Hypoxia promotes glycogen synthesis and accumulation in human ovarian clear cell carcinoma

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Abstract. Ovarian clear cell carcinoma (OCCC) has several significant characteristics based on molecular features that are distinct from those of ovarian high-grade serous carcinoma. Cellular glycogen accumulation is the most conspicuous feature of OCCC and in the present study its metabolic mechanism was investigated. The amount of glycogen in cells cultured under hypoxia increased significantly and approximately doubled after 48 h (P<0.01) compared to that under normoxic conditions. Periodic acid-Schiff positive staining also demonstrated intracellular glycogen storage. Western blot analysis revealed that HIF1α, which was overexpressed and stabilized under hypoxic conditions, led to an increase in the levels of cellular glycogen synthase 1, muscle type (GYS1), and conversely to a decrease in inactive phosphorylated GYS1 at serine (Ser) 641. Additional increases were observed in both protein phosphatase 1, which dephosphorylates and thereby induces GYS1 enzyme activity, and glycogen synthase kinase 3 beta (GSK3β) phosphorylated at Ser9, which is inactive on phosphorylation of GYS1 and subsequently induces its enzyme activity. By contrast, the level of PYGM-b decreased. These results indicated that the glycogen accumulation under a hypoxic environment resulted in the promotion of glycogen synthesis, but did not lead to inhibition of glycogen degradation and/or consumption. Under hypoxic conditions, HAC2 cells showed activation of the PI3K/AKT pathway caused by a mutation in exon 20 of PIK3CA, encoding the catalytic subunit p110α of PI3K. The resulting activation of AKT (phosphoSer473) also plays a role as a central enhancer in glycogen synthesis through suppression of GSK3β via phosphorylation at Ser9. Hypoxia decreased the cytotoxic activity of cisplatin and doxorubicin to various degrees. In conclusion, the hypoxic conditions together with HIF1 expression and stabilization increased the intracellular glycogen contents and resistance to the anticancer drugs.

Introduction

Ovarian cancer still remains the most lethal form of gynecologic malignancy. In Japan, ovarian cancer is the eighth leading cause of death from cancer in woman, but a steady upward trend has become evident in recent years. Ovarian clear cell carcinoma (OCCC) has a higher incidence in Japan at 24.7% of all epithelial ovarian cancers (1), than in North America and Europe (1-12%) (2-5). OCCC has clinical characteristics and pathologic significance based on molecular features that are distinct from high-grade serous carcinoma of the ovary (6). Unfortunately, some studies reported that OCCC may be relatively resistant to standard platinum-based chemotherapy (7-11). It is necessary to improve our understanding of pathobiologic characteristics of OCCC in order to develop new therapeutic strategies and administer current treatment effectively.

The intracellular storage of glycogen is one of the most conspicuous pathodiagnostic features of OCCC (6). Clarification of the mechanism of these metabolic characteristics may yield a new strategy to overcome the disease. Several studies have considered the mechanism of facilitated glycogen accumulation of OCCC (12,13), but a conclusion has not yet been reached. Some studies have found that the glycogen accumulation in several kinds of tumor cells was accelerated under hypoxic conditions (14-18). Under such a hypoxic environment, hypoxia-inducible factor 1 (HIF1) is expressed and functions as the main factor regulating the metabolism of the cell (19). HIF1 is a heterodimeric transcription factor that consists of an O2-regulated HIF1α and a constitutively expressed HIF1β subunit (20). HIF1α is unstable in well-oxygenated tissues owing to ubiquitin-mediated proteasomal degradation, but rapidly becomes stable under hypoxic conditions (19). Two kinds of hydroxylation enzyme (prolylhydroxylase and asparaginylhydroxylase) act in the functional maintenance of HIF1α. To initiate the degradation process in well-oxygenated cells, HIF1α is hydroxylated on a proline residue at 402 (Pro402) and/or Pro564 in the oxygen-dependent degradation domain (ODDD) by HIF-prolyl-4-hydroxylase (PHD) enzyme family, mainly PHD2, which comprises HIF1 downstream expressing genes (21). Prolyl-hydroxylated HIF1α is bound and ubiquitinylated by the von Hippel-Lindau tumor suppressor protein (pVHL), which recruits an E3-ubiquitin ligase that targets HIF1α for proteasomal degradation (22). HIF1 regulates the transcription of over hundreds of genes that...
encode proteins involved in every aspect of cancer biology, including cell immortalization and stem maintenance, genetic instability, glucose and energy metabolism, vascularization, autocrine growth factor signaling, invasion and metastasis, immune evasion, and resistance to chemotherapy and radiation therapy (23). It is well known that HIF1 promotes catabolism for energy production in many kinds of tumor cells mainly under an anaerobic microenvironment, which is the so-called Warburg effect (19). The present study demonstrated that hypoxia induced and enhanced cellular accumulation of glycogen in a human PIK3CA-mutation-positive OCCC cell line, HAC2, via stabilization rather than overexpression of HIF1 protein and also determined the manifestation of chemo-resistance under such conditions. Prevention and treatment strategies based on the metabolic properties of the OCCC were determined.

