Collecting sufficient quantities of primary neural crest cells (NCCs) for experiments is difficult, as NCCs are embryonic transient tissue that basically does not proliferate. We successfully induced NCCs from human induced pluripotent stem cells (iPSCs) in accordance with a previously described method with some modifications. The protocol used in this study efficiently produced large amounts of iPSC-derived NCCs (iPSC-NCCs). Many researchers have recently produced large amounts of iPSC-NCCs and used these to examine the physiological properties, such as migratory activity, and the potential for medical uses such as wound healing. Immunological properties of NCCs are yet to be reported. Therefore, the purpose of this study was to assess the immunological properties of human iPSC-NCCs. Our current study showed that iPSC-NCCs were hypoimmunogenic and had immunosuppressive properties in vitro. Expression of HLA class I molecules on iPSC-NCCs was lower than that observed for iPSCs, and there was no expression of HLA class II and costimulatory molecules on the cells. With regard to the immunosuppressive properties, iPSC-NCCs greatly inhibited T cell activation (cell proliferation and production of inflammatory cytokines) after stimulation. iPSC-NCCs constitutively expressed membrane-bound TGF-β, and TGF-β produced by iPSC-NCCs played a critical role in T cell suppression. Thus, cultured human NCCs can fully suppress T cell activation in vitro. This study may contribute to the realization of using stem cell-derived NCCs in cell-based medicine.

Keywords: eye, neural crest cells, immunosuppression, T cells

Introduction

Neural crest cells (NCCs) are vertebrate-specific, transient, multipotent migratory stem cells that give rise to a wide range of cell and tissue types throughout the body [1]. It has been reported that corneal endothelium [2], gingivae [3,4], oral mucosal progenitor cells [5], and part of the mesenchymal stem cells (MSCs) [6] are derived from NCCs and exhibit immunosuppressive properties in vitro. However, corneal endothelial cells also express PD-L1 (B7-H1) and TGF-β, which inhibit T cell proliferation and induce regulatory T cells [7,8]. Gingivae express the Fas ligand, which then induces T cell apoptosis [3]. Oral mucosal progenitor cells can inhibit lymphocyte proliferation in mixed lymphocyte cultures [5]. Numerous studies have illustrated the ability of human MSCs to inhibit T cell proliferation in vitro and promote the expansion of regulatory T cells [9–13]. Moreover, MSC infusion has been shown to be an effective treatment of steroid-refractory acute graft-versus-host disease after bone marrow transplantation [14,15] and is now being used in clinics. Human induced pluripotent stem cell (iPSC)-derived MSCs also have immunosuppressive properties and can prolong pancreatic islet survival after islet transplantation in streptozocin-induced diabetic mice [16]. Thus, NCC-derived differentiated cells/tissues exhibit immunosuppressive properties.

Previous reports have described NCC induction protocols from human iPSCs [17–20]. NCCs derived from iPSCs (iPSC-NCCs) were reported to have a potential use in wound healing of diabetic neuropathy [21] and tendon repair [22] in vivo. However, there are yet to be any studies that have investigated the interaction of NCCs with specific immune cell subtypes. It was reported that iPSC-NCCs were effective in treating congenital aganglionic megacolon (Hirschsprung’s disease) [23,24]. Human iPSC-NCCs have been shown to migrate and differentiate into functional neurons in aganglionic human gut tissue in vitro [23]. Human iPSC-NCCs restore mature ganglia and neurons in human intestinal organoid-derived tissue-engineered small
IMMUNOSUPPRESSION OF IPS-NEURAL CREST CELLS

INTRODUCTION

Li et al. suggested that iPSC-NCCs may be an attractive tool that can be used for studying the pathogenesis of gastrointestinal disorders, as they represent a potentially ideal cell source for enteric neural transplantation treatments [23]. The multipotency of the NCCs suggests a promising role for iPSC-NCCs as a source for cell-based therapy. When the iPSC-NCCs are clinically applied, it is important to investigate their immunological properties.

Therefore, the purpose of this study was to assess the immunological properties of iPSC-NCCs in vitro. After successfully inducing NCCs from human iPSCs in accordance with a previous method with some modifications [25,26], we used the iPSC-NCCs as effectors and immune cells as targets in vitro to evaluate the immunosuppression by NCCs.

