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Control of hypoxia-induced tumor cell adhesion by cytophilic human catalase

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Abstract

Hypoxia-induced reactive oxygen species (ROS)-mediated expression of a variety of genes in endothelial cells has been suggested to be involved in abnormal cell adhesion. To prevent this by accelerated binding of catalase to endothelial cells, human catalase (hCAT), an enzyme catalyzing the decomposition of hydrogen peroxide, was fused with three repeats of arginine-glycine-aspartic acid peptide or nona arginine peptide at the C-terminal to obtain hCAT-(RGD)3 and hCAT-R9, respectively. Human CAT and its derivatives were expressed in yeast *Pichia pastoris* and purified. The specific activity and secondary structure of hCAT-(RGD)3 and hCAT-R9 were close to those of hCAT, but these derivatives showed higher binding to mouse aortic vascular endothelial cell line MAEC than hCAT, indicating that they are cytophilic derivatives. Hypoxic treatment of MAEC increased the intracellular ROS level, the binding of mouse melanoma cells, and the activity of transcription factors, hypoxia inducible factor-1 and nuclear factor-κB. hCAT-(RGD)3 or hCAT-R9 efficiently inhibited these changes compared with hCAT. These results indicate that cytophilic hCAT-(RGD)3 and hCAT-R9 are effective in inhibiting hypoxia-induced tumor cell adhesion to endothelial cells.

Keywords: recombinant human catalase; abnormal cell adhesion; reactive oxygen species; hypoxia; free radicals
Introduction

Vascular endothelium is a monolayer lining the interior surface of blood vessels, which forms an interface between blood and tissue. Dynamically responding to changes in environmental factors, such as oxygen tension, cytokines and growth factors, endothelial cells play a pivotal role in maintaining physiological homeostasis [1].

Abnormal cell adhesion to vascular endothelium is observed in a wide range of pathological situations, including arteriosclerosis, thrombosis and tumor metastasis [2-4]. Under these pathological conditions, local blood flow is impaired to induce microenvironmental changes that lead to the imbalance of oxygen demand and supply, and resulting hypoxia [5]. Endothelium that has experienced hypoxia can be the ‘soil’ for tumor metastasis, because cells, including endothelial cells, increase the transcription of a variety of genes for adaptation to hypoxia [4]. An increasing body of evidence suggests that ROS play a central role in controlling hypoxia-induced transcription [6, 7]. Therefore, eliminating ROS in endothelial cells can be an efficient approach to controlling abnormal cell adhesion to endothelium. Catalase is an antioxidant enzyme that catalyzes the degradation of hydrogen peroxide, ROS with a long half-life. However, previous studies have demonstrated that catalase poorly interacts with cells [8], so that any approaches to increasing its binding to cells could increase its therapeutic potential in a wide range of disease conditions.

A variety of molecules, including integrins and other adhesion molecules, are
expressed on the surface of endothelial cells [9]. In addition, negative charge of cell surface provides a vast surface area for binding of cationic compounds [9]. In this study, two types of cytophilic human catalase (hCAT) derivatives were newly designed: one is hCAT fused with an arginine-glycine-aspartic acid (RGD) peptide, which is a well-known ligand for αvβ3 integrin, and the other is one fused with a nona arginine (R9) peptide, a cationic peptide. hCAT and its derivatives were expressed in *P. pastoris*, and their activity to inhibit hypoxia-induced tumor cell adhesion was examined.
Materials and Methods

Cell culture

A mouse aortic vascular endothelial cell line (MAEC), a gift from Professor Ichiro Saito (Department of Pathology, Tsurumi University School of Dental Medicine, Yokohama, Japan), was cultured as previously reported [10]. B16-BL6/Luc [11] and colon26/Luc [12], clones of murine melanoma B16-BL6 and murine colon carcinoma colon26 that stably express the firefly luciferase, were grown as reported previously. A murine Lewis lung cancer (LLC) was cultured as reported previously [13]. To establish LLC/Luc, LLC was transfected with pCMV-Luc [14] using Lipofectamine 2000 (Invitrogen, Carlsbad, CA, USA) according to the manufacturer’s instruction. Cell culture under hypoxic conditions (8 % oxygen) was carried out using an AnaeroPack (Mitsubishi Gas Chemical Co., Tokyo, Japan).

