Stem Cells, Tissue Engineering, and Hematopoetic Elements

Adipose Injury–Associated Factors Mitigate Hypoxia in Ischemic Tissues through Activation of Adipose-Derived Stem/Progenitor/Stromal Cells and Induction of Angiogenesis

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Based on the analysis of exudates from injured adipose tissue, we prepared a mixture containing the injury-associated growth factors at the same proportion as the exudates, named adipose injury cocktail (AIC). We hypothesized that AIC induces a series of regenerating and angiogenic processes without actual wounding. The purpose of this study is to elucidate the therapeutic potentials of AIC. AIC preferentially activated adipose-derived stem/progenitor/stromal cells (ASCs) to proliferate, migrate, and form networks compared with vascular endothelial cells, whereas vascular endothelial growth factor did not induce mitogenesis or chemotaxis in human ASCs. Each component growth factor of AIC was differently responsible for the ASC activation. AIC-treated ASCs tended to differentiate into adipocytes or vessel-constituting cells rather than into other cell types. In ischemic adipose tissues of mice, induced by either a surgical intervention or diabetes, AIC administration enhanced proliferation, especially of CD31−/CD34+/H11545 ASCs, and mitigated tissue hypoxia by increasing capillary density and reducing fibrogenesis. These results suggest that AIC may have therapeutic potentials for various ischemic/hypoxic conditions by inducing adipose remodeling and neovascularization through activation of ASCs and other cells. Treatment with AIC has many advantages over cell-based therapies regarding morbidity, cost, and physical risks and may be used as an alternative therapy for improving tissue oxygen.

Adipose tissue turns over very slowly and contains a diversity of cell types, including adipocytes, stromal cells, vascular endothelial cells (ECs), pericytes, and resident blood-derived cells.1,2 Vascular stromal cells isolated from adipose tissue have been shown to contain multipotent cells called adipose-derived stem/progenitor/stromal cells (ASCs).3 The adipose progenitors were recently identified in perivascular locations, and this generated a hypothesis about their identity with pericytes.4–7 ASCs contribute to adipose tissue turnover by providing new-generation cells, and they also play predominant roles during the remodeling process that compensate for apoptotic or degenerative changes.8,9 In diverse models of ischemic tissues, ASCs were shown to be potential therapeutic tools owing to their ability to promote angiogenesis by providing ECs and/or releasing angiogenic growth factors.9–15

In regenerative medicine, stem or progenitor cells need to be activated with a microenvironment optimized for each purpose. By supplying conditions and factors involved in inflammation or tissue remodeling/repair, we can provide microenvironments that evoke regenerative events, including cell proliferation, migration, and differentiation. Therefore, tissue pretreatments, eg, intentional wounding, have been occasionally used for preparing microenvironments to induce tissue remodeling. For example, skin resurfacing procedures, such as laser or chemical peeling, are destructive but induce skin renewal and resolve various cutaneous problems. As another example, ectopic osteogenesis in skeletal muscle

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could be experimentally induced by overexpression of bone morphogenetic protein only when muscle ischemia and regeneration were induced by free grafting. Thus, tissue wounding is considered to be a potent method for preparing microenvironments to be suitable for tissue engineering and regenerative therapies. In this study, instead of tissue wounding, we used injury-associated soluble factors to mimic such regenerative microenvironments.

The repair/remodeling process of injured adipose tissue is controlled by spatiotemporal actions of multiple growth factors secreted from the extracellular matrix and various types of cells, including dying cells. We previously showed that after surgical injury to human adipose tissue, different growth or inflammatory factors were secreted from the injured tissue in each wound-healing phase (coagulation, inflammation, and proliferation). In the coagulation stage (within 1 day after wounding), basic fibroblast growth factor (bFGF), epidermal growth factor (EGF), platelet-derived growth factor (PDGF), and transforming growth factor-β (TGF-β) were released, but all of them were detected in decreasing amounts in the following stages.

In this study, we hypothesized that the soluble factors released in the initial phase triggered a series of subsequent inflammatory and regenerating responses and that administration of the factors in vivo or in vitro would reproduce at least part of the microenvironment associated with adipose injury without actual wounding. We prepared a mixture of the factors, termed adipose injury cocktail (AIC), which comprised the four major early-stage factors (bFGF, EGF, PDGF, and TGF-β) combined in the proportions found in early wound exudates. We also hypothesized that AIC may improve tissue oxygen tension by activating quiescent ASCs and/or ECs, initiating a series of adipose-repairing events and regenerating adipose and vasculature. To examine the therapeutic potential of AIC, we further performed in vivo experiments to evaluate the effects of AIC on acutely induced ischemic adipose tissue, chronically ischemic adipose tissue with diabetes, and unperturbed adipose tissue.