MATERIALS AND METHODS

Culture of iPSCs and the induction of NCCs

Human iPSC lines 253G1 and 201B7, which are generated from adult skin fibroblasts [27,28], were obtained from the Riken BioResource Center (Ibaraki, Japan). The iPSCs were cultured under feeder-free and serum-free conditions with AK03N medium (Ajinomoto, Tokyo, Japan) in six-well culture plates coated with iMatrix-511 (Nippi, Tokyo, Japan) as previously described [29]. The iPSCs were subcultured once a week, with the medium changed daily. The iPSCs were differentiated into NCCs according to a previously described method [25,26] with some modifications. Briefly, after the undifferentiated iPSCs reached confluence in culture, the iPSCs were dissociated with Accutase (Millipore, Temecula, CA) and seeded first on a 3-D culture plate (EZSPHERE®; AGC Techno Glass, Shizuoka, Japan) [30] at a density of 2×10⁵ cells/mL in an embryoid body (EB) induction medium, which was composed of DMEM/F12 (Sigma-Aldrich, St. Louis, MO) supplemented with 1% N-2 (Gibco, Grand Island, NY), 2% MACS® NeuroBrew-21 (Miltenyi Biotec, Auburn, CA), 2 mM t-glutamine (Sigma-Aldrich), 20 ng/mL recombinant human epidermal growth factor (EGF: PeproTech, Rocky Hill, NJ), 1% penicillin/streptomycin (Gibco), and 10 μM Y-27632 (Wako, Osaka, Japan). The spent medium was changed once on EB culture day 4, with uniformly sized EBs forming in 7 days. Subsequently, the EBs were replated on a noncoated 10 cm dish in NCC induction medium, DMEM/F12 with 1% N-2, 2 mM t-glutamine, 1% penicillin/streptomycin, 20 ng/mL EGF, 20 ng/mL recombinant human fibroblast growth factor-basic (bFGF: Wako), and 5 μg/mL heparin (Sigma-Aldrich). The culture dish was left untouched for the first 3 days to allow the EBs to attach to the culture dish and the induced NCCs to migrate from the attached EBs, after which the spent medium was changed on days 4, 8, 11, 13, and 14 in conjunction with the replating of the EBs. NCC induction was completed by day 14. The iPSC-NCCs were used for subsequent experiments.

This study was performed in accordance with ethics protocols approved by the Keio University Ethics Committee and the Institutional Review Board at RIKEN Center for Developmental Biology. Informed consent was obtained from all healthy volunteers.

Flow cytometric analysis

After washing, iPSC-NCCs and iPSCs were harvested and resuspended at a density of 5×10⁵ cells/mL. Specimens were then incubated in PE, FITC, or PE/Cy5 conjugated primary antibody or PI solution (Dojindo, Kumamoto, Japan; catalog #341-07881) at 4°C for 30 min and washed thrice. Flow cytometric analysis was performed using the CantoTM II flow cytometer (BD Biosciences, San Jose, CA), with the data analyzed using the FlowJo software (Tree Star, Ashland, OR). The primary antibodies included PE-conjugated anti-human SSEA-4 (eBioscience, San Diego, CA; catalog #12-8843), PE-conjugated anti-human CD271 (BioLegend, San Diego, CA; catalog #345106), PE/Cy5-conjugated anti-human CD49d (BioLegend; catalog #304306), FITC-conjugated anti-human HLA class I (Sigma-Aldrich; catalog #F5662), PE-conjugated anti-human β2-microglobulin (BioLegend; catalog #316306), FITC-conjugated anti-human HLA-DR, DP, DQ (BD Biosciences; catalog #555558), FITC-conjugated anti-human CD80 (B7-1: eBioscience; catalog #11-0809), PE-conjugated anti-human CD86 (B7-2: eBioscience; catalog #12-0862), PE-conjugated anti-human CD274 (PD-L1: BioLegend; catalog #12-5983), PE-conjugated anti-human CD273 (PD-L2: BD Pharmingen, San Jose, CA; catalog #558066), and the appropriate isotype-matched controls. Expression of TGF-β2 in iPSC-NCCs and iPSCs (control) was assessed by flow cytometry. Cell-surface staining (no intracellular staining) was performed for the expression of TGF-β2 using these cells. After human Fc block staining, these cells were stained with anti-human TGF-β2 abs (Abcam, Cambridge, United Kingdom; catalog #ab66045) at 4°C for 30 min. iPSC-NCCs were then stained with Alexa Fluor-488-labeled anti-rabbit IgG (2nd abs: Life Technologies, Carlsbad, CA; catalog #A11008) at 4°C for 30 min, and the cells were evaluated by fluorescence-activated cell sorting (FACS) analysis.

Immunohistochemistry

Cells were fixed with methanol for 15 min at −30°C, permeabilized with 0.2% Triton X-100-PBS (Sigma-Aldrich) at room temperature for 15 min, and blocked in a solution of PBS containing 1% bovine serum albumin (Sigma-Aldrich), followed by an overnight incubation in primary antibody solutions at 4°C. After three washes in 0.05% Tween-20-PBS (Sigma-Aldrich), cells were incubated with Alexa fluoresecently conjugated secondary antibodies and 5 μg/mL 4′,6-diamidino-2-phenylindole dihydrochloride (DAPI: Invitrogen, Carlsbad, CA) at room temperature for another 60 min. After three washes in PBS, labeled cells were imaged with a fluorescence microscope (Olympus, Tokyo, Japan).

