Angiopoietin-1-expressing adipose stem cells genetically modified with baculovirus nanocomplex: investigation in rat heart with acute infarction

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Abstract: The objective of this study was to develop angiopoietin-1 (Ang1)-expressing genetically modified human adipose tissue derived stem cells (hASCs) for myocardial therapy. For this, an efficient gene delivery system using recombinant baculovirus complexed with cell penetrating transactivating transcriptional activator TAT peptide/deoxyribonucleic acid nanoparticles (Bac-NP), through ionic interactions, was used. It was hypothesized that the hybrid Bac-NP_Ang1 system can efficiently transduce hASCs and induces favorable therapeutic effects when transplanted in vivo. To evaluate this hypothesis, a rat model with acute myocardial infarction and intramyocardially transplanted Ang1-expressing hASCs (hASC-Ang1), genetically modified by Bac-NP_Ang1, was used. Ang1 is a crucial pro-angiogenic factor for vascular maturation and neovasculogenesis. The released hAng1 from hASC-Ang1 demonstrated profound mitotic and anti-apoptotic activities on endothelial cells and cardiomyocytes. The transplanted hASC-Ang1 group showed higher cell retention compared to hASC and control groups. A significant increase in capillary density and reduction in infarct sizes were noted in the infarcted hearts with hASC-Ang1 treatment compared to hASC or the untreated group. Furthermore, the hASC-Ang1 group showed significantly higher cardiac performance in echocardiography (ejection fraction 46.28% ± 6.3%, P < 0.001 versus control, n = 8) than the hASC group (36.35% ± 5.7%, P < 0.01, n = 8), 28 days post-infarction. The study identified Bac-NP complex as an advanced gene delivery vehicle for stem cells and demonstrated its potential to treat ischemic heart disease with high therapeutic index for combined stem cell-gene therapy strategy.

Keywords: combined stem cell-gene therapy, baculovirus, nanoparticle, myocardial therapy, angiogenesis, tissue engineering

Introduction
A promising therapeutic approach currently under intensive clinical trials is mesenchymal stem cell (MSC) therapy for congestive heart diseases, which depends on several corroborated mechanisms, such as myocyte formation and neovascularization to improve cardiac function and attenuate ventricular remodeling.1,2 Despite promising initial results, such clinical application remains limited due to logistic, economic, and timing issues when harvesting autologous stem cells from elderly patients. Moreover, it has been reported that MSCs obtained from elderly donors and patients with diabetes or ischemic heart disease have a significantly reduced capacity for neovascularization, proliferation, and differentiation potential.3
Delivering proangiogenic proteins, such as vascular endothelial growth factor (VEGF) and angiopoietin-1 (Ang1), using MSC-based gene therapy approaches are currently being employed in recent studies as an alternative strategy to promote myocardial angiogenesis and regeneration.\(^{4-6}\) However, the mammalian gene delivery vectors widely used in such studies have various drawbacks and severe safety concerns.\(^{7-9}\) These vectors are prone to integration into coding regions of transcriptionally active genes, raising concerns about gene silencing and insertional mutagenesis. On the other hand, nonviral gene delivery systems are much safer for potential clinical applications, but are currently limited by low transfection efficiency and highly transient nature of expression with suboptimal level of transgene expression.\(^{12}\) The potential of this new system for direct gene therapy with minimal cytotoxicity, including MSCs as shown in earlier studies, and reports confirm the safety of transplanting Bac-engineered MSCs into immunocompetent animals for cell-based gene therapy.\(^{9,11}\)

The present study aimed to develop a hybrid nanodelivery system to genetically modify MSCs efficiently, utilizing the complementary strengths of Bac, such as relatively high transduction efficiency and easy scale-up, and transactivating transcriptional activator (TAT)/deoxyribonucleic acid (DNA) nanoparticles (NPs), such as low immunogenicity. TAT peptide sequence, obtained from the protein transduction domain of human immunodeficiency virus-1 responsible for nuclear import of human immunodeficiency virus genome, was modified here by incorporating histidine and cysteine residues for enhanced DNA transport, efficient cell penetration, cell vesicle escape, and transgene expression.\(^{12}\) The potential of this new system for direct gene therapy with cardiomyocytes has recently been demonstrated.\(^{13}\) Here, it was hypothesized that MSC transduction efficiency can also be significantly enhanced by this new system where the negatively charged Bac is coupled with positively charged endosomolytic histidine rich TAT peptide/DNA NPs, both carrying the transgene. For this, a unique pool of MSCs located within the adipose tissue was used, called adipose tissue-derived stem cells (ASCs), mainly because of their practical availability and pro-angiogenic, immunomodulatory, and other unique properties.\(^{14,15}\) Based on the positive efficacy data in disease-relevant animal models, ASCs have recently entered into their first clinical trial.\(^{16}\) In a recent intriguing study, Metzele et al reported that human ASCs (hASCs) can fuse to newborn rat heart cells to form new cardiomyocytes with several nuclei, which can beat when maintained in a culture environment.\(^{17}\) ASCs possess a natural ability to secrete VEGF, which overexpresses under hypoxic conditions in ischemic tissues.\(^{18}\)

It has been reported that VEGF in cooperation with another angiogenic growth factor, Ang1, promotes significant neovascularization, and their combined action leads to a formation of mature and functional vasculature.\(^{19,21}\) Thus, it was hypothesized that the secretion of Ang1 along with naturally releasing VEGF from the genetically modified ASCs, together with their inherent transdifferentiation abilities to cardiomyocytes, can induce a superior synergistic therapeutic effect for myocardial regeneration therapy. Additionally, acute myocardial infarction also induces a high circulating endogenous serum VEGF state.\(^{22}\) With these rationales, Ang1 complementary DNA-carrying recombinant Bac and TAT NPs were generated, a self-assembled binary nanocomplex was prepared by hybridizing the two, and its efficiency to express functionally active Ang1 was determined using optimized transduction protocol with hASCs. The in vivo efficacy of the formulated nanobiohybrid (Bac-NP) system for combined stem cell-gene therapy applications was also evaluated using human Ang1 (hAng1)-expressing genetically modified hASCs for myocardial therapy using an immunocompetent infarcted rat heart model. A schematic presentation of the entire procedure is demonstrated in Figure 1.

