Generation and manipulation of human iPSC-derived platelets

Naoshi Sugimoto1 · Koji Eto1,2

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Abstract
The discovery of iPSCs has led to the ex vivo production of differentiated cells for regenerative medicine. In the case of transfusion products, the derivation of platelets from iPSCs is expected to complement our current blood-donor supplied transfusion system through donor-independent production with complete pathogen-free assurance. This derivation can also overcome alloimmune platelet transfusion refractoriness by resulting in autologous, HLA-homologous or HLA-deficient products. Several developments were necessary to produce a massive number of platelets required for a single transfusion. First, expandable megakaryocytes were established from iPSCs through transgene expression. Second, a turbulent-type bioreactor with improved platelet yield and quality was developed. Third, novel drugs that enabled efficient feeder cell-free conditions were developed. Fourth, the platelet-containing suspension was purified and resuspended in an appropriate buffer. Finally, the platelet product needed to be assured for competency and safety including non-tumorigenicity through in vitro and in vivo preclinical tests. Based on these advancements, a clinical trial has started. The generation of human iPSC-derived platelets could evolve transfusion medicine to the next stage and assure a ubiquitous, safe supply of platelet products. Further, considering the feasibility of gene manipulations in iPSCs, other platelet products may bring forth novel therapeutic measures.

Keywords Clinical application · Megakaryopoiesis · Thrombopoiesis

Introduction
In 2006, induced pluripotent stem cells (iPSCs) were established from mouse fibroblasts by the introduction of four reprogramming factors, Oct3/4, Sox2, Klf4 and c-Myc [1]. In the next year, the establishment of human iPSCs was reported by the same or similar combination of reprogramming factors [2–4]. Analogous to embryonic stem cells (ESCs), iPSCs bear pluripotency and self-renewal capacity, in other words, they can proliferate indefinitely in vitro and have the capacity to differentiate into any cell type, recapitulating the development process in vivo (Fig. 1). Because iPSCs can be established from somatic cells such as skin fibroblasts or blood cells, they bear none of the ethical issues associated with embryo use and can be essentially established from any individual. In addition, iPSCs can be edited for their genes by CRISPR/Cas9 and other technologies. These characteristics have encouraged their application to disease remodeling in vitro and regenerative medicine [5, 6] (Fig. 1).

The versatility of iPSCs is applicable to the field of hematopoiesis. One way is to recapitulate hematopoietic diseases, ranging from monogenic congenital diseases or myeloproliferative diseases (MPD) to multifactorial hematopoietic malignancies and bone marrow failures, by using iPSCs established from patients [7, 8]. These models will contribute to uncovering the unknown pathogenesis or drug testing, thus leading to novel treatment modalities for the diseases. Another way is the clinical application of the hematopoietic cells differentiated from iPSCs, such as platelets, erythrocytes, lymphocytes and macrophages [5, 6]. This review will focus on the current status of the development, application and manipulation of iPSC-derived platelet-like particles (iPSC-PLTs), which has been leading the field of regenerative medicine using iPSC-derived blood cells and has reached the first-in-human clinical trial.
Expectations in manufacturing blood products from iPSCs

The primary treatment for patients with severe anemia or thrombocytopenia is the transfusion of blood products [9, 10]. Patients with primary hematopoietic failures and malignancies are those that require transfusion the most. Normal hematopoiesis is decreased while myelosuppressive treatments such as chemotherapy, radiotherapy and hematopoietic stem cell transplantation (HSCT) are applied. Non-hematopoietic cancers can also require transfusions due to chemotherapies or the invasion of malignant cells into the bone marrow. Other conditions that require transfusions include massive hemorrhage by trauma or surgery, the consumption and loss of platelets through extracorporeal circulation in cardiovascular surgeries using artificial heart and lung apparatuses, and respiratory failures using extracorporeal membrane oxygenation (ECMO).

