ISL1 Protein Transduction Promotes Cardiomyocyte Differentiation from Human Embryonic Stem Cells

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Abstract

Background: Human embryonic stem cells (hESCs) have the potential to provide an unlimited source of cardiomyocytes, which are invaluable resources for drug or toxicology screening, medical research, and cell therapy. Currently a number of obstacles exist such as the insufficient efficiency of differentiation protocols, which should be overcome before hESC-derived cardiomyocytes can be used for clinical applications. Although the differentiation efficiency can be improved by the genetic manipulation of hESCs to over-express cardiac-specific transcription factors, these differentiated cells are not safe enough to be applied in cell therapy. Protein transduction has been demonstrated as an alternative approach for increasing the efficiency of hESC differentiation toward cardiomyocytes.

Methods: We present an efficient protocol for the differentiation of hESCs in suspension by direct introduction of a LIM homeodomain transcription factor, Islet1 (ISL1) recombinant protein into the cells.

Results: We found that the highest beating clusters were derived by continuous treatment of hESCs with 40 μg/ml recombinant ISL1 protein during days 1-8 after the initiation of differentiation. The treatment resulted in up to a 3-fold increase in the number of beating areas. In addition, the number of cells that expressed cardiac specific markers (cTnT, CONNEXIN 43, ACTININ, and GATA4) doubled. This protocol was also reproducible for another hESC line.

Conclusions: This study has presented a new, efficient, and reproducible procedure for cardiomyocytes differentiation. Our results will pave the way for scaled up and controlled differentiation of hESCs to be used for biomedical applications in a bioreactor culture system.

Introduction

Cardiomyocytes derived from human embryonic stem cells (hESCs) potentially offer large numbers of cells for biomedical and industrial applications. Current protocols for differentiation of cardiomyocytes from hESCs are time consuming, have low yield, and lack reproducibility (for review see, ref [1]). However, for the applicability of these cells in biomedicine it is necessary to produce sufficient numbers of functional cardiomyocytes or their progenitors. This requires the development of large-scale expansion of hESCs and their controlled differentiation protocols. In recent years technologies for the suspension expansion of hESCs and application of bioreactors have been introduced [2,3,4,5]. For example, we recently expanded hESCs as carrier-free suspension aggregates for an extended period of time [6].

On the other hand, the differentiation of cardiomyocytes from hESCs has progressed rapidly through a growth factor-mediated approach. Although the efficiency of differentiation protocols has increased over time, a desirable efficiency has not been attained by these methods. It has been shown that the forced expression of instructive transcription factors such as Tbx5 and Nkx2.5 successfully increased the differentiation efficiency toward cardiomyocytes [7,8].

There is strong evidence that cardiomyocyte specification and differentiation is controlled by transcription factors such as the LIM-homeodomain transcription factor, Islet 1 (ISL1). ISL1 is a

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marker of myocardial lineage during mammalian cardiogenesis and marks a common population of progenitors in the heart that can differentiate into cardiomyocytes, smooth muscle, and endothelial cells [9,10]. It has been demonstrated that approximately 97% of cells within the outflow tract, 92% of cells within the right ventricle, 63% of cells within the left atria, 70% of cells within the right atria, and approximately 20% of cells within the left ventricle of a normal heart are ISL1-positive. Thus, two-thirds of the cells within the entire heart originate from ISL1-positive progenitor cells [11]. It has also been shown that ISL1 is required for survival, proliferation, and migration of progenitor cells into the cardiac tube [12]. Cells differentiated from Isl1 knockdown ESCs have shown severely reduced beating frequencies and compromised expression of cardiac sarcomeric genes (Myh6, Myh7, Mlc2a, and Mbi2c). On the other hand, over-expression of Isl1 during spontaneous differentiation of mouse ESCs into EBs resulted in a higher expression level of cardiac muscle genes compared with the control. A 2-fold over-expression of Isl1 led to a 25% increase in the number of cardiac cells [13], and the expression level of Nkx2.5 (a cardiovascular progenitor marker) increased after over-expression of Isl1 in hESCs [14]. These and other data have proven that ISL1 acts at the top of a cascade of cardiac transcription factors in the myocardial lineage [12].

Although these reports represent a critical step forward in determining the potential of ISL1 in cardiac differentiation, genetic alteration of cells continues to raise safety concerns due to transgene reactivation and insertional mutagenesis [15]. Ultimately, derivation of cardiomyocytes without viral integration is essential for the generation of safe cells for therapeutic applications.