Materials and methods

Cell culture, hypoxic exposure and DFO treatment. The human OCCC cell line, HAC2, was kindly provided by Dr M. Nishida (National Organization of Kansumiga Medical Center). HAC2 was cultured in RPMI-1640 supplemented with 10% heat-inactivated fetal bovine serum (Invitrogen, USA). Normoxic cells were maintained conventionally at 37°C in 5% CO2 incubator (~21% O2 concentration). Hypoxic cells were exposed to hypoxic conditions of 1% O2, 5% CO2, and 94% N2 at 37°C for up to 48 h using an MIC-101 modular incubator chamber (Billups-Rothenberg, Inc., USA). Hypoxia was consistently monitored by an OX-02G gas monitor (Riken Keiki, Japan). To ensure the precise expression of HIF1α, deoxygenase (DFO, Wako, Japan), which is a potent hypoxia-inducing iron-chelating material, was added to the culture medium. For DFO treatment, the cells were incubated for relatively short time of 6 h in medium with 50 μM DFO because of the severe cytotoxic effect. Human ovarian cancer cell line, 2008 was provided by Dr S.B. Howell (UCSD, San Diego, USA). A2780 and OV1063 were obtained from European Collection of Cell Cultures (UK) and American Type Culture Collection (Rockville, USA), respectively. Human OCCC cell lines (JHOC5, JHOC7 and JHOC9) were obtained from Riken Bioreource Center (Tsukuba, Japan).

Quantitative assays of glycogen. After washing the cultured cells with cold 20 mM Tris-HCl, pH 7.4, with 0.15 M NaCl, (Tris-buffered saline, TBS), glycogen was extracted with 30% KOH from the cells and the extracts were boiled at 100°C for 15 min, followed by precipitation with 100% ethyl alcohol. After centrifugation (16,000 x g for 10 min) at 4°C, the pellet was re-suspended in 70% ethyl alcohol and re-centrifuged. The resultant glycogen precipitate was dissolved in 0.1 M sodium acetate. Glycogen was measured spectrophotometrically (infinite M200PRO, Tecan, Switzerland) using Glycogen Assay Kit (Bio Vision, USA). The results were expressed in μg of glycogen per mg of protein.