**Materials and Methods**

**Human Cell Isolation and Culture**

Human liposuction aspirates were obtained from 12 healthy female donors [mean ± SD age, 39.3 ± 3.4 years; mean ± SD body mass index (calculated as weight in kilograms divided by height in meters squared), 22.6 ± 0.7] undergoing elective liposuction of the abdomen or thighs. Each patient provided informed consent. The protocol was approved by the ethical committee of University of Tokyo School of Medicine before the procedure. Human ASCs (hASCs) were isolated from the aspirated fat as described previously. Briefly, the aspirated fat tissue was washed with PBS and was digested at 37°C in PBS containing 0.075% collagenase for 30 minutes on a shaker. Mature adipocytes and connective tissue were separated from pellets by centrifugation (800 × g for 10 minutes). The cell pellets were resuspended, filtered through 100-μm mesh, plated at a density of 5 × 10^6 nucleated cells/100-mm dish, and cultured at 37°C in an atmosphere of 5% CO_2 in humid air. The culture medium was Dulbecco's modified Eagle's medium (Nissui Pharmaceutical, Tokyo, Japan) containing 10% fetal bovine serum (FBS). Primary cells were cultured for 7 days and were defined as passage 0. The medium was replaced every 3 days. Cells were passaged every week by trypsinization. hASCs at passages 1 to 3 were used in the experiments.

Isolation and culture of human umbilical vein ECs (HUVECs) were performed according to the methods described previously. Samples were collected immediately after delivery. The umbilical cord was separated from the placenta by clipping both ends and was irrigated with 1% iodine/PBS solution. To eliminate iodine, the intracelal space was rinsed with M199 medium and filled with 0.25% trypsin in PBS. Both ends were again clipped, followed by incubation for 10 minutes at 37°C. Then, the intracelal space was rinsed with endothelial basal medium (EBM; Cambrex, Walkersville, MD), and cells were collected. The cells were centrifuged at 450 × g for 5 minutes, attached to 100-mm plastic dishes, and cultured with EBM containing 2% FBS. Primary cells were cultured for 7 days and were defined as passage 0. HUVECs at passages 3 to 5 were used in the experiments.

**Proliferation Assay**

In this study, the mixture of four injury-associated growth factors (PDGF, TGF-β, bFGF, and EGF) composed the AIC. The final concentration of AIC was 2.5 ng/mL PDGF, 1 ng/mL TGF-β, 10 ng/mL bFGF, and 2 ng/mL EGF (all from Wako Pure Chemicals, Osaka, Japan) mixed in an appropriate solution (EBM containing 0% or 2% FBS in vitro or PBS in vivo); the proportion of the four factors was determined in a previous study, and the concentrations were determined based on a preliminary experiment. hASCs or HUVECs were plated in a six-well plate at 2 × 10^4 cells per well in EBM containing 2% FBS with or without AIC (control). The cell numbers were counted after 3, 6, and 9 days in culture using a cell counter (NucleoCounter; ChemoMetec, Allerod, Denmark). 5-Bromo-2-deoxyuridine (BrdU) incorporation assays were performed using the APC BrdU Flow Kit (BD Biosciences, San Jose, CA). Cells were cultured for 48 hours in EBM containing 2% FBS with or without AIC, followed by BrdU labeling for 2 hours at a final concentration of 10 μmol/L. Flow cytometry was then performed using an LSR II flow cytometry system (BD Biosciences).

**Migration Assay**

hASCs and HUVECs were resuspended (2 × 10^5 cells in 200 μL of serum-free EBM) and seeded into the upper chamber of a cell migration apparatus that had a floor perforated with 8-μm pores (Transwell; Corning Life Sciences, Lowell, MA). The lower chamber was filled with 600 μL of serum-free EBM with or without AIC or with 100 ng/mL...
vascular endothelial growth factor (VEGF). To evaluate the dose-dependent effects of AIC, a medium containing 10 times the concentration of AIC (10 × AIC) and diluted AIC media (0.1 ×, 0.01 ×, and 0.001 × AIC) were used. Furthermore, we tested different combinations of two or three of the four AIC factors (incomplete AIC). The cells were then allowed to migrate for 4 hours at 37°C. Then they were fixed in 100% methanol for 15 minutes and stained with 0.1% Giemsa solution for 10 minutes. Cells remaining on the bottom of the upper chamber were removed using a cotton swab. Cells that migrated through the pores were counted under a light microscope.

**Induced Differentiation of Cultured Cells**

After culturing in EBM containing 2% FBS with or without AIC for 3 days, hASCs were differentiated into adipogenic, chondrogenic, and osteogenic lineages. For adipogenic differentiation, cells were incubated for 21 days in adipogenic medium (Dulbecco’s modified Eagle’s medium containing 10% FBS, 0.5 mmol/L isobutyl-methylxanthine, 1 μmol/L dexamethasone, 10 μmol/L insulin, and 200 μmol/L indomethacin). Adipogenic differentiation was visualized with oil red O staining. For quantitative analysis of lipid droplets, Nile red fluorescence was measured by AdipoRed (Cambrex) excitation at 485 nm and emission at 535 nm using a fluorescent plate reader (DTX 880 Multimode Detector; Beckman Coulter, Fullerton, CA). For chondrogenic differentiation, cells were incubated for 21 days in Dulbecco’s modified Eagle’s medium containing 1% FBS supplemented with 6.25 mg/mL insulin, 10 ng/mL TGF-β1, and 50 nM of ascorbate-2-phosphate. Chondrogenic differentiation was visualized with Alcian blue staining. For quantitative analysis, a micromass culture system was used, as reported previously. Cells were plated in a 15-mL tube and were cultured in the chondrogenic medium for 21 days. Then, the diameter of a micromass was measured. For osteogenic differentiation, cells were incubated for 21 days in Dulbecco’s modified Eagle’s medium containing 10% FBS supplemented with 0.1 mM of dexamethasone, 50 mM of ascorbate-2-phosphate, and 10 mM of β-glycerophosphate (Nacalai Tesque, Kyoto, Japan). Osteogenic differentiation was visualized with von Kossa staining. For quantitative analysis of total calcium, calcium deposition was evaluated based on the ortho-cresolphthalein complexone method using the Calcium C-Test Wako Kit (Wako Chemicals, Osaka, Japan) according to the manufacturer’s instructions.