Protein transduction has been shown to be an alternative approach for the over-expression of a desired gene in the absence of genetic manipulation [16]. However, because of eukaryotic cell membrane structure the directed intracellular delivery of proteins is less efficient in these cells. A significant exception to this rule is the application of protein transduction domains (PTDs), also known as cell-penetrating peptides (CPPs) that are capable of transporting cargo across the membrane and delivering biologically active proteins inside the cell. The initial discovery of CPPs originated from the observation that the HIV TAT transactivator could translocate across the plasma membrane by its 11 basic amino acids (residues 47–57), the TAT PTD. It has been shown that TAT has a higher efficiency for protein delivery into the cells when compared to other PTD signals [16,17,18]. The positive charges allow the protein to interact with lipid rafts in a negatively charged membrane and overcome the cell membrane barrier by different mechanisms, including macropinocytosis [19,20,21].

Recent studies have demonstrated that protein transduction of transcription factors can stimulate over-expression of its genes and initiate the specific pathway that is needed for differentiation toward a particular cell fate. By transduction of Tat-PDX-1 protein into hESCs, insulin protein production was induced [22]. In another experiment, Stock et al. succeeded in doubling the efficiency of olgodendroglial differentiation of mouse ESC-derived neural stem cells by Nkx2.2 protein transduction [23].

In this study, we successfully applied a Tat-base protein transduction system to deliver the Tat-ISL1 protein into hESCs with the intent to improve the cardiomyocyte differentiation rate under a suspension culture condition. We have demonstrated that the application of Tat-ISL1 increased the differentiation of cardiomyocytes (2–3 folds) without genetic modification.

Materials and Methods

Cloning of ISL1 cDNA

Total RNA from hESC-derived cardiac precursor cells was extracted using TRIzol reagent (Invitrogen, CA, USA) and treated with RNase-free Dnase (Invitrogen, Carlsbad, CA, USA). Reverse transcription was performed under the conditions recommended by the manufacturer using SuperScriptIII reverse transcriptase (Invitrogen, Carlsbad, CA, USA) and Oligo DT primer. Next, the Isl1 fragment was amplified by PCR using (p5x) DNA polymerase (Invitrogen, Carlsbad, CA, USA) and cloned into the pENTR-D/TOPO Gateway entry vector according to the supplier’s instructions (Invitrogen, Carlsbad, CA, USA). PCR forward and reverse primers were 5’-CACCTGCGAGCGCCGAGGGG3’ and 5’-TTAAGCCGCTCGAGTTTG3’, respectively. Forward primers included the 4 base pair sequences (GACCC) necessary for directional cloning on the 5’ end of the forward primer.

Construction of pDest17/ISL1 expression vector

cDNA from the pENTER D-TOPO/ISL1 entry clone was transferred into the pDest17 Gateway expression vector using an LR clonase recombination according to the manufacturer’s instructions (Invitrogen, Carlsbad, CA, USA). The expression vector was transformed to E. coli strain BL21 (DE3; Novagen, Madison, WI, USA) by the heat shock method according to the supplier’s manual (User Protocol TB09 Rev. F 0104). The sequence of Isl1 was verified by DNA sequencing.

Recombinant fusion protein expression and purification

For recombinant fusion protein expression, the selected clones were grown until the OD600 reached 0.8. Recombinant fusion protein expression was then induced by the addition of isopropyl-d-thiogalactopyranoside (IPTG). The expressed His6-TAT-ISL1 fusion proteins (rISL1) were purified by immobilized metal affinity chromatography (IMAC) and eluted with 8 M urea (pH 3.5), then desalted by Tris (5 mM) that contained 50% glycerol and maintained at -20°C until use. Identical volumes of elution fractions were mixed with 1/5 volume of 5x loading buffer [1 M Tris–HCl (pH 6.8), 10% w/v SDS, 0.05% w/v bromophenol blue, 50% glycerol, and 200 mM β-mercaptoethanol], heated at 95°C for 5 min and then analyzed by SDS-PAGE on a 12% (w/v) separating gel. This was followed by staining with 0.1% Coomassie brilliant blue (CBB) R-250. CBB tainted protein bands of interest were excised from the SDS-PAGE gel and samples analyzed by a Matrix-assisted laser desorption/ionization-tandem time of flight mass spectrometry (MALDI TOF/TOF MS).