Sodium dodecylsulphate-polyacrylamide gel electrophoresis (SDS-PAGE) and Western blot analysis. After washing once with ice-cold TBS, cells were harvested, lysed with SDS-PAGE sample buffer (50 mM Tris-HCl, pH 6.8, 2% SDS, 10% glycerol, 0.05% bromophenol blue) and immediately boiled at 100°C for 5 min. Lots of each sample (50 μg) were subjected to SDS-PAGE on 10% polyacrylamide gel and electrophoresed. The fractionated proteins were transferred to PVDF filters (Bio-Rad, USA), then blocked with blocking buffer (TBS, 0.05% Tween-20, 0.1% casein, 0.1% gelatin, and 0.02% NaN3), and the filters were incubated for 17 h at 4°C with the respective primary antibodies. Primary antibodies used were murine monoclonal antibodies against human HIF1α, Vimentin (VIM), E-cadherine (CDH1), VHL, heat shock protein (HSP70, HSP90, plasminogen activator inhibitor 1 (PAI), extracellular signal regulated kinase 1/2 (ERK1/2), phosphatase and tensin homolog (PTEN), P21(WAF1) (WAF1) (BD Biosciences, Japan), ERK1/2 diphosphorylated at Thr183 and Tyr185 (p-ERK1/2), β-actin, α-tubulin (Sigma-Aldrich Japan, Japan), CD73 (Abcam, Japan), rabbit monoclonal antibodies against glycogen synthase 1, muscle type (GYSI), glycogen synthase kinase 3β phosphorylated at Ser9 [p-GSK3β(ser9)], Notch1, cMET, protein phosphatase 1 (PPP1A), mammalian target of rapamycin phosphorylated at Ser248 (p-mTOR) (Epitomics Inc., USA), rabbit polyclonal antibodies against protein phosphatase 1, regulatory subunit 3C (PPP1R3C), pyruvate dehydrogenase kinase 1 (PDK1), prolyl-4-hydroxylase 2 (PDH2), carbonic anhydrase 9 (CA9), FK506-binding protein 38 (FKBP38), glucose transporter isoform 1 (GLUT1), glycogen phosphorylase, muscle type, dephosphorylated form (PYGM-b), matrix metalloproteinase 14 (MMP14), m-TOR (Abcam), monocarboxylate transporter 4 (MCT4) (Sigma-Aldrich Japan), and hexokinase 2 (HK2) (Santa Cruz Biotechnol. Inc., USA), AKT, AKT phosphorylated at Ser473 (p-AKT), GYS1 phosphorylated at Ser641 (p-GYS1) (Cell Signaling Technology Inc., USA). After 17-h incubation at 4°C, the filters were further incubated with horseradish peroxidase-conjugated secondary antibodies against mouse or rabbit IgG (1:5000, Cell Signaling Technology) for 2 h at 37°C. The signal was detected by the ImmunoStar LD chemiluminescence system (Wako), and the images were captured by Light Capture II (Atto, Japan). Digital images of bands were analyzed by the NIH Image J, 1.45 software.

Morphologic examination. Cells cultured using the Lab-Tek chamber slide (Thermo Fisher Scientific, USA) under hypoxic or normoxic conditions were fixed with 10% formalin in TBS, washed with TBS and dehydrated by graded concentrations of ethyl alcohol. For tumor tissues transplanted in nude mice, the resected tissue samples were washed once with cold TBS and fixed with 10% formalin in TBS, then routinely processed and embedded in paraffin. Both fixed cultured cells and depafrinized tissue sections were stained by hematoxylin and eosiin and periodic acid-Schiff (PAS) with or without diastase digestion.

RNA extraction and RT-PCR. After culturing the cells, total RNA was extracted using the RNasy kit (Qiagen, Japan), and the concentration was determined by a spectrophotometer (Hitachi, Japan). Reverse transcription was carried out using the PrimeScript II 1st Strand cDNA Synthesis Kit (Takara, Japan). cDNAs were then amplified using GoTaq Green Master Mix (Promega, USA) with a pair of gene-specific primers. Primer sequences for the analyzed genes are shown in Table I.

Xenotransplantation of HAC2 cells into nude mice and preparation of tissue extracts. HAC2 cells (5x10⁶) were transplanted into the subcutaneous tissue of the lateral abdominal wall of...
male nude mice (BALB/c nu/nu, 5-weeks old, Clea, Japan) under specific pathogen-free conditions. Eleven weeks after the transplantation, tumor tissues were removed and homogenized. The harvested tumor tissues were rinsed immediately with cold TBS. For RNA preparation, tumor tissue fragments were homogenized in Buffer RLT Plus (Qiagen) with 1% 2-mercaptoethanol and the total RNA was isolated by the same procedure as described in the above section. For protein preparation, some tumor fragments were homogenized in 50 mM Tris-HCl, pH 7.5, 0.1% SDS, 1 mM EDTA, 1% TritonX-100, 1% deoxycholate, 1 mM NaF, 1 mM NaVO₃, 1 mM NaF, protease inhibitor cocktail and phosphatase inhibitor cocktail 1, 2 (Sigma-Aldrich). After incubation for 20 min on ice followed by centrifugation (16,000 x g for 20 min) at 4°C, SDS-PAGE sample buffer (5-fold concentration) was added to the supernatant and boiled. All animal studies were performed in conformity with the Guidelines on Animal Experimentation of the Jikei University containing rules of ethics on animal experimentation.