Fortunately, many countries have well-established systems in which blood products manufactured from healthy blood donors are supplied to patients in need of blood transfusions (Table 1). However, the percentage of young blood donors is decreasing in developed countries due to declining birthrates and aging populations [11, 12]. Elderly people have higher incidences of hematological diseases such as leukemia and myelodysplastic syndromes (MDS) along with cancers and cardiovascular disease. Furthermore, owing to advances in medical care, the aforementioned aggressive treatments that require transfusions are being increasingly applied. Thus, there is concern that a supply shortage of blood products will occur in the future [11, 12].

In particular, platelet products are stored at room temperature to best maintain their function, but still have a short shelf life of only four or five days based on their loss of function and the risk of bacterial growth. Therefore, platelet products must be continuously supplied to meet demand. Moreover, supply can be significantly affected by extraneous factors such as holidays, inclement weather and pandemics,

| Table 1  | Characteristics of donor-derived platelets and iPSC-PLTs
|-----------------|-----------------|
| Donor-derived platelets | iPSC-PLTs |
| Established massive supply system with high safety profile | Early clinical trial phase |
| Supply shortage risk | Potential donor-independent stable supply |
| 4–5 days shelf life | Infection-free |
| Ageing society | Certified pathogen-free source |
| Contamination | GMP-grade processing |
| Bacteria flourish | Autologous, HLA-homologous or HLA-null products for allo-PTR |
| Test window period | Uniform quality and amenable to gene manipulation |
| Emerging infections | Currently high cost |
| Alloimmune platelet transfusion refractoriness (allo-PTR) | |

Fig. 1 Applications of induced pluripotent stem cells (iPSCs). iPSCs are established from somatic cells such as skin fibroblasts and blood cells by the introduction of the reprogramming genes OCT3/4, SOX2, KLF4 and c-MYC or other combinations. iPSCs can be expanded indefinitely and induced to differentiate into any cell type in vitro. Cells differentiated from iPSCs are expected to contribute to disease remodeling in vitro, which can be used to clarify the pathogenesis and develop novel drugs and therapeutics, and to regenerative medicine as cell therapies.
like the current COVID-19 pandemic [13]. The risk of supply shortages is especially higher for patients with rare blood types who require matched donors to avoid alloimmune rejection [12, 14]. In addition, although improved testing and collection procedures have made them rare, blood-borne infection and bacterial contamination cannot be completely eliminated partly due to the time window of the infection tests and emerging infections for which reliable tests are unavailable [15].

As an alternative to these donor-dependent blood product supply systems, biosynthesized blood products have been created by introducing molecules into liposomes and other platform materials to mimic the function of red blood cells or platelets [16–18]. These products are expected to have advantages in emergency situations with limited logistics, such as trauma at battlefields, owing to better preservation capacity, but it is especially difficult to recapitulate the delicate and complicated functional properties of native platelets.

Another approach is to derive the desired cells from iPSCs, which would realize the supply of bona fide blood cells independent of blood donors (Table 1). In addition, iPSC-derived blood products can be devoid of infection by aseptic manufacturing from the source cells, which are assured of being pathogen-free. From the perspective of iPSC-based regenerative medicine, since platelets and erythrocytes are non-nucleated cells, there is essentially no concern about their tumorigenesis, and the risk from nucleated cells that contaminate the product can also be dealt with by irradiation, which is generally performed for transfusion preparations. The differentiation of hematopoietic cells may be relatively easy compared with other cell types because cells can be cultured in suspension and do not require three-dimensional tissue construction. However, difficulties lie in the development of a production system that can provide $10^{11}$ platelets or $10^{12}$ red blood cells and a quality equivalent to current blood products. In addition, cost is a concern: the cost for red blood cells is approximately US $150–450 per pack and for platelets is $400–1000 per pack in developed countries [19].