**Methods and materials**

**Cell culture and preparation of Bac gene delivery complex**

Human ASCs (n = 1 male donor) were obtained from Invitrogen Life Technologies (Carlsbad, CA) and cultured in Gibco® Dulbecco’s Modified Eagle Medium (Invitrogen) supplemented with 10% fetal bovine serum (FBS, Gibco, Grand Island, NY). The cells were routinely maintained as stationary cultures in 75 cm\(^2\) tissue culture flasks (Corning, MA) and incubated at 37°C in a controlled environment with an air atmosphere of 5% carbon dioxide. Human umbilical vein endothelial cells (HUVECs) (ScienCell Research Laboratories, Carlsbad, CA) were cultured and expanded on tissue culture flasks according to the supplier’s instructions. They were cultured in endothelial cell medium (ScienCell) supplemented with 5% FBS and placed in an incubator containing 5% carbon dioxide at 37°C. H9c2 myogenic cell line, derived from embryonic rat heart...
ventricles, was obtained from American Type Culture Collection (CRL-1446; Manassas, VA) and cultured in Dulbecco’s Modified Eagle Medium supplemented with 10% FBS. The cells were routinely maintained as stationary cultures in flasks, incubated at 37°C in a controlled environment with 5% carbon dioxide. SF9 insect cells (Invitrogen) were maintained at 27°C in SF-900™ III serum free medium (Invitrogen). The cells were maintained in exponential growth phase and subcultured twice per week. Larger volumes were prepared in shaker flasks (Erlenmeyer; Corning Life Sciences, Lowell, MA) which were agitated at 120 rpm in an incubator shaker as mentioned in earlier studies.\textsuperscript{11}

Generation of LacZ and Ang1 gene-carrying recombinant Bac and their subsequent hybridization with NPs to form

**Figure 1** Schematic representation of the overall scheme: generation of recombinant baculovirus (Bac\textsubscript{Ang1}), preparation of hybridized baculovirus with TAT/DNA nanoparticles, in vitro human adipose tissue-derived stem cell transduction, and in vivo investigation using direct intramyocardial transplantation of ASC-Ang1 using a rat model with acute myocardial infarction. Transmission electronic microscopic pictures of baculovirus, nanoparticle, and baculovirus-nanoparticle complex shown in subsets confirm the formation of the nanocomplex (in white arrows).  

**Abbreviations:** ASC-Ang1, angiopoietin-1-expressing adipose tissue-derived stem cells; Bac\textsubscript{Ang1}, angiopoietin-1-carrying baculovirus; Bac-NP\textsubscript{Ang1}, angiopoietin-1-carrying baculovirus-nanoparticle complex; DNA, deoxyribonucleic acid; ELISA, enzyme-linked immunosorbent assay; NP\textsubscript{Ang1}, angiopoietin-1-carrying nanoparticles; TAT, transactivating transcriptional activator.
Bac-NP<sub>Ang1</sub> and Bac-NP<sub>LacZ</sub> were performed using protocol as mentioned in earlier work. 13

Detection of hAng1 expressed by the transduced hASCs: enzyme-linked immunosorbent assay (ELISA) and immunofluorostaining

In order to evaluate the transduction efficiency of Bac-NP on hASCs and compare it with free Bac and NP gene system, NP<sub>Ang1</sub>, Bac<sub>Ang1</sub>, and Bac-NP<sub>Ang1</sub> at multiplicity of infection (MOI; defined as plaque forming units per cell) of 200 were used. hASCs were seeded in six-well plates at 0.5 × 10<sup>6</sup> cells/well and incubated overnight at 5% carbon dioxide and 37°C. Following this, an appropriate volume of the nanocomplexes suspended in phosphate buffered saline (PBS; Life Technologies, Burlington, ON, Canada) was added to each well and incubated for 4 hours. Following this, the transduction solution was replaced with fresh media and grown in a cell culture incubator. Conditioned media were collected every alternate day for 21 days and stored at −80°C for Ang1 ELISA (R&D Systems, Inc, Minneapolis, MN) analysis using standard procedure provided by the manufacturer. 13 To detect the Ang1 expressed within the transduced cells, in another set of experiments, hASCs transduced with NP<sub>Ang1</sub>, Bac<sub>Ang1</sub>, and Bac-NP<sub>Ang1</sub> or nontreated controls were grown on glass microscope slides for 96 hours. After washing with PBS, the cells were fixed with −20°C methanol for 10 minutes followed by immuno-staining as mentioned elsewhere. 23 Briefly, after blocking for 1 hour with 10% donkey serum (Santa Cruz Biotechnology, Santa Cruz, CA), the cells were incubated overnight at 37°C with 1:50 dilution of goat anti-hAng1 (Santa Cruz Biotechnology) primary antibodies. On the second day, the cells were thoroughly washed with wash buffer. The cells were incubated with donkey anti-goat immunoglobulin G-tetramethyl rhodamine isothiocyanate (Santa Cruz Biotechnology) with 1:200 dilutions for 1 hour. The proportions and intensities of tetramethyl rhodamine isothiocyanate-positive hASCs, as seen under fluorescence microscope (Eclipse TE2000U; Nikon Corporation, Tokyo, Japan), gave a qualitative idea of the relative amount of cellular Ang1 expressed due to transgene delivery by the different delivery systems.