Expression and purification of recombinant human catalase derivatives

The pCMV6-Entry vector containing the genomic Homo sapiens catalase (NM_001752) cDNA was purchased from OriGene Technologies Inc. (Rockville, MD, USA). Oligonucleotides corresponding to the amino acid sequence RGDRGDRGD (5’-AGAGGTGATAGAGGCGACAGAGGTGAC-3’) or RRRRRRRRRR (5’-AGAAGAAGGAGAAGGAGAAGGAGAAGGAGAAGGAGA-3’) were purchased from Greiner Bio-One Co. Ltd. (Tokyo, Japan). For the expression of glutathione S-transferase
(GST)-fusion proteins in the cytoplasm of *P. pastoris*, pNT401 vector was first constructed from pIB2 vector [15]; DNA sequence encoding GST was PCR-amplified and cloned immediately downstream of glyceraldehyde-3-phosphate dehydrogenase promoter within the parental vector. A PCR fragment of human catalase cDNA was digested with *Bgl*II and *Xba*I and inserted into *BamH*I-*Spe*I site of pNT401 expression vector. To construct plasmid vectors expressing hCAT-(RGD)3 or hCAT-R9, the DNA sequence corresponding to the C-terminal 11 amino acid residues (SHLAAREKANL) of human catalase, which contains a peroxisomal sorting signal [16, 17], was replaced with that corresponding to each peptide.

The plasmid constructs were linearized with *Sal*I. *P. pastoris* yeast SMD1168 was transformed by electroporation at 1.5 kV, 25 μF, 200 Ω. Minimal SD agar plates (2 % glucose, 0.5 % ammonium sulfate, 0.17 % yeast nitrogen base w/o amino acids and ammonium sulfate, 2 % agar) were used for the selection of *P. pastoris* transformants. Single colonies were picked and inoculated into 5 mL of YPD medium (1 % yeast extract, 2 % peptone, 2 % glucose) and grown at 30 °C in a bio-shaker with shaking at 250 rpm for 36 hr. A 2.5 mL aliquot of this suspension was inoculated to 500 mL of YPD medium in 3 L flask, and the culture was grown for 24 hr. The cells were harvested with 2000 ×g centrifugation, and disrupted using a French Press at 2,000 psi. The clarified supernatants of cell lysates containing hCAT, hCAT-(RGD)3 or hCAT-R9 were loaded onto a glutathione sepharose column (GE Healthcare, U.K.), and these
proteins were eluted by applying PreScission protease (GE Healthcare) in 50 mM Tris-HCl solution (pH 7.5). After concentration by ultrafiltration, the solvent of each elute was changed to water. The purified catalase derivatives were stored at -20 °C until use.

**Determination of catalase specific activity**

Protein concentration was quantified using a BCA Protein Assay Kit (Pierce, Rockford, IL, USA). Each catalase derivative (0.04 µg) was added to 25 mM phosphate buffer (pH 7.0) containing 10 mM hydrogen peroxide, and the decomposition of hydrogen peroxide was monitored spectrophotometrically at 240 nm. Samples were used without any separation of active (heme-containing) and inactive (heme free) forms. Bovine liver catalase (50,400 U/mg protein; Sigma Aldrich, USA) was used as a standard. Each sample was measured in five replicates and their averages were employed.