Gel shift assay

To investigate the interaction of rISL1 with DNA, we applied a gel shift assay with some modifications. Genomic DNA was extracted from the hESCs. A total of 50 μg of extracted DNA was digested by Bam HI enzyme for 2 h at 37°C. We incubated 5 μg native purified rISL1 with 2.5, 5, and 10 μg of digested DNA for 1 h at 37°C. Formaldehyde was added to the reaction and allowed to incubate for 5 min in order to cross-link reacted DNA and proteins. To terminate the cross-linking reaction, 1 M glycine was used. In order to check for any nonspecific interactions between the DNA and protein, some of the bacterial proteins were allowed to remain in the final elution. Samples were then analyzed with SDS-PAGE.
Suspension cell culture and differentiation protocol

Roya5 H5 and Royan H6 hESC lines [24] were used in this study. Suspension culture of hESCs was performed according to a recently published protocol [6]. Briefly, cells were treated with 10 mM ROCK inhibitor Y-27632 (Sigma-Aldrich, Y0503) 1 h prior to dissociation from Matrigel. Cells were then washed by Ca2+- and Mg2+-free phosphate buffered saline (PBS; Gibco, 21600-051) and incubated with 0.05% trypsin at 37°C for 4–5 min. Dissociated cells were transferred into non-adhesive bacterial plates (60 mm; Griner, 628102) at 15×10^4 viable cells/ml in hESC medium that had been conditioned on mouse embryonic fibroblasts (MEFs) [25], which contained 10 mM ROCK inhibitor. After 2 days, half of the medium was replaced by the hESC medium conditioned on MEFs. The medium was changed every other day.

Differentiation of the cells into cardiomyocytes in suspension was performed according to the Laflamme et al. protocol [26] with some modifications. Briefly, 6-day old spheres were treated by 100 ng/ml Activin A for 1 day in RPMI medium (Gibco, 51800-055) supplemented with 2% B27 without vitamin A, followed by 4 days of 10 ng/ml BMP4. At day 5, the spheres were plated on gelatin-coated plates in RPMI/B27 medium without cytokines. Beating clusters were observed 5 days post-plating. In the rISL1 treated group the recombinant protein was added from days 1–8 after initiation of differentiation induction. All experiments with hESCs were performed under supervision of the Institutional Review Board and Institutional Ethical Committee of Royan Institute.

Stability and penetration of the rISL1 protein

To analyze the stability and penetration of recombinant proteins, 40 μg/ml rISL1 and elution buffer (as control) were added to the aggregated hESC differentiation media one day after differentiation initiation. Cells and culture media were collected after 2, 6, 12, 24, 36, and 48 h. The quantity of rISL1 in the cell extracts and media were analyzed by Western blot and qRT-PCR as described below.

Penetration was further confirmed by immunostaining analysis of adherent and aggregated hESC colonies treated with 40 μg/ml rISL1 protein and elution buffer (as control) for 2 h. Cells were washed 3 times by PBS/tween to ensure the removal of all rISL1 proteins that were loosely bound to the cell surfaces. The penetration of rISL1 was then investigated using anti-ISL1 and TAT antibodies.

**ISL1-GFP reporter assay**

A 4kb fragment that contained the *Isl1* promoter was isolated from human genomic DNA extracted from Royan H5 using the Expand Long Template PCR System (Roche, 10201179). The PCR forward primer was 5’ CATGCAAGATCTAATCGTCTGTTCTCGGTAG 3’ and the reverse was 5’ CTGATATTAAAGGGCTGTCTGGCAG 3’. The isolated fragment was then cloned into the pIRE2-EGFP vector. For transfection hESCs, cells were plated at 200,000 cells/6 cm diameter dish, and transfected 24 h after. 3 μg of plasmid DNA was mixed with 4 μl of X-tremeGENE 9 DNA Transfection Reagent (Roche, 06 356 787 001) and 300 μl of DMEM/F12 for 15–30 minutes and then applied to cells for a total volume of 1 ml of hESCs culture medium. After 48 h, the medium was replaced by fresh media that contained 100 μg/ml G418 (active concentration). After 1–2 weeks, transfected colonies took up and cultured. In order to examine the ability of rISL1 to induce its own gene expression, we treated undifferentiated aggregated ISL1-GFP cells with 40 μg/ml rISL1 and elution buffer (as a control); cells were examined after 5 days for GFP expression by flow cytometry.

**Western blotting**

Proteins were separated by 12% SDS-PAGE electrophoresis at 100 V for 2 h using a Mini-PROTEAN 3 electrophoresis cell (Bio-Rad, Hercules, CA, USA) and transferred to a PVDF membrane by wet blotting (Bio-Rad, Hercules, CA, USA). Membranes were blocked for 1 h with 5% BSA and incubated for 1.5 h at room temperature (RT) with the respective primary antibodies (anti-ISL1 (Abcam, ab86472, 1:5000) and anti-β-TUBULIN 49 (Millipore, P07437, 1:5000)). At the end of the incubation period membranes were rinsed 5 times (15 min each) with PBS-tween-20 (0.05%) and incubated with the peroxidase-conjugated secondary antibody [anti-mouse (Millipore, 1:6000)], as appropriate for 1 h at RT. The blots were visualized with Sigma detection reagents (Sigma, G9107) and films were scanned with a densitometer (GS-800, Bio-Rad, Hercules, CA, USA). Quantification of immunological signals was performed by Image Master software. The volume of each band was analyzed by dividing the volume percent of ISL1 by the housekeeping gene (ISL1/β-TUBULIN49) in order to assure uniformity of the protein amounts loaded on the gels.