Cell proliferation and viability assay

For the cell proliferation assay, 2 × 10<sup>4</sup> HUVEC cells/well were seeded in triplicate for each sample in 96-well plates. After 8 hours of culturing, the cells were washed twice with PBS and 200 μL of conditioned media from nontransduced hASCs, Bac-NP<sub>LacZ</sub>-transduced hASCs, Bac-NP<sub>Ang1</sub>-transduced hASCs, and Bac-NP<sub>Ang1</sub>-transduced hASCs supplemented with anti-hAng1 antibodies were added to the corresponding set of wells. After 96 hours, absorbance was measured at 490 nm using CellTiter 96<sup>®</sup> AQueous Non-Radioactive Cell Proliferation Assay (Promega, Fitchburg, WI) in a Victor3 Multi Label Plate Counter (Perkin Elmer, Montreal, QC, Canada). 24 In a similar way, cardiomyocyte cell viabilities under oxidative stress in groups treated with different conditioned media were measured as described later in the study, using the same assay. The experiment was performed in triplicate.

Wound healing assay with HUVECs

In order to check the wound healing potential of released Ang1, HUVECs were seeded into 24-well plates and grown to confluency. After 24 hours of serum starvation (1% FBS), lesions were made in the monolayer using a cell scraper. 13 Cells were rinsed with PBS, and then incubated with the hASC cardiomyocytes from different experimental groups (NP<sub>Ang1</sub>, Bac<sub>Ang1</sub>, and Bac-NP<sub>Ang1</sub>) for 24 hours. Cardiomyocytes from untransduced hASCs were used as control. To confirm the beneficial effect of Ang1 in particular, cardiomyocytes from Bac-NP<sub>Ang1</sub> group were preincubated with anti-Ang1 neutralizing antibody (R&D Systems; 1 μg/mL) for 30 minutes before being added to the wells. Cells were fixed with 4% paraformaldehyde (Sigma-Aldrich, St Louis, MO) after 24 hours and the number of cells which had moved across the starting scratched lines were measured for all groups. Three fields were analyzed for each well at 200x magnification under bright field settings of the microscope mentioned earlier.

Apoptosis assay using cardiomyocytes

H9c2 cells were seeded in 96-well microtiter plates at a density of 2 × 10<sup>4</sup> cells/well and cultured overnight. The media was replaced with cardiomyocytes from different treatment groups and oxidative stress was induced by adding 200 μM hydrogen peroxide to the media as described elsewhere. 25 After 6 hours, the apoptotic cells were detected by tracking the loss of mitochondrial membrane potential using MitoCapture<sup>TM</sup> Apoptosis Detection Kit (BioVision, Inc, Mountain View, CA) cell staining cationic dye, according to the manufacturer’s protocol, which fluoresces differently in healthy and apoptotic cells under fluorescence microscope (488-nm and 543-nm excitation filters). The red emission of the dye is due to a potential-dependent aggregation in the mitochondria reflecting normal membrane potential and the green fluorescence detects the monomeric form of MitoCapture, appearing in the cytosol after mitochondrial membrane depolarization. 26 For each
experimental group, total cell number was counted and related to the number of cells that displayed fluorescence. In a similar way, after oxidative stress of 6 hours, the cell viabilities were also checked using CellTiter 96 assay as mentioned above.

**In vivo studies in myocardially infarcted rat model**

Immunocompetent female Lewis rats (200–250 g; Charles River Laboratories, Senneville, QC, Canada) were used. All procedures were in compliance with the Guide for the Care and Use of Laboratory Animals (National Institutes of Health, publication 85-23) and the Guide to the Care and Use of Experimental Animals of the Canadian Council on Animal Care. Female Lewis rats were anesthetized. The rats were intubated with an 18-gauge catheter and mechanically ventilated (Harvard Ventilator 683; Harvard Apparatus, St Laurent, QC, Canada) at 80 breaths/minute. Anesthesia was maintained with 3% isoflurane. A left thoracotomy was performed through the fourth intercostal space to expose the left ventricle. The left coronary artery was ligated 2 mm from its origin with a 7-0 polypropylene suture (Ethicon, Inc, Somerville, NJ) using standard protocol. The ischemic myocardial segment rapidly became identifiable through its pallor. Fifteen minutes after ligation of the artery, intramyocardial injections were given. In the control group with no hASC (n = 8), 300 µL of the culture medium was divided into three equal periinfarct left ventricular intramyocardial injections using a 27 gauge needle. In the hASC group (n = 8), the same procedure was repeated using culture medium containing 3 × 10⁶ hASCs, while the group hASC-Ang1 (n = 8) received the same number of cells transduced with Bac-NPang1 (MOI: 200) suspended in culture medium.

To detect the transgene expression in transplanted cells, three animals underwent the same experimental procedure as described above, followed by injection of three million cells transduced with Bac-NP <sup>LacZ</sup> (MOI: 200). Three days post-transplantation, LacZ expression was detected in histological samples using a standard staining LacZ staining procedure.

**Detection of transplanted male hASCs in female rat hearts: polymerase chain reaction (PCR) analysis for human Y chromosome**

In order to detect the transplanted hASCs from the male donor in heart tissue, samples from groups with no hASC (n = 6), hASC (n = 6), and hASC-Ang1 (n = 6) were taken. After 3 days and 28 days post-injection, three animals from each group were sacrificed and heart tissue samples were snap frozen. PCR analysis was done on the extracted DNA samples to confirm the survival of the implanted gender-mismatched cells in the hearts using standard procedure followed in earlier studies. Briefly, equal amounts of DNA were extracted from peri-infarct portions of the ventricular heart in each group and PCR analysis was done to confirm the survival of the implanted gender-mismatched cells in the hearts 3 days and 28 days post-operation. Genomic DNA was purified using DNeasy (QIAGEN, Valencia, CA) according to the manufacturer’s instructions, and the presence of living male cells in female hearts was confirmed by targeting a specific microsatellite sequence within the Y chromosome (DYS390). The primer pairs used were forward primer 5’TATATTTTACACATTITTG GCC3’ and reverse primer 5’TGACAGTA AAAATGAACACATTGC3’ with product length of 250 base pairs.

