Differentiation of dermis-derived multipotent cells into insulin-producing pancreatic cells in vitro

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AIM: To observe the plasticity of whether dermis-derived multipotent cells can differentiate into insulin-producing pancreatic cells in vitro.

METHODS: A clonal population of dermis-derived multipotent stem cells (DMCs) from newborn rat with the capacity to produce osteocytes, chondrocytes, adipocytes and neurons was used. The gene expression of cultured DMCs was assessed by RNA microarray using rat RGU34A gene expression probe arrays. DMCs were further cultured in the presence of insulin complex components (Insulin-transferrin-selenium, ITS) to observe whether DMCs could be induced into insulin-producing pancreatic cells in vitro.

RESULTS: DNA microarray analysis showed that cultured DMCs simultaneously expressed several genes associated with pancreatic cell, neural cell, epithelial cell and hepatocyte, widening its transcriptomic repertoire. When cultured in specific induction medium containing ITS for pancreatic differentiation, DMCs were shown to produce insulin-positive cells detected by immunohistochemistry.

CONCLUSION: Our data indicate that dermal multipotent cells may serve as a source of stem/progenitor cells for insulin-producing pancreatic cells.

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INTRODUCTION

Organ-specific stem cells possess plasticity that can permit differentiation along new lineages and have significant implications for future therapies of diabetes. The production of endocrine pancreas and insulin-secreting beta cells from adult nonpancreatic stem cells and hepatic oval stem cell has been demonstrated[12]. Zulewski et al. also showed that pancreatic islets contained a heretofore unrecognized distinct population of cells that expressed the neural stem cell-specific marker nestin. The nestin-positive islet-derived progenitor (NIP) cells are a distinct population of cells that reside within pancreatic islets and may participate in the neogenesis of islet endocrine cells[2]. According to this result, liver stem cells have been proved to differentiate into pancreatic islet-like cells[3]. Dermis is a highly accessible tissue source for adult stem cells. Nestin-positive skin-derived stem cells have been isolated from dermis by Toma et al[4]. In our previous study, we isolated a clonal population of dermal multipotent cells by their adherence to tissue culture plastic (termed as plastic adherent DMCs) from newborn rat dermis. These cells showed the differentiation capacity to produce nestin-positive cells[5]. In this study, we aimed to investigate whether plastic adherent DMCs had the differentiation capacity to produce insulin-producing pancreatic cells in vitro.

MATERIALS AND METHODS

Materials

A clonal population of dermis-derived multipotent stem cells (DMCs) was used in the experiments. The clonal population of DMCs were isolated and identified from primary dermal cells of male newborn Wistar rat as previously described[5]. All tissue culture reagents, including Iscove’s Modified Dulbecco’s Medium (IMDM), ITS (insulin-transferrin-selenium), epidermal growth factor, basic fibroblast growth factor, were purchased from Sigma (St. Louis, CA, USA). Fetal bovine serum (FBS) was purchased from Hyclone (Logan, UT, USA). The rat RGU34A gene expression probe arrays were purchased from Affymetrix (Santa Clara, CA). Mouse anti-insulin monoclonal antibody was purchased from Sigma (St. Louis, CA, USA). Horseradish peroxidase (HRP)-labeled goat anti-mouse IgG antibody was purchased from Boster (Wuhan, China).

Methods

Cell culture The DMCs were cultured in IMDM medium containing 10 mL/L fetal bovine serum and 100 U/mL penicillin and 10 µg/mL streptomycin. Cultures were maintained at 37 °C in a humidified atmosphere containing 5% CO₂. For differentiation induction, DMCs were transferred into specific inducing medium containing 10 µg/mL of keratocyte growth factor (KGF), 20 ng/mL of epidermal growth factor (EGF), 10 mmol/L of nicotinamide, and 1 mg/mL of ITS. The differentiated cells were examined for the expression of insulin by immunohistochemistry.

DNA Microarray analysis Total RNA from cultured DMCs was isolated using QIAGEN’s RNeasy total RNA isolation kit (Rneasy, Qiagen) following the manufacturer’s instructions, and quantified. Transcript profiling was conducted by means of rat RGU34A gene expression probe arrays, containing 8 799 probe sets, interrogating primarily annotated genes. The experiment was conducted according to the recommendations of the manufacturer (Affymetrix GeneChip expression analysis manual). The resulting “data-file” was processed further using the microarray analysis suite 5 software package (Affymetrix). According to the statistical expression analysis algorithm, the presence of a gene within a given sample was determined at a detection P value of <0.05 and was graded as absent (A), marginal (M), or present/positive (P).

Immunocytochemistry Immunocytochemistry study with peroxidase-labeled streptavidin biotin method was performed to detect the expression of insulin in differentiated cells. Before
detection, the cells were washed three times for 20 min with phosphate-buffered saline (PBS, PH = 7.2) to exclude the pollution of exogenous insulin in the culture medium. After washing, the cells were blocked for 30 min with 100 mL/L normal goat serum in PBS and were incubated with mouse anti-insulin antibody (1:100) at 4 °C for 24 h. Incubation at room temperature with anti-mouse secondary antibody and avidin-biotinylated peroxidase complexes was performed for 2 h. The specimens were washed for 15 min with 0.01 mol/L PBS between all steps. The reaction product was developed with 0.5 g/L 3'3-diaminobenzine tetrahydrochloride (DAB). After staining, the result was observed microscopically. PBS was used as a substitute for primary antibody as negative control.