**Analyses of gene mutations.** Exons 9 and 20 of the PIK3CA gene encoding phosphatidyl inositol-3, 4, 5-trisphosphate 3-kinase catalytic subunit α isoform (PIK3 p110α, EC2.7.1.153) and VHL in HAC2 cells were amplified using PCR. Purification of cDNA and the PCR method were as described in the above section. The primers for PIK3CA were the same as described previously (24), briefly, exon 9: forward, 5'-CCAGAGGGGA AAAATGACAA-3'; reverse, 5'-ACCTGACTCCATCAATGAAA-3'. Exon 20: forward, 5'-TGTAGCAGATTGATCAT CG-3'; reverse, 5'-AGATTGTGAGAGATCCAATCC-3'. Primers for VHL were as follows: forward, 5'-GAAGACT TCG-3'; reverse, 5'-AATTGTGTGGAAGATCCAATCC-3'.

Each RT-PCR product was electrophoresed on 1.5% agarose gel (Wako) in Tris-acetate buffer, 1.26 μM ethidium bromide. Amplified PCR products were purified with Wizard SV Gel and PCR Clean-Up System (Promega). They were subsequently subjected to direct sequencing PCR with BigDye terminator v3.1/1.1 cycle sequencing reagents (Applied Biosystems, USA) and were analyzed with the Applied Biosystems 3730xl DNA Analyser (Applied Biosystems) with basic BLAST in NCBI. When a mutation(s) was detected by sequencing analysis, genetic analysis was repeated at least twice from the step of genomic DNA amplification.

**Table 1. PCR primer.**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer</th>
<th>Reverse primer</th>
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<tr>
<td>HNF1B</td>
<td>5'-CCCAGACCAAGCGCCGTTTT-3'</td>
<td>5'-ACTGTCTTGTTGAATTCTGCGGAGG-3'</td>
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<tr>
<td>HIF1A</td>
<td>5'-GAAAGCGCCAGCAGCTCCTCAAG-3'</td>
<td>5'-TGGGTAGGAGATGGAGATGCG-3'</td>
</tr>
<tr>
<td>ABCB1</td>
<td>5'-GACTGAGCTGGAGTGGAAG-3'</td>
<td>5'-CCACCAAGAGCTGTTGTC-3'</td>
</tr>
<tr>
<td>ABCC1</td>
<td>5'-TGGAACAAGCTGAGCTGAG-3'</td>
<td>5'-CACAGTGCAGGAACCAGA-3'</td>
</tr>
<tr>
<td>GCLC</td>
<td>5'-TGAGATTTAAAGGCCGCTCTT-3'</td>
<td>5'-TGGCATTAAACTCCCTCCT-3'</td>
</tr>
<tr>
<td>TOP2A</td>
<td>5'-GTTTACTGGGCTAAGCGGA-3'</td>
<td>5'-CACGACCAACCGTACCGAT-3'</td>
</tr>
<tr>
<td>TOP2B</td>
<td>5'-AAACAATGTCAGCAAGCTT-3'</td>
<td>5'-CCCACAGCGCCTCTGAG-3'</td>
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**Cytotoxicity assay.** After washing once with calcium-free Dulbecco's phosphate buffered saline [PBS(-)], cells were detached with 0.125% trypsin and 0.02% EDTA in PBS(-). The cells (2x10⁴ cells/well) were seeded into a 48-well culture plate (Greiner Bio One, Germany) containing 500 μl of medium with various concentrations of cisplatin (CDDP, Sigma-Aldrich Japan) or doxorubicin (DXR, Sigma-Aldrich Japan) and incubated continuously for 96 h under conditions of normoxia or hypoxia. To prepare cells cultured under hypoxic conditions, the cells were precultured for 48 h under hypoxic conditions as stated above and then detached in a hypoxic chamber before assay. After 96-h incubation with test materials under respective hypo- and normo-oxygen conditions, 40 μl of Cell Titer 96 Aqueous MTS solution (Promega) was added to each well and the plates were incubated for additional 3-h period under normoxic conditions. Absorbance at 490 nm for each well was measured using a microtiter plate reader (infinite M200PRO). The cell viability was expressed as the fraction of surviving cells relative to untreated controls and the 50% growth inhibition concentration (IC₅₀) was calculated.