**Scar area analysis: myocardial infarct size**

Twenty-eight days post-operation, rats were deeply anesthetized and sacrificed by rapid excision of the heart. The excised hearts were immediately soaked in cold saline to remove excess blood from the ventricles and fixed in neutral-buffered 4% formalin. Paraffin embedded samples were sectioned at 5 µm, and Masson’s trichrome staining (Diagnostic BioSystems, Pleasanton, CA) was performed to delineate scar tissue (blue color) from the total area of myocardium. Masson’s trichrome-stained section images were analyzed by ImageJ 1.41 software (National Institutes of Health, Bethesda, MA). Infarct area, epicardial and endocardial length of infarction, and ventricular and septal wall thickness were calculated and expressed as a percentage.

**Immunohistochemistry for detecting neovascularization**

Neovascularization was evaluated by analyzing capillary and arteriole density in the periinfarct area. For this, immunohistochemical staining was performed with anti-PECAM (Santa Cruz Biotechnology) for identification of endothelial cells and antismooth muscle α-actin (Santa Cruz Biotechnology) for tracing the smooth muscle cells, as described elsewhere. Briefly, for measurement of capillary density, five fields in the periinfarct area were imaged with 200× magnification and average numbers of capillaries <10 µm in diameter were counted. Capillary density was quantified as (mean total PECAM-positive microvessels/mm² using three tissue sections spanning the periinfarct tissue region of each animal. Similarly, arteriole densities were quantified as (mean total smooth muscle α-actin-positive microvessels/mm²).
Transthoracic echocardiography

Transthoracic echocardiography was performed on all surviving animals in rat groups treated with hASC (n = 8), hASC-Ang1 (n = 8), and untreated control (n = 8) on day 3, day 14, and day 28 post-infarction. Echocardiograms were obtained with a commercially available system (Titan®; SonoSite, Inc, Bothell, WA) equipped with a 15-MHz transducer. After sedating the animals with 2% isoflurane, echocardiography was performed according to the American Society of Echocardiology leading-edge method.27 Briefly, parasternal long- and short-axis views were obtained with both M-mode and two-dimensional images. Left ventricular end-diastolic and end-systolic diameters were measured with M-mode tracings between the anterior and posterior walls from the short-axis view just below the level of the papillary muscles of the mitral valve. Two images on average were obtained in each view and averaged over three consecutive cardiac cycles. Left ventricular end-diastolic volume was calculated as $7.0 \times \frac{\text{left ventricular end-diastolic diameter}^3}{(2.4 + \text{left ventricular end-diastolic diameter})}$, and left ventricular end-systolic volume as $7.0 \times \frac{\text{left ventricular end-systolic diameter}^3}{(2.4 + \text{left ventricular end-systolic diameter})}$. Ejection fraction was estimated as (left ventricular end-diastolic volume – left ventricular end-systolic volume)/left ventricular end-diastolic volume.

Statistical analysis

Quantitative variables are presented as mean ± standard deviation from independent experiments as described in the figure legends. Statistics were performed using two-way analysis of variance by Bonferroni’s multiple comparison post hoc test. All statistical analyses were performed with GraphPad Prism (v 5.0; GraphPad Software, Inc, La Jolla, CA). $P < 0.05$ was considered significant.

Results

Bac-Ang1-transduced hASCs can efficiently release hAng1

Released Ang1 from the transduced hASCs was analyzed at protein level by hAng1 ELISA. As shown in Figure 2A–C, hAng1 expression was detected from day one post-transduction until day 21 with regular intervals for all groups (NP_Ang1, Bac_Ang1, and Bac-NP_Ang1). The highest expression was seen on day four with the gene delivery systems. However, highest expression was achieved with Bac-NP_Ang1 (18.5 ± 1.2 ng/10^6 cells), which stayed consistently higher than other groups all throughout the experimental period. Moreover, cellular Ang1 expression, demonstrated by tetramethyl rhodamine isothiocyanate-stained cells 96 hours post-transduction, was shown to be highest in the Bac-NP_Ang1-treated group as compared to NP_Ang1 and Bac_Ang1 (Figure 2D–G). ELISA and immunostaining analysis confirmed that Bac-NP_Ang1 was a superior gene delivery vehicle than free Bac_Ang1 and NP_Ang1 alone, and was able to express higher amounts of hAng1 transgene at both extracellular and intracellular level.

Biologic activity of the secreted hAng1 on vascular endothelial cells

To confirm the functionality of the expressed hAng1, HUVEC proliferation assay was performed using the conditioned media from Bac-NP_Ang1-transduced hASCs and compared with that of control Bac-NP_LacZ-transduced hASCs and untransduced hASCs. As reported in Figure 3A, the samples with high Ang1 concentrations, ie, from cardiomyocytes of Bac-NP_Ang1-transduced hASCs, significantly increased the proliferation rate of HUVECs ($2.67 \times 10^4$ cells compared to $1.85 \times 10^4$ cells in Bac-NP_LacZ cardiomyocytes, $1.89 \times 10^4$ in control nontransduced cardiomyocytes, and $1.59 \times 10^4$ cells in cardiomyocytes with antibody – $2 \times 10^4$ cells were taken as the initial density).

Furthermore, the ability of cardiomyocytes (containing secreted hAng1) to increase HUVEC wound healing in a monolayer was tested. As depicted in Figure 3B and C, stimulation of wounded HUVEC monolayer with cardiomyocytes from Bac-NP_Ang1 (33.3% ± 2.0%) induced a significant reduction of wound compared with the unstimulated control (16.8% ± 1.4%) and cardiomyocytes from Bac-NP_LacZ (16.1% ± 4.2%). Thus, consistent with mitotic assay results, Bac-NP_Ang1 showed highest healing potential, indicating that efficient overexpression of Ang1 is one of the determining factors for inducing substantial biological effects. Preincubation of cardiomyocytes with neutralizing anti-Ang1 antibodies completely inhibited Bac-NP_Ang1 cardiomyocyte-induced wound healing, clearly suggesting that chemotactic signals from hAng1 are essential for this effect.