Primary antibodies and their working dilutions were as follows: mouse anti-human nerve growth factor receptor (NGFR: Advanced Targeting Systems, San Diego, CA; catalog #AB-N07, 1:200 dilution) and mouse anti-human transcription factor AP-2 alpha (TFAP2A: Developmental Studies Hybridoma Bank, Iowa City, IA; catalog #3B5, 1:200 dilution). The secondary antibodies used were the corresponding Alexa-fluorescent labeled antibodies (Life Technologies, catalog #341-07881) at 4°C for 30 min and washed thrice. Immunohistochemistry was performed using the High Pure RNA Isolation Kit (Roche, Basel, Switzerland) appropriate isotype-matched controls. Expression of TGF-β2 on iPSC-NCCs and iPSCs (control) were also assessed by immunohistochemistry using the same antibodies as those in the FACS analysis.

Quantitative real-time polymerase chain reaction

Total RNA from iPSC-NCCs and control iPSCs were purified using the High Pure RNA Isolation Kit (Roche, Basel, Switzerland).
TGF-β10, Master (Roche) was performed using the LightCycler chain reaction (qRT-PCR) with the LightCycler 480 Probe manufacturer’s instructions. Quantitative real-time polymerase for single-standard cDNA using the Transcriptor First Strand cDNA Synthesis Kit (Roche) in accordance with the manufacturer’s instructions. Quantitative real-time polymerase chain reaction (qRT-PCR) with the LightCycler® 480 Instrument II (Roche). Table 1 lists the primer sequences. The qRT-PCR was performed by denaturation at 95°C for 10 min, followed by 45 cycles of denaturation at 95°C for 10 s, annealing at 60°C for 30 s, and extension at 72°C for 1 s. All amplifications were performed in triplicate, with β-actin used as a normalizer of the RNA content. A negative control with no cDNA was also used for each analysis.

### Adipogenic, chondrogenic, and osteogenic differentiation

To validate whether iPSC-NCCs maintain multipotency, iPSC-NCCs were plated at a density of 5 × 10^5 cells in 35 mm dishes. Cells were fed every 2 days by completely replacing the medium with NCC induction medium (3–5 days). At 100% confluence, we performed adipogenic differentiation by treating the iPSC-NCCs with hMSC Adipogenic BulletKit (Lonza, Basel, Switzerland; catalog #PT-3004) for 3 weeks according to the manufacturer’s protocol, followed by staining with oil red O (Muto Pure Chemicals, Tokyo, Japan; catalog #40491) to detect lipids. A pellet culture system was used to evaluate the capacity of iPSC-NCCs to differentiate into chondrocytes. iPSC-NCCs were applied to a pellet at a density of 5 × 10^5 cells and cultured in hMSC Chondro BulletKit (Lonza; catalog #PT-3003). The medium was changed every 3–4 days. After 4 weeks in culture, the induced cartilage tissue was fixed with 4% formaldehyde, sliced, and stained with 1% Alcian Blue (Amresco, Solon, OH; catalog #0298). To investigate the osteogenic differentiation capacity of iPSC-NCCs, confluent iPSC-NCCs in 35 mm dishes were treated with hMSC Osteogenic BulletKit (Lonza; catalog #PT-3002). The medium was changed every 2–3 days. After 2–3 weeks, the cells were stained with Alizarin Red S (Nacalai Tesque, Kyoto, Japan; catalog #01303-52), rinsed with water, and examined by microscopy.

### T cell-NCC assay of cell proliferation

Human peripheral blood mononuclear cells (PBMCs) were obtained from the blood of five healthy donors by Lymphoprep (Stemcell Technologies, Vancouver, BC) density gradient centrifugation in accordance with the manufacturer’s instructions. PBMCs were incubated in RPMI 1640 (Nacalai Tesque) containing 10% FBS supplemented with human recombinant interleukin-2 (rIL-2; BD Biosciences). To assess immunosuppressive effects of iPSC-NCCs in vitro, we used a mixed lymphocyte reaction (MLR) test, with iPSCs used as a negative control and iPSC-derived retinal pigment epithelial (iPSC-RPE) cells used as a positive control. iPSC-RPE cells were established in accordance with a previously reported method [31,32]. PBMCs were categorized into four groups as follows: PBMC monoculture (PBMC only: 3 × 10^6 cells/well), PBMC coculture with iPSC-NCCs (1 × 10^6 cells/well), PBMC coculture with iPSCs, and PBMC coculture with iPSC-RPE cells. After 120 h of incubation, PBMCs were collected and analyzed by flow cytometry of the PBMC fraction. In the T cell suppression assay with iPSC-NCCs, anti-human CD3 antibody (1 μg/mL, Ancell, Bayport, MN; catalog #144-020) and anti-human CD28 antibody (1 μg/mL, BD Pharmingen; catalog #555725) were used for the T cell stimulation. After 72 h of incubation, PBMCs were collected and analyzed by flow cytometry.

For the TGF-β block assay, we used TGF-β RI Kinase Inhibitor VI (SB431542; Sigma-Aldrich; catalog #S4317). After adding 1, 5, and 10 μM of SB431542 to the PBMC coculture with iPSC-NCCs, PBMCs were then incubated in RPMI1640 medium plus rIL-2 and stimulated with anti-CD3/CD28 antibodies. PBMCs were categorized into three groups as follows: PBMC monoculture, PBMC coculture with iPSC-NCCs, and PBMC coculture with iPSC-RPE cells plus SB431542.

