

Characterization and Angiogenic Potential of Human Neonatal and Infant Thymus Mesenchymal Stromal Cells

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Key Words. Thymus gland • Mesenchymal stromal cells • Cardiac surgery • Neonatal • Angiogenesis

ABSTRACT

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on February 23, 2015.

©AlphaMed Press

http://dx.doi.org/

Received October 22, 2014:

1066-5099/2015/\$20.00/0

10.5966/sctm.2014-0240

accepted for publication January

22, 2015; published Online First

Resident mesenchymal stromal cells (MSCs) are involved in angiogenesis during thymus regeneration. We have previously shown that MSCs can be isolated from enzymatically digested human neonatal and infant thymus tissue that is normally discarded during pediatric cardiac surgical procedures. In this paper, we demonstrate that thymus MSCs can also be isolated by explant culture of discarded thymus tissue and that these cells share many of the characteristics of bone marrow MSCs. Human neonatal thymus MSCs are clonogenic, demonstrate exponential growth in nearly 30 population doublings, have a characteristic surface marker profile, and express pluripotency genes. Furthermore, thymus MSCs have potent proangiogenic behavior in vitro with sprout formation and angiogenic growth factor production. Thymus MSCs promote neoangiogenesis and cooperate with endothelial cells to form functional human blood vessels in vivo. These characteristics make thymus MSCs a potential candidate for use as an angiogenic cell therapeutic agent and for vascularizing engineered tissues in vitro. STEM CELLS TRANSLATIONAL MEDICINE 2015;4:339–350

INTRODUCTION

Mesenchymal stromal cells (MSCs) isolated from bone marrow, a tissue with regenerative abilities [1, 2], are being evaluated for degenerative and acquired diseases of tissues and organs (e.g., the heart) that may have relatively limited intrinsic regenerative potential [3]. The human thymus is a vascularized organ that plays a pivotal role in T-lymphocyte development, and its durability in providing a robust acquired immune response throughout the lifespan may be partially attributed to its ability to regenerate after injury, infection, or partial surgical resection. A brisk angiogenic phase that involves the participation of MSCs supports thymus regeneration [4, 5]. The in vivo regenerative and proangiogenic function of resident MSCs in the thymus makes them a potential candidate as a cell therapeutic agent in ischemic diseases of other tissues.

According to the Society of Thoracic Surgeons National Database, severe congenital heart disease requiring surgical correction in the neonatal and infant period occurs in >10,000 patients each year in the U.S. Partial or subtotal thymus resection is required to expose the heart and great vessels during many of these surgical procedures. We and others have identified this normally discarded tissue as a plentiful source of MSCs obtained by enzymatic digestion [6, 7]. However, the use of dissociative enzymes and subsequent washes are significant additional steps for clinical translation of these MSCs into patients and may yield numerous other thymus cell types, such as thymic epithelial cells.

The objective of this investigation was twofold. First, we characterized MSCs isolated by explant culture of mechanically fragmented human neonatal and infant thymus tissue. Second, we investigated the in vitro and in vivo proangiogenic qualities of explant culture-isolated human neonatal thymus MSCs. Our results indicate that discarded human neonatal thymus tissue is a source of MSCs with angiogenic potential.

MATERIALS AND METHODS

Cell Isolation and Culture

All studies were performed under an approved protocol from the University of Michigan institutional review board. After informed consent was given by the parents, discarded thymus tissue from infant heart operations was mechanically minced and subjected to explant culture. Experimental details are shown in the supplemental online data. Unless specified otherwise, all experiments were performed with cells from passages 3–9. Downloaded from http://stemcellstm.alphamedpress.org/ by Janko Mrkovacki on April 3, 2015

Thymus MSC Isolation Yield Estimate

Thymus tissue fragments from 4 patients were weighed and placed in 35-mm tissue culture dishes for explant culture (3–4 fragments per patient). After 10 days of culture, tissue fragments were removed, and the number of cells in each dish was counted. The average number of cells per gram of tissue was then calculated.

CFU-F Efficiency and Limiting Dilution Assay

All colony-forming unit (CFU) assays were performed with passage 7 thymus MSCs from 8 patients. Experimental details of CFU efficiency (CFE) determination are shown in the supplemental online data. Colonies with >50 cells were counted and recorded.

Fibroblastic CFU (CFU-F) prevalence was also estimated by a limiting dilution assay (LDA) [8]. Experimental details are shown in the supplemental online data. CFU-F frequency was then calculated using L-Calc software (Stemcell Technologies, Vancouver, BC, Canada, http://www.stemcell.com).

Thymus MSC Growth Kinetics

Growth kinetics of thymus MSCs isolated from four patients were studied with a cumulative population-doubling analysis. Experimental details are shown in the supplemental online data.

Flow Cytometric Analysis

Surface markers of passage 4–9 thymus MSCs isolated from 6 patients were characterized by flow cytometry using antibodies against CD29, CD44, CD45, CD90, CD105, CD73, CD166, CD49e, CD56, STRO-1, CD271, SSEA-4, HLA-ABC, HLA-DR, and nestin. Experimental details are shown in the supplemental online data.