**Immunofluorescence staining**

For immunofluorescence staining, differentiated cells were washed twice in PBS and fixed in 4% paraformaldehyde for 20 min at RT. For permeabilization, 0.2% (v/v) Triton X-100 was used for 20 min. Nonspecific antibody binding was blocked for 30 min at 37°C with 10% heat-inactivated serum. Cells were incubated overnight at 4°C with the appropriate primary antibodies: GATA4 (1:200, Santa Cruz, SC-1237); MHC (1:200, Abcam, Ab15); ISL1 (1:200, Abcam, Ab86472); NKX2.5 (1:200, R&D Systems, AF2444); DESMIN (1:200, Abcam, Ab8470); ACTININ (1:200, Abcam, Ab75805); and TAT (1:200, Cell Application Inc., CB0883). Cells were then washed twice with PBS/0.1% Tween-20 for 3 min and incubated with the appropriate secondary antibody in PBS. Fluorescence-conjugated secondary antibodies, goat anti-mouse IgG FITC (1:200, Sigma-Aldrich, F0906), goat anti-rabbit IgG FITC (1:200, Sigma-Aldrich, F1262), goat anti-mouse IgG Alexa 488 (1:100, Invitrogen, Q10110MP), and rabbit anti-goat IgG FITC (1:200, Sigma-Aldrich, F7367) were used as appropriate for 1 h at 37°C. After two washes with PBS+0.1% Tween-20 for 5 min, cells were counterstained with DAPI (Sigma-Aldrich, D4176) and analyzed with a fluorescent microscope (Olympus, IX71).

**Flow cytometric analysis**

In order to quantify cardiac protein expression, cells were dissociated by trypsinization and centrifuged for 5 min at 1500 rpm to remove cell debris. After supernatant removal, cell viability was determined by trypan blue exclusion. Cells were then washed twice in PBS and fixed in 4% paraformaldehyde for 30 min at 4°C. For permeabilization, 0.1% (v/v) Triton X-100 was used for 10 min. Nonspecific antibody binding was blocked for 30 min at RT with 10% heat-inactivated serum. The cells were incubated overnight at 4°C with the appropriate primary antibodies followed by 45 min at 37°C for the secondary antibody. Antibodies used were the same as those for immunofluorescence staining in addition to CONNEXIN43 (1:200, Abcam, Ab62689) and cTnT (1:200, Abcam, Ab64623). For each analysis, 1–5×10^5 cells were used per sample. Flow cytometric analysis was performed with a BD-FACS Calibur flow cytometer (Becton Dickinson, San Jose, CA, USA). The experiments were replicated
Quantitative reverse transcriptase PCR (qRT-PCR)

Gene expression was assessed by qRT-PCR for genes of interest. The PCR mix in each well included 10 μl of SYBR® Premix Ex Taq™ II (RR081Q, Takara Bio, Inc.), 6 μl dH2O, 1 μl each of the forward and reverse primers (5 pmol/μl), and 2 μl of single strand cDNA (16 ng/μl) in a final reaction volume of 20 μl. Primer sequences are given in Table S1. PCR was performed on a Rotor-Gene™ 6000 Real-Time PCR System (Corbett Life Science) using the following program of 95°C for 10 min (stage 1) and 95°C for 10 s, 60°C for 20 s, and 72°C for 20 s (stage 2), for 40 cycles.

qRT-PCR was conducted using 3 biologically independent replicates. Thermal conditions for all genes were the same. The annealing temperature was 60°C. Amplification specificity was verified by the melting curve method. Relative gene expression was calculated by the ΔΔCT method [27]. Target genes normalized by the reference gene Gapdh; CT did not vary under different experimental conditions when equal amounts of RNA were used. PCR efficiencies for different primer pairs were ~1 as determined by a standard curve method on serially diluted templates. All data are represented as log2-linear plots.

Statistical analysis

All quantitative experiments including Western blot, qRT-PCR, and flow cytometry were performed using 3 biologically independent replicates. Significant differences between groups were examined by the student’s t-test. P<0.05 was considered statistically significant. Data were presented as mean±SD.