**Quantitative Real-Time RT-PCR**

Extraction of total RNA from homogenized mouse inguinal adipose tissue was performed using the RNeasy Mini Kit (QIAGEN Inc., Valencia, CA), followed by reverse transcription. We amplified cDNA in 40 cycles using a SYBR Green PCR Master Mix and an ABI 7700 sequence detection system (both from Applied Biosystems, Foster City, CA). Expression levels were calculated by the comparative CT method with ribosomal 18S RNA as an internal control. The primer sequences used in real-time PCR analysis were as follows: CD31: 5′-TTGAGCCTCACCAAGAGAACGGA-3′, 5′-AATCCAGGAATCGGCTGTCTTCTCT-3′; Flk-1: 5′-AGGGCATTGAGTCCAATTACACA-3′, 5′-AGACCATGTGCTGTCTTCTCA-3′; CD34: 5′-AGACTCAGGAAAGGCAATGTA-3′, 5′-GCCACCACATGGTCTTGCTGAA-3′; and 18SrRNA: 5′-TCACTTCTCGATGTAGTCGCCC-3′, 5′-TCTTGGATTGTGATGCCTTCT-3′.

*In Vitro Angiogenesis (Network Formation) Assay*

Network formation was assessed for hASCs and HUVECs. Matrigel (BD Biosciences) was poured into 96-well plates (50 μL per well) and polymerized (30 minutes at 37°C). After incubation in EBM containing 2% FBS with or without AIC for 24 hours as a pretreatment, 5000 cells resuspended in 50 μL of EBM containing 2% FBS without AIC were plated on Matrigel and cultured for 6 hours. Branching formation was observed with phase microscopy, and the length of cytoplasmic extensions per field was calculated. Dose-dependent effects of AIC on network formation were also evaluated as described in the migration assay. To distinguish the two types of cells co-cultured on Matrigel, hASCs and HUVECs were stained with DiO and CM-Dil (Invitrogen-Molecular Probes, Carsibad, CA), respectively, in advance (overnight at 37°C). For staining the networks formed on the Matrigel, cells were fixed with 4% paraformaldehyde, blocked with 10% normal goat serum with PBS containing 1% bovine serum albumin for 30 minutes at room temperature, and incubated with Alexa Fluor 488–conjugated isoelectin or anti-von Willebrand factor (vWF) antibody overnight at 4°C. After 1 hour of incubation with a secondary antibody (only for immunostaining), Hoechst 33342 (Invitrogen) staining was performed, and the cells were observed using a confocal microscope system.

**Mouse Nonischemic and Ischemic Models of Inguinal Adipose Tissue**

All animal care was in accordance with institutional guidelines. Eight-week-old ICR mice or db/db mice were anesthetized with pentobarbital (50 mg/kg weight).

**Intact (Nonischemic) Model**

A 3-mm incision was made in the inguinal region, and the subcutaneous adipose pad was identified. After the measurement of tissue partial pressure of oxygen (pO2), AIC (0.2 mL of PBS containing 12.5 nmg/mL PDGF, 5 ng/mL TGF-β, 50 ng/mL bFGF, and 10 ng/mL EGF; 5× AIC used in vitro) was injected into the fat pad via a 29-gauge needle-attached syringe (Myjector; Terumo, Tokyo, Japan). As a control, 0.2 mL of PBS was injected into the contralateral fat pad. At designated time points (before injection and at 1, 2, and 4 weeks; n = 10 mice per time point), tissue pO2 was measured, and the fat pad was harvested. The samples were evaluated using whole-mount staining and capillary density measurements.
Acute Ischemia Model

A linear 1-cm incision was made in the inguinal region, and the \( p_O_2 \) of the subcutaneous fat pad was measured. The nutrient vessels of the fat pad arising from the femoral artery and its branch to the adductor muscles were exposed and ligated. Small communicating vessels to the skin from the fat pad were left intact. Immediately after the ligation, \( p_O_2 \) was measured again, and then AIC or PBS was injected into the adipose tissue as described for experiment 1. At designated time points (before injection and at 1, 2, and 4 weeks; \( n = 10 \) mice per time point), tissue \( p_O_2 \) was measured and the fat pad was harvested. The samples were evaluated using whole-mount staining, measurements of capillary density and fibrosis area, and real-time PCR.

Chronic Ischemia Model in db/db Mice

The subcutaneous inguinal adipose tissue of db/db mouse was identified after a small incision. \( p_O_2 \) was measured, and AIC or PBS was injected as described for experiment 1. At designated time points (before injection and at 1, 2, and 4 weeks; \( n = 5 \) mice per time point), tissue \( p_O_2 \) was measured and the fat pad was harvested. The samples were evaluated using whole-mount staining, capillary density measurements, histologic analysis, and flow cytometric analysis.

