌연변이와 5-ALA 형광발현과의 관계를 연구해 보기로 한다.

본 연구를 위해 렌티바이러스 벡터를 이용하여 IDH1/2 야생형 (U87MG-IDH1WT, U87MG-IDH2WT) 또는 IDH1/2 돌연변이형 (U87MG-IDH1R132H, U87MG-IDH1R172K)이 계속적으로 발현하는 악성뇌교종 세포인 U87 MG 세포를 구축하였다. 이 연구에서 IDH1/2 돌연변이형 (U87MG-IDH1R132H, U87MG-IDH1R172K)에서 5-아미노레블린산 (5-ALA)의 대사 및 프로토포피린 IX (PPIX) 축적이 지연되는 것을 확인하였으며, 액체 크로마토그래피- 질량분석기 (LC-MS)를 통해 5-아미노레블린산 (5-ALA)에 노출되었을 때, 크렘스 회로 (TCA cycle)와 관련된 대사물질들의 변화를 확인하였다. 그 결과 Heme 분해의 반응속도결정단계에서 보조인자로 작용하는 NADPH가 IDH1 돌연변이형 (U87MG-IDH1R132H)에서 낮게 발현하는 것을 발견하였다. 이것은 과량의 5-아미노레블린산 (5-ALA)을 대사시키기 위해서 많은 양의 NADPH가 필요로 하는
데, IDH1 돌연변이형 (U87MG-IDH1<sub>R132H</sub>)에서는 NADPH의 양이 적으므로, 5-ALA를 과량 투여했을 때 일시적으로 5-ALA 형광이 증가하게 된다는 가설을 뒷받침해주었다. 또한 악성 뇌교증 환자 조직에서도 대사물질 분석을 통해 이 가설을 증명할 수 있었다.

결론적으로, 이 연구를 통해 WHO grade 3 악성 뇌교증에서 증가된 5-ALA의 형광과 IDH1 돌연변이는 연관이 있고, IDH1 돌연변이를 가진 종양조직에서 낮게 발현하는 NADPH가 증가된 형광과 관련이 있음을 보여주었다.

주요어 : 형광유도수술, 뇌종양, 교모세포종, IDH1, 5-ALA, NADPH
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