Results and Discussion

Direct protein transduction into the cells both in vitro and in vivo is an efficient alternative to genetic manipulation, which leads to the production of safe cells required for cell therapy. Using this method, the concentration and duration of proteins in the cells can be easily controlled [28,29].

Figure 1. Daily qRT-PCR analysis in aggregate differentiation of hESCs. Undifferentiated aggregates of hESCs were treated by Activin A for 1 day and then for 4 days by BMP4. At day 5, the aggregates were plated without cytokines. The data show the maximum expression of the mesenendodermal marker, Brachyury, one day after Activin A treatment (day 2 after differentiation initiation). By continuing differentiation with BMP4 for the next 4 days Isl1, a marker of precardiac mesoderm, and Actinin were reached to their highest expression level. Isl1 expression was remained at high level for the next 3 days and by decreasing its expression, Meis2, a cardiac progenitor marker showed its maximum expression and after that, other cardiac progenitor genes, Gata4, Nkx2.5 and Tbx5 reached to their highest expression level respectively. Finally, the expression of MHC and cTnT, which are structural cardiomyocytes markers, got to maximum level (Fig. 1). These data shows that 3-dimensional structures of the cells are very important for cardiac differentiation and aggregated differentiation method enhances cardiac differentiation and functionality. Target genes were normalized by the reference gene Gapdh. The relative expression was calculated by dividing the normalized target gene expression of the treated sample with that of the undifferentiated state (day 0). All data represented as log2-linear plots. All data are statistically significant otherwise marked with "ns" (P>0.05).

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ISL1 Transduction Promotes hESC-Cardiomyocytes

**Figure Legend**

**A**
- DNA ladder: 50 kDa, 40 kDa, 30 kDa, 20 kDa, 10 kDa, 5 kDa, 3.75 kDa, 2 kDa
- 6X His, attB1, TAT, NLS, ISL1, attB2

**B**
- Gel electrophoresis of DNA samples
- Lanes: M, T, E1, E2

**C**
- Dilution buffer: 5 µg, 10 µg, 50 µg

**D**
- 50 kDa, 40 kDa DNA ladder
- DNA samples: rISL1 5 µg, rISL1 10 µg
- DNA+ samples: 3.75 µg, 5 µg, 7.5 µg, 10 µg

**E**
- Culture medium:
  - W/o cells + rISL1
  - Conditioned medium on cells + rISL1
  - Cell lysate of rISL1 treated
  - Cell lysate of vehicle group

**F**
- Graph showing percent volume over time:
  - rISL1 Treated: 2h, 6h, 12h, 24h, 36h, 48h
  - Control

**G**
- Graph showing log expression in rISL1 treated cells:
  - Expression in untreated cells:
    - 2h, 6h, 12h, 24h, 36h, 48h

**H**
- Flow cytometry histograms:
  - TAT/DAPI, Adherent cells
  - ISL1/DAPI, Adherent cells
  - ISL1/DAPI, Aggregate cells

**I**
- Fluorescent images:
  - TAT/DAPI, Adherent cells
  - ISL1/DAPI, Adherent cells
  - ISL1/DAPI, Aggregate cells
We have demonstrated that the transduction of rISL1 protein enhanced hESCs differentiation into the beating cardiomyocytes phenotype.

Establishment of a cardiac differentiation protocol

hESCs were expanded in feeder and serum-free conditions on Matrigel-coated plates. hESCs (RoyaH5) were induced to differentiate under adherent conditions and as aggregates in the absence of bFGF, with the addition of Activin A for one day and BMP4 for four days, and subsequently in growth factor-free medium in the presence of 2% B27, as described previously [26]. In adherent conditions none of the colonies were able to beat during 40 days of culture (Fig. S1A). RT-PCR analysis of differentiated cells during days 0–14 showed the onset of the expression of cardiac markers, *Brachyury*, *Isl1*, *Mef2c*, *Tbx5*, *Gata4*, *Nkx2.5*, *αMHC*, *βMHC*, and *MLC2v* (Fig. S1B). This showed that despite morphological changes and expression of some cardiac genes in the differentiated cells they are not functional, therefore they were unable to beat. In aggregate conditions, hESCs initially expanded in suspension aggregates as previously described [6]. Next, 6-day aggregates were induced to differentiate as described above and plated on gelatin-coated plates. In this way, 20 ± 2.5% of the aggregates showed evidence of beating 5 days post-plating. qRT-PCR analysis of aggregate differentiation of hESCs showed the maximum expression of the myocardial marker, *Brachyury*, one day after Activin A treatment (day 2 after differentiation initiation) (Fig. 1). By continuing differentiation with BMP4 for the next 4 days *Isl1*, a marker of precardiac mesoderm, and *Actinin* were reached to their highest expression level. *Isl1* expression was remained at high level for the next 3 days and by decreasing its expression, *Mef2c*, a cardiogenic progenitor marker showed its maximum expression level. After that, other cardiogenic progenitor genes, *Gata4*, *Nkx2.5* and *Tbx5* reached to their highest expression level respectively. Finally, the expression of *MHC* and *cTnT*, which are structural cardiomyocytes markers, got to maximum level (Fig. 1). These data showed that in aggregate differentiation, the 3-D structure of cells enhanced cardiac differentiation and functionality.