Measurements of \( p_O_2 \) in Inguinal Adipose Tissue

Tissue \( p_O_2 \) was measured using an oxygen electrode (200 \( \mu \)m in diameter) and an oxygen monitor (Eiko Kagaku, Tokyo, Japan). The oxygen electrode was directly inserted into mouse inguinal adipose tissue, and an indifferent electrode was inserted into the abdominal subcutaneous space; the system was allowed to equilibrate for 15 to 20 minutes before every measurement.

Whole-Mount Staining

Visualization of living adipose tissue was performed as described previously. Briefly, the adipose tissue was minced into 3-mm pieces and was incubated with the following reagents for 30 minutes: BODIPY 558/568 (Invitrogen-Molecular Probes) to stain adipocytes, Alexa Fluor 488–conjugated isoelectin to stain ECs, and Hoechst 33342 to stain all nuclei. The sample was then washed and directly observed using a confocal microscope system.

Histologic and Immunohistochemical Analyses

Harvested adipose tissue samples were cleaned in PBS, zinc fixed (Zinc Fixative; BD Biosciences), and paraffin embedded. Specimens were sliced into 5-\( \mu \)m sections and were stained with H&E to examine histologic changes. Azan staining was performed to assess tissue fibrosis. To evaluate capillary density, the sections were incubated for 30 minutes with a staining solution containing Alexa Fluor 488–conjugated isoelectin and Hoechst 33342 for the same purpose, immunostaining with goat anti-mouse CD31 antibody (BD Biosciences) was also performed. Photographic images of the stained sections were acquired using a fluorescence microscope equipped with a camera. Capillary density was expressed as the number of microvessels observed per optical field averaged across three to four images acquired from each section. The sections were also immunostained with rabbit anti-mouse perilipin antibody (Sigma-Aldrich, St. Louis, MO) to visualize viable adipocytes. The images acquired using the fluorescence microscope were evaluated for the average size of viable adipocytes and their distribution.

Flow Cytometry

Mouse inguinal adipose tissue was minced into 1-mm pieces and digested at 37°C in PBS containing 0.075% collagenase for 60 minutes on a shaker. After centrifugation (800 \( \times g \) for 10 minutes), the cell pellets (stromal vascular fraction) were resuspended, filtered through a 100-\( \mu \)m mesh, and counted for the total number of cells. Cells were stained with the following monoclonal antibodies conjugated to fluorochromes: anti-mouse CD31-phycoerythrin (PE; BD Biosciences), anti-mouse CD34–fluorescein isothiocyanate (eBioscience, San Diego, CA), and anti-mouse CD45-PE Cy7 (Beckman Coulter, Fullerton, CA). Then, multicolor flow cytometric analyses were performed using an LSR II flow cytometry system. We counted the total number of CD45+/CD31+ cells (vascular ECs) and CD45+/CD31+/CD34+ cells (ASCs) contained in the stromal vascular fraction.

Primary cultured hASCs and HUVECs were examined for surface marker expression using flow cytometry. The following fluorochrome-conjugated monoclonal antibodies were used: anti-CD31–PE, anti-CD34–PE, anti-CD45–PE, anti-CD90–PE, anti-CD105–PE, anti-CD146–PE, anti-CD117–PE (all from BD Biosciences), anti–Fk–1–PE (R&D Systems, Minneapolis, MN), anti-CD133–PE (Miltenyi Biotech, Gladbach, Germany), and anti–vWF–PE (Abcam, Cambridge, UK). Cells were incubated with each antibody for 30 minutes and then were analyzed using an LSR II flow cytometry system. Gates were set on the basis of staining with isotype controls so that no more than 0.1% of cells were detected with control antibodies.

Statistical Analysis

Results are expressed as mean ± SEM The unpaired t-test was used to evaluate the differences between two groups, except in the comparison of \( p_O_2 \) among different mouse inguinal adipose tissues, for which the paired t-test was applied. The level of significance was set at \( P < 0.05 \).

Results

AIC Promotes the Proliferation of hASCs and HUVECs

Because AIC administration was intended to simulate adipose tissue injury, we first investigated the re-
sponse to AIC in vitro of two primary cellular components of adipose tissue, ASCs and ECs. Using fluorescence-activated cell sorting analysis, we confirmed in advance that there were no contaminating ECs in isolated hASC populations (see Supplemental Figure S1 at http://ajp.amjpathol.org).

AIC significantly promoted proliferation of hASCs and HUVECs (Figure 1, A and B; also see Supplemental Figure S2 at http://ajp.amjpathol.org). Although the promotion was observed earlier in HUVECs than in hASCs, HUVECs lost the proliferative effects by day 9 and detached from the dishes. To examine the synergism of the AIC factors, we performed BrdU incorporation assays with incomplete AIC (different combinations with three factors contained in AIC). The proliferative effect of incomplete AIC on hASCs was maintained despite the lack of any one of the four growth factors in AIC (Figure 1C). This suggests that the four factors acted synergistically in promoting hASC proliferation. All the growth factors of AIC showed proliferative effects on ASCs except TGF-β, and EGF had the strongest effect of the four growth factors (see Supplemental Figure S3A at http://ajp.amjpathol.org).