For the Ki-67 FACS analysis, PBMCs were fixed and permeabilized with 70% ethanol at −30°C for 1 h, followed by washing thrice using cell staining buffer (BioLegend). Subsequently, the cells were intracellularly stained with PE-conjugated anti-Ki-67 antibody (BioLegend; catalog #350504), APC-conjugated anti-CD3 (BioLegend; catalog #300412), APC-conjugated anti-CD4 (eBioscience; catalog #17-0049), anti-CD8 (eBioscience; catalog #17-0088), anti-CD11b (Miltenyi Biotec; catalog #130-098-088), or anti-CD56 antibodies (BioLegend; catalog #304604).

### Transwell culture assays

Transwell chambers with a 0.3 μm pore size membrane (Corning Costar, Cambridge, MA) were used to separate the

<table>
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lymphocytes physically from the iPSC-NCCs. In the transwell assay, PBMCs \((1.0 \times 10^6 \text{ cells/well})\) were added to the upper wells of a transwell plate. PBMCs were cultured in the presence or absence of iPSC-NCCs \((1.0 \times 10^5 \text{ cells/well})\). Cultures were plated in triplicate and incubated for 3 days. After incubation, PBMCs were collected and stained with PE-conjugated anti-Ki-67 antibody and APC-conjugated anti-CD3 antibody. T cell proliferation (Ki-67 FACS) was assessed by flow cytometry.

**Cytokine measurements**

Concentrations of IFN-\(\gamma\) in the supernatant of the PBMC monoculture, PBMCs cocultured with iPSCs, and iPSC-NCCs were examined using ELISA in accordance with the manufacturer’s manual (R&D Systems, Minneapolis, MN). Each result was evaluated statistically, with \(P<0.05\) defined as being significant. In addition, concentrations of IFN-\(\gamma\) in the supernatant of the PBMC monoculture, PBMC coculture with iPSC-NCCs, and PBMC coculture with iPSC-NCCs plus SB431542 were also examined.

For the FACS analysis, double staining of CD4 and IFN-\(\gamma\) was performed to determine the proportion of the CD4\(^+\) IFN-\(\gamma\) expressing T cells. Intracellular staining was performed to determine the expression of IFN-\(\gamma\). Before staining for IFN-\(\gamma\), PBMCs were prestimulated with T cell stimulation materials (CytoStim; Miltenyi Biotec) for 1 h, and then cells were incubated for an additional 4 h in the presence of a protein transport inhibitor (BD GolgiStop\textsuperscript{TM}; BD Biosciences). Cells were then treated with an intracellular staining material (BD Cytofix/Cytoperm Kits, BD Pharmingen). After human Fc block staining, these T cells were stained with APC-labeled anti-human CD4 ab (Miltenyi Biotec) and PE-labeled anti-human IFN-\(\gamma\) (R&D Systems) at 4\(^\circ\)C for 30 min. PBMCs were also stained with PE-labeled isotype control abs (mouse IgG) at 4\(^\circ\)C for 30 min.

**DNA microarray**

For comparing the gene expression between iPSCs and iPSC-NCCs, after the cells were harvested, the total RNA of iPSCs (\(n=2\)) and iPSC-NCCs (\(n=2\)) was extracted using a High Pure RNA Isolation Kit (Roche). Whole Human Genome Oligo Microarray (4*44K, Agilent Technologies, Santa Clara, CA) analysis was then performed by Filgen (Aichi, Japan). Briefly, RNA quantity and quality were measured by NanoDrop ND-1000 (NanoDrop Technologies, Santa Clara, CA). RNA integrity was assessed by standard denaturing agarose gel electrophoresis. Sample labeling and array hybridization were performed according to the Agilent One-Color Microarray-Based Gene Expression Analysis protocol. We registered these data in the GEO database (Accession No. GSE109136).

Next, the gene expressions between inflamed and normal conditions of iPSC-NCCs were compared. For the inflamed condition, iPSC-NCCs \((1.0 \times 10^6 \text{ cells/well})\) were incubated in the presence of PBMC mix (MLR; \(3.0 \times 10^6 \text{ cells/well}\)) in a transwell culture plate. To avoid contamination with iPSC-NCCs, PBMCs were incubated in an upper well of the transwell plate. For the normal condition, iPSC-NCCs were incubated in the absence of PBMCs. For both conditions, iPSC-NCCs were incubated in RPMI1640 medium. After incubation for 120 h, the total RNA of iPSC-NCCs (inflamed condition, \(n=2\)) and iPSC-NCCs (normal condition, \(n=2\)) was extracted. The procedures were performed as previously described. We also registered these data in the GEO database (Accession No. GSE116110).

**Statistical analysis**

All experiments were repeated at least thrice, and data are expressed as the mean± standard deviation (SD). A two-tailed \(t\)-test was used for all of the statistical analyses. Results were considered significant when \(P<0.05\).