Expression pattern of ISL1 during differentiation

*ISL1* is necessary for the proliferation and survival of ISL1+ progenitor cells and the inhibition of further cardiac differentiation, therefore a decrease in its expression level is necessary before ISL1+ progenitor cells can differentiate into cardiac cells [11,12]. It is important to define the optimum time for addition and removal of the rISL1 protein in order to achieve the highest number of ISL1+ progenitor and cardiac cells, respectively. To address this, we have analyzed the expression pattern of *Isl1* during the first 15 days of differentiation using qRT-PCR. *Isl1* expression was detected after Activin A treatment and reached a maximum level at days 7–8 (Fig. 1). Therefore, we decided to add rISL1 protein to the differentiation medium from days 1–8 after the initiation of differentiation. In this condition, rISL1 treatment was started from the first day of *Isl1* gene expression and none of the *Isl1* expressing cells were missed. We did not continue rISL1 treatment after day 8, because it inhibited further differentiation.

Generation of cell-permeable rISL1 protein

A 522 bp of the *Isl1* native sequence was amplified from human cardiac precursor cell mRNA with specific primers and its DNA sequence was confirmed by sequencing. The right orientation of the primary cloning of *Isl1* was demonstrated by a PCR that utilized T7 forward and *Isl1* reverse primers for the pENTER D-TOPO/*Isl1* entry clone. A fusion protein (rISL1 protein) that consisted of the TAT transduction domain for protein transduction, a N-terminus histidine tag for protein purification, and the *Isl1* protein was then generated using a bacterial pDest17/*Isl1* expression vector (Fig. 2A). Since we used the *Isl1* native sequence for protein expression, there was no exogenous nuclear localization signal included. To produce rISL1 protein in the bacterial host, we used the pDest17 expression vector system which is one of the most frequently employed ways to efficiently and effectively synthesize heterologous proteins in prokaryotic cells. This system possesses the characteristics of having an exceptionally strong promoter allowing high-level production of recombinant proteins. The rISL1 protein (42 kDa) was successfully purified from the bacterial expression system as demonstrated by SDS-PAGE and Western blot analysis using anti-ISL1 antibody (Figs. 2B and C). The identity of the expressed protein was also confirmed by mass spectrometry (data not shown). We confirmed that the rISL1 protein was bound to digested genomic DNA by using a gel shift assay (Fig. 2D) which showed that not only the purified protein was rISL1, but that it was functional and had the ability to bind with DNA in vitro.