Biologic activity of the secreted hAng1 on cardiomyocytes

In order to assess whether the secreted hAng1 have a protective effect on cardiomyocytes under oxidative stress, the treated H9c2 cells from different groups underwent cell viability studies after 6 hours oxidative stress using hydrogen peroxide. The data in Figure 4A show that cardiomyocytes from Bac-NP_Ang1 were able to significantly reduce cell death
compared to Bac-NP<sub>LacZ</sub> and Bac-NP<sub>Ang1</sub>-antibody (1.82 ± 0.02 versus 1.04 ± 0.04 versus 0.93 ± 0.08), taking a normalized value of the cell number in the untreated control group as 1.0.

As positive control, cells with no oxidative stress were taken which showed a viability amount of 1.82 ± 0.02.

Furthermore, the antiapoptotic effect of the released hAng1 on cardiomyocytes under oxidative stress was examined and quantified by MitoCapture cell staining (Figure 4B and C). Cultivation of H9c2 cardiomyocytes under hydrogen peroxide-induced oxidative stress for 6 hours provoked a strong apoptotic response with 60.7% ± 3.7% of apoptotic cells in the untransduced control cardiomyocyte group. Addition of cardiomyocytes from Bac-NP<sub>Ang1</sub>, hASCs strongly suppressed cardiomyocyte apoptosis (24.1% ± 2.9%) in comparison to other groups, which did not show any significant effect.

**hASCs can survive and efficiently express their transgene in vivo**

To analyze whether the xenotransplanted genetically modified hASCs were able to survive and express the transgene in the rat heart, hearts (n = 3) were myocardially infarcted.
and injected with hASCs transduced with Bac-NP<sub>LacZ</sub> in the left ventricular region, as described earlier. Three days post-transplantation, the rat hearts were harvested and ventricular tissue samples were stained with X-gal in order to trace the transplanted LacZ-expressing hASCs in the peri-infarct region of the heart. Figure 5A confirms that the transplanted cells were able to survive and express the transgene in the heart.

Higher retention of implanted hASC-Ang1 in rat myocardium

After confirming transplant survival and transgene expression, the effect of Ang1-expressing hASCs compared to normal hASCs on cardiac cellular transplant retention was further explored. Three groups (no hASC [n = 6], hASC [n = 6], and hASC-Ang1 [n = 6]) were taken as mentioned earlier.
Three days and 28 days post-injection, three animals from each group were sacrificed for PCR analysis.

DNA was extracted from the injected sites of the heart tissues to detect the presence of Y chromosome of transplanted male hASC using standard PCR. The gel electrophoresis of the PCR products shows a clear distinction in band intensities of hASC-Ang1 and normal hASCs (Figure 5B). The hASC-Ang1 group showed around 1.72 times higher retention of viable transplanted cells in the heart in comparison to normal hASCs (0.55 times versus...
0.32 times, taking initial 3-day band intensity as 1.0; \( P < 0.05 \) as quantified by relative band intensities using ImageJ software. This semiquantitative analysis indicates that a much higher number of cells were able to survive the transplantation when they were genetically modified to express Ang1 protein.

**Bac-Ang1 hASCs can significantly attenuate scar area formation in infarcted heart**

Macroscopic views of Masson’s trichrome stained heart sections are shown in Figure 6A. In both the control and cell treated groups, positively stained fibrous infarct areas were clearly observed in the heart 28 days after myocardial infarction.
Thin infarcts and left ventricular wall with dilated left ventricular cavity were observed in the control hearts. On the other hand, the hearts in the cell treated groups had significantly lesser infarct areas (Figure 6B: 13.0% ± 3.1% for hASC-Ang1, 31.5% ± 5.0% for hASCs, and 38.0% ± 3.3% for control; P < 0.01) and higher left ventricular wall thickness (Figure 6C: 2.53 ± 0.19 mm for hASC-Ang1, 1.44 ± 0.24 mm for hASCs, and 0.91 ± 0.07 mm for control; P < 0.001) than the control heart. A total of 24 rats were analyzed for the determination of infarct size (no hASC control [n = 8], hASCs [n = 8], and hASC-Ang1 [n = 8]). Infarct size and wall thickness in the left ventricles at the section of the middle point between ligation and apex were measured as described elsewhere.22 There was also a significant improvement in the group treated with hASC-Ang1 compared to hASCs group (P < 0.001) with respect to the percentage left ventricle infarct area and left ventricle infarct wall thickness.

Ang1-expressing hASCs induce angiogenesis and arteriogenesis

Reporting the significant attenuation of scar area with Ang1-expressing hASCs in the last section, attempts were made to understand whether it was only the paracrine effect of the hASCs or a combinatorial effect of hASC-released Ang1 and paracrine factors which was responsible for the improvement. For this, the neovasculature formation in the periinfarct area was assessed by detecting angiogenesis and arteriogenesis densities (Figure 7). Indeed, a significant improvement was seen in angiogenesis in the hASC and hASC-Ang1 groups compared to the control (201 ± 25/mm² for hASC-Ang1, 174 ± 14/mm² for hASCs, and 141.5 ± 6/mm² for control; P < 0.01). Moreover, the hASC-Ang1 group showed significantly higher capillary density compared to hASC. Similar results were obtained with arteriole density in the hASC-Ang1 group (12.25 ± 2.4/mm² for hASC-Ang1, 9.5 ± 1.8/mm² for hASCs, and 6.5 ± 1.32/mm² for control; P < 0.05), with the hASC-Ang1 group showing significantly higher arteriole density compared to hASC.