**Results**

**Differentiation and identification of iPSC-derived NCCs**

In the first step, we established human iPSC-derived NCCs. iPSC-NCCs were successfully induced from human iPSCs after 3 weeks of differentiation. During the differentiation, the cells gradually changed into a dendritic morphology (Fig. 1A). Flow cytometry staining revealed that the iPSCs were positive for stage-specific embryonic antigen-4 (SSEA4), which is a pluripotency marker, while the iPSC-NCCs were negative for SSEA4 (Fig. 1B). iPSC-NCCs highly expressed CD271 (NGFR) and CD49d (both NCC markers), compared with iPSCs (Fig. 1C). iPSCs expressed only CD271 and not CD49d. Immunofluorescence staining revealed that the iPSC-NCCs were positive for neural crest markers, NGFR and TFAP2A (Fig. 1D). In addition, qRT-PCR showed that iPSC-NCCs expressed low levels of the pluripotency markers, Nanog and Lin28A. The expression of the neural crest markers, NGFR, snail family transcriptional repressor 2 (SNAI2), sex determining region Y-box 10 (SOX10), and TFAP2A, were higher in the iPSC-NCCs versus the iPSCs (Fig. 1E).

In addition, we successfully induced adipocytes, chondrocytes, and osteocytes from iPSC-NCCs (Fig. 1F). We validated the NCC induction protocol in two different iPSC cell lines (201B7, 253G1), with the results being the same in both cell lines.

**Expression of HLA and costimulatory molecules on iPSC-NCCs**

To confirm the immunogenicity in iPSC-NCCs, we examined the expression of HLA and costimulatory molecules on the iPSC-NCCs. At first, we confirmed that analyzed cells (iPSCs and iPSC-NCCs) were >99% negative for PI (which stains the dead cells). Our results showed that iPSC-NCCs exhibited a lower expression of HLA class I, but had no expression of HLA class II (Fig. 2A). Similarly, the iPSC-NCCs had a lower expression of \(\beta2\)-microglobulin, which forms HLA class I, compared to the iPSCs (Fig. 2A). Furthermore, there was a statistically significant difference between the iPSCs and iPSC-NCCs (Fig. 2B). In addition, we found no difference in the costimulatory molecule expression between the iPSCs and iPSC-NCCs (Fig. 2C). For example, iPSC-NCCs, as well as iPSCs, did not express CD80 (B7-1), CD86 (B7-2), CD274 (PD-L1), or CD273 (PD-L2) on their surface. In contrast, monocytes (positive control: from a healthy donor) clearly expressed these...
FIG. 1. Differentiation of NCCs from human iPS cells. (A) Bright-field image of iPS-NCCs. Scale bars, 100 µm. (B) Expression of SSEA4 in iPSCs and iPS-NCCs by flow cytometry analysis. iPSCs are positive, and iPS-NCCs are negative for SSEA4. Red histogram; isotype control. (C) In the FACS analysis, iPSCs are negative for CD49d and slightly positive for CD271 expression. iPS-NCCs are clearly positive for CD49d and CD271. Numbers in the FACS histograms indicate double positive cells. (D) Expression of NCC marker NGFR and TFAP2A. Immunocytochemistry analysis shows that iPS-NCCs are positive for NGFR and TFAP2A. Cell nuclei were counterstained with DAPI. Scale bars, 100 µm. (E) Expression of NCC markers: qRT-PCR data showing the expression of NGFR, SNAI2, SOX10, TFAP2A, Lin28A, and Nanog in iPS-NCCs (n = 3) relative to iPSCs (n = 3). NGFR, SNAI2, SOX10, and TFAP2A are significantly upregulated in iPS-NCCs, while Lin28A and Nanog are significantly downregulated in iPS-NCCs when the detection of mRNA is compared in these cells. *P < 0.05, **P < 0.01, when comparing the two groups. (F) Characterization and differentiation into adipocytes, chondrocytes, and osteocytes of iPS-NCCs. iPS-NCCs were differentiated into adipocytes. In adipocyte-differentiated cells, the accumulation of oil red O-stained lipid drops was observed. Scale bar, 200 µm. iPS-NCCs were differentiated into chondrocytes. The pellet was verified by Alcian blue staining of cartilage proteoglycans. Scale bar, 500 µm. iPSC-NCCs were differentiated into osteocytes and exhibited enhanced calcium deposition by alizarin red S staining. Scale bar, 200 µm. iPSC, induced pluripotent stem cell; NCC, neural crest cell; qRT-PCR, quantitative real-time polymerase chain reaction.
FIG. 2. Expression of HLA class I, class II, and costimulatory molecules on iPSC-NCCs. (A) For the flow cytometry analysis, HLA class I expression of iPSC-NCCs is lower compared with the iPSCs. Expression of β2-microglobulin of the iPSC-NCCs is lower compared with iPSCs. Expression of HLA class II is not detected in either of the cells. MFI, mean fluorescence intensity. (B) MFI of HLA class I and β2-microglobulin was also examined. Data are the mean ± SD of three experiments. * P < 0.01, comparing two groups. (C) Expressions of CD80 (B7-1), CD86 (B7-2), CD274 (PD-L1), and CD273 (PD-L2) failed to be detected in the iPSCs and iPSC-NCCs. Red histogram; isotype control.
costimulatory molecules (data not shown). Thus, these results suggest that iPSC-NCCs have low immunogenicity.