AIC Induces hASC Migration but Not HUVEC Migration

The migration assay showed that AIC induced hASC migration but not HUVEC migration (Figure 1D). In contrast, VEGF induced HUVEC migration but not hASC migration. Medium containing 10× AIC promoted the migration of both cell types to the same degree as 1× AIC (Figure 1D), whereas the dilution experiments showed that media containing <0.1× AIC had no significant effects on the migration of either hASCs or HUVECs (see Supplemental Figure S4A at http://ajp.amjpathol.org).

Incomplete AIC had a reduced migration effect on hASCs, except for the combination lacking bFGF (Figure 1E). These results suggest that three growth factors, PDGF, TGF-β, and EGF, worked synergistically in promoting hASC migration, although an increased synergistic effect of all four factors cannot be completely ruled out. Of the three necessary factors, EGF seemed to be the most influential for inducing hASC migration.

AIC-Treated hASCs Show a Higher Capacity for Adipogenic Differentiation and a Lower Capacity for Chondrogenic and Osteogenic Differentiation

Human ASCs were experimentally induced to differentiate into adipocytes, chondrocytes, or osteocytes, regardless of AIC treatment (Figure 2A). Quantitative analyses showed that hASCs pretreated with AIC produced significantly more lipid droplets with the induced adipogenic differentiation compared with controls. On the other hand, AIC-treated hASCs formed significantly smaller micromasses after chondrogenic differentiation, and significantly less calcium was deposited after osteogenic differentiation than control (Figure 2B).

AIC-Treated hASCs Form Complex Capillary-Like Networks

Network formation assay on Matrigel showed that AIC-pre-treated hASCs formed a complex capillary-like network structure, whereas untreated hASCs showed cell sprouting and formed only closed polygon-like structures (Figure 3A). AIC-treated hASCs formed networks more quickly and intricately than did HUVECs, which formed orderly networks regardless of AIC pretreatment. Supplementation of VEGF
did not affect the network length of hASCs or HUVECs, except when HUVECs had been pretreated with AIC (Figure 3B). The effects of AIC on hASC network formation were dose dependent in a range of $0.001 \times 10^{-3}$ to $1 \times 10^{-3}$ AIC, although $1 \times 10^{-3}$ and $10 \times 10^{-3}$ AIC showed equivalent efficacy (see Supplemental Figure S4B at http://ajp.amjpathol.org). The assessments using incomplete AIC showed that the growth factors had synergistic effects on hASC network formation. Among the four factors, bFGF had the strongest promoting effect; the absence of bFGF significantly suppressed hASC network formation. In addition, any combination of two growth factors including bFGF or bFGF alone showed a significant increase in total network length compared with control (Figure 3C; also see Supplemental Figure S3B at http://ajp.amjpathol.org). The networks formed by AIC-treated hASCs were positive for lectin (Figure 3D) and expressed vWF at the levels observed in HUVECs, whereas the formations of control hASCs were nearly negative for vWF (Figure 3E). Furthermore, we detected a small number of CD31-expressing cells in the networks formed by AIC-treated hASCs (see Supplemental Figure S5 at http://ajp.amjpathol.org). These indicated that AIC pretreatment contributed to ASCs’ differentiation into ECs. In the co-culture assay with HUVECs and AIC-treated hASCs on Matrigel, the two types of cells collaboratively formed networks. Both cells were equally positive for vWF, suggesting that AIC-treated hASCs were similar to ECs (see Supplemental Figure S6 at http://ajp.amjpathol.org).

AIC Effect Quicker Recovery of $pO_2$ and Higher Capillary Density in a Surgically Induced Ischemic Adipose Tissue

To investigate the in vivo effects, AIC was injected into ischemic or nonischemic subcutaneous adipose tissue in mice. When AIC or vehicle was administered into unperturbed inguinal adipose tissue of healthy mice, AIC in-

![Figure 3](http://ajp.amjpathol.org)
creased capillary density and resulted in a higher mean ± SEM pO₂ value (61.5 ± 3.5 mm Hg, n = 9) than in controls (51.9 ± 3.6 mm Hg, n = 9) at 1 week, but there was no significant difference between the two groups at 2 and 4 weeks (see Supplemental Figures S7 and S8 at http://ajp.amjpathol.org). By surgical intervention (ligation of the nutrient vessels; Figure 4A) for the ischemic models, mean ± SEM tissue pO₂ was reduced from 61.4 ± 1.0 mmHg (n = 134) to 18.1 ± 1.0 mmHg (n = 64). One week later, the AIC-treated ischemic adipose tissue showed significantly improved mean ± SEM pO₂ (44.6 ± 2.9 mmHg, n = 9) compared with the vehicle-injected contralateral ischemic adipose tissue (29.2 ± 2.1 mmHg, n = 9) (Figure 4B). Whole-mount histologic analysis showed a visible increase in adipose capillaries as early as 1 week after AIC treatment (Figure 4C), and the capillary density was significantly higher in the AIC-treated group throughout the investigation period (Figure 4D; see also Supplemental Figure S8 at http://ajp.amjpathol.org).