### Hematopoiesis in vivo vs. hematopoiesis in vitro from iPSCs

Since the 1990s, several groups have achieved the hematopoietic differentiation of mouse ESCs [20–22] and human ESCs [23–28]. Soon after human iPSCs were established in 2007, the hematopoietic differentiation of iPSCs from healthy donors and patients were reported for many blood cell lineages [7, 29–32]. One issue with these pluripotent stem cells (PSCs), however, is that the differentiated hematopoietic cells tend to have primitive or early definitive of embryonic/fetal phenotype, not the definitive adult phenotype [33]. Primitive hematopoiesis occurs in the yolk sac and gives rise to primitive HSC-like cells, which differentiate into primitive erythroid cells, macrophages and megakaryocytes. Then, for a brief period, yolk sac gives rise to erythromyeloid progenitors (EMPs) which differentiate to both erythroid and myeloid cells, among which tissue-resident macrophages persist into adulthood [34]. Following these limited spectrums of hematopoiesis, definitive bona fide HSCs originate from the hemogenic endothelium in the aorta-gonad-mesonephros (AGM) during mid-gestation, then migrate to the fetal liver, spleen and finally the bone marrow, to differentiate to all adult-type hematopoietic cells throughout life [33]. Conventionally, HSCs are postulated to differentiate to multipotent progenitor cells that differentiate through a branching fashion to common myeloid progenitor cells (CMPs) and then to megakaryocyte/erythroid bipotent progenitor cells (MEPs) and finally megakaryocytes. However, megakaryocyte differentiation from HSCs has been recently revisited, with studies reporting megakaryocyte-biased HSCs and the direct differentiation of megakaryocyte progenitors from HSCs that bypasses CMPs and MEPs [35, 36].

The primitive or embryonic/fetal nature of iPSC-derived hematopoiesis may be the reason for the several limitations in differentiating certain hematopoietic cells. First, the induction of HSCs has not been successful unless the HSCs were collected from the bone marrow of mice in which teratomas were formed by transferring iPSCs [37, 38] or by the forced expression of seven transcription factors (ERG, HOXA5, HOXA9, HOXA10, LCOR, RUNX1 and SPI1) in iPSC-derived hemogenic endothelial-like cells [39]. Second, the myeloid lineage is dominant, and lymphoid hematopoiesis have been less well reconstituted. Thus, most in vitro iPSC hematopoietic disease models are myeloid types, such as MPD, acute myeloid leukemia (AML), MDS and bone marrow failures [8]. Thirdly, the dominant types of hemoglobin in iPSC-derived erythrocytes are neonatal (HbE) or fetal (HbF), not adult-type HbA [33, 40–42]. Fourthly, the primitive or embryonic/fetal nature could also be the cause for the low yield of iPSC-PLTs from iPSC-derived megakaryocytes, which are currently differentiated from hematopoietic progenitor cells (HPCs), but not definite HSCs, since neonatal megakaryocytes have a higher capacity for expansion but lower polyploidy and platelet generation compared to the adult type [43]. In addition, fetal/neonatal platelets are suggested to be hyporeactive compared to adult-type platelets, although they survive longer, which may also become the nature of platelets derived in vitro from iPSCs [44, 45].
Development of iPSC-PLTs

Success and limitations in the direct differentiation of platelets from HSCs and PSCs in vitro

The derivations of erythrocytes and platelets from iPSCs have been both pursued, but the derivation of red blood cells is very much behind in terms of clinical application due to the scale and immature phenotype. As noted above, one magnitude more red blood cells is needed for transfusions compared with platelets \(10^{12} \text{ vs. } 10^{11}\). In addition, only one red blood cell can be derived from one erythroblast through enucleation, while 800 to 2,000 platelets are fragmented from one megakaryocyte in vivo. Furthermore, with current methods, iPSC-derived erythroblasts do not complete enucleation or globin switching to adult-type HbA [40–42].

iPSC-PLTs have reached clinical trial, but to do so, several issues had to be overcome. The first was obtaining enough megakaryocytes, the mother cell of all platelets (Table 2). As megakaryocytes mature, they become large and polyploid through endomitosis, store platelet-specific granules and develop the demarcation membrane system (DMS) [46]. In steady state, mature megakaryocytes in the vascular niche of the bone marrow extend their proplatelets into the bone marrow sinusoid, where blood flow tears off proplatelets or smaller fragments, i.e. platelets, which are released into the systemic circulation [47].