Pluripotency Gene Expression Analysis

Expression of pluripotency-related genes *Oct-4*, *Sox-2*, and *Nanog* was determined in thymus MSCs (passage 5 or 6, n = 5) relative to human induced pluripotent stem cells using quantitative polymerase chain reaction (qPCR). Experimental details are shown in the supplemental online data.

Multilineage Differentiation

The ability of thymus MSCs isolated from three patients to differentiate into osteogenic, adipogenic, and chondrogenic lineages when cultured in specific differentiation media was investigated. Experimental details are shown in the supplemental online data.

CD248 Expression

Expression of CD248 (endosialin) on neonatal human thymus MSCs was performed by immunofluorescent staining. Experimental details are shown in the supplemental online data.

Two-Dimensional Angiogenesis Assay

A two-dimensional (2D) in vitro assay was performed to investigate whether thymus MSCs, with or without human umbilical vein endothelial cells (HUVECs), induced tube formation. Experimental details are shown in the supplemental online data.

Multicellular Spheroid Generation

HUVEC, thymus MSC, and HUVEC plus thymus MSC spheroids used in the three-dimensional (3D) angiogenesis assay and in vivo experiments were created by hanging drop culture. Experimental details are shown in the supplemental online data.

Fluorescent Labeling of Cells and Spheroid Sprouting Time Course Study

HUVECs and thymus MSCs were labeled with the vital cell dyes PKH26 and PKH67, respectively, according to the manufacturer's directions (Sigma-Aldrich, St. Louis, MO, https://www. sigmaaldrich.com). Nuclei were stained with Hoechst 33342 (Molecular Probes, Life Technologies; Thermo Fisher Scientific, Waltham, MA, http://www.thermofisher.com/en/home.html). After completion of the labeling, spheroids were generated as described above and imaged consecutively at 0, 24, and 48 hours after placement in fibrin hydrogel with a confocal microscope. Independent experiments were performed in triplicate.

3D Angiogenesis Assay

Fibrin hydrogel was generated in each well of a 24-well plate, as described, followed by the addition of 75 spheroids per well prior to polymerization to ensure that spheroids were embedded within the hydrogel. There were three spheroid groups: HUVECs, HUVECs plus thymus MSCs, and thymus MSCs. After fibrinogen polymerization, basal EGM-2 was added to each well. Spheroids were then incubated overnight and imaged at \times 100 using an inverted phase contrast microscope (20 spheroids per group). Images were digitally acquired and then analyzed using NeuronJ plugin (Erik Meijering, http://www. imagescience.org/meijering/software/neuronj/) for ImageJ software (NIH, Bethesda, MD, http://imagej.nih.gov/ij/). Primary sprouts emanating from each spheroid were measured by tracing from the base to the furthest tip. Branches from primary sprouts were measured from their origin to the tip. Cumulative branch length and total number of branches were then calculated for each spheroid. Spheroids located along the edge of the wells or in close proximity to each other were excluded from image analysis. Independent experiments were performed in triplicate. In separate experiments, we explored the effects of modifying the ratio of cell types in combination cell spheroids with thymus MSC/HUVEC ratios of 1:3, 1:1, and 3:1 on spheroid sprouting.

Angiogenic Gene Expression Analysis

To gain insight into the findings demonstrated by the spheroid and monolayer sprouting assays, we evaluated differential gene expression for *VEGFA*, *bFGF*, and *HIF-1* α in the different groups and culture conditions using qPCR. After 48 hours, fibrin gels containing spheroids or cell monolayers were digested with nattokinase (50 fibrinolytic units per milliliter, NSK-SD; Japan Bio Science Laboratory Co., Ltd, Osaka, Japan, http://jbsl-net.com/ english/index.html) [9] prior to RNA isolation and qPCR. The effects of modifying the ratio of cell types in combination cell spheroids with thymus MSC/HUVEC ratios of 1:3, 1:1, and 3:1 on angiogenic gene expression were also investigated. Experimental details are shown in the supplemental online data and supplemental online Table 1.

Enzyme-Linked Immunosorbent Assay

Vasculogenic and angiogenic growth factor (VEGFA and bFGF) levels in supernatants were measured using enzyme-linked immunosorbent assay kits (VEGFA kit: Invitrogen, Life Technologies, Thermo Fisher Scientific; bFGF kit: Abcam, Cambridge, MA, http://www.abcam.com), according to the manufacturer's directions. To investigate the effects of the combinations of HUVECs and thymus MSCs on growth factor production, HUVEC, thymus MSC, and HUVEC plus thymus MSC spheroids were generated and embedded in fibrin gel in a 12-well plate (500 spheroids per well), and supernatants were collected after 48 hours of culture for analysis. Independent experiments were performed in triplicate.