**Measurement of circular dichroism**

Circular dichroism (CD) spectra were recorded using a JASCO-820 type spectropolarimeter (JASCO, Tokyo, Japan) at 4 °C. For calculation of the mean residue ellipticity [θ], the molecular mass of the catalase derivatives was assumed to be as 240 kDa [8]. Far-UV (wavelength 200-250 nm) CD spectra were recorded at a protein concentration of 15 µM in Tris buffer (50 mM Tris-HCl (pH 7.5), 150 mM NaCl).
**Interaction of catalase derivatives with endothelial cells**

MAEC seeded on 24 well plates at a density of $1.0 \times 10^5$ cells/well were exposed to Medium 199 (Gibco, USA) containing one of the catalase derivatives at a concentration of 100 U/mL for 30 min at 4 °C or 37 °C. Then, the cells were washed with PBS and lysed with a lysis buffer (0.05 % Triton-X, 2 mM EDTA, 0.1 M Tris-HCl (pH 7.8)). The catalase activity of cell lysates was determined as described above.

**Intracellular ROS production under hypoxic conditions**

Intracellular ROS production was detected using CM-H$_2$DCFDA (Invitrogen). MAEC were treated with 5 μM CM-H$_2$DCFDA for 1 hr, and then were cultured under hypoxic conditions of 8 % oxygen. The fluorescence intensity was measured using a 1420 Multilabel counter (PerkinEimer, USA) at an excitation wavelength of 485 nm and an emission wavelength of 530 nm.

**Measurement of luciferase activity**

Cells were lysed with the lysis buffer. Twenty microliters of the lysate was mixed with 20 μL luciferase assay buffer (Picagene, Tokyo Ink, Tokyo, Japan) and the produced light was measured using a luminometer (Lumat LB 9507, EG&G Berthold, Bad Wildbad, Germany).
**Cell adhesion assay**

To examine the effect of hydrogen peroxide on tumor cell adhesion, MAEC (1.0×10⁵ cells/well) were seeded on 24 well plates, added with hydrogen peroxide at a final concentration of 0 to 100 µM, and incubated for 6 h. Then, tumor cells (B16-BL6/Luc, LLC/Luc and colon26/Luc) at a density of 1.0×10⁴ cells/well were added into each well containing the MAEC monolayer and incubated for 3 hr. Then, non-attached cells were washed out with PBS, and the remaining adherent cells were lysed with the lysis buffer. The luciferase activity of the lysates was measured as described above. Separately, MAEC were incubated under normoxic or hypoxic conditions for 6 hr. Then, tumor cells were added and their adhesion was examined as above.

**Transcription factor dependent reporter gene assay**

MAEC seeded on 24 well plates at a density of 1.0×10⁵ cells/well were transfected with pHRE-Luc [18], pNF-κB-Luc, or pAP-1-Luc (Stratagene, La Jolla, CA, USA) using Lipofectamine 2000 as described above. After incubation under hypoxic or normoxic conditions for 6 hr, cells were lysed with the lysis buffer. The luciferase activity of the cell lysates was measured as described above.

**mRNA quantification**
Total RNA was isolated from MAEC (2.0×10^5 cells) after incubation under hypoxic or normoxic conditions for 6 hr, and the mRNA expression of intercellular adhesion molecule-1 (ICAM-1) and vascular cell adhesion molecule-1 (VCAM-1) in MAEC was estimated by real-time quantitative PCR using a LightCycler system (Roche Diagnostics, Mannheim, Germany). The mRNA expression of the target genes was normalized to the mRNA level of GAPDH. The primer sequences for RT-PCR were as follows, ICAM-1: forward 5’-GGGAATGTCACCAGGAAT-3’, reverse 5’-TCCTGAGCCTTCTGTAACTTGT-3’, VCAM-1: forward 5’-CGGCATATACGAGTGTGAATC-3’, reverse 5’-ATCCCGATGGCAGGTTATT-3’, GAPDH: forward 5’-TCTCCTCGACTTCAACA-3’, reverse 5’-GCTGTAGCCGTATTCATTGT-3’.