Interestingly, the lung has been postulated to produce platelets or fragment to smaller platelets based on the observation that megakaryocytes reside in the lung and that the number of efferent platelets is more than of afferent platelets, whereas for megakaryocytes the opposite is true [48]. Although one report questions the lung as a platelet source [49], a recent in vivo imaging study of mouse bone marrow captured images of the vasculature with megakaryocytes trapped in the lung and spleen extending proplatelets and eventually releasing platelets as they do in the bone marrow [50]. The study further suggested that the lung also harbors HSCs and HPCs, which may migrate out of the lung to the bone marrow under thrombocytopenia and relative stem cell deficiency in the bone marrow. Meanwhile, our in vivo live imaging study of the mouse bone marrow observed that, under inflammatory conditions, megakaryocytes release platelets through a “rupture” fashion by an IL-1α- and caspase-3-dependent mechanism [51]. In the thrombopoietin (TPO)-dependent steady state, alpha and beta-1 tubulins are expressed in a regulated fashion to form proplatelets, but in the rupture state, the expression of tubulins is dysregulated, resulting in possibly diminished plasma membrane stability. Interestingly, our collaborator also reported that an activated form of tyrosyl-tRNA synthetase (YRSACT) functions to enhance megakaryopoiesis and platelet production in TPO-independent fashion, most likely under stress conditions [52].

Although megakaryocytes can be collected through bone marrow aspiration, they are not considered a source for the production of platelets ex vivo (Table 2). The collection procedure is invasive, and megakaryocytes are normally only about 50–150/μL, or less than 0.1% of nucleated cells in the bone marrow, which range from HSCs to fully differentiated hematopoietic-lineage cells [53, 54]. Because megakaryocytes cannot expand in vitro, it is unrealistic to prepare enough of them from the bone marrow for ex vivo production. HSCs, the origin of all adult-type hematopoietic cells, mostly reside in the bone marrow, but they can be isolated from cord blood without invasiveness or from peripheral blood under specific conditions such as mobilized conditions that reduce the invasiveness compared with the bone marrow (Table 2). Significantly owing to the discovery of the key cytokine TPO [55–58], HSCs can be differentiated into megakaryocytes in vitro [59–61]. However, the number of obtainable HSCs is small \(10^6\) order, and their complete expansion has not been achieved, thus they cannot provide a sufficient number of megakaryocytes for clinical application [62, 63].

PSCs, such as ESCs and iPSCs, offer a solution because they can expand limitlessly in vitro and have the potency to

| Progenitor | Origin | Collection | Amount in source | Source availability | Ethical issues |
|------------|--------|------------|------------------|--------------------|---------------|
| Megakaryocyte | Bone marrow | Invasive | Few | (N.C.) | (N.C.) |
| Hematopoietic stem cell | Cord blood | Non-invasive | Low | Donation-dependent (Limited) | Donation-dependent (Limited) |
| Embryonic stem cell (ESC) | Established from embryo | Non-invasive | Expandable | Donation-dependent (Very limited) | High |
| Induced pluripotent stem cell (iPSC) | Established from somatic cells | Minimally invasive | Expandable | Anyone | Few |

Bold indicate disadvantages; italics indicate advantages
N.C.: not considered
differentiate into essentially any kind of cell type, including hematopoietic lineages (Table 2). ESCs were successfully differentiated into megakaryocytes that could generate platelets in vitro [27, 64–69], an approach later replaced with iPSCs [30, 70] (Table 2). There are several ways to differentiate megakaryocytes from iPSCs. Mimicking embryogenesis, iPSCs are differentiated stepwise to mesodermal, HPC-like cells, and then to megakaryocytes by 2D cultivation using extracellular components [70] or embryoid bodies under 3D cultivation [71]. For the mesoderm and HPC differentiation, humoral factors such as BMP4, VEGF, bFGF, and WNT activators are added. For the megakaryocyte differentiation, TPO and SCF are indispensable but other humoral factors are also required. Feeder cells can also support HPC and megakaryocyte differentiation [30]. Based on these early successes, the amplification of iPSCs to large quantities and then their differentiation to megakaryocytes to derive platelets was considered [70]. However, the differentiation efficiency from iPSCs to megakaryocytes was low. We estimated that this technique would require hundreds or more liters of culture media supplemented with expensive cytokines and other substances for a long culture process, thus demanding high labor and unacceptable cost.