In Vivo Assessment of Thymus MSC-Induced Angiogenesis

The care of animals was in accordance with institutional guidelines. Constructs (3 per group) containing 500 spheroids (HUVECs, thymus MSCs, and HUVECs plus thymus MSCs) or no spheroids (control group) were made in 96-well plates with 200 μ l of fibrin hydrogel and then were implanted subcutaneously in the dorsal region of NOD/SCID mice (Charles River Laboratories International, Wilmington, MA, http://www.criver.com). Constructs were explanted 14 days after implantation, fixed with 4% paraformaldehyde, serially dehydrated in ethanol solutions, infiltrated with xylene, and then embedded in paraffin. Paraffin blocks were sectioned at 5- μ m thickness, mounted on glass slides, and then stained with hematoxylin and eosin. The vessel density within and adjacent to the constructs (between the panniculus carnosus muscle layer and the superficial aspect of the construct) was determined. Details on vessel density quantification are shown in the supplemental online data. To determine the origin of blood vessels identified within and adjacent to the construct, immunohistochemical analysis was performed using a monoclonal antibody specific for human CD31 (Dako, Carpinteria, CA, http:// www.dako.com) along with the appropriate secondary antibodies. Cell nuclei were counterstained with hematoxylin. To determine the possibility of natural killer (NK) cell involvement in the observed angiogenesis surrounding and within the construct, a separate group of HUVEC plus thymus MSC constructs were explanted on day 7 (when early infiltrate would have been expected) and evaluated for CD335 (BioLegend, San Diego, CA, http://www.biolegend.com), a NK cell marker.

Statistical Analysis

Data were expressed as mean \pm SD. Statistical differences between multiple groups were determined using one-way analysis of variance (ANOVA) with post hoc Tukey honestly significant difference test. Two-way ANOVA with post hoc Tukey honestly significant difference test was used to analyze angiogenic gene expression and vessel density data. Significance was defined as a *p* value <.05.

RESULTS

Characterization of Explant Culture Isolated Neonatal Human Thymus MSCs

Thymus MSCs were isolated from 10 different neonates and infants (aged 2–150 days; mean age: 51 \pm 61 days) who carried a diagnosis of hypoplastic left heart syndrome, D-transposition of

the great arteries, pulmonary atresia, complete atrioventricular septal defect, aortic arch hypoplasia, tetralogy of Fallot, and pulmonary artery sling by using an explant culture method. Only thymus MSCs from patients with an absence of known chromosomal abnormalities (n = 8) were evaluated. Discarded thymus tissue was mechanically minced under sterile conditions in the operating room (Fig. 1A, 1B). After 5–10 days of culture, elongated cells with fibroblastic morphology migrated from thymus tissue onto the culture surface (Fig. 1C). An average of (1.95×10^5) \pm (2.71×10^5) plastic-adherent cells with MSC morphology per gram of thymus tissue was isolated by using this explant culture method (n = 4).

Clonogenicity of thymus MSCs (all between passages 4–9) was evaluated by CFE and CFU-F frequency by limiting dilution assays. Two weeks after plating thymus MSCs (n = 7), we observed a CFE of 20%–57% (Fig. 1D, 1E). Using the LDA method (n = 9), we found that CFU-F frequency was an average of 1 in 166 (range: 1 in 31 to 1 in 467).

We evaluated the growth kinetics of thymus MSCs isolated from four patients. Thymus MSCs displayed exponential growth in nearly 30 population doublings (Fig. 1F). Average doubling time for these thymus MSCs during this period was 2.85 \pm 0.54 days.

Human neonatal thymus MSCs from three different patients differentiated into adipogenic, osteogenic and chondrogenic lineages (Fig. 2A–2D). By flow cytometric analysis, human neonatal thymus MSCs from five different patients were nearly all positive for CD44, CD90, CD29, CD105, CD73, and CD166 and negative for CD45, HLA-DR, CD56, STRO-1, SSEA-4, and nestin (Fig. 2E). Approximately 1%–3% of thymus MSCs were also negative for HLA-ABC.

Human MSCs are also known to express pluripotency genes during early passages, and that may contribute to their regenerative abilities [10–13]. We investigated this characteristic in thymus MSCs (n = 5, passage 5 or 6) with qPCR and found that thymus MSCs expressed low levels of *Oct-4*, *Nanog*, and *Sox-2* gene expression (supplemental online Fig. 1). All lines that were evaluated consistently expressed low levels of *Oct-4*, whereas there was more variability in the expression of *Nanog* and *Sox-2*.

Proangiogenic Characteristics of Thymus MSCs

Because CD248-positive thymus MSCs have been demonstrated to contribute to angiogenesis during murine thymus regeneration [5], we determined whether human thymus MSCs also expressed this surface marker. With immunofluorescence staining, we found that human thymus MSCs (n = 6) were all positive for CD248 (supplemental online Fig. 2).

We evaluated the in vitro proangiogenic characteristics of thymus MSCs combined with HUVECs using a conventional 2D monolayer method and a quantitative 3D spheroid sprouting assay because 3D culture of MSCs has been demonstrated to potentiate their proangiogenic effects and may be more representative of their in vivo behavior [14, 15].