**Suppression of the proliferation of inflammatory immune cells by iPSC-NCCs**

We examined whether established iPSC-NCCs have immunosuppressive effects in vitro. For this assay, we used the MLR test. In this experiment, iPSCs and iPSC-RPE cells were used as controls. Compared to a mix of PBMCs without NCCs, our results showed that iPSC-NCCs inhibited the proliferation of PBMCs (Fig. 3A). In contrast, iPSCs failed to suppress the proliferation of PBMCs, while iPSC-RPE cells strongly inhibited the PBMC proliferation. Compared to using only the PBMC mix, the PBMC mix plus iPSC-NCCs significantly suppressed CD4+ helper T cells, CD8+ cytotoxic T cells, CD11b+ monocytes/macrophages, and CD56+ natural killer (NK)/NKT cells (Fig. 3B). In addition, iPSC-NCCs did not increase the proliferation of PBMCs stimulated with anti-human CD3 and anti-CD28 antibodies (data not shown). Thus, these results suggest that iPSC-NCCs have low immunogenicity.

**Ability of iPSC-NCCs to suppress MLR**

To clarify the role that cell contact plays in the inhibition of T cell proliferation by NCCs, cells were separately cultured in culture plates. Transwell cell insert membranes, which contained PBMCs plus anti-human CD3 and anti-human CD28 antibodies, were placed in the wells. In the PBMC-NCC cocultures, T cell proliferation was significantly suppressed in the absence of the transwell membrane (Fig. 5A). However, iPSC-NCCs failed to suppress the T cell proliferation as contact was prohibited by the transwell membrane. There was a statistically significant difference between iPSC-NCCs and iPSC-NCCs plus cell insert (Fig. 5B). These results imply that cell contact molecules are essential for suppression of activated T cells by iPSC-NCCs. Thus, NCCs can exclusively suppress T cell activation through immunosuppressive cell surface molecules.

**Survey for candidate immunosuppressive molecule(s) by iPSC-NCCs using microarray analysis**

Based on the immunosuppressive properties of the iPSC-NCCs that appeared to be associated with the cell-to-cell contact molecules, we hypothesized that cell surface molecules on iPSC-NCCs were responsible for the suppression. To determine possible candidate molecules, we examined iPSC-NCCs using a microarray assay. For the assay, we prepared human iPSC-NCCs and control iPSCs. GeneChip analysis showed that there was downregulation of the mRNA expression of HLA class I and class II in iPSC-NCCs compared with control iPSCs (Fig. 6A). These data indicated that iPSC-NCCs had poor immunogenicity after differentiation from iPSCs. Furthermore, these findings were confirmed by the flow cytometry results. Among the T cell related immunosuppressive factors, iPSC-NCCs highly expressed mRNA for TGF-β1 and especially TGF-β2, compared to iPSCs (Fig. 6B). Other factors such as FASL, PTGES, PTGES2, CD274, PDCD1LG2, IL10, NOS1, NOS2, NOS3, CD80, CD86, IDO1, IDO2, and TGF-β3 were not involved in the expression of iPSC-NCCs. We also examined how gene expression of iPSC-NCCs changes during the inflamed condition. Similar to previous results by GeneChip analysis, mRNA for TGF-β1 and TGF-β2 in iPSC-NCCs was highly expressed during the inflamed condition, as well as the normal culture condition (Supplementary Fig. S2). These data suggest that NCCs can express and produce these immunosuppressive factors even under inflammatory conditions.

FACS, immunostaining, and qRT-PCR were used to evaluate the expression of TGF-β2 and confirm the expression of TGF-β on iPSC-NCCs. As shown in Fig. 7A and B, iPSC-NCCs clearly expressed TGF-β2 on their surface (= membrane-bound TGF-β2). In addition, qRT-PCR also showed that iPSC-NCCs significantly expressed mRNA for TGF-β2 (Fig. 7C). Based on these findings, we focused on TGF-β as a candidate immunoregulatory factor that suppresses T cells.
FIG. 4. Capacity of iPSC-NCCs to suppress activation of T cells. (A) PBMCs (a healthy donor) in the presence of anti-CD3 and anti-CD28 antibodies were cocultured with iPSC-NCCs for 3 days. After 3 days, the PBMCs exposed to iPSC-NCCs were harvested for Ki-67 FACS analysis. Numbers in the FACS dot plots indicate double-positive cells (e.g., CD3-Ki-67). These data are representative of three experiments. (B) Percentages of the proliferating T cells (double-positive cells in A) were also examined. Data are the mean ± SD of three experiments. * P < 0.05, compared to the two groups. NS, not significant. (C) FACS density plots represent the expression of IFN-γ on CD4+ T cells in the presence of iPSC-NCCs or control iPSCs. PBMCs were stained with anti-CD4 and anti-IFN-γ. Numbers in the density plots indicate the percentage of cells double positive for CD4/IFN-γ. These data are representative of three experiments. (D) Percentages of the IFN-γ secreting CD4+ T cells (double-positive cells in C) were also examined. Data are the mean ± SD of 3 experiments. * P < 0.05, compared to the two groups. NS, not significant. (E) Concentrations of IFN-γ in supernatant (MLR plus iPSC-NCCs) were examined by human IFN-γ ELISA. We also collected control samples: MLR without iPSC-NCCs and MLR plus iPSCs. Data are the mean ± SD of three ELISA determinations. * P < 0.01, compared to the two groups. NS, not significant.
Capacity of iPSC-NCCs to suppress T cell activation in the TGF-β block assay