**Protein concentration.** The protein concentration was measured by the DC protein assay kit (Bio-Rad). Bovine serum albumin was used as the standard.

**Statistical determination.** When indicated, data were given as means ± SD with n indicating the number of experiments. One-way analysis of variance and t-tests were applied for the statistical analysis, as appropriate, using the StatMate software (ATMS, Japan). Values were considered significant at P<0.05.

**Results and Discussion**

**Expression of proteins under hypoxia.** Under hypoxic environment, HAC2 cells showed slight but non-significant growth retardation as compared with that under normoxic condition. HAC2 cells expressed the hepatocyte nuclear factor-1β (Fig. 1), a representative gene expressed in OCCE, as reported previously (25). Under the culture condition of low oxygen, mRNA of HIF1α in HAC2 cells was somewhat overexpressed at certain points of time (Fig. 1), and protein was stabilized at increased levels as compared to that of normoxic cells. Expressed and stabilized HIF1α simultaneously promoted transcription of many genes crucial for cellular response to hypoxia including cell survival.
angiogenesis, cell motility and invasion as shown (Fig. 2A). HIF1 also regulates the expression of the genes involved in glucose metabolism. HIF1 downstream targets, such as GLUT1, HK2, PDK1, MCT4, were significantly increased in terms of levels of proteins under hypoxia compared to normoxia (Fig. 2B).

The stability and function of HIF1 are mainly regulated by PHD-driven hydroxylation of ODDD, followed by pVHL-caused ubiquitinylation and proteasomal degradation (21). In the present study, the VHL gene from HAC2 cells did not manifest any genetic mutations in the DNA sequence and was compatible with that from normal cells as reported previously (26). Furthermore, pVHL expression without loss of function in HAC2 cells did not show any difference between cells cultured under normoxic and hypoxic conditions (Fig. 2C). It is well known that both somatic and germline abnormalities of the VHL gene play a pivotal role in the carcinogenesis of renal clear cell carcinomas (27-32) but have not been reported in OCCC (0/74 ovarian samples in COSMIC) (6). Our present study also demonstrated that a VHL mutation did not appear in OCCC, different from renal clear cell carcinoma. In contrast, in the initial 24 h after hypoxic culture, the occurrence was noted of continuous 48-h overexpression and stabilization of PHD2, which is a HIF1 downstream gene and the main HIF1α regulator via hydroylation of ODDD (33,34), and also of transient downregulation of FKBP38, a key regulator in the stabilization and function of PHD2 by binding, followed by proteasomal degradation (35-37) (Fig. 2C). Since the function of PHD2 is downregulated via molecular interaction between FKBP38 and PHD2 protein, HIF1α protein stability is prolonged by the inhibition of PHD2, to which the

Figure 1. HAC2 and other OCCCs expressed various levels of HNF1β and HIF1. (A) Each cell line cultured under normoxia was lysed and mRNAs were extracted for RT-PCR followed by agarose gel electrophoresis. (B) HAC2 cells were cultured under normoxia for 48 h or hypoxia for 24 and 48 h, or normoxia for 6 h with DFO. Cells cultured under each condition were lysed. mRNA for RT-PCR and protein for SDS-PAGE followed by Western blotting were obtained. HAC2, JHOC5, JHOC7 and JHOC9 were OCCC in nature. β-actin and α-tubulin were loading controls of protein.