Significantly improved cardiac function is detected in hASC-Ang1-treated infarcted heart

To investigate if the reduction of scar area after stem cell therapy results in improved heart function, the ejection fraction was measured using echocardiography. The hASC-Ang1 group showed a significantly higher ejection fraction compared to the control (60.0 ± 3.0% for hASC-Ang1, 55.0 ± 3.5% for hASCs, and 50.0 ± 2.0% for control; P < 0.001). These results suggest that the improvement in cardiac function is due to the regenerative effect of hASC-Ang1 on myocardial tissue.
fraction in a rat model of myocardial infarction at different time periods (days three, 14, and 28) was monitored. Ejection fraction of all groups was around 30% on day three post-infarction, indicating successful acute myocardial infarction in all groups. As presented in Figure 8, there were significant improvements in ejection fraction in groups treated with hASC-Ang1 and hASCs compared to the control group on day 14 and day 28, although there were no significant differences between the groups on day three post-infarction (P > 0.05). In addition, the hASC-Ang1 treated group showed a significantly greater increase in ejection fraction as compared to hASC-treated groups on day 14 (44.78% ± 6.14% versus 37.1% ± 4.9%; P < 0.01) and day 28 (46.28% ± 6.3% versus 36.35% ± 5.77%; P < 0.01). Thus, analysis using echocardiographic ejection fraction data suggests that hASC-Ang1 can be a better alternative to hASCs to improve cardiac function after acute myocardial damage.

**Discussion**

Cardiovascular and ischemic heart disease is a major health concern in both developing and industrialized countries. Although different gene therapy approaches are being employed to treat the disease, clinical trials showed limited results suggesting the need for a better gene therapy strategy. It has been demonstrated in small animals, that ASCs induce wound healing effects in damaged ischemic

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**Figure 7** Angiogenesis and arteriogenesis in the peri-infarct area. Immunohistological staining of CD31 to detect endothelial cells in (A) no hASC, (B) hASC, and (C) hASC-Ang1 groups. Immunohistological staining of smooth muscle α-actin for smooth muscle cells in (D) no hASC, (E) hASC, and (F) hASC-Ang1 groups. Quantification of (G) capillary and (H) arteriole density. 

**Notes:** Data are expressed as mean ± standard deviation. One-way analysis of variance: (G) F = 24.86, P < 0.0001 and (H) F = 18.48, P < 0.0001 (treatment groups). Statistically significant differences between groups compared to control no hASC are indicated as ***P < 0.001; **P < 0.01; *P < 0.05. Significant difference between hASC and hASC-Ang1 is indicated by †.

**Abbreviations:** hASC, human adipose tissue-derived cell; hASC-Ang1, angiopoietin-1-expressing human adipose tissue-derived cell; ctrl, control.
tissues mainly through proangiogenic paracrine mechanism followed by neovascularization, local recruitment of progenitor cells, and exhibition of cardiomyocyte-like properties.\(^{17,30}\) Moreover, ASCs secrete VEGF which has a favorable impact on myocardial neovascularization, tissue perfusion, and contractile performance. In addition, a larger number of ASCs can be easily harvested using noninvasive standard fat extraction techniques, which is of immense importance under clinical settings because of their ready availability.\(^{14,18}\) Most importantly, as allogeneic hASCs can be easily obtained from young healthy donors, they could be of great value in elderly patients with advanced diseases, where their own ASCs could be dysfunctional.

The initial step involved the development of a recombinant Bac carrying Ang1 and assessing its potential for direct gene therapy for efficient myocardial neoangiogenesis as reported in previous work.\(^{13}\) As the next step, the present study further explored its potential for stem cell-based gene therapy, apart from direct gene therapy. The present study reports for the first time, the angiogenic potential of this newly formulated hybrid nanodelivery system composed of vascular gene-carrying recombinant Bac armed with noncovalently conjugated TAT/DNA NPs for stem cell-based gene therapy. As stem cells are known to have a natural potential for cardiovascular therapies, complementing them with rapid Ang1 overexpression can further improve their efficacy. The present work illustrates this hypothesis in a rat model with acute myocardial infarction. Moreover, this is the first study to report the potential of recombinant Bac to genetically modify ASCs. In addition, the work also demonstrates the potential of genetically modified ASCs for successful xenotransplantation applications.

Using a hybrid model of TAT/DNA\(^{\text{Ang1}}\) NPs coupled to Bac\(^{\text{Ang1}}\) for ASC transduction under optimized conditions (Figure S1), significantly higher gene transfer efficiency and gene carrying capacity was achieved than free Bac\(^{\text{Ang1}}\) or NP\(^{\text{Ang1}}\).
alone (Figure 2 and Figure S2). Figure S2 demonstrates that Bac-NP, with Bac MOI of 200 and N/P ratio (defined as the molar ratio of amino groups/phosphate groups) of 3 of TAT/DNA NP, was optimal so as to achieve the highest transduction efficiency with no cytotoxic effects (Figure S3). Data also confirm that gene transfer efficiency can be significantly enhanced by this hybrid nanoformulation of Bac and NP, as compared to their individual efficiencies. In vitro analysis showed that Bac-NP-transduced cells were able to express their transgene for at least 3 weeks (Figure 2C), which is comparable to other experimental results. This temporal expression of the delivery system is beneficial in many cases, particularly for angiogenesis where the expression of the therapeutic protein ceases once its job is done. Further studies need to be done to comprehend the exact mechanism of such a gene delivery system. Rapid attachment of cationic NPs to a negatively charged cell surface followed by efficient intracellular gene entry to the nucleus by Bac may be a probable way. Optimizing the pH condition and transduction time may further enhance the gene delivery efficiency of the Bac-NP system.

