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Kyoto University
Two differential flows in a bioreactor promoted platelet generation from human pluripotent stem cell-derived megakaryocytes

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Abstract, 209 words; Main text, 2083 words, 6 figures, 1 table

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Abstract (209 words)

Induced pluripotent stem cell (iPSC) technology enables us to investigate various potential iPSC-based therapies. Although the safety of iPSC-derivation has not been completely validated, anucleate cells, such as platelets or erythrocytes, derived from iPSCs are promising targets. However, the efficiency of \textit{in vitro} platelet generation from megakaryocytes (MKs) under static culture conditions is lower than is seen \textit{in vivo}. Here we demonstrate the proof-of-concept by a two-dimensional flow culture system that enabled us to increase platelet yield from human embryonic stem cell (hESCs) or iPSC-derived MKs using a biomimetic artificial blood vessel system. The bioreactor was composed of biodegradable scaffolds with ordered arrays of pores made to mimic \textit{in vivo} bone marrow through salt-leaching. Within the system, two flows in different directions in which the angle between the directions of flow is 60° but not 90° contributed to suitable pressure and shear stress applied to MKs to promote platelet generation. Generated platelets derived from hESCs or hiPSCs through the bioreactor with 60° angle revealed intact integrin αIIbβ3 activation after agonist stimulation. Collectively, our findings indicate that two flows in different directions of two-dimensional flow culture may be a feasible system for \textit{in vitro} generation of platelets from pluripotent stem cells, i.e. iPSC-derived MKs, in numbers sufficient for transfusion therapy.
Introduction

Platelets are essential for hemostasis and thrombosis, and also play important roles in wound repair, inflammatory reactions, angiogenesis, lymph-vessel separation, and tumor metastasis (1). In humans, approximately $1 \times 10^{11}$ platelets are produced each day, presumably through the cytoplasmic fragmentation of megakaryocytes (MKs) and/or formation of proplatelets, elongated pseudopods that extend from mature MKs and release platelets (2). Platelets production requires the commitment of hematopoietic stem cells (HSCs)/hematopoietic progenitor cells (HPCs) to the MK lineage, followed by proliferation of the progenitors and terminal differentiation. Ultimately, immature MKs develop a unique membrane complex called the demarcation membrane (DM) system, which is thought to serve as a membrane reservoir for platelet biogenesis through both direct cytoplasmic rupture and elongation of proplatelets (3). Both of those processes are controlled by the orchestrated activities of numerous signaling molecules (3). A variety of gene-targeted mouse models have shed light on the roles played by the transcriptional factors involved in platelet formation, and have contributed to the characterization of the linkage between the machinery of MK maturation and platelet release (3,4,5,6,7). However, the actual machinery of platelet release (platelet biogenesis) from MKs has not yet been characterized (8).

The establishment of mouse (9) and human embryonic stem cells (hESCs) (10) and human induced pluripotent stem cells (hiPSCs) (11,12) were crucial advances in the area of regenerative medicine. Notably, hiPSC-based platelet generation is a particularly fascinating therapy model, in part because blood donors are becoming increasing scarce in most industrial countries, including Japan (13). In addition, because platelets are anucleate, they can be
subjected to gamma irradiation prior to their transfusion, which would prevent the occurrence of oncogenesis or teratoma formation due to the presence of residual undifferentiated hiPSCs or hiPSC-derived cells. Finally, for patients requiring repeated transfusions, it is important that the platelets are human leukocyte antigen (HLA)-matched (13); thus an HLA-identical donor could provide the origin cells for creating iPSCs that contribute to a sustained platelet supply.

Here, by primarily using human ESCs as a gold standard of reference (14, 15), we describe a two-dimensional flow culture that recapitulates bone marrow in the view of “blood flow” application and may be a feasible system for *in vitro* generation of platelets from human pluripotent stem cell-derived MKs in numbers sufficient for transfusion therapy.
Methods