Penetration and stability of the rISL1 protein

Cellular uptake and stability of the rISL1 protein was confirmed by Western blot analysis of cell lysates from control or rISL1-treated hESCs (Fig. 2E). Our results showed a higher abundance of ISL1 in the rISL1-treated group compared to the control, which
Figure 3. Optimization of the effect of rISL1 protein on hESCs. (A) To evaluate the effect of discontinuous (2 h/day) or continuous rISL1 protein addition on hESCs (Royan H5) differentiation, cells were treated continuously or discontinuously from days 1–8 post initiation of differentiation. *isl1 qRT-PCR analysis of differentiated cells at day 8 showed higher significant endogenous isl1 expression in hESCs in the continuous protocol. Thus, continuous treatment was applied in the next steps. * : P < 0.05. (B) To determine the best concentration of rISL1 protein for cardiac differentiation, cells were treated with four different concentrations of recombinant protein: 10, 20, 30, and 40 μg/ml in continuous treatment of hESCs during days 1–8 after initiation of differentiation. During differentiation, cells that were treated by 10 and 20 μg/ml rISL1 protein were morphologically similar to hematopoietic and endothelial progenitors, while the 30 and 40 μg/ml concentrations showed cardiomyocyte and
muscular appearances. It seems that 30 and 40 μg/ml rISL1 protein are better concentrations for cardiac differentiation. *P<0.05 (C) qRT-PCR analysis of differentiated cells at day 8 by different concentrations of rISL1 also showed that 40 μg/ml of the rISL1 protein induced more endogenous *Isl1*, but less *Me2c* and *Nkx2.5* expressions. *P<0.05 (D) Schematic diagram of the differentiation protocol by the addition of rISL1 protein (40 μg/ml), which was added after induction with Activin A (days 1–8). qRT-PCR analysis of endogenous *Isl1* expression in hESCs demonstrated that treated cells expressed higher significant endogenous *Isl1* than the untreated control. *P<0.05 (E) The percentage of beating clusters in continuous treatment of hESCs by 40 μg/ml rISL1 protein during days 1–8 after differentiation initiation in comparison with the control (vehicle-treated) group. The percentage of beating clusters in the rISL1-treated group was significantly higher than the untreated group at day 14 after plating (75±10% vs. 20±2.5%). *P<0.05 (F) rISL1 treatment resulted in a 3.2±0.5 fold increase in the number of beating areas in comparison with untreated control group, rISL1 also caused a 2.2±0.4 fold increase in the other hESC line, Royan H6, which shows the reproducibility of this protocol for another hESC line. *P<0.05 (G) In order to assess the expression of cardiac-specific genes, we collected samples at 3 stages: day 3 after plating (the day of rISL1 removal); day 14 after plating (day of maximum beating); and day 20 after plating (day that beating decreased and cells were mature) by qRT-PCR in two hESC lines. Target genes were normalized by the reference gene Gapdh. The relative expression was calculated by dividing the normalized target gene expression of treated hESCs with rISL1 protein and elution buffer (as control) with that of the undifferentiated state (day 0). All data are statistically significant in comparison with undifferentiated state (day 0) otherwise marked with “ns” (ns: P>0.05). a: P<0.05 in comparison with control group (elution buffer treated group). All data were represented as log2-linear plots.

Based on these results, we decided to add rISL1 protein into the differentiation media every other day during medium replacement. Cell penetration of the rISL1 protein was further confirmed by immunostaining analysis of adherent and aggregated cells using both anti-TAT and ISL1 antibodies. Two hours following transduction of hESCs, most cells were positive for the labeled TAT or ISL1 protein (Fig. 2F). rISL1 proteins were detected around the nucleus in adherent and aggregated cells, which suggested the ability of the recombinant protein to penetrate deep inside aggregated cells (Fig. 2F). This finding was consistent with the prevailing view that TAT can promote cellular uptake via endocytosis [30,31,32].

Defining rISL1 treatment conditions

Previous studies have demonstrated that discontinuous or continuous addition of recombinant protein into cell culture media is also an important factor that should be considered [23,33]. In order to optimize protein transduction, the cells were either discontinuously (2 h/day) or continuously (from days 1–8) treated by rISL1. qRT-PCR analysis of differentiated cells at day 8 showed higher endogenous *Isl1* expression in hESCs in the continuous protocol (P<0.05, Fig. 3A). The following experiments were performed by continuous protein treatment.

To study the dose dependency of rISL1 transduction, hESCs were exposed to different concentrations of the purified protein (10, 20, 30, and 40 μg/ml) in the continuous treatment of hESCs during days 1–8 after differentiation initiation. We observed that the concentration greater than 40 μg/ml was lethal (data not shown). The differentiating cells at concentrations of 10 and 20 μg/ml of the rISL1 protein were morphologically similar to hematopoietic and endothelial progenitors, while 30 and 40 μg/ml rISL1 protein showed cardiomyocytes and muscular appearances (Fig. 3B). According to qRT-PCR analysis, 40 μg/ml rISL1 protein induced more endogenous *Isl1* and less expression of *Me2c* and *Nkx2.5* (Fig. 3C). These data were consistent with previous data which has shown that ISL1 marks a common population of progenitors in the heart that can differentiate into cardiomyocytes, smooth muscle, and endothelial cells [9,10]. It seems that different levels of ISL1 protein direct cells towards specific lineages. However, more experiments are needed to find the exact amount of ISL1 expression required for each lineage differentiation.