**Statistical analysis**

Differences were statistically evaluated by one-way ANOVA followed by the Tukey-Kramer multiple comparison test, and the level of statistical significance was set at P<0.05.
Results

Characterization of catalase derivatives

Figure 1 shows the schematic representation of human catalase derivatives: hCAT, hCAT-(RGD)3 and hCAT-R9. Three constructs were expressed in *P. pastoris*, and purified as described in the Materials & Methods. Figure 2A shows the SDS-PAGE patterns of the human catalase derivatives. Each sample showed a single band of about 60 kDa, indicating that the catalase derivatives were adequately purified. The far-CD spectra of hCAT-(RGD)3 and hCAT-R9 were similar to that of hCAT (Figure 2B), suggesting that the fusion of peptides hardly changed the secondary structure of human catalase. Figure 2C shows the time-dependent decomposition of hydrogen peroxide by the human catalase derivatives. The activities of hCAT, hCAT-(RGD)3 and hCAT-R9 were 20,000, 19,500 and 24,300 units/mg protein, respectively, indicating that the fusion hardly reduced the specific activity of human catalase. These results are in a good agreement with the previous finding that the modification of C-terminus of catalase hardly influenced its activity [19].

Next, the cellular association of catalase derivatives in MAEC was examined at 37 °C or 4 °C. The activity to degrade hydrogen peroxide in cellular fractions was used to estimate the amount of catalase derivatives associated with cells. Cells untreated with any catalase derivative had a very low activity of $0.85 \pm 0.003$ mU/10^5 cells. The activity of cells was increased to $94.2 \pm 14.3$ mU/10^5 cells when cells were incubated
with hCAT for 30 min at 37 °C, indicating hCAT is associated with cells. The activity of cells treated with hCAT-(RGD)3 or hCAT-R9 at 37 °C was significantly higher (217 ± 31 and 261 ± 9 mU/10^5 cells, respectively) than that treated with hCAT. At 4 °C, a temperature at which internalization is essentially blocked, the activity of cells added with hCAT-R9 was significantly lower than that at 37 °C. On the other hand, the activities of cells added with hCAT or hCAT-(RGD)3 were not significantly different between the temperatures. These results suggest that the mechanisms of increased interaction of catalase derivatives with MAEC are different between hCAT-(RGD)3 and hCAT-R9.

Effect of catalase derivatives on intracellular ROS generation in MAEC under hypoxic conditions

The levels of intracellular ROS in MAEC were monitored using an ROS-sensitive probe. The fluorescence intensity increased significantly at 3 to 9 hr after exposure to hypoxia (Figure 4). Treatment of cells with hCAT before exposure to hypoxia slightly reduced the level of intracellular ROS in MAEC. Greater reduction was observed in MAEC treated with hCAT-(RGD)3 or hCAT-R9. The addition of 10 U/mL hCAT-(RGD)3 or hCAT-R9 was more effective in inhibiting the increase of the intracellular ROS than the addition of 100 U/mL hCAT (Figure 4). These results suggest that intracellular ROS generation in MAEC is induced soon after exposure to hypoxic stress, and can be
efficiently eliminated by catalase derivatives, especially by hCAT-(RGD)3 and hCAT-R9, in a dose-dependent manner.