Cells and Reagents

The KhES-3 human ESC clone (Institute for Frontier Medical Science, Kyoto University, Kyoto, Japan) was used upon the approval by the Minister of Education, Culture, Sports, Science, and Technology of Japan (MEXT) (14). In addition, The TkDA3-4 hiPSC clone was used (15). The mouse C3H10T1/2 cell line was purchased from the RIKEN Bio-Resource Center (Tsukuba, Ibaraki, Japan) and was cultured as described previously (14,15). Human umbilical cord endothelial cells (HUVECs) were purchased from KURABO (Tokyo, Japan). Human vascular endothelial growth factor (VEGF), human thrombopoietin (TPO) and human stem cell factor (SCF) were all from R&D Systems (Minneapolis, MN). Heparin was from Ajinomoto Pharmaceuticals Co., LTD (Tokyo, Japan). The following antibodies and stains were obtained from BD Biosciences (Franklin Lakes, NJ): allophycocyanin (APC)-conjugated anti-CD41a, FITC-conjugated anti-CD41a (GPIX), anti-CD42b (GPIbα) and Hoechst 33342 (blue). DAPI (Vector Laboratories, Burlingame, CA) and Hoechst 33342 were used for nuclear staining in MKs. FITC-conjugated PAC-1 antibody (BD Biosciences, San Diego, CA) was used for platelet activation studies as described (14, 15).

Hematopoietic differentiation of human ESCs or iPSCs

Differentiation of human ESCs or iPSCs into hematopoietic cells was performed as described previously (14,15). In brief, small clumps of ESCs or iPSCs (<100 cells treated with PBS containing 0.25% trypsin (Invitrogen, Carlsbad, CA), 1 mM CaCl2 (Sigma-Aldrich, St. Louis,
MI) and 20% Knock-out Serum Replacement (KSR, Invitrogen [Life technology Japan, Tokyo]) were transferred onto mitomycin (Sigma)-treated or irradiated C3H10T1/2 cells and co-cultured in hematopoietic cell differentiation medium (Iscove’s Modified Dulbecco’s Medium [IMDM, Life technology Japan, Tokyo] with 15% fetal bovine serum and VEGF as described previously (14,15), which was replaced every 3 days. On days 14-15 of culture, the HPCs within ES-(14) or iPS-Sacs (15) were collected, transferred onto fresh mitomycin-treated or irradiated feeder cells, and further cultivated in differentiation medium supplemented with 50 ng/ml TPO, 50 ng/ml SCF and 25 U/ml heparin sodium, as described (14,15). The medium was refreshed every 3 days, and non-adherent cells were collected on day 20 of culture and transferred to a bioreactor.

**Flow cytometric analysis of platelets**

Washed platelets were prepared as described previously (14,15). The resultant platelet pellets were resuspended with staining medium and stained with anti-human CD41a-APC and CD42b-PE for 30 min at room temperature. The platelets were then diluted in 200 µl of staining medium and analyzed by flow cytometry (FACSARia, BD biosciences, San Diego, CA). Platelet numbers were counted using True Count Beads (BD Biosciences).

**Construction of a bioreactor to obtain platelets from human iPSC-derived MKs**

The fabrication of a bioreactor to produce platelets entailed the following steps. Using photo-resist SU8-3050 (Nippon Kayaku, Tokyo, Japan), a microfluidic device was constructed by centrifugation on a silicon wafer at 5000 rpm for 30 s. The device was then exposed after
applying a mask using a mask aligner, developed in developer and rinsed with de-ionized water. Polydimethylpolysiloxane (PDMS) was then solidified on the fabricated mold at 60°C in an oven. Finally, the PDMS was released from the mold and bonded to a glass plate.

**Experimental procedure (dissemination of cells, culture and counting platelets)**

Sterilization of the bioreactor was achieved through treatment with ethanol for 30 min irradiation with UV light. Platelet generation in the bioreactor was observed in a connected circulation system on the stage of a confocal microscope (FV1000, Olympus, Tokyo, Japan) at 37°C. The process of platelet generation from MKs was visualized using a confocal microscope system, and the images were recorded in the hard disc of a connected computer. Flow pressure was determined by flow speed and the medium collected, including the cell populations in the reservoir. After adding ACD medium to the culture medium (1:10 dilution), the collected samples were centrifuged at 900 rpm for 10 min (slow acceleration). Finally, the cell samples were stained for cell surface molecules specific for MKs/platelets and analyzed by flow cytometry.