Bac surface modification by chemical coupling and electrostatic interactions is recently being exploited in different ways for enhanced gene delivery. It has been demonstrated that galactosylated polyethylenimine/DNA hybridized Bac improves gene delivery to hepatoblastoma and lung carcinoma cells. Coating of Bac using positively charged polymer polyethylenimine is also shown to significantly enhance their in vivo gene therapy potential compared to uncoated viruses. Recently Chen et al demonstrated that genetically modified Bac generated by fusing the protein transduction domain of human immunodeficiency virus TAT protein with VP39, the major capsid protein, resulted in improved transduction of various mammalian cells. Similarly, the present study demonstrates that surface modifying Bac with TAT/DNA NPs can also improve gene transfer efficiency. But, to achieve significant therapeutic effects, specifically with stem cells, maintenance of their natural functionalities is of immense importance. The present study confirmed this by successfully differentiating the genetically modified hASCs to adipogenic and osteogenic lineages (Figure S4). In addition, data revealed that Bac-NP-mediated hAng1 expression from hASCs were biologically active as indicated by their enhanced mitotic and chemotactic activities on HUVEC (Figure 3). The study also confirmed that the released Ang1 possesses antiapoptotic and protective effects on cardiomyocytes (Figure 4). This supports the findings reported by Wang et al who reported that Ang1 can protect cardiomyocytes from oxidative stress-induced apoptosis.

In addition to efficient transgene expression, retention of viability of transplanted cells at the infarct site is an important parameter that determines the myocardial therapeutic index of the study. This is because higher retention of viable cells directly correlates with enhanced secretion of paracrine factors and better scope for cellular cardiomyoplasty. Previous works have aimed to address this problem using polymeric microcapsules to reduce the biologic and mechanical loss of implanted cells in the continuously beating heart. In an interesting study, Liu et al demonstrated that Ang1 can protect MSC against serum deprivation and hypoxia-induced apoptosis. Detection of a significantly higher amount of human Y chromosome-positive hASCs in recipient female rat hearts treated with hASC-Ang1 compared to hASC, 28 days post-implantation, confirmed these findings. Most importantly, hASC-Ang1 treatment resulted in enhancement of vascular density (Figure 7), thereby significantly improving cardiac function (Figure 8) compared to hASC and control groups; although cardiomyogenic transdifferentiation of grafted cells was not detected in any of the groups. Moreover, histological findings confirmed the reduction of fibrosis in the infarct area (Figure 6), which is an important indicator for improved heart function since late reperfusion of infarcted vascular beds attenuates left ventricular remodeling including infarct expansion. Measurement of blood flow using cardiac magnetic resonance imaging-based techniques can give a better real time picture on the progress of such myocardial regenerative process post cell transplantation. Although preclinical findings exemplify the beneficial effects of such combined adipose stem cell–gene therapy treatment, further in vivo studies are needed to elucidate the precise mechanism and to investigate its prolonged effects on cardiac function; these will help determine whether one-time cell delivery is a viable option or whether multiple cell transplantations are required to achieve a long term desirable outcome.

**Conclusion**

From a clinical standpoint, the study promises major advancements where ASCs could be easily harvested and mass produced well in advance, genetically modified to express biologically active therapeutic proteins, and stored for immediate off-the-shelf use on any patient without delay after an acute myocardial infarction.

In summary, it was demonstrated that a hybrid nanodelivery complex of Bac and NPs can take advantage of the unique features associated with the two individual vector systems and exhibit enhanced gene transfer efficiency, supported by in vitro and in vivo experimental demonstrations. It was illustrated that
genetically-engineered hASCs, using a hybrid nanocomplex, enhanced the transplant retention in a myocardially infarcted rat model and induced a significant favorable impact on tissue perfusion and contractile performance, which have been corroborated with vasculogenesis. Taken together, the current findings confirm the feasibility of a combined stem cell-based gene therapy for ischemic heart diseases using a novel, biologically safe nanobiobohybride gene delivery system.

Acknowledgments
The authors gratefully acknowledge the assistance received from the Canadian Institutes of Health Research (MOP #64308) to S Prakash, and the Natural Sciences and Engineering Research Council of Canada to S Prakash and D Shum-Tim. A Paul acknowledges the Alexander Graham Bell Post Graduate Scholarship and Michael Smith Foreign Study Award – Doctoral from the Natural Sciences and Engineering Research Council of Canada.

Disclosure
The authors report no conflicts of interest in this work.

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Supplementary information

Characterization of baculovirus (Bac)-nanoparticle (NP) hybrid complex

The particle size and zeta potential of the NP, Bac, and Bac-NP hybrid particles were measured by the technique of electrophoretic laser Doppler anemometry using a Zeta Potential Analyzer (ZetaPlus; Brookhaven Instruments Corporation, Holtsville, NY). ZetaPlus Particle Sizer Software (v 4.11; Brookhaven) was used to determine the size distribution of the particles and Zeta Potential Analyzer software (v 3.57; Brookhaven) was used for zeta potential analysis. Both particle size and zeta potential were measured for three independent preparations and each measurement was obtained after taking the average of the three runs. Transmission electron microscopy (TEM) was used to obtain the size characterization. The NPs were suspended in 1x phosphate buffered saline and analyzed on CM200 FEG-TEM (Royal Philips Electronics, Markham, ON, Canada). Results are presented in Figure S1.

The scanning electron microscope (SEM; S4700 FEG-SEM Hitachi, Oakville, ON, Canada) and atomic force microscope (AFM; Digital Instruments, Palo Alto, CA) photomicrographs in Figure 2A and B confirm the formation of NP by transactivating transcriptional activator (TAT) and DNA complexation. The formed NPs were further studied by TEM and zeta potential analyzer, as demonstrated in Figure S1C–F. The Bac-NP complex was first characterized by measuring the zeta potential of the nanocomplex with laser Doppler electrophoresis (Figure S1C). At its natural pH 6.8, Bac is negatively charged with zeta potential of −6.5 ± 1.4 mV. At physiological pH 7.4, Bac had a zeta potential of −12.8 ± 3.1 mV. On the other hand, prepared TAT/DNA NPs showed high positive charge of 26.5 ± 3.2 mV at N/P ratio of 3. The positively charged TAT/DNA NPs, upon conjugation with the negatively charged Bac, formed positively charged (5.1 ± 2.7 mV) Bac-NP hybrid nanocomplexes.