Gene expression analyses of the whole inguinal adipose tissue revealed that AIC treatment induced up-regulation of CD34 mRNA at 1 week, followed by up-regulation of Flk-1 and CD31 (Figure 5A). The CD34 up-regulation might suggest the ASC proliferation promoted by AIC because CD34⁺/CD31⁻ cells in a living adipose tissue are regarded as ASCs. In the adipose repair/remodeling processes, fibrous scar tissue was also formed. Histologic assessment revealed that ischemia induction and vehicle administration caused a significant increase in the fibrous area throughout the 4-week experimental period (Figure 5B). AIC treatment significantly suppressed the fibrogenesis at 2 and 4 weeks.

**AIC Induces Angiogenesis and Adipogenesis in Chronically Ischemic Adipose Tissue of Diabetic Mouse**

The adipose tissue of diabetic obese mice showed significantly lower mean ± SEM pO₂ (45.5 ± 1.5 mmHg, n = 36) than that of nondiabetic mice (61.4 ± 1.0 mmHg, n = 134). Administration of AIC also caused an elevation of pO₂ in the chronically ischemic adipose tissue accompanying diabetes; the pO₂ level was significantly higher at 1 and 4 weeks in AIC-treated adipose tissue compared with untreated adipose tissue and vehicle-treated counterparts (Figure 6A). AIC treatment was also associated with a significant increase in capillary density compared with control; this was observed as early as 1 week and up to 4 weeks (Figure 6B and C; see also Supplemental Figure S8 at http://ajp.amjpathol.org).

Flow cytometric analysis showed that AIC-treated adipose tissue included a much larger number of adipose-derived proliferating (CD45⁻/BrdU⁺) cells than did the control tissue (Figure 7A). Among the proliferating cells, CD31⁺/CD34⁺ cells (ASCs) and CD31⁻/CD34⁺ cells (ECs) were main populations and composed 53.1% and 32.4%, respectively. Next, we performed immunohistologic staining with perilipin. We observed viable small-sized adipocytes (<25 μm in diameter), suggesting ongoing adipogenesis, and they increased in number over time in...
AIC-administered adipose tissue (Figure 7, B and C). Capillaries were frequently observed around the small-sized adipocytes.

Thus, local administration of AIC mitigated hypoxia of diabetic adipose tissue. Our results suggest that AIC-responsive cellular events contributed to angiogenesis and adipogenesis during the remodeling process and improved tissue oxygen tension.

Discussion

Of the four AIC factors, bFGF and TGF-β are generally released from injured extracellular matrix or dying cells, whereas EGF, PDGF, and TGF-β are released from activated platelets on bleeding. Recently, it was also demonstrated that EGF is released from apoptotic ECs and activates antiapoptotic response in mesenchymal stem cells. AIC does not contain either VEGF or hepatocyte growth factor (HGF), which are considered to be major proangiogenic factors. In migration and capillary-forming assays, VEGF preferentially affected ECs compared with hASCs. VEGF and HGF are released from hASCs in some conditions, including hypoxia, and may not be essential activators for ASCs. Indeed, VEGF and HGF protein levels increased in the later phase of adipose wound healing.

We found that AIC significantly promoted hASCs to proliferate, migrate, and form networks. Especially, the promotion of ASCs to form networks was one of the most impressive effects of AIC in this study. AIC-treated hASCs were able to form complex networks, which were positive for vWF and lectin, by themselves. Untreated hASCs expressed pericyte markers, such as α-smooth muscle actin and CD140b, yet their expression was suppressed by AIC treatment (data not shown). These results suggest that AIC induced hASCs at least partly to differentiate into cells of endothelial lineage. As for mesenchymal differentiation capacities, AIC-treated hASCs showed a higher capacity for adipogenic differentiation and a lower capacity for chondrogenic or osteogenic differentiation compared with controls. Thus, in response to AIC, hASCs seemed...
to preferentially differentiate into adipocytes and/or vascular cells. This would explain a lot about the physiologic manifestations of adipose tissue remodeling observed after injury.

We performed preliminary experiments to evaluate the extent of endothelial differentiation of hASCs placed in diverse environments (eg, in cultures with bFGF or inflammatory cytokines, in various major media, on major extracellular matrix, in hypoxic conditions, and in co-cultures with HUVECs), and we found that no single factor or simple combination could induce a remarkable tendency to differentiate the endothelial lineage of hASCs (data not shown), although antibodies against bFGF and PDGF were reported to affect tube formation by ASCs.25 Therefore, various factors and their complicated interplay seemed to be essential. Indeed, in some previous reports, superior results for angiogenesis have been obtained through the application of multiple factors than through the introduction of a single factor, eg, paired combinations of bFGF and PDGF,26 VEGF and bFGF,28,29 VEGF and PDGF,30 or PDGF and TGF-β,31 or a triple combination of bFGF, PDGF, and VEGF.32 Our findings with incomplete AIC with either two or three growth factors suggest that PDGF, EGF, and TGF-β contributed to hASC migration and that bFGF played a pivotal role in network formation, although these effects were likely augmented by the synergism of all of the factors. Although the reproducible methods for inducing hASC

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**Figure 6.** AIC effects on diabetic adipose tissue with chronic hypoxia, part I. A: Time course of adipose tissue $\rho_{O_2}$ in db/db mice ($n = 5$). B: Whole-mount histologic analysis of living adipose tissue. Harvested, unfixed tissue samples were stained with BODIPY (adipocytes; green), lectin (ECs; red), and Hoechst 33342 (nuclei; blue). Scale bars = 50 μm. C: Histologic sections were stained with lectin (ECs; red) and Hoechst 33342 (nuclei; blue), and capillary density was calculated ($n = 6$). *$p < 0.05$, **$p < 0.01$. Scale bars = 100 μm. Error bars represent SEM.