**Statistics**

We evaluated results by using an unpaired t-test for significance.
**Results and Discussion**

Observations in living mice made using two-photon confocal microscopy revealed that platelet release occurs within both the capillaries and sinusoids of the bone marrow (8). Within human bone marrow, sinusoidal capillaries are the discontinuous type, with wall thicknesses less than 30 µm and containing numerous pores. One approach to reconstituting the structure of such vessels *in vitro* is to use a sheet-like scaffold with multiple pores (16). We constructed a structure using the salt-leaching method as shown in Figure 1A. The contents of the solution used for the salt-leaching are shown in Table 1.

Using the constructed sheet-like porous structure, we designed the bioreactor to recapitulate mainly two functions of capillary blood vessel for platelet production: (i) one function that is a porous structure of endothelial cells for fixing MKs on the outside of blood vessel wall (17), and, (ii) another function that is a platelet production of MKs by applying the shearing effects of blood flow. Within the bioreactor, the membrane scaffold for platelet release from MKs was composed of PDMS. Moreover, the scaffold was completely covered by HUVECs where adjacent chamber with a layer of HUVECs and a formed porous barrier in between appeared 30 µm. By adjusting the speed of flow through the structure, different levels of shear stress were generated, which were calculated using equation (A), where $Q$ is flow speed [ml/h], $\mu$ is the viscosity of the medium [Pa/s], $a$ is the width of the duct [5.0 mm], and $b$ is the height of duct [0.3 mm].

$$\tau = \mu \frac{6Q}{ab^2} \quad (A)$$
In the system, the medium flows past MKs on HUVECs in two directions (pressure flow and main flow), such that the pressure flow is perpendicular to the main flow; the main flow applies shear stress to the MKs, while the pressure flow applies pressure for MK fixation to the each slit. Thus we estimated that exposed shear stress was occurred from 0.14-37.5 dyne/cm² onto MKs within chamber and iPSC-derived MKs were applied into the regions of the pores, as shown in the left panel of Figure 1B, although it is not ruled out that the flow was turbulence.

In addition, we expected that vascular cell adhesion molecule-1 (VCAM-1) on HUVEC might promote platelet release from MKs because of the reports that integrin α4β1 expression in MKs is crucial for platelet production in vivo in mouse models (reference 18) or that the binding between integrin α4β1 in MKs and VCAM-1 promoted proplatelets, a hallmark of platelet generation, from cultured MKs in vitro (19).

Our bioreactor system enabled us to visualize real-time changes to MKs during culture. Time-lapse confocal microscopy revealed that applied iPSC-derived MKs grew on HUVECs within the bioreactor (Figure 2A), and the numbers of adherent MKs were greater under weak flow conditions (0.14 dyne/cm²) (Figure 2B). We applied 1.2 x 10⁵ MKs totally into the bioreactor and approximately 5.2-5.6 x 10⁴ MKs (average=5.42 x 10⁴, Figure 2A) or 5.8-6.5 x 10⁴ MKs (average=6.2 x 10⁴, Figure 2B) were counted in the absence or presence of flow respectively. When we compared the final yield of platelets between the static culture (no flow) in a conventional culture dish (14) and our new bioreactor system with flow, as shown in Figure 3A, we found that platelet generation from human ESC-derived MKs tended to be better in the flow system (Figure 3B). Moreover, time-lapse imaging of the platelet production from MKs under flow revealed that increasing the shear stress by adjusting flow speed (1.0 ml/hr)
stimulated large-proplatelet (90 min in Figure 4), followed by cytoplasmic fragmentation from ESC-derived MKs (92 min in Figure 4). However, the increase in platelet production was not significant (Figure 3B).

To improve platelet yield, we designed a new bioreactor using two flows with an angle between the main and pressure flows. The schematic in Figure 5A shows the newer design of a bioreactor, in which the angle between the directions of flow is 60°. The platelets were generated from MKs placed between the slits. We again applied 1.2 x 10^5 MKs for each experiment within a newer bioreactor. Under these conditions, the numbers of platelets produced from human ESCs were as much as 3.6 times higher than in static culture (mean count: 19822 under static vs. 71755 under flow, Figure 5B, from independent 4 experiments, p<0.05). These results using the same improved reactor system were confirmed by using human iPSC-derived MKs (2 experiments, data not shown). Purity of CD41a+ or CD42b+ platelet populations was also examined by flow cytometer and there was no significant difference between static condition and bioreactor (Figure 5C and 5D).