Figure 2. HIF1-downstream (A and B) and HIF1-regulating (C) gene products. Cells cultured under each condition described in Fig. 1B were lysed and then SDS-PAGE, followed by Western blotting were performed. For details, see Materials and methods.
increased FKBP38 molecules are bound. The rapid recovery by 48 h under hypoxic conditions and the unchanged level with DFO treatment in this study, suggested that FKBP38 is not a HIF target, as previously reported (37). These results indicated that the early and immediate compensatory bio-responses of HAC2 cells against sudden and severe hypoxic changes were almost the same as those described in previous reports (33,36).

A decreased level of HSP90 and slight but detectable increased level of HSP70 were additionally found (Fig. 2C). It has been reported that HSP70/HSP90 axis is another mechanism used to stabilize HIF1 and the downregulation of HSP70 and upregulation of HSP90 induced destabilization and degradation of the HIF1 protein by means of their chaperon mechanism (38,39), independently from pVHL-ubiquitin system. These results suggest that the hypoxic environment in our experiment was sufficient.

Hypoxia-induced glycogen accumulation. HAC2 cells exposed to hypoxia or normoxia were lysed and their glycogen contents were measured at different points of time. The intracellular amount of glycogen under hypoxic conditions was increased significantly after 24 h (641.4±33.4 µg/mg protein, P<0.05, n=4), and had approximately doubled after 48 h (1252.9±239.3 µg/mg protein, P<0.01, n=4) as compared to that under normoxic condition (533.2±16.15 µg/mg protein, n=4) (Fig. 3A). As shown from the cytologic examination (Fig. 3B), detectable PAS-positive, diastase-soluble granules in HAC2 cells indicated a marked increase in the cellular glycogen concentration under hypoxia. Histopathologic observations revealed that HAC2 cells transplanted into nude mice formed an adenocarcinoma-like glandular structure lining several large hobnail type tumor cells and had an irregular shape with bizarre nuclei. These cells also showed intracellular PAS-reactive, diastase-soluble glycogen accumulation in a large number of the tumor cells (Fig. 3B) with significantly elevated levels of HIF1α protein, as mentioned above in the biochemical analysis section (Fig. 1). These histologic findings were compatible with the histopathologic feature of OCCC and confirmed that the HAC2 cell line was established from OCCC tumor tissue.

Expression of several proteins related to glycogen metabolism under hypoxia. To confirm whether hypoxia modulated enzymes were involved in glycogen synthesis and/or degradation in HAC2 cells, the expressions of some enzymes related to glycogen metabolism, such as GYS1, p-GSK3β(Ser9), PPP1A, PPP1R3C and PYGM-b, were examined under respective normoxic and hypoxic conditions. Protein level of GYS1 was increased under hypoxic conditions. Conversely, the level of p-GYS1 in ser641 protein was diminished (Fig. 4A and B). GSK3β is known as a key regulator in the function of GYS1 via phosphorylation and consequent inactivation of GYS1 activity (40). In spite of the omission of a direct assay of glycogen synthase enzyme activity in this study, a previous report demonstrated that the glycogen synthase activity increased in parallel with GYS1 protein accumulation (17). Phosphorylation at Ser9 of GSK3β inhibits its own enzyme activity (41) and in the present study, p-GSK3β(Ser9) increased under hypoxia (Fig. 4C). Moreover, PPP1A and its regulatory subunit PPP1R3C were slightly but significantly elevated when HAC2 cells were cultured under hypoxic conditions. Conversely, the level of p-GYS1 in ser641 protein was diminished (Fig. 4A and B). GSK3β is known as a key regulator in the function of GYS1 via phosphorylation and consequent inactivation of GYS1 activity (40). In spite of the omission of a direct assay of glycogen synthase enzyme activity in this study, a previous report demonstrated that the glycogen synthase activity increased in parallel with GYS1 protein accumulation (17). Phosphorylation at Ser9 of GSK3β inhibits its own enzyme activity (41) and in the present study, p-GSK3β(Ser9) increased under hypoxia (Fig. 4C). Moreover, PPP1A and its regulatory subunit PPP1R3C were slightly but significantly elevated when HAC2 cells were cultured under hypoxia (Fig. 4A and B). It is well known that PPP1A catalyzes dephosphorylation of GYS1 and activates its enzyme activity, and additionally catalyzes dephosphorylation and inactivation of PYGM, contributing to further stimulation of glycogen deposition (42). In the present study, the level of PYGM-b was diminished (Fig. 4A and B) and this indicated that glycogen accumulation resulted in facilitated glycogen deposition.
synthesis, but not in the inhibition of glycogen degradation and/or consumption. These facts partly explained why the promotion of glycogen accumulation resulted in synthesis in the HAC2 cells under hypoxia.