To determine whether TGF-β is the major factor responsible for inhibiting T cell proliferation by iPSC-NCCs, human TGF-β RI kinase inhibitor (SB431542) was added to the coculture of PBMCs and iPSC-NCCs. The FACS analysis showed that iPSC-NCCs failed to suppress activated T cells in the presence of 10 μM of SB431542 (Fig. 8A). Moreover, there was a statistically significant difference between the PBMC-NCC cocultures and the cultures with 10 μM of SB431542 (Fig. 8B). Dose-dependent effects of SB431542 were observed, although there was no statistically significant difference among the doses (Supplementary Fig. S3). SB431542 had no influence on the cell viability of iPSC-NCCs, and SB431542 itself did not increase the PBMC proliferation (data not shown). In addition, ELISA revealed similar results for IFN-γ, that is, supernatants from PBMC-NCC cocultures in the presence of SB431542 contained significantly higher levels than that observed for the cocultures without this inhibitor (Fig. 8C). These results demonstrate that the blockers of TGF-β RI are able to neutralize the suppression of NCCs. Taken together, our results indicate that TGF-β produced by NCCs can be used to suppress bystander T cells in vitro.

Discussion

While purification of primary NCCs is possible, obtaining a sufficient quantity for use in experiments can be very
difficult. Although Stemple et al. succeeded in isolating primary NCCs from a rat embryo [33], the amount of NCCs that can be collected from an embryo is limited as the neural crest is an embryonic transient tissue. Moreover, the collection of human primary NCCs is ethically impossible. However, as the protocol used in our current study was able to efficiently produce a large amount of induced NCCs, this made it possible to use NCCs in our present experiments. As we were able to obtain sufficient cell quantities, we were able to successfully examine the immunological properties using human iPSC-NCCs. Moreover, as iPSC-NCCs have poor immunogenicity, this means that they can potentially be used for T cell suppression. However, to achieve this suppression, TGF-β produced by NCCs is an important factor in achieving these immunosuppressive properties.

The neural crest plays an important role in the development of the whole human body. Thus, developmental disorders associated with the neural crest, which are referred to
as neurocristopathy, can affect the entire body. Neurocristopathy disorders include Hirschsprung’s disease, CHARGE syndrome, and Treacher-Collins syndrome, among others. Eye related complications such as ocular coloboma and heterochromia iridis are often known to arise from such diseases [34]. Furthermore, these complications include not only the above congenital diseases but also degenerative diseases of neural crest-derived tissues such as keratoconus, Fuchs dystrophy, and adult-onset primary open-angle glaucoma [1]. Thus, NCCs induced from neurocristopathy-

**FIG. 7.** Expression of membrane-bound TGF-β2 on iPSC-derived NCCs. (A) Detection of membrane-bound TGF-β2 on iPSC-NCCs by flow cytometry analysis. We also prepared iPSCs as a control. These cells were stained with anti-human TGF-β2 antibodies. Blue histograms represent isotype control staining. (B) Detection of TGF-β2 in iPSC-NCCs by immunostaining. iPSC-NCCs, but not control iPSCs, clearly expressed TGF-β2 on their surface. Cell nuclei were counterstained with DAPI. Scale bars, 100 μm. (C) iPSC-NCCs or control iPSCs were harvested and examined for expression of TGF-β1, β2 mRNA by qRT-PCR. Results indicate the relative expression of these molecules (ΔΔCt: control iPSCs = 1.0).
FIG. 8. Effect of TGF-β produced by iPSC-NCCs on suppression of T cell activation. (A) PBMCs plus anti-CD3 and anti-CD28 antibodies were cocultured with iPSC-NCCs in the presence of 10 μM of SB431542 and PBMC-NCCs without SB431542 (middle panel) and evaluated by Ki-67 FACS analysis. Numbers in the histogram indicate double-positive cells. These data are representative of three experiments. (B) Percentages of proliferating T cells (the double-positive cells in A) are examined. Data are the mean ± SD of three experiments. *P < 0.05, compared to the two groups. NS, not significant. (C) Concentrations of IFN-γ in supernatant are examined by human IFN-γ ELISA. We collected samples of PBMCs only and PBMC plus iPSC-NCCs with or without SB431542. Data are the mean ± SD of three ELISA determinations. *P < 0.05, comparing two groups. NS, not significant.
specific iPSCs might be an attractive tool for studying the pathogenesis of neurocristopathies. Okuno et al. reported that iPSC-NCCs from CHARGE syndrome patients exhibited defective migratory activity [35].