In order to confirm the functionality by collected platelets through second generation of a bioreactor in Figure 5A, agonist-induced integrin activation in platelets was examined. The PAC-1 binding, the activated form of an integrin αIIbβ3 after agonist stimulation showed intact functionality by hESC- or iPSC-derived platelets (Figure 6), confirming the similarity to the previous studies obtained under static conditions (14, 15).

Using a novel 3-dimensional bioreactor, which enabled us to regulate shear stress through application of flow in two directions, we enhanced platelet generation from MKs by subjecting the cells to the appropriate levels of shear stress. Within this system the shear stress created by
the main flow could be regulated by the pressure flow. Furthermore, it appears that an angle of 60° between the main and pressure flows is optimal.

This novel conception has the potential to influence a new era of regenerative medicine through the use of hESCs or hiPSCs. In an effort to better understand the underlying molecular mechanisms of platelet release and production from MKs, we are attempting to mimic previously visualized \textit{in vivo} platelet production (8), which showed evidence of shear stress dependence. Recently, several groups reported \textit{in vitro} platelet generation under conditions in which high shear stress was produced utilizing one-directional flow (18, 21). On the other hand, our proposed bioreactor makes use of flow in two directions. We suggest that the flow applying pressure to the cells stimulates MK maturation on HUVECs, while the flow applying shear stress stimulates platelet generation. In addition, it appears that the angle between those two-directional flow a key for improving production efficiency as exemplified by three times difference in comparison between Figures 3 and 5). This two-direction flow strategy is seemingly a feasible approach to the future development of bioreactors able to supply platelets derived from human pluripotent stem cells in numbers sufficient for transfusion.
References


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**Author Contributions**
Y.N. and R.T. designed the experiments and performed experiments. S.N., H.E., and T.D. prepared samples. M.N. prepared materials. F.A. and T.F. supervised and designed the construction of a bioreactor. H.N. provided valuable discussion. K.E. wrote and edited the manuscript.

**Conflict of Interest**

H. Nakauchi and K. Eto are scientific advisory board of Megakaryon Co. Ltd. without salary. Others have no conflict of interest.
Figure legends.

Figure 1. Fabrication process of a sheet-like scaffold and Medium circulation model of bioreactor.

(A) (a) Produce a wax (made of toluenesulfonamide) model. (b) Wax model was dipped into PVA (Polyvinylalcohol) solution and pulled up with a dip coater (pull-up velocity: 1 mm/s). Dip coating of PVA onto the wax model was repeated 6 times, and each coating was followed by drying at room temperature for 30 minutes. (c) After an interval of 12 hours, the PVA-coated wax model was soaked in acetone. Because of the insolubility of PVA and solubility of wax in acetone, the wax was selectively dissolved, leaving the structure of PVA. (d) The PVA model then was coated with the polymer and NaCl particle solution using the dip coater (velocity: 3 mm/s). To coat the PLCL (poly (L-lactide-co-epsilon-caprolactone)) membrane as uniformly as possible, dip coating of polymer solution onto the PVA model was repeated three times in one direction and three times in the opposite direction by setting the model upside down. (e) Dip coated PLCL of the second layer in the same way as (d). (f) Each coating was followed by drying at room temperature for 2 minutes. The PLCL-coated PVA models then were soaked in deionized (DI) water followed by dissolution of PVA and the NaCl microparticles. After PVA and the NaCl microparticles were completely removed from the PLCL, the salt-leached PLCL scaffold was dried for 24 hours at room temperature.

(B) Culture within the reactor system was performed at 37°C in a CO₂ incubator. Circulation was achieved using a PFA tube connected to a roller pump to produce flow. The left side schematic shows the cross-sectional view of bioreactor. Human umbilical vascular endothelial cells (HUVECs) were pre-applied for stable adhesion to the micropores in the scaffold 2 days
before the megakaryocytes (MKs) were applied.

**Figure 2.** Confocal microscope images of megakaryocytes  (A) Static culture.  (B) Flow culture in which the shear stress produced with the reactor system depicted in Figure 2 was 0.14 dyne/cm². Nuclei were stained with DAPI (blue) and CD41a was stained with anti-human CD41a-FITC (green). Low-magnification images (left top) shows only DAPI staining. Bars, 50 µm.