**Adhesion of tumor cells to MAEC**

Next, we investigated whether hypoxia induces the adhesion of tumor cells to endothelial cells. Preincubation of MAEC with hydrogen peroxide significantly increased the adhesion of B16-BL6/Luc, colon26/Luc and LLC/Luc to MAEC (Figure 5A). The number of cells bound to MAEC tended to increase depending on the concentration of hydrogen peroxide from 25 to 100 µM. Separately, MAEC were incubated under hypoxic or normoxic conditions, and then adhesion of tumor cells was examined. Again, an increased binding of tumor cells to MAEC was observed under hypoxic conditions compared with under normoxia (Figure 5B). Then, the effects of catalase derivatives on the hypoxia-induced adhesion of B16-BL6/Luc were examined by treating MAEC with catalase derivatives at a concentration of 1 or 100 U/mL. hCAT reduced the hypoxia-induced cell adhesion (Figure 6). Under the same condition, hCAT-(RGD)3 and hCAT-R9 showed a greater inhibitory effect on the hypoxia-induced tumor cell adhesion than hCAT. Significantly lower numbers of tumor cells were bound to MAEC treated with hCAT-(RGD)3 or hCAT-R9 (p<0.05 compared with the same concentration of hCAT). The effects of catalase derivatives on tumor cell adhesion under normoxic conditions were less apparent than those under hypoxia, suggesting that
catalase derivatives inhibit hypoxia-induced, hydrogen peroxide-mediated tumor cell adhesion under hypoxic conditions.

Effect of catalase derivatives on transcription factor activity in MAEC under hypoxic conditions

To investigate the mechanism of inhibitory effect of catalase derivatives on hypoxia-induced cell adhesion, the activity of transcription factors HIF-1, NF-κB and AP-1 was estimated using pHRE-Luc, pNF-κB-Luc and pAP-1-Luc, respectively. The addition of hydrogen peroxide to cells increased the luciferase activity in cells (data not shown), suggesting that the transcription factors are activated by the addition of hydrogen peroxide. Culture of MAEC under hypoxic conditions induced significant activation of HIF-1, NF-κB and AP-1 (Figure 7). The activation of HIF-1 and NF-κB was inhibited by catalase derivatives, especially by hCAT-(RGD)3 or hCAT-R9. Strikingly, pre-treatment of 100 U/mL of hCAT-R9 almost completely inhibited the activation of HIF-1, NF-κB and AP-1. In accordance with the results of tumor cell adhesion (Figure 6), the effects of catalase derivatives under normoxic conditions were less significant than those under hypoxic conditions.

mRNA expression of adhesion molecules under hypoxic conditions

Finally, the changes in mRNA level of cell adhesion molecules that have reported to be
induced by HIF-1 or NF-κB [20] were examined using RT-PCR. The expression of

*ICAM-1* was induced in MAEC under hypoxia (Figure 8), but that of *VCAM-1* was

hardly changed (data not shown). The increase in the mRNA level of *ICAM-1* was

reduced by all catalase derivatives examined.
Discussion

Antioxidant enzymes including catalase have been believed to be promising therapeutic agents for a variety of human diseases related to oxidative stress, but the poor cell affinity of these enzymes has been a major obstacle in developing them as agents to protect cells and tissues from oxidative damage [8]. Recent studies have demonstrated that targeted or sustained delivery of bovine catalase by chemical modification is a therapeutic approach for ROS-mediated diseases, such as hepatic ischemia/reperfusion injury, cisplatin-induced nephrotoxicity and tumor metastasis [21-24]. However, chemical modification of protein, in general, has some disadvantages, such as productivity and uniformity of the products, compared with genetic modification from a viewpoint of clinical application. Furthermore, heterologous proteins, such as bovine catalase, can be immunogenic, so that the use of human catalase instead of bovine origin will reduce the possibility of antibody production after administration to patients [25]. In this study, we selected genetic engineering technique for modification of human catalase.