**Establishment of expandable megakaryocytic cell lines through genetic modifications**

Genetic manipulation technology has enabled the induction of megakaryocytes through various ways (Fig. 2a). One approach is the genetic manipulation of iPSCs or HPCs to establish expandable megakaryocyte cell lines. Based on the finding that the transient activation of c-MYC is critical for efficient megakaryopoiesis from human iPSCs [30], we established immortalized megakaryocyte progenitor cell lines (imMKCLs) by the sequential introduction of c-MYC, BMI-1, and then BCL-XL during the differentiation process of iPSC-derived HPCs into megakaryocyte progenitors (Fig. 2b) [72]. Overexpressing c-MYC promotes proliferation, while BMI1 and BCL-XL suppress cell senescence and cell death, respectively. These three transgenes were incorporated into a doxycycline (DOX)-inducible gene expression vector to control their expression. In the presence of DOX, the expression of the transgenes is induced to endow imMKCLs to vigorously proliferate (DOX-ON). Then, when switched to DOX-free media, the expression of the three diminishes, which turns imMKCL from the proliferation state to maturation state, and platelets are released three days later (DOX-OFF) (Fig. 2b). We were able to expand imMKCLs effectively in static or rocking-motion type bioreactors [73]. Another group used a single-use bioreactor to expand commercial megakaryocytes through automated gas exchange and orbital shaking, which might be also useful [74].

Meanwhile, other groups have sought to activate transcription factors that determine hematopoiesis and megakaryopoiesis to induce megakaryocytes (Fig. 2a). Ono et al. reported the conversion of fibroblasts into megakaryocytes by introducing a set of three genes: NF-E2, MAFG and MAFK [75]. Later, Pulecio et al. reported similar transdifferentiation by a set of six genes: GATA2, RUNX1, GATA1, TAL-1, LMO2 and c-MYC [76]. Using another overexpression approach but with a different
concept, Moreau et al. established an expandable megakaryocyte cell line, forward programmed MKs (fopMKs) [71]. Based on a screening of candidate megakaryocyte-specific transcription factor genes, GATA1, FLI1 and TAL1 were chosen to be introduced into PSCs two days before the differentiation into megakaryocytes. fopMKs can be cultivated in the absence of feeder cells and highly expanded in the presence of TPO and SCF. Then, by changing the cytokines to TPO and IL-1α, fopMKs mature to release platelets.

imMKCL and fopMKs can be cryopreserved, thawed upon demand, expanded and then induced to produce platelets upon switching to the maturation stage (Fig. 3). The issue of the low differentiation efficiency from iPSCs to megakaryocytes is removed, and the number of days required for the production is reduced. In principle, a clone that is confirmed for safety and high platelet productivity can be banked in advance for master cells, the source cells to start production. Then, after thawing, these cells would be subject to a standard operating procedure (SOP) that complies with good manufacturing practice (GMP) to stably produce platelets ex vivo with guaranteed clinical-grade quality (Fig. 3).

Besides genetic manipulation, Tozawa et al. reported that a human adipose tissue-derived mesenchymal stromal/stem cell line (ASCL) can produce platelets upon megakaryocyte induction (Fig. 2a) [77]. Interestingly, because of the unique ability of these ASCLs to produce endogenous TPO through transferrin stimulation, TPO does not need to be added to the media. Meanwhile, Patel et al. reported a scaled-up induction of cryopreservable megakaryocytes from human cord blood HSCs through an optimized culture condition [78]. While these induced megakaryocytes have not produced platelets at the clinically required quantity and quality, the findings may contribute to improved ex vivo platelet production.