**Figure 3.** Original model of the bioreactor and comparison of platelet yield (static vs. reactor) using the bioreactor system.

(A) The bioreactor was 25 mm x 28 mm and contained numerous circuits for flow. The termination of each circuit was a pore (~8 µm) formed by a pocket containing a single MK. MKs were stimulated to mature by the pressure flow, while platelet release was stimulated by the main flow. (B) Numbers of released platelets were determined based on CD41a⁺CD42b⁺ counts per reservoir (over a period of 12 h). Black bars show the platelet counts obtained under static conditions using the protocol described in ref. 14. Gray bars show the platelet counts obtained using the first generation bioreactor. Number of platelets from static condition equalized 1.0. The input was 1.2 x 10⁵ mature MKs in both conditions.

**Figure 4.** Confocal micrographs showing the time course of a platelet release from a MK (green: CD41a, blue: Hoechst). Nuclei were stained with Hoechst.
**Figure 5.** Schematic of an improved bioreactor with two-directional flow (second type of a bioreactor) and comparison of platelet yields (static vs. reactor).

(A) The illustration shows the design of a bioreactor with two-directional flow in which the angle between the directions of the main and pressure flow is 60°. Platelets were generated from MKs placed between the slits. The slits were 16 μm wide with 4 μm in between.

(B) Numbers of released platelets were determined based on hESC-derived CD41a⁺CD42b⁺ platelet population counts per reservoir (over a period of 12 h). Black bars show the platelet counts obtained under static conditions using the protocol described in ref. 14. Gray bars show the platelet counts obtained using second generation of a bioreactor. Average number of platelets from static condition equalized 1.0. The input was 1.2 x10⁵ mature MKs in both conditions. Experiments were mean+/−s.e.m. (n=4).

(C) Representative dot plots of platelets in flow cytometer, derived from human ESCs through second generation of a bioreactor (right). Same gate in side scatter (SSC) and forward scatter (FSC) of fresh human platelets was used for ESC-derived platelets in flow cytometer (left). This gate was re-analyzed for detection of CD41a (integrin αIIb) and CD42b (GPIbα).

(D) Purity of CD41a⁺ or CD41a⁺CD42b⁺ platelet population of gated population depicted in the left panel of Figure 5C (SSC and FSC) was also evaluated as comparison between static culture condition and bioreactor. Black column indicates static culture and silver column indicates bioreactor. In each column, white bar is merged as CD41a⁺CD42b⁺ population.

**Figure 6.** Integrin activation of human ESC- or iPSC-platelets. Integrin activation in ESC-platelets (KhES-3) (A) and in iPSC-platelets (TkDA3-4)(B). The binding of PAC-1
(indicative of platelet activation) to individual platelets was quantified in the absence and presence of 50 μM ADP, 0.5 Unit human Thrombin, or 200 nM phorbol myristate acetate (PMA) using flow cytometry. Mean fluorescence intensity (MFI) of bound PAC-1, obtained from CD42b^+ fraction was evaluated (experiments were independently performed twice).
Figure 1 Nakagawa et al.

A

(a) Wax  (b) PVA  (c) NaCl Particle  (d) PLCL  (e) PLCL  (f) Pore

B

Porous Scaffold

Flow to Reservoir

CO₂ Incubator

Bio-Reactor

Medium tank

PFA Tube

Damper

Valve

Manometer

Roller Pump
Figure 2 Nakagawa et al.
Figure 4 Nakagawa et al.
Figure 5 Nakagawa et al.

A

Outlet port
Pressure flow channel
Slit
Main flow channel
MK Supply port
Outlet port
Pressure flow

B

Relative number of CD41a+CD42b+ platelets

Static
Bioreactor

p<0.05

D

Purity of CD41a+ (%)

Static
Bioreactor

N.S.

C

Peripheral blood

hESC-platelets from bioreactor
Figure 6 Nakagawa et al.

A

KhES3

PAC1 (MFI)

0 500 1000 1500

No Agonist ADP Thrombin PMA

B

TkDA3-4

PAC1 (MFI)

0 500 1000 1500

No Agonist ADP Thrombin PMA
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<th>Solution 2</th>
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