It is well known that HUVECs cultured on Matrigel form a tubule network after overnight culture [16]. We chose to use fibrin hydrogel instead of Matrigel to eliminate the effects of the angiogenic growth factors contained in the latter. We observed that HUVECs cultured on fibrin hydrogel did not manifest any evidence of tubulogenesis at 48 hours of culture (Fig. 3A). Thymus MSCs tended to cluster together and form multicellular masses when cultured on fibrin hydrogel (Fig. 3B). The combination of thymus



Figure 1. Discarded human neonatal thymus tissue is a source of mesenchymal stromal cells (MSCs). (A): Discarded human neonatal thymus tissue during pediatric cardiac surgery. (B): Minced thymus tissue prior to plating. (C): Cells migrating from thymus tissue fragments during explant culture at 10 days. (D): Clonogenicity of thymus MSCs at 2 weeks (representative of 7 donors). (E): Colony-forming efficiency of thymus MSCs. (F): Averaged cumulative population doubling of thymus MSCs (*n* = 4) over 9 weeks of culture. Abbreviation: CFU-F, fibroblastic colony-forming unit.

MSCs and HUVECs on fibrin hydrogel yielded an interconnected honeycomb tubule network (Fig. 3C).

Multicellular spheroids embedded in fibrin hydrogel were used for the 3D assessment of angiogenesis. Spheroids composed only of HUVECs did not manifest much sprouting at 16 hours (Fig. 4A). Spheroids composed of only thymus MSCs demonstrated variable sprouting that was dependent on the donor (Figs. 4B-4F). However, spheroids with the combination of HUVECs plus thymus MSCs formed extensive, radiating, tubule-like structures within 16 hours, regardless of donor source of thymus MSCs (Fig. 4G, 4H). The morphology of the sprouts was also distinct in that the combination HUVEC plus thymus MSC spheroids had the most complex branching pattern compared with the thymus MSC-only group (Fig. 4I). At 72 hours, sprouts from HUVEC plus thymus MSC spheroids were more extensive and appeared to anastomose with those from adjacent spheroids (Fig. 4J). We had arbitrarily chosen a 3:1 ratio of HUVECs to thymus MSCs (3:1) because we speculated that endothelial cells (ECs) would outnumber MSCs in vivo based on separate measures of MSCs and ECs in bone marrow [17–19]. We also repeated these sprouting experiments with a HUVEC/thymus MSC ratio of 1:3 and 1:1 and found no significant differences in branch number, and the 1:1 ratio group demonstrated longer total branch length (supplemental online Fig 3).

To understand the contribution of the different cell types to spheroid sprout formation, HUVECs and thymus MSCs were labeled with vital dyes PKH67 and PKH26, respectively. Fluorescent labeling in the combination group of HUVECs plus thymus MSCs revealed that after spheroid formation, MSCs composed the core of the spheroid, whereas the HUVECs remained in the periphery (Fig. 5). After allowing the spheroids to sprout, the thymus MSCs migrated to form the leading edge of the sprouts, whereas the HUVECs composed the core of the spheroid. At 8 hours, many of these sprouts originated from the thymus MSCs, with HUVECs colocalized to a subset of the bases of these sprouts (Fig. 5). At 24 hours, the sprouts had increased in number, complexity, and length. Again, many of the sprouts were composed of thymus MSCs; however, a larger subset also contained HUVECs, which extended further out from the base compared with those observed at the 8-hour time point (Fig. 5).

Thymus MSC Angiogenic Gene and Protein Expression

To gain further insight into the angiogenic sprouting seen in 3D versus 2D culture conditions and the effects of cell composition (HUVECs, thymus MSCs, and HUVECs plus thymus MSCs) on spheroid sprouting, we investigated changes in expression for genes known to play a role in angiogenesis and vasculogenesis: VEGFA, bFGF, and HIF-1 α . Culturing HUVECs in spheroids resulted in a decrease in expression of all studied genes compared with monolayer culture. However, placing thymus MSCs in spheroid form had varying effects on the genes (Fig. 6A). For VEGFA, expression decreased when thymus MSCs were cultured in spheroids (Fig. 6A). For *bFGF* and *HIF-1* α , expression increased from two to five times when culturing thymus MSCs in spheroid form (Fig. 6A). In monolayer and spheroid culture, thymus MSCs had similar levels of VEGFA expression. Relative to thymus MSCs in spheroid form, we noted that combining HUVECs with thymus MSCs resulted in an increase in VEGFA expression; however, in this combination of cell types, the expression of *bFGF* or *HIF-1* α did not change or slightly decreased (Fig. 6A). Increasing the thymus MSC content of spheroids resulted in an increase in VEGFA, bFGF, and HIF-1 α gene expression (supplemental online Fig 3).



Figure 2. Characterization of human neonatal thymus mesenchymal stromal cells (MSCs) isolated by explant culture. **(A)**: Oil Red O staining representing adipogenic differentiation after differentiation with adipogenic-differentiation media. **(B)**: FABP-4 expression on thymus MSCs after differentiation with adipogenic-differentiation media. **(C)**: Osteocalcin expression on thymus MSCs after differentiation media. **(D)**: Aggrecan expression on thymus MSCs after differentiation with chondrogenic-differentiation media. **(A)**: Flow cytometric profile of thymus MSCs. All results are representative of three independent experiments. **(E)**: Flow cytometric profile of thymus MSCs. All results are representative of thymus MSCs isolated from five to seven patients. Scale bar = 50 μ m.