Previous studies have reported that enhanced delivery of biologically active proteins to cells is possible by using peptides that have high affinity with cells, including RGD peptide and oligoarginine peptide [26-28]. RGD motifs are found in many extracellular matrix proteins, and are recognized by and bound to integrins, a family of cell adhesion receptors expressed on the surface of various cells [29]. On the
other hand, oligoarginine peptide is a short repeat of basic amino acid arginine, and it has been reported not only to bind to negatively charged cell membranes, but also to initiate macropinocytosis, a special form of endocytosis, which facilitates the internalization of surface-bound compounds [30]. In order to increase the affinity of catalase with endothelial cells, the C-terminus of catalase was fused with (RGD)3 peptide or R9 peptide. It has been reported that three repeat of RGD has higher affinity with integrins than monomeric RGD [29, 31], so that the three repeat of the peptide was used for the construction of a catalase derivative containing RGD peptide. Both hCAT-(RGD)3 and hCAT-R9 showed an increased affinity for MAEC, but their cellular distribution was suggested to be different each other (Figure 3); hCAT-(RGD)3 could be bound to cell surface, whereas a part of hCAT-R9 was considered to be internalized during the 30-min incubation. R9 peptide fused to C-terminus of catalase might initiate the active cellular uptake mechanism as suggested in a previous report [30]. Although cellular distribution of these two types of catalase derivatives can be different each other, both eliminated intracellular ROS in MAEC, and inhibited the activation of HIF-1 and NF-κB under hypoxic condition. Because hCAT-(RGD)3 or hCAT-R9 was estimated more than 10 times effective than hCAT, the cytophilic nature of hCAT-(RGD)3 and hCAT-R9 is highly efficient for the removal of hydrogen peroxide generated in endothelial cells under hypoxic conditions. Two types of cytophilic catalase derivatives showed somewhat different intracellular distribution, but their inhibitory effects against
tumor cell adhesion were comparable. These results would be explained by the fact that hydrogen peroxide easily passes through cellular membranes.

The present study demonstrated that the cytophilic catalase derivatives have high binding affinity to MAEC. However, these results do not mean that their inhibition of ROS-mediated events in MAEC is the only mechanism for inhibiting tumor cell adhesion. The intracellular ROS level in tumor cells would be increased by several mechanisms [32]. In addition, the interaction with MAEC would further increase the ROS level in tumor cells. Therefore, cytophilic catalase derivatives bound to MAEC would reduce the ROS level in tumor cells which are interacting with MAEC.

In the design of catalase derivatives, the C-terminus of catalase was selected as the site for fusion with (RGD)3 peptide or R9 peptide. An X-ray structural analysis verified that the C-terminal region of human catalase is exposed on the surface of catalase molecule, and is distant from the active site [33]. In addition, human catalase contains a peroxisomal targeting signal in the C-terminal 11 amino acid sequence [16, 17]. These findings suggest that a peptide fused at the C-terminus is extended outside the protein, and is able to interact with integrins or negatively charged compounds on cell surface, which does not interfere with enzymatic activity of catalase. In fact, the activity and secondary structure of human catalase derivatives were not different from those of wild-type human catalase. These properties were quite contrast to those of chemically modified bovine catalase with an overall positive charge [22].
Tumor cells detached from primary tumors would be distributed to a variety of downstream organs via vascular or lymphatic pathways [34]. The adhesion to endothelial cells is the first step in the metastatic processes for tumor cells that have entered the blood circulation. It has been known that surgical removal of primary tumors increases the chance of tumor metastasis through a variety of mechanisms, including the release of tumor cells from primary tumors into circulation. Administration of cytophilic catalase derivatives before or during surgical removal of primary tumors could reduce the incidence of tumor metastasis through the inhibition of tumor cell adhesion as observed in the present studies using cultured cells.

In conclusion, we have demonstrated that the fusion of (RGD)3 peptide or R9 peptide to the C-terminus of catalase is an efficient method of increasing the cytophilic activity of catalase. Both hCAT-(RGD)3 and hCAT-R9 efficiently inhibited hypoxia-induced cell adhesion through controlling intracellular oxidative stress in MAEC under hypoxic conditions. These results suggest that newly designed cytophilic human catalase derivatives can be effective therapeutic agents for controlling hypoxia-induced tumor cell adhesion.
Acknowledgement

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Abbreviations

ROS, reactive oxygen species; hCAT, human catalase; hCAT-(RGD)3, human catalase fused with RGDRGDRGD; hCAT-R9, human catalase fused with RRRRRRRRR; CD, circular dichroism; NF-κB, nuclear factor κB; HIF-1, hypoxia inducible factor-1; AP-1, activator protein-1; HRE, hypoxia response element; ICAM-1, intercellular adhesion molecule-1; VCAM-1, vascular cell adhesion molecule-1; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; RT-PCR, reverse transcriptional polymerase chain reaction; P. pastoris, Pichia pastoris; CM-H2DCFDA, 5-(and-6)-chloromethyl-2',7'-dichlorodihydrofluorescein diacetate, acetyl ester
Reference


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Figure legends

Fig. 1: Schematic representation of human catalase derivatives.