HAC2 cells exposed to hypoxic condition showed the activation of AKT and MAP kinase (ERK1/2). In OCCC, like numerous other cancers (43,44), somatic mutations in the PIK3CA gene have been common and highly frequent (24,45,46). The PIK3CA gene, encoding the catalytic subunit p110α of PI3K, is located on chromosome 3q26.3 and mutation and/or amplification of this locus has been shown to increase PI3K activity (43,47). HAC2 cells used in the present study have a distinct PIK3CA mutation in exon 20 of the H1047R (A3140G) substitution (Fig. 5A), and activation of its downstream AKT and MAP kinase pathways was noticeable (Fig. 5B). Since PIK3CA gene mutation causes the continuous activation of AKT, it was not surprising in HAC2 cells that the phosphorylated AKT at Ser473 was elevated under hypoxic culture conditions as well as DFO treatment, moreover, the stabilized HIF1 (48,49) was further activated by PI3K/AKT signaling pathway, one of the main mechanisms of HIF1 activation, as reported previously (50). Additionally, it is well known that AKT regulates glycogen synthesis through phosphorylation at Ser9 followed by inactivating the GSK3β. When the cells are exposed to hypoxic conditions, time-dependent biphasic HIF1α stabilization and activation by PI3K/AKT/GSK3β signaling pathways has been proposed but this explanation remains controversial (50). Our present results supported part of the initial response reported in previously findings (50), although the hypoxic treatment of the cells with a different type of PIK3CA mutation was for a longer time compared with that in the previous report (50). In the present study, downregulation in p-GSK3β(Ser9) was promoted and the level of GSY1 in the dephosphorylated form was consequently increased and probably enhanced glycogen synthesis.

In contrast, apparent phosphorylated ERK1/2 at Thr183 and Tyr185 was noted only in cells exposed to hypoxic culture conditions and there were no detectable signals in cells treated by DFO because of the extremely short time of exposure and the effect of severe toxicity of DFO on the cells (Fig. 5B). Treatment of the target cells with DFO usually has been used to shift the cell metabolism in order to mimic hypoxic conditions (51), instead of culturing the cells under ‘real’ oxygen-regulated hypoxia. In the present study, however, several discrepancies were noticed, and the reasons for such differences between the two conditions were not resolved at this time. The MAP kinase signaling pathway is concerned with almost all cellular responses, including not only acceleration, but inhibition of the cell metabolism and functions under a variety of physiologic and pathologic states. Therefore, HAC2 cells exposed to hypoxia may evoke multipotential functions in their character.

HIF1-mediated epithelial-mesenchymal transition (EMT) may have occurred simultaneously in this experiment. Multifunctional GSK3β regulates the E-cadherin transcriptional repressor, Snail, as described in a previous report (52), via phosphorylation regulation. In the present study, E-cadherin was downregulated time-dependently by hypoxia treatment of the cells and several EMT-related gene families were activated under the experimental hypoxia (Fig. 2A and B). Our previous and other reports have indicated that the three dimensional (3D)
culture provided a cellular architecture similar to that of living organisms and closely mimicked the living hypoxic microenvironment without any specific modification and/or treatments, such as cell culturing under anaerobic condition or cell exposure to hypoxia-inducible reagents such as DFO (53-56). The 3D culture system may be a highly useful tool for investigating metabolic changes in cells under hypoxia without treatment, and further study is necessary to clarify whether or not some metastatic properties expressed in the OCCC without anoikis are based on EMT or MET.