RESULTS

Biological properties of DMCs

Most adherent DMCs were of spindle-shaped cells (Figure 1). Phenotype analysis showed that DMCs were negative for some lineage-specific surface markers, including pan-cytokeratin, cytokeratin19, factor VIII, CD31, CD45, CD34, α-smooth muscle actin (α-SMA), desmin, collagen II and nestin, but were positive for CD59, CD90, CD44, vascular cell adhesion molecule-1 (VCAM-1) and intercellular adhesion molecule-1 (ICAM-1). The doubling time of cultured DMCs was about 40 h. Following induction, DMCs had the capacity to differentiate into cells with phenotypic characteristics of osteocytes (alkaline phosphatase activity and alizarin red staining), adipocytes (oil red staining), chondrocytes (collagen II and Alcian blue staining) and neurons (nestin and NF-200 staining) in specific induction media.

Figure 1 Morphology of cultured DMCs. (Inverted microscope, ×100).

Gene expression profile of plastic-adherent DMCs

The gene expression of DMCs was assessed by DNA microarray analysis and the harbinger of a potential cell type was viewed by simultaneous expression of a panel of lineage-related genes. The result revealed that the cultured DMCs simultaneously expressed genes associated with pancreatic cells (Table 1). DMCs also simultaneously expressed transcripts for epithelial cell, neural cell, and hepatocyte as well (Table 1). These transcripts were developmentally related with pancreatic tissue and further widened the transcriptomic repertoire of DMCs. To confirm the data by DNA microarray, several genes were further tested by RT-PCR and the result was consistent with the result from microarray (data not shown).

Figure 2 Differentiation of DMCs into insulin-producing cells. A: After induction in the medium containing ITS for 4 wk, cells were positive for insulin; B: Negative control (SP, ×200).

Table 1 DMC gene lists (partial) associated with diverse cellular lineages

| Cell lineage       | Representative examples of associated genes                                                                                                                                 |
|--------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| A: Pancreatic cell | Pancreatic eukaryotic initiation factor 2 alpha-subunit kinase (PEK), pancreatic secretary trypsin inhibitor-like protein (PSTI), pancreatic islet cDNA Rattus norvegicus cDNA similar to rapamycin-binding protein FKBP13, insulin II gene, IRS-1 mRNA for insulin-receptor, mRNA for glucose-dependent insulinotrophic polypeptide |
| B: Neural cell     | Neural visinin-like Ca2+-binding protein type 3, neuron-specific enolase, neuronal nitric oxide synthase, GABA receptor-associated proteins, NCAM, brain-derived neurotrophic factor (BDNF), syntaxin 7, glial fibrillary acidic protein (GFAP), adrenomedullin, myelin protein SR13 (growth-arrest-specific Gas-3 homolog), neurotomin; secretogranin II, synapse-associated protein 102, latentin, kexin-like protease PC7A, N-methyl-D-aspartate (NMDA) receptor |
| C: Epithelial cell | Epithelial membrane protein 1, epithelial cell transmembrane protein antigen precursor, epidermal growth factor precursor, heparin-binding EGF-like growth factor, mucin, parathyromosin and thy mosin 4 and 10 |
| D: Hepatocyte      | Liver glycogen phosphorylase enzyme, liver IL-6 receptor ligand binding chain, liver specific transcription factor L-F-B, liver nuclear protein P47, liver al-Fucosidase, hepatocyte nuclear factor 3α (HNF-3beta) |

Expression of a single gene can’t define the phenotype of a particular cell type. Simultaneous expression of a panel of lineage-related genes in single isolated cell was viewed as the harbinger of a potential cell type. Representative examples of genes corresponding to each cell lineage are outlined in the above table.
**Differentiation of DMCs into insulin-producing pancreatic cells**

DMCs were further tested for their capacity to produce pancreatic cells in the inducing medium containing ITS. The morphology of DMCs were changed into epithelial cells-like cells when cultured in medium containing ITS for 2 wk and immunohistochemistry showed that a proportion of cells (less than 10%) were positive for insulin when cultured in medium containing ITS for 4 wk (Figure 2). The negative control cells showed no positive staining for insulin. This result further indicated that DMCs could undergo differentiation to form insulin-producing cells.

**DISCUSSION**

Replacement of the insulin-producing pancreatic islet ßcells represents the ultimate treatment for type I diabetes. Recent advances in islet transplantation underscore the urgent need for developing alternatives to human tissue donors, which are scarce. The generation of insulin-producing cells from adult stem cells is one possible approach. Recent studies have suggested that there are closely developmental relationship between the nestin positive progenitors and the pancreatic cells. In addition to liver stem cells and neural stem cells, multipotent adult stem cells from the dermis of mammalian skin. Nat Cell Biol 2001; 3: 778-784.

Nevertheless, the efficiency of adult stem cells to differentiate into insulin-producing cells in vitro is low at present and expansion with large scale is needed for application. Genetic manipulation in tissue culture may be a choice and it is possible that new insights into endocrine pancreas development will lead to the manipulation of progenitor cell fate towards the ß cell phenotype of insulin production, storage and regulated secretion. If successful, these approaches could lead to widespread cell replacement therapy for type I diabetes.

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