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**Figure 7.** AIC effects on diabetic adipose tissue with chronic hypoxia, part II. A: Flow cytometric analysis of freshly isolated stromal vascular fraction from diabetic adipose tissue. The histograms of CD45$^-$ cells represent the proportion of BrdU$^+$ cells in vehicle-treated tissue (left) or in AIC-treated tissue (right). B: Immunohistochemical staining images. The sections were stained with perilipin (viable adipocytes; green), lectin (ECs; blue), and Hoechst 33342 (nuclei; blue). Perilipin-negative adipocytes surrounded by macrophages (so-called crown-like structure; white arrows), suggesting degenerated adipocytes, were occasionally observed. The proportion of perilipin-positive small-sized adipocytes (yellow arrowheads), suggesting ongoing adipogenesis, was counted and calculated ($n = 4$). *$p < 0.05$. Scale bars = 100 μm. Error bars represent SEM.
differentiation into smooth muscle cell lineages have been reported previously. ASC differentiation protocols into ECs currently remain to be established. These results provide some insights into establishing an efficient, standard protocol for this purpose.

After confirmation of AIC potentials in vitro, we targeted the development of a therapy for mitigating tissue hypoxia; the overall results of animal studies indicated that AIC had promising therapeutic effects. When administered in normoxic adipose tissue (mean \( \pm \) SEM \( pO_2 \) = 61.4 \( \pm \) 1.0 mmHg), the capillary density significantly increased, but the adipose tissue \( pO_2 \) did not significantly change. This was possibly because each tissue has an individual maximum tissue \( pO_2 \) that is restricted by the arterial \( pO_2 \). When administered into hypoxic adipose tissue induced by surgical intervention, the AIC treatment significantly increased capillary density and accelerated the recovery of tissue \( pO_2 \). Furthermore, AIC contributed to the inhibition of ischemia-induced fibrogenesis, probably by accelerating favorable tissue regeneration/remodeling, as suggested by a previous study. A key contribution of AIC in the repair process is likely angiogenesis; AIC administration activated ASCs, increased the number of ECs, increased capillary density, and, consequently, accelerated tissue \( pO_2 \) recovery. In chronically hypoxic adipose tissue of diabetic mice, the most influential effects among the three experimental models were observed. The subcutaneous adipose tissue in diabetic obese mice exhibited low \( pO_2 \), as reported previously, presumably as a result of increased adipocyte size and lower capillary density; in other words, because of imbalance of adipogenesis and angiogenesis. Decreased capillary density in obese adipose tissue might underlie low \( pO_2 \) and chronic adipose inflammation, which increased the risk of cardiovascular disease. The present results suggest that AIC has therapeutic potential as a proangiogenic tool against chronic ischemia and inflammation in obese adipose tissue and may also contribute to reducing the risk of metabolic syndrome. A recent study suggested that aggregate formation and delivery may be another way to induce the therapeutic potentials of ASCs in diabetic mice, although it is cell therapy.

This study focused on adipose tissue as a target tissue. To our knowledge, adipose tissue generally shows considerably high tissue \( pO_2 \) compared with other tissues or organs and supports the vascularity of overlying skin. Intact adipose tissue exhibited \( pO_2 \) >50 mmHg, whereas other tissues were reported to exhibit \( pO_2 \) <30 mmHg; eg, the \( pO_2 \) values of brain, spleen, and thymus were reported to range from 5 to 20 mmHg. Tissue \( pO_2 \) is affected by arterial \( pO_2 \), blood flow, and the cellular oxygen consumption rate. It is known that adipose tissue is more vulnerable to ischemia than skin, and necrosis of subcutaneous adipose tissue, known as deep tissue injury, can seriously affect the vascularity of overlying skin. Therefore, it is practical to target adipose tissue for angiogenesis in clinical settings. For example, AIC could be used to treat intractable skin ulcers with ischemia or hypoxia, such as diabetic or de-cubitus ulcers, or to improve vascularity in ischemic limbs with artery occlusive diseases.

Angiogenic therapy with AIC has several therapeutic advantages. First, AIC treatment is a minimally invasive therapy without any particular donor site morbidity and the need for cell preparation and manipulation, which are costly and raise risk issues. Second, a single administration of AIC is likely to be sufficiently effective, as shown in this study; AIC was the initiator, but presumably not a main player, in the remodeling process. Third, AIC can be prepared at a low cost. Because EGF, PDGF, and TGF-\( \beta \) are released from activated platelets and are well contained in the platelet-rich plasma of patients, a solution similar to AIC can be prepared by supplementing platelet-rich plasma with bFGF, although the therapeutic effects of this substitute remain to be determined. Recombinant bFGF protein is an approved therapeutic product (Fiblast Spray; Kaken Pharmaceutical Co. Ltd., Tokyo, Japan) in Japan; it is topically used to accelerate wound healing in chronic skin ulcers.