Figure 3. Thymus mesenchymal stromal cells (MSCs) cooperate with human umbilical vein endothelial cells (HUVECs) to form a network in a two-dimensional angiogenesis assay. (A): Monolayer appearance of HUVECs after 48 hours of culture on fibrin hydrogel. (B): Thymus MSCs clustered together after 24 hours of culture on fibrin hydrogel. (C): Combining HUVECs with thymus MSCs (2:1) resulted in the appearance of interconnected tubules at 24 hours. Scale bars = 100 μ m. Results are representative of two independent experiments.

Monolayer and spheroid VEGFA and bFGF gene expression changes were confirmed by measuring protein levels in supernatants (n = 3) (Fig. 6B, 6C). We found that HUVEC spheroids had undetectable VEGFA levels, whereas supernatant VEGFA levels in both thymus MSC spheroids and HUVEC plus thymus MSC spheroids were significantly higher. Supernatant VEGFA levels were not different in HUVEC plus thymus MSC spheroids compared with thymus MSC spheroids (Fig. 6B). Supernatants from HUVEC spheroids contained low levels of bFGF. Thymus MSC spheroids had significantly higher levels of bFGF in the supernatant compared with HUVEC spheroids. The combination of HUVEC plus thymus MSC spheroids yielded the greatest concentration of bFGF in

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Figure 4. Thymus MSCs cooperate with HUVECs to promote tubule formation in a spheroid three-dimensional (3D) angiogenesis assay. (A): Spheroids composed of HUVECs and embedded in fibrin hydrogel did not manifest any sprouting after 16 hours. (**B**–**F**): Spheroids composed of thymus MSCs isolated from different donors manifested varying degrees of tubule formation at 16 hours. (**G**, **H**): Spheroids composed of HUVECs plus thymus MSCs (regardless of donor) all demonstrated extensive branching tubule formation at 16 hours. Scale bars = 100 μ m. (**I**): Quantitative analysis of 3D angiogenesis assay with HUVEC plus thymus MSC spheroids demonstrating the greatest cumulative branch length and total number of branches (*n* = 20 spheroids per group). (**J**): Spheroids composed of HUVECs plus thymus MSCs demonstrated complex, anastomosing branches at 72 hours. Results are representative of three independent experiments. Abbreviations: HUVEC, human umbilical vein endothelial cell; MSC, mesenchymal stromal cell.

the supernatant (p < .05 vs. supernatants from both HUVEC spheroids and thymus MSC spheroids) (Fig. 6C).

Thymus MSCs Promote In Vivo Vascularization

We evaluated the function of spheroid angiogenic sprouts by implanting fibrin hydrogel constructs containing no cells or spheroids composed of HUVECs, thymus MSCs, and HUVECs plus thymus MSCs subcutaneously in NOD-SCID mice. On explant, control constructs did not appear to manifest any adjacent host reaction, whereas the constructs with spheroids made of thymus MSCs or HUVECs plus thymus MSCs incited an obvious host vascular response (Fig. 7A–7D).

Histologically, controls demonstrated cellular (but avascular) infiltrate from the surrounding tissue. No blood vessels were detectable within the control constructs, and the region adjacent to the construct did not appear to have increased vascularity (Fig. 7E).

Constructs with spheroids from all groups shared several characteristics. First, the central portions of the construct demonstrated "ghosts," or empty pockets, that were indicative of the prior locations of spheroids (Fig. 7F). There were also spheroids within the more interior portions of the constructs that manifested varying degrees of cell death, suggesting that the cells were subjected to ischemia (Fig. 7F). However, spheroids were generally present at the surface of the construct, suggesting that an adequate level of tissue partial pressure of oxygen was present to

sustain the survival of these cells. All spheroid groups had an increased cell number and vascularity adjacent to the construct. This increase in vascularity was specifically identified between the panniculus carnosus muscle layer and the more superficial aspect of the construct, whereas no blood vessels were identified deep in the construct (Fig. 7F). This finding suggests that implanted constructs attracted host blood vessels from more superficial (and vascular) regions of the mouse dermis. Any vascularization within a construct occurred at its surface, consistent with the findings of others [20].

Constructs composed of spheroids made of only HUVECs attracted host blood vessels adjacent to the construct but had few vessels penetrate the construct (Fig. 7F). Constructs composed of thymus MSC-only spheroids demonstrated increased cellularity and vascularity surrounding and penetrating these constructs (Fig. 7G). Constructs containing spheroids with both HUVECs and thymus MSCs also demonstrated an increase in adjacent tissue cellularity and vascularity and had the greatest vessel density within the construct (Fig. 7H, 7I). In some regions adjacent to constructs with thymus MSC-only or HUVEC plus thymus MSC spheroids, we noted significant erythrocyte extravasation (data not shown).