The DNA sequence corresponding to the C-terminal 11 amino acids (SHLAAREKANL) of human catalase, which contains peroxisomal targeting signals, was replaced with oligonucleotides coding RGDRGDRGD or RRRRRRRRR to obtain hCAT-(RGD)3 or hCAT-R9, respectively. The fusion genes were inserted into the pNT401 expression vector and expressed in *P. pastoris*.

Fig. 2: Characteristics of human catalase derivatives.

(A) Coomassie brilliant blue stained SDS-PAGE gel. Lane 1, Precision Protein Unstained Marker (Bio-Rad); Lane 2, soluble fraction of cell extracts from *P. pastoris*; Lane 3; purified extracts from *P. pastoris* using glutathione sepharose column. (B) Far-CD spectra of catalase derivatives. The protein concentration was 15 mM in Tris buffer. Key: (solid line), hCAT; (broken line), hCAT-(RGD)3; and (dotted line), hCAT-R9. (C) Time-dependent decomposition of hydrogen peroxide by catalase derivatives. The concentration of remaining hydrogen peroxide was measured spectrophotometrically at 240 nm. Key: (solid line), hCAT; (broken line), hCAT-(RGD)3; and (dotted line), hCAT-R9.

Fig. 3: Catalase activity in MAEC after incubation with catalase derivatives.
The enzymatic activity of catalase to degrade hydrogen peroxide in MAEC was measured. Key: (closed bar), hCAT; (hatched bar), hCAT-(RGD)3; (open bar), hCAT-R9. Results are expressed as the mean ± SD of three independent measurements. The catalase activity of untreated MAEC was 0.85 ± 0.003 mU/10^5 cells. *p<0.05 compared with the hCAT-treated group; † p<0.05 compared with the same sample at 37 °C.

**Fig. 4: Effect of catalase derivatives on ROS level in MAEC.**

Intracellular ROS level was monitored using an ROS-sensitive probe CM-H2DCFDA. After exposed to solution containing catalase derivatives for 30 min, MAEC were treated with 5 µM CM-H2DCFDA for 1 hr. Cells were cultured under hypoxic conditions (8 % oxygen). The fluorescence intensity was measured (excitation 485 nm, emission 530 nm), and the ratio to the initial value was plotted against the incubation period as relative fluorescent intensity. Key: (▼) normoxia, untreated; (▽) hypoxia, untreated; (●) hypoxia, hCAT; (△, ▲) hypoxia, hCAT-RGD; (□, ■) hypoxia, hCAT-R9. Concentrations of the derivatives added were 10 (open symbols) or 100 U/mL (closed symbols). Results are expressed as the mean of 24 independent measurements. *p<0.05 compared with the hypoxia, untreated group (▽). † p<0.05 compared with the 100 U/mL hCAT-treated group (●).

**Fig. 5: Effect of hydrogen peroxide or hypoxia on tumor cell adhesion to MAEC.**
(A) Effect of hydrogen peroxide on tumor cell adhesion to MAEC. MAEC were cultured in M199 containing 0-100 µM hydrogen peroxide for 6 hr, and then co-cultured with B16-BL6/Luc, LLC/Luc or Colon26/Luc for 3 hr. Cell adhesion was evaluated by measuring the luciferase activity of B16-BL6/Luc, LLC/Luc and Colon26/Luc cells bound to MAEC. Results are expressed as mean ± SD of three independent measurements. *p<0.05 compared with 0 µM hydrogen peroxide-treated group. (B) Effect of hypoxia on tumor cell adhesion to MAEC. Results are expressed as mean ± SD of three independent measurements. *p<0.05 compared with the normoxia group. Key: (Closed bar), B16-BL6/Luc; (hatched bar), LLC/Luc; (open bar), colon26/Luc.