Increasing cardiac differentiation using rISL1 protein

Based on the above mentioned experiments, we continuously added 40 μg/ml of rISL1 protein into differentiation medium from days 1–8 after differentiation initiation. The effect of rISL1 protein on the expression of endogenous *Isl1* was analyzed using qRT-PCR at 1, 2, 3, 5, and 8 days after differentiation initiation. Our results showed that treated cells expressed higher endogenous *Isl1* than the untreated control (P<0.05, Fig. 3D). We further continued differentiation to obtain beating clusters. The beating areas appeared at day 5 post-plating of aggregates and the percent of beating areas were significantly higher in rISL1-treated cells compared to the control (Fig. 3E). The difference was more pronounced at 14 days after plating when the percent of beating areas reached 75±10% in rISL1-treated cells compared to 20±2.5% in the control when more than 1000 embryoid bodies were assessed in each group (Fig. 3E). Therefore, rISL1 treatment resulted in a 3.2±0.05 fold increase in the number of beating areas (Fig. 3F). In order to check reproducibility of this protocol the same experiments were performed using another hESC line, Royan H6. Our data indicated that rISL1 treatment could also cause a 2.2±0.4 fold increase in the number of beating areas in Royan H6. Temporal expression of cardiac genes showed the highest levels of *Isl1*, *Me2c*, *Hand1*, *Nkx2.5*, *Actinin*, *MHC*, *cTnT*, *MLC2a*, and *MLC2v* at day 14 in both hESC lines, Royan H5 and Royan H6 (Fig. 3G). Our data showed that *MLC2v* expression increased in the rISL1-treated groups while *MLC2a* decreased. These results suggested that differentiated cells were directed toward ventricular cardiomyocytes. This observation was consistent with previous reports in which approximately 92% of cells within the right ventricle and about 20% of cells within the left ventricle of a normal heart were ISL1-positive [11].

Immunofluorescence staining of differentiated cells showed expressions of NKK2.5, GATA4, ISL1, ACTININ, DESMIN, and MHC (Fig. 4A). By flow cytometric analysis, higher expressions of cTnT, CONNEXIN 43, ACTININ, and GATA4 were detected (P<0.05, Fig. 4B). Based on the *Isl1* gene expression profile, rISL1 was added to the cell culture media from days 1–8 of differentiation, when the expression level of endogenous *Isl1* was first detected (day 1) and reached its maximum level (day 8). The addition of rISL1 with
Figure 4. Marker analysis of hESC-derived cardiomyocytes. Cardiomyocytes were dissociated at differentiation day 14 post plating, cultured for an additional 2 days, and assessed by (A) immunofluorescence staining and (B) flow cytometry for cardiac-specific markers. Flow cytometric analysis showed significant increases in the levels of cTnT, CONNEXIN43, ACTinin, and GATA4 protein expression (P<0.05). The experiment was performed for at least 3 independent biological replicates. *: P<0.05.
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BMP4 to the cell culture media may also enhance its effect. It has been demonstrated that ISL1 promoted BMP expression (such as BMP6 and BMP7), and the expression level of BMP4 was reduced in ISL1 mutant cells [11]. It is likely that rISL1 increases the number of beating cells through enhancing the expression of the gap junction protein Connexin40 (a major protein in the conduction system) by BMP signaling. It has been shown that BMP signaling is necessary for the expression of T-box transcription factors [34]. Connexin40 is one of the direct downstream targets of the T-box transcription factors, which play an important role in the conduction system [35].

Taken together, these data indicate that direct delivery of the transcription factor ISL1 by protein transduction enhanced the cardiomyocyte differentiation in hESCs in vitro.

Conclusions

In this study we showed that under cell culture conditions purified rISL1 protein was stable for at least 48 h. When the protein was added to hESCs cultures, it efficiently penetrated into the cells and enhanced the differentiation of the two hESC lines into cardiac cells up to 3-fold.

This approach may pave the way for the scaled up expansion of hESCs as carrier-free suspension aggregates for an extended period of time. It provides a controlled environment for a homogeneous culture and simplifies the handling and controlling of the differentiation of hESCs, which are required for their applications in bioreactor culture systems and cell therapy.

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Another advantage of this method is the lack of genetic manipulation of hESCs which may decrease the risk of their application in cell therapy. In conclusion, our data indicate that by addition of rISL1 protein into the differentiation medium we have successfully produced large numbers of functional cardiomyocytes that can be easily applied in drug discovery or cell therapy. However, further research is necessary to additionally increase the efficiency of differentiation using this method.

Supporting Information

Figure S1 Adherent differentiation and gene expression of hESCs (Rown H5) by the Laflamme et al. protocol [26]. (A) D0: Undifferentiated colony of hESCs. D1: One day after addition of Activin A. D5: Four days after treatment by BMP4. D15: Fifteen days after differentiation initiation. (B) RT-PCR data of daily expression of cardiac genes after differentiation induction. (TIF)

Table S1 Primers and reaction conditions used in real time RT-PCR analysis. (DOC)

Author Contributions

Conceived and designed the experiments: GHS NA HB. Performed the experiments: HF MY. Analyzed the data: GHS NA HB. Contributed reagents/materials/analysis tools: HR BAM FF ZG MA. Wrote the paper: HB GHS NA.

ISL1 Transduction Promotes hESC-Cardiomyocytes
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