In addition, iPSC-NCCs can also be used for various applications. For example, the generation of sufficient iPSC-NCC numbers can be used to perform drug screening assays. Moreover, iPSC-NCCs can be utilized as a cell source for improving pathological neurocristopathy conditions. Li et al. and Schlieve et al. also recently reported that iPSC-NCCs differentiated into neurons in human or engineered gut tissues [23,24]. As a result, the transplantation of iPSC-NCCs might make it possible to restore the bowel movements of Hirschsprung’s disease patients.

The simplicity and high reproducibility of the NCC induction protocols can potentially reveal molecular functions and mechanisms of neural crest originated disorders. If so, this could provide an important insight for novel cell-based medicine. In our current study, we succeeded in inducing NCCs and were able to examine the immunological properties of the iPSC-NCCs. Study results demonstrated that while iPSC-NCCs had a lower expression of HLA class I, they had no expression of HLA class II or positive costimulatory molecules (eg, CD80/CD86) against T cells, thereby indicating that iPSC-NCCs exhibit poor immunogenicity.

The MLR test in this study demonstrated that iPSCs failed to inhibit PBMC proliferation. Lu et al. previously reported that in contrast to human skin fibroblasts, which significantly induced allogeneic T cell activation and proliferation, human iPSCs had low or negligible immunogenicity, were able to induce IL-10-secreting regulatory T cell, and failed to induce T cell activation and proliferation [36]. In contrast, iPSC-NCCs globally inhibited PBMC proliferation. Since a low level of HLA molecules theoretically increases the susceptibility to attacks by NK cells, this is an important pathway for defending against pathologic microorganisms under normal conditions [37]. The MLR test, however, revealed that iPSC-NCCs did not activate NK cell proliferation. When PBMCs were stimulated with anti-CD3/CD28 agonistic antibodies, iPSC-NCCs inhibited CD4+ and CD8+ T cell proliferation in cocultures. Moreover, iPSC-NCCs inhibited IFN-γ expressing CD4+ T cells. Thus, not only do iPSC-NCCs suppress CD4+ T cell proliferation but also they inhibit helper T cell function (= suppression of Th1 cells). As per our MLR results (Fig. 3), iPSC-NCCs can also inhibit the IFN-γ secretion from potential producers (CD8+ T cells, CD11b+ monocytes/macrophages, and CD56+ NK/NKT cells), as well as CD4+ T cells.

Our transwell culture results suggested that cell-to-cell contact was critical in order for iPSC-NCCs to inhibit T cell proliferation. And perhaps more importantly, the iPSC-NCCs constitutively expressed membrane-bound TGF-β2. In addition, SB431542 (human TGF-β RI kinase inhibitor VI) prevented the T cell suppression mediated by iPSC-NCCs. This finding demonstrated that iPSC-NCCs suppressed T cell activation through membrane-bound TGF-β. In fact, TGF-β is a powerful mediator of inflammatory immune cell suppression [38]. Based on these findings, it has been assumed that iPSC-NCCs are able to suppress T cell activation by delivering membrane-bound TGF-β to the migrating T cells that are targeted through the TGF-β/TGF-β receptor interactions.

NCCs have immunological properties, which are similar to the corneal endothelial cells, and are derived from the neural crest. Since neural crest derivatives may already have developmentally acquired immunological properties during the neural crest stage, when the differentiation advances from the neural crest, it is then possible that more progressive immunological properties can be gained. Zhao et al. succeeded in inducing corneal endothelial cells from human iPSCs through NCCs [39]. In future studies, we hope to establish iPSC-derived corneal endothelial cells, after which, we will then compare the immunological properties between the NCCs and the corneal endothelial cells.

In conclusion, we successfully induced NCCs from human iPSCs. Using iPSC-NCCs, this makes it possible to determine the molecular functions of NCCs and the pathogenesis of neurocristopathy. iPSC-NCCs are hypoimmunogenic cells that can potentially suppress immune cells, including T cells. TGF-β produced by NCCs plays an important role in T cell suppression. iPSC-NCCs have been reported to have a potential for wound healing [21,22] and enteric neural transplantation treatment [23,24]. The current results may help lead to the development of a new treatment of neurocristopathies. In addition, if iPSC-NCCs are clinically used in humans, it is likely that they will cause less or no inflammation and exhibit optimum wound healing, as they exhibit a low immunogenicity and have the potential for inflammatory immune cell suppression. We believe that the current study findings could contribute to realization of the use of iPSC-NCCs in cell-based medicine.

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Author Disclosure Statement

No competing financial interests exist.

References