To determine the origin of the new blood vessels identified within the constructs, we performed human-specific CD31 staining. We found that only constructs with spheroids containing HUVECs plus thymus MSCs (n = 3) contained human-specific CD31 luminal structures that contained erythrocytes, suggesting



Figure 5. Thymus MSCs initiate the spheroid sprouting response in vitro. HUVECs and thymus MSCs were labeled with PKH26 and PKH67 prior to spheroid formation by overnight hanging drop culture and then embedded in fibrin hydrogel. Spheroids were then imaged with a confocal microscope at the indicated time points. As seen at 0 hour, spheroids with HUVECs and thymus MSCs formed core-shell bodies with MSCs forming the interior core of the spheroid. Spheroids manifested sprouting at 8 hours, with thymus MSCs composing the majority of sprouts and HUVECs colocalized to the bases of a subset of sprouts. At 24 hours, spheroids had more extensive branches, all composed of thymus MSCs and a subset containing HUVECs as well. Scale bars for 0- and 8-hour images = 50 μ m. Scale bars for 24-hour images = 100 μ m. Results are representative of three independent experiments. Abbreviations: HUVEC, human umbilical vein endothelial cell; MSC, mesenchymal stromal cell.

the formation of functional human vessels that integrated with the host's vasculature (Fig. 7J). Human CD31-positive vessel density from five sections of these HUVEC plus thymus MSC constructs was determined to be 20.9 \pm 10.6 vessels per square millimeter. We did not find any positive CD335 NK cells within or around day 7 explants (supplemental online Fig. 4).

DISCUSSION

One aim of this study was to isolate MSCs from discarded thymus tissue using a nonenzymatic, explant culture technique. Digestive enzymes may damage and/or alter cells and complicate the generation of a cell therapeutic product under good manufacturing practice conditions [21, 22]. Similar to our approach in this study with discarded thymus tissue, others have described the efficient isolation of adipose stem cells by explant culture of lipoaspirate tissue fragments [23–25]. It was not the purpose of our study to optimize MSC isolation from discarded thymus tissue using a manual fragmentation method, and this may explain the variability in the yield that we observed with different samples. However, compared with published results of the isolation of MSCs from adipose tissue, our method of isolating MSCs from thymus tissue appears to provide a comparable yield [26].

We have extended the prior characterization of MSCs from discarded neonatal human thymus tissue isolated by an enzymatic method [6, 7]. In this study, we demonstrated that explant culture-isolated thymus MSCs share many characteristics with bone marrow-derived MSCs. The prevalence of thymus MSC CFU-Fs is comparable to that determined for bone marrow and adipose tissue MSCs [27, 28]. The average thymus MSC doubling was also comparable to those of MSCs isolated from other tissues [29]. Various types of MSCs have been shown to express pluripotency genes [10–12]. We demonstrated that passage 5 and 6 thymus MSCs exhibited low levels of *Oct-4*, *Nanog*, and *Sox-2* gene expression and were also multipotent. Thymus MSCs shared several surface marker characteristics with bone marrow MSCs but also, like MSCs from other tissues, were distinct in that they lacked some of the markers that have been used to identify a subset of bone marrow MSCs such as STRO-1 [30–32]. Unlike amniotic fluid MSCs, neonatal thymus MSCs did not express SSEA-4 [32].

Organ and tissue regeneration requires growth of new tissue or repair of damaged tissue that contributes relevant, organspecific, physiological function. The regenerative capacity of human organs varies considerably. Although heart, brain, and kidneys appear to have limited regenerative potential after injury, the preadolescent thymus gland possesses an extraordinary ability to regenerate itself after injury and stress. Fundamental to any significant regenerative response of thick tissues and to the engineering of viable thick tissues is the requirement of angiogenesis and vasculogenesis; therefore, regenerative medicine strategies should include components that boost neovascularization [33–37].

We found that thymus MSCs possess proangiogenic characteristics in vitro and in vivo. Although bone marrow MSCs form a tubule network when cultured on Matrigel [38, 39], thymus MSCs tended to cluster together to form microcellular aggregates on fibrin hydrogel. This different behavior may be attributable to



Figure 6. Increased angiogenic gene and protein expression in thymus MSCs. HUVECs and thymus MSCs were cultured as monolayer or in spheroids for 48 hours prior to RNA isolation for quantitative polymerase chain reaction analysis. **(A)**: Fold change in expression for *HIF-1* α , *VEGFA*, and *bFGF* relative to HUVECs cultured in spheroids. In general, monolayer and spheroids with thymus MSCs had significantly increased expression of all studied genes. Culturing thymus MSCs in spheroids enhanced the expression of these angiogenic genes. For each gene, all comparisons between groups were significant by two-way analysis of variance (ANOVA) and post hoc Tukey test except those comparisons indicated by an asterisk. **(B, C)**: Levels of VEGFA and bFGF in supernatants from spheroid cultures in **(A)** were determined by enzyme-linked immunosorbent assay. VEGFA levels were increased in spheroids containing thymus MSCs, with no apparent effect by culturing with HUVECs. There was no difference in VEGFA levels between thymus MSCs. Results are representative of three independent experiments. Comparisons were performed by one-way ANOVA with post hoc Tukey test. Abbreviations: 2D, two-dimensional; 3D, three-dimensional; HUVEC, human umbilical vein endothelial cell; MSC, mesenchymal stromal cell.

the composition and biomechanical properties of the fibrin hydrogel [40]. When HUVECs and thymus MSCs were cultured together, a honeycomb tubule network rapidly developed on fibrin hydrogel, suggesting that the addition of HUVECs altered the clustering and contractile behavior of the thymus MSCs.