Fig. 6: Effect of catalase derivatives on adhesion of melanoma cells to MAEC.

MAEC were exposed to solution containing catalase derivatives for 30 min. Cells were cultured in hypoxia for 6 hr, and then co-cultured with B16-BL6/Luc for 3 hr. Cell adhesion was evaluated by measuring the luciferase activity of B16-BL6/Luc cells bound to MAEC. Results are expressed as mean ± SD of three independent measurements. Key: (hatched bar), 1 U/mL; (open bar), 100 U/mL. *p<0.05 compared with the hypoxia, untreated group; #p<0.05 compared with the hCAT-treated group at the same dose; §p<0.05 compared with the normoxia, untreated group.

Fig. 7: Effect of catalase derivatives on transcription factor activity in MAEC
under hypoxia.

MAEC were transfected with pHRE-Luc (A), pNF-kB-Luc (B), or pAP-1-Luc (C). MAEC were exposed to solution containing catalase derivatives for 30 min. After incubation under hypoxic conditions for 6 hr, cells were lysed with lysis buffer. Each transcription factor activity was determined by measuring luciferase activity of the cell lysates. Results are normalized to the value of the normoxia, untreated group and are expressed as mean ± SD of three independent measurements. Key: (hatched bar), 1 U/mL; (horizontal striped bar), 10 U/mL; (open bar), 100 U/mL. *p<0.05 compared with the hypoxia, untreated group; #p<0.05 compared with the hCAT-treated group at the same dose; §p<0.05 compared with the normoxia, untreated group; †p<0.05 compared with the 100 U/mL hCAT-treated group.

Fig. 8: Effect of catalase derivatives on mRNA expression of ICAM-1 in MAEC under hypoxic conditions.

Total RNA was isolated from MAEC after incubation under hypoxic conditions for 6 hr, and the mRNA expression of ICAM-1 in MAEC was estimated by RT-PCR. The mRNA expression of ICAM-1 was normalized to that of GAPDH, and the value was then normalized to the value of the normoxia, untreated group. Results are expressed as mean ± SD of three independent measurements. *p<0.05 compared with the hypoxia, untreated group.
Fig. 1
Fig. 2

Panel A: Gel electrophoresis images of hCAT, hCAT-(RGD)₃, and hCAT-R9 with molecular weight markers.

Panel B: Wavelength absorption data showing changes in absorbance over time.

Panel C: Time-course of absorbance changes at 240 nm.
Fig. 3

Catalase activity (mU/10^5 cells)

- 37°C
- 4°C

* indicates significant difference
† indicates another significant difference
Fig. 4
Fig. 5

A

Cell adhered (% of cells added)

0 20 40 60 80

0 μM 25 μM 50 μM 100 μM

B

Cell adhered (% of cells added)

0 20 40 60 80 100

Normoxia Hypoxia
Fig. 6

**Graph Description:**

- **X-Axis:** Cells adhered (% of cells added)
- **Y-Axis:** Number of cells adhered
- **Legend:**
  - CAT
  - Hypoxia

**Data Points:**

- **Normoxia:**
  - Control
  - CAT
  - CAT-RGD

- **Hypoxia:**
  - Control
  - CAT
  - CAT-RGD

**Significance:**

- *: Statistically significant difference
- #: Additional significance marker

**Comparison:**

- Normoxia shows lower adhered cells compared to Hypoxia.
- CAT-RGD treatment shows increased adherence compared to CAT treatment.
Fig. 8