There are also some unresolved issues that suggest limitations of AIC therapy. AIC administration is not expected to show remarkable effects in tissues with acute wounds because the release of AIC components is a natural response to injury and hemorrhage and the beneficial effects unlikely increase with excessive administration of AIC, as shown by our results with 10x AIC. In addition, AIC would not have large effects in ASC-deficient (or resident stem/progenitor cell–deficient) sites, such as irradiated tissues, because many AIC effects are likely mediated by the activation of ASCs. In those cases, the administration of AIC-treated ASCs or the combined administration of ASCs and AIC may be required to achieve sufficient therapeutic effects.

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Supplemental Figure S1. Flow cytometric analysis of primary isolated hASCs (n = 3) and HUVECs (n = 3). In fluorescence-activated cell sorting plots, solid gray histograms represent cells stained with the fluorescent-conjugated antibody of interest, and solid red lines represent cells stained with fluorescent-conjugated isotype control antibody. None of the hASCs expressed CD31, vWF, Flk-1, or CD117, which were expressed by HUVECs, whereas all of the hASCs expressed CD90, which was not expressed by HUVECs, suggesting the absence of contamination of vascular ECs in the ASC population.
Supplemental Figure S2. Inverted microscopic images of hASCs and HUVECs cultured with or without AIC for 5 days. Scale bars = 200 μm.
Supplemental Figure S3. The effects of each growth factor included in AIC on proliferation and network formation of hASCs.  

**A:** The proliferative effect of each single factor on hASCs was examined using BrdU incorporation assays \((n = 3)\). \(^*P < 0.05\).  

**B:** Effect of each single factor on network formation by hASCs \((n = 3)\). \(^*P < 0.05\). Error bars represent SEM.
Supplemental Figure S4. Dose-dependent effects of AIC on migration and network formation of hASCs and HUVECs. **A**: Effects of the concentrated AIC medium (10× AIC) and diluted AIC media (0.1×, 0.01×, and 0.001× AIC) on migration of hASCs (n = 4) and HUVECs (n = 4). *0×* indicates control medium (no AIC). *P < 0.05. **B**: Effects of the concentrated AIC medium (10× AIC) and diluted AIC media (0.1×, 0.01×, and 0.001× AIC) on network formation by hASCs (n = 3) and HUVECs (n = 3). *P < 0.05. NS, not significant. Error bars represent SEM.
Supplemental Figure S5. Immunohistochemical analysis of networks formed by hASCs and HUVECs. Most HUVECs on Matrigel were positive for CD31 (green) regardless of AIC pretreatment. A small number of AIC-treated ASCs were positive for CD31 (green), whereas none of the control hASCs were positive for CD31. All nuclei were stained with Hoechst 33342 (blue). Scale bars = 50 μm.
Supplemental Figure S6. The Matrigel co-culture system of AIC-treated hASCs and HUVECs. A: AIC-treated hASCs were labeled with DiO (green) and seeded with Dil-labeled HUVECs (red). Scale bars = 200 μm. B: AIC-treated and DiO-labeled hASCs (green) and unlabeled HUVECs. The networks were immunostained for vWF (red). Scale bar = 100 μm.
Supplemental Figure S7. AIC effects on nonischemic adipose tissue. A: \( pO_2 \) of the inguinal adipose pad \((n = 10)\). To investigate the in vivo effects of AIC on nonischemic subcutaneous adipose tissue, AIC or vehicle was injected into the inguinal subcutaneous adipose pad of mice. At 1 week, the AIC-treated adipose tissue showed a higher \( pO_2 \) value than did the vehicle-treated control, but there was no significant difference between the two groups at 2 or 4 weeks. *\( P < 0.05 \). B: Whole-mount histologic analysis of the adipose tissue. Harvested living tissue samples were stained with BODIPY (adipocytes; green), lectin (ECs; red), and Hoechst 33342 (nuclei; blue). An increased number of infiltrating nucleated cells and lectin-positive cells were observed in the interstitial spaces between adipocytes at 1 week in both groups, and they seemed to decrease in number earlier in the AIC-treated group than in controls. Scale bars = 50 \( \mu m \). C: Quantification of capillary density \((n = 10)\). Paraffin-embedded tissue was stained with lectin (ECs; red) and Hoechst 33342 (nuclei; blue). A significantly increased number of capillaries compared with the control were observed in AIC-treated adipose tissue at 1, 2, and 4 weeks. Scale bars = 100 \( \mu m \). *\( P < 0.05 \); **\( P < 0.01 \). Error bars represent SEM.
Supplemental Figure S8. Microscopic images of ischemic and nonischemic murine adipose tissues after administration of AIC or vehicle. Each section was stained with CD31 (red) and Hoechst 33342 (blue). A: Nonischemic adipose tissue. B: Surgically induced ischemic adipose tissue. C: Chronically ischemic adipose tissue of diabetic mice. In all cases, AIC treatment increased the number of CD31-positive vascular structures. Inflammatory cells infiltrating interstitial spaces and around dead adipocytes were also stained with CD31, especially in the surgically induced ischemia model. Scale bars = 100 μm.