We also evaluated the angiogenic characteristics of thymus MSCs using an established 3D sprouting assay [41-43] because 3D culture is known to enhance the regenerative abilities of MSCs and more closely approximates the in vivo setting [44-47]. In multicellular spheroids, HUVECs and thymus MSCs are in a 3D, highcell-density aggregate during initial formation, facilitating direct contact and paracrine cell interactions. These cell interactions can be homotypic (MSC/MSC and EC/EC) or heterotypic (EC/ MSC). Although it is clear that thymus MSC spheroids had widespread upregulation of gene expression because of homotypic interactions, the slightly decreased upregulation in angiogenic gene expression in the combination of HUVEC plus thymus MSC spheroids suggested that HUVECs may temper this response (heterotypic interactions) or that this may be a dilutional effect of the HUVEC mRNA that decreases the thymus MSC mRNA signal during qPCR. The decrease in angiogenic gene expression caused by an increase in the ratio of HUVECs to thymus MSCs in spheroids is consistent with either explanation. Regulation of MSCs by endothelial cells in vasculogenesis and angiogenesis has not been delineated in detail and is an area of needed research.

Lee et al. described large core-shell bodies (5,000 cells per spheroid) composed of HUVECs and bone marrow MSCs, with MSCs forming the core and HUVECs forming the shell [48]. These investigators used a two-step process that took 48 hours to force a core-shell body arrangement, and the sprouting from these spheroids was seen first at day 3 after embedding in Matrigel. Our method used overnight hanging drop culture to generate spheroids that demonstrated long sprouts in fibrin hydrogel within 16 hours. The improved sprouting may have been secondary to the difference in angiogenic properties of thymus versus bone marrow MSCs, and allowing thymus MSCs and HUVECs to comingle with each other may have maximized contact and interactions between both cell populations before they sorted into a core-shell formation. The improved sprouting we observed may also be explained by the differences in surrounding matrix and their biophysical properties, which can affect the sprouting response [40, 49].

Recruited supporting cells stabilize nascent angiogenic sprouts [50–53]. Placing thymus MSCs and HUVECs together in spheroids eliminates the recruitment step, potentially explaining the rapid sprouting observed in the heterotypic cell combination



Figure 7. Thymus mesenchymal stromal cells (MSCs) incite angiogenesis in vivo. Fibrin constructs without spheroids (control) or with 500 spheroids with 600 human umbilical vein endothelial cells (HUVECs) per spheroid, 200 thymus MSCs per spheroid, or 600 HUVECs plus 200 thymus MSCs per spheroid were generated (n = 3 per group) and were implanted subcutaneously for 14 days in NOD-SCID mice. Explanted constructs were photographed (edges traced in A–D) and processed for histology. (A): Controls did not manifest local reaction. (B): HUVEC constructs appeared avascular. (C): Thymus MSC constructs were integrated and caused increased adjacent vascularization. (D): HUVEC plus thymus MSC constructs were integrated and surrounded by a host vascular response and appeared to have vessels within. (E–H): Construct with adjacent cellularity and vascularity between panniculus carnosus muscle layer (*) and construct. "Ghost" (†) of the prior locations of spheroid and necrotic spheroid (‡) were present in internal regions of all constructs with spheroids. (G): Thymus MSC constructs had minimal adjacent vascularization. Thymus MSC constructs promoted the greatest adjacent response, whereas HUVEC plus thymus MSC constructs had minimal adjacent vascularization. Thymus MSC construct. (J, K): Immunohistochemical staining with human-specific CD31 monoclonal antibody revealed that only constructs with HUVEC plus thymus MSCs constructs. (J, K): Immunohistochemical staining with blood cells. Scale bar = 20 μ m. Abbreviations: C, controls; H, human umbilical vein endothelial cell constructs; T, thymus mesenchymal stromal cell construct.

spheroids. It has been speculated that the relative availability of supporting cell-secreted growth factors and their diffusion distance from supporting cells to endothelial cells affects the robustness of tubule formation; therefore, minimizing the distance between endothelial and supporting cell, as in spheroids, may promote microvascularization [49]. In this study, we also found that thymus MSCs may lead the angiogenic sprouting response; this is a new angiogenesis and vasculogenesis paradigm because tip endothelial cells are thought to lead the formation of new sprouts [54–56].

Because of the small diameter of the spheroids that we used, which was well within the oxygen diffusion distance, hypoxia was not expected; however, $HIF-1\alpha$ gene expression was upregulated when thymus MSCs were cultured in spheroid form versus monolayer, suggesting a hypoxic response. One possible explanation of this finding is that the oxygen consumption rate for thymus MSCs may be high (or increased during spheroid culture) such that the hypoxia is present at the core of these relatively small spheroids. The combination of thymus MSCs and HUVECs in spheroids demonstrated a decrease in supernatant VEGFA levels. This may represent consumption or binding of VEGFA by HUVECs and/or negative regulation of MSCs by HUVECs and represents an area of future study.