To reconfirm the successful formation of the Bac-NP complexes, particle sizes of each complex were measured. Free Bac had an average size of 238 ± 10 nm, whereas free TAT/DNA NP showed an average size of around 72 ± 4.6 nm. The Bac-NP hybrid complexes had an average size of 480 ± 18.2 nm. This significant increase in size of Bac-NP particles, compared to that of free Bac and NPs, indicates the efficient production of the nanobiohybrid complexes, generated by strong electrostatic interactions of Bac with the NPs.

In order to look for the morphological evidence for successful conjugation of the budded Bac particles with the NPs, TEM was used. Electron micrographs showed the well-dispersed NPs (Figure S1D). On coming in contact with free Bac (Figure S1E; the rod-shaped particles with a length of 200–250 nm), there was an instant virus-NP complex formation by the negatively charged Bac with the positively charged NP, as indicated in Figure S1F to form the hybridized Bac-NP complex. The images of Bac-NP complexes also confirm the proper retention of the typical rod-shaped morphological appearance and envelope structure of Bac, suggesting that Bac were able to sustain their morphological integrity even after hybridization with NPs.

Optimization of viral dose for Bac-NP mediated human adipose tissue-derived stem cell (hASC) transduction

In order to achieve maximum transduction, the effect of multiplicity of infection (defined as plaque forming units per cell) and their combinatorial effects on the hybrid Bac-NP system were optimized. For this, multiplicity of infection ranging from 100 to 400 and N/P ratio of 3 were used. Initially, hASCs were seeded in six-well plates at 0.5 × 10⁴ cells/well and incubated overnight at 5% carbon dioxide and 37°C. Following this, an appropriate volume of transduction solutions from different experimental groups (LacZ-carrying Bac, LacZ-carrying NP, Bacnull-LacZ-carrying NP, LacZ-carrying Bac, LacZ-carrying NP, LacZ-carrying Bac-NPnull), suspended in phosphate buffered saline, was added to each well according to varied multiplicity of infection, and incubated for 4 hours at 25°C. Bacnull and NPnull represent delivery systems carrying DNA with no gene of interest. The wells were replenished with fresh media and grown 37°C in carbon dioxide incubator. After 24 hours, the cells were stained with X-gal to detect the transduced LacZ-expressing cells. Results are presented in Figure S2.

Bac-NP has no toxic effects on hASCs: cytotoxicity assay

For the cytotoxicity assay, 2 × 10⁴ hASCs/well were seeded in triplicate for each sample in 96-well plates. After culturing overnight, the cells were washed twice with phosphate buffered saline. NP, Bac, and Bac-NP particles suspended in culture media (200 µL) were added to the corresponding set of wells. Nontransduced cells were used as control. After 24 hours, absorbance of each well were measured at 490 nm using CellTiter® 96 AQueous Non-Radioactive Cell Proliferation Assay (Promega, Fitchburg, WI) in a plate reader. The percentage of viable cells in different experimental
Figure S1 Characterization of the Bac-NP hybrid nanocomplex. (A) Scanning electron microscope photograph of transactivating transcriptional activator/deoxyribonucleic acid nanoparticles with subset showing a magnified image. (B) Atomic force microscope photograph of NPs demonstrating their surface topography. (C) Zeta potential of free Bac at pH 6.8 (pH of insect cells media), free Bac at pH 7.4 (pH of mammalian cell culture), free NP (phosphate buffered saline: pH 7.4), and hybrid Bac-NP nanocomplexes at pH 7.4, with N/P ratio of 3. The complexes were prepared from 1 µg deoxyribonucleic acid (complexed with transactivating transcriptional activator) per 10⁹ plaque forming units Bac. Transmission electron microscope images of (D) NP with N/P ratio of 3, (E) Bac, and (F) Bac-NP suspended in phosphate buffered saline.

Notes: Arrows indicate NPs hybridized on Bac surface. Scale bar: 100 nm.

Abbreviations: Bac, baculovirus; Bac-NP, baculovirus-nanoparticle complex; NP, nanoparticle.
groups was quantified. All the experiments were performed in triplicate. Results are presented in Figure S3.

**Bac-NP transduced hASCs retain their differentiation potential**

Bac-NP-transduced hASCs were then seeded in a 24-well plate at a high confluency of $6 \times 10^4$ cells/well. Nontransduced cells were used as control. After 24 hours, the medium was replaced with either adipogenic or osteogenic differentiation medium (Invitrogen Life Technologies, Carlsbad, CA). Osteogenic differentiation was assessed using Alizarin Red S staining (Invitrogen) after a 21-day period of induction towards this lineage. After a 15-day period, adipogenic differentiation was evaluated by LipidTOX™ red neutral lipid staining (Invitrogen). Results are presented in Figure S4. The presence of calcium deposits following osteogenic induction, as well as lipid vacuoles following adipogenic induction, is indicative of the transduced hASCs’ ability for multilineage differentiation.
Figure S4 Adipogenic and osteogenic differentiation of baculovirus-nanoparticle complex-transduced human adipose tissue-derived stem cells. Transduced adipose tissue-derived stem cells were cultured in either adipogenic or osteogenic differentiation media. Adipogenic differentiation was assessed via LipidTOX™ Red neutral lipid staining of lipid vacuoles and 4′,6-diamidino-2-phenylindole staining of nucleus: (i) transduced differentiated and (ii) nontransduced differentiated. Osteogenic differentiation was determined by Alizarin Red staining of calcium deposits: (i) transduced differentiated and (ii) nontransduced differentiated. Notes: Arrows show the cell differentiated area. Results confirm that human adipose tissue-derived stem cells retain their multilineage differential potential even after baculovirus-nanoparticle complex transduction.