Hypoxia altered sensitivity to some anticancer agents of HAC2 cells. OCCC has been demonstrated to show chemoresistance (7-11), hence its sensitivity under hypoxic conditions to CDDP, which has been used routinely in gynecologic cancer clinics, was analyzed and then compared to that under normoxic conditions. Cell viability assessed by MTT assay indicated that the IC_{50} of CDDP increased under hypoxic conditions (IC_{50} 2.35±0.25 µM, n=5) as compared to normoxia (IC_{50} 1.51±0.07 µM, n=5) (P<0.001), suggesting that known and/or unknown mechanisms of resistance to CDDP under hypoxia may be upregulated and induced (Table II). It is of interest that the hypoxic environment did not attenuate the drug efficacy of DXR (IC_{50}: hypoxia/normoxia = 0.307±0.019, n=5/0.314±0.018 µM, n=5) (Table II). The expression of some genes related to drug-resistant phenotypes was analyzed and the mRNA expression levels of ABCB1, ABCC1, γ-glutamylcysteine synthetase (GCLC) and DNA topoisomerase IIα, β (TOP2A, B) did not show any significant difference between hypoxia and normoxia (Fig. 6), although ABCB1 is reported to be one of the HIF target downstream genes (57) and its gene product, P-glycoprotein (Pgp), actively transports its typical substrates (DXR) outside the cells (58). This result indicated that ABCB1 gene expression might not be sufficient to show an effect in the relatively short time of

### Table II. Cytotoxicity of cisplatin and doxorubicin on HAC2 cells.

<table>
<thead>
<tr>
<th>Drug</th>
<th>Normoxia (µM)</th>
<th>Hypoxia (µM)</th>
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<tr>
<td>Cisplatin</td>
<td>1.51±0.07 (n=5)</td>
<td>2.35±0.25 (n=5)</td>
</tr>
<tr>
<td>Doxorubicin</td>
<td>0.307±0.02 (n=5)</td>
<td>0.314±0.02 (n=5)</td>
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*Data are presented as the means ± SD of 50% growth inhibition concentration (IC_{50}). P-values compared to normoxia, ^P<0.001; ^NS, P>0.05.*
Hypoxia promotes glycogen accumulation in OC. 48-h culture. DXR, an anthracycline anticancer agent, exhibits cytotoxicity that intercalates DNA and induces TOP1-mediated strand breaks (59). A recent convincing and meaningful report (60) proposed evidence that HIF1 binds to hypoxia-responsive elements (HREs), which are cis-acting DNA sequences containing the consensus binding site to mediate HIF-inducible and -dependent transcription; moreover, anthracyclines disrupt the binding of HIF1 to HREs in vitro and probably in vivo. Another study (61) reported, however, that neither DXR nor daunorubicin inhibited the expression of HIF1-dependent genes, and that only aclacinomycine B inhibited expression. It is known that aclacinomycine B, a member of the anthracycline anticancer drug family, is not a substrate or is at most a weak substrate for Pgp (62-64). Inhibitory effect of the induction of anthracyclines on HIF-downstream genes is controversial, because the discrepancy might have been caused by the use of another experimental model of cells with unknown and various expression levels of ABCB1 and/or the ABCB family and related biomaterials including GSH. In the present study, DXR cytotoxicity under hypoxia did not increase, but in contrast, decreased slightly by a non-significant level as compared with that under normoxia. The result might have been off set by other GS-X pump molecules in the ABCB family in HAC2 cells under hypoxia having actively transported the DXR outside the cells, in spite of the inhibitory effect of DXR in HIF1 regulating transcription. Another mechanism to reduce the cytotoxicity of DXR might be that under a well-conditioned hypoxic environment, HIF1-proteins translocate into the nuclei, bind and show fully functional transcriptional activity, which may lead to competitive inhibition of the binding of DXR to DNA under such conditions. The reduced level of DXR-cytotoxicity under hypoxia was based on this competitive binding inhibition of DXR to HREs. Anticancer efficacy of DXR on ovarian cancer chemotherapy will have to be re-considered strategically again. CDDP also showed reduced cytotoxicity under hypoxia. Since CDDP exhibits cytotoxicity that intercalates the DNA strands, a slight but efficient reduction on CDDP cytotoxicity may have a considerable effect similar to that of DXR on HREs. This phenomenon requires further study to clarify the mechanism.

Acknowledgements

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