The thymus gland involutes after infection, injury, and chemotherapy [57]. Resolution of thymic stress initiates a potent regenerative response in the thymus gland. Underlying the regenerative potential of the thymus gland is a brisk angiogenic response [4, 5]. This response has been investigated in rodents but has been poorly characterized in humans [4, 5]. Sepsis during the neonatal period has been demonstrated to cause involution [58]. CD248-positive MSCs significantly contributes to the angiogenic response of thymus regeneration in mice [5]. In this study, we demonstrated that thymus MSCs expressed CD248, a cellsurface receptor associated with cell adhesion and migration [59, 60]. CD248 has also been demonstrated to play a role in peripheral lymph node and splenic expansion and regeneration [61, 62]. The significance of CD248 in the injury and regeneration of other tissues has not been defined, although human bone marrow MSCs, rat adipose MSCs, and blood vessel-associated mural cells express CD248, and an anti-CD248 antibody has been shown to inhibit tube formation in bone marrow MSCs [63, 64].

We found that subcutaneously implanted constructs with homotypic or heterotypic multicellular spheroids attracted host dermal blood vessels, with thymus MSC spheroids inciting the greatest response; however, the combination of HUVECs plus thymus MSCs yielded the greatest vascularization within the construct. This indicates that the cooperation of both cell types promoted the angioinvasion of the adjacent host dermal blood vessels. Furthermore, only the combination of HUVECs plus thymus MSCs yielded the formation of functional human blood vessels, indicating that the importance of the interactions between the two cell types in vasculogenesis. An alternative explanation for the increase in surrounding vascular density is host NK cellmediated vascular changes because this immune cell is intact in NOD-SCID mice. We performed immunohistochemistry on earlier (day 7) explants to determine the presence of NK cells (CD335) but did not identify any within or around the construct, making NK cell-mediated increase in vascular density less likely.

Although MSCs, with or without endothelial cells, have already been used by others to promote vascularization of implanted constructs and ischemic tissues in both preclinical and clinical settings [47, 65–69], the optimal MSC yielding the maximal therapeutic angiogenic response has yet to be determined. Results presented in this study and elsewhere suggest that thymus MSCs may be a promising candidate for therapeutic use [6, 7]. Given the low expression human leukocyte antigen class I on a subset of neonatal human thymus MSCs, as shown in this study and elsewhere [7, 70], use of these MSCs may also yield advantages in the allogeneic setting (e.g., use in older patients with ischemic heart disease), in which it is more likely to complement the advantages of neonatal MSCs over adult MSCs [71–73]. However, improvements in isolation yield are needed to make clinical use of thymus MSCs feasible.

Other sources of MSCs and stem cells that may have therapeutic potential in neonates with heart disease include amniotic fluid, placenta, umbilical cord, and Wharton's jelly [32, 74–78]. MSCs have also been isolated from aborted fetuses [79]. Similar to human neonatal thymus MSCs, fetal MSCs also had trilineage differentiation capacity. The doubling time for fetal MSCs was slightly less than what we observed in human thymus MSCs, but the CFE was greater in human neonatal thymus MSCs. Like neonatal thymus MSCs, fetal MSCs express pluripotency-related genes such as *Oct-4* and *Nanog* [80, 81]. Full characterization and detailed comparison in relevant disease models are needed to determine which MSC type will yield the optimal therapeutic benefit.

CONCLUSION

MSCs are present in large numbers in discarded neonatal and infant thymus tissue and can be isolated by an explant culture method. Explant culture-isolated thymus MSCs demonstrate the classical characteristics of bone marrow MSCs. Thymus MSCs also possess proangiogenic qualities and deserve further consideration as a potential cell therapeutic agent to promote tissue regeneration and repair and for use in strategies to vascularize engineered tissues.

ACKNOWLEDGMENTS

This work was supported by the University of Michigan, Department of Cardiac Surgery. We thank Kaihong Wu for his technical assistance with cell culture.

AUTHOR CONTRIBUTIONS

S.W.: conception and design, collection and/or assembly of data, data analysis and interpretation, manuscript writing; L.M.: collection and/or assembly of data, data analysis and interpretation, manuscript writing; S.J.: collection and/or assembly of data; J.W. and R.W.: collection and/or assembly of data, data analysis and interpretation; R.G.O.: provision of study material or patients, manuscript writing, financial support; M.-S.S.: conception and design, provision of study material or patients, collection and/or assembly of data, name data analysis and interpretation; R.G.O.: provision of study material or patients, manuscript writing, financial support; M.-S.S.: conception and design, provision of study material or patients, collection and/or assembly of data, data analysis and interpretation, manuscript writing, final approval of manuscript.

DISCLOSURE OF POTENTIAL CONFLICTS OF INTEREST

The authors indicated no potential conflicts of interest.

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Characterization and Angiogenic Potential of Human Neonatal and Infant Thymus Mesenchymal Stromal Cells

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Stem Cells Trans Med 2015, 4:339-350. doi: 10.5966/sctm.2014-0240 originally published online February 23, 2015

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