**Development of drugs that contribute to megakaryocyte maturation and platelet biogenesis**

TPO and SCF are an indispensable combination of cytokines for the in vitro differentiation of iPSCs to megakaryocytes [70–72]. Concerns for using recombinant TPO are its high cost and immunogenicity. Indeed, a clinical trial of recombinant TPO for patients with thrombocytopenia was discontinued due to the formation of antibodies against TPO in general, leading to even lower platelet count than before the administration [79, 80]. In response, TPO mimetics have been developed for clinical use [81]. We developed TA-316, a low-molecular compound with effects higher than current TPO mimetics and comparable to recombinant TPO in vitro [82]. These TPO mimetics could be used as an alternative to recombinant TPO for platelet production ex vivo. Other cytokines such as interleukin-1 (IL-1) and IL-6 have also been used [83].

There have been many substances and conditions reported to enhance megakaryopoiesis and thrombopoiesis. Chemotactic factors secreted from endothelial and other cells in the bone marrow niche induce megakaryocyte migration and proplatelet formation. These include the secreted molecules stromal-derived factor 1 (SDF-1), sphingosine 1-phosphate (S1P) and chemokine ligand 5 (CCL5, or RANTES) [84–86]. WNT family proteins are also involved in megakaryocyte proliferation, maturation and proplatelet formation [87]. Nicotinamide increases the polyploidization and proplatelet formation by increasing TP53 activity [88]. Src-family kinases inhibit megakaryocyte proliferation and differentiation and either a Src inhibitor or a multi-kinase inhibitor induces polyploidization [89, 90]. Rho A and the downstream Rho-associated protein kinase (ROCK) contributes to actin function, and ROCK inhibitors have been reported to enhance megakaryocyte polyploidization and proplatelet formation [91–94]. A myosin inhibitor showed a similar effect [95]. A bromodomain and extraterminal motif protein (BET) inhibitor enhanced megakaryocyte...
maturation through the inhibition of c-MYC and enhancement of GATA1 expression [70]. Aurora B kinase is indispensable for megakaryocyte endomitosis, but its inhibitor has been reported to increase megakaryocyte ploidy, size and granularity [96, 97]. Aryl hydrocarbon receptor (AhR) antagonists are known to enhance HSC expansion [98–100] as well as increase the ploidy of megakaryocytes and the proportion of platelet-producing megakaryocyte precursors during in vitro differentiation [101]. In addition, hypercholesterolemia correlates with increased platelet biogenesis and function [102, 103]. Finally, hypoxia is postulated to exist in the bone marrow and reported to increase platelet production in vitro [104], and mild hyperthermia of 39 °C is also reported to enhance platelet production in vitro [105].

The combinations of different chemicals have been tested by many groups to enhance megakaryocyte maturation and eventual platelet production. For instance, Avanzi et al. found that the combination of nicotinamide, an Src inhibitor and Aurora B inhibitor was effective [96]. Feng et al. incorporated hypoxia, hyperthermia as well as a ROCK inhibitor and BET inhibitor [70]. In an in vivo megakaryocyte infusion setting, pre-infusion treatment with a Src inhibitor enhanced the platelet yield, but not a ROCK inhibitor or an Aurora kinase B inhibitor [97]. Through an assessment of relevant substances or drugs, we found that the combination of a ROCK inhibitor and an AhR antagonist enables platelet generation without feeder cells for imMKCL cultivation [73]. For further high-throughput screening of drugs that enhance megakaryocyte maturation, we developed an imMKCL with a fluorescent reporter that correlates with the expression of a maturation-specific protein [106].

During the in vitro generation of platelets, the metalloproteases ADAM17 and ADAM10 are activated at 37 °C, the temperature at which cultivation is suitable. ADAM17 and ADAM10 cleave GPIbα (CD42b) and GPVI on the surface of platelets, respectively [107, 108], rendering the platelets to become like those in thrombasthenia, a series of diseases with significantly reduced platelet functionality due to mutations in genes including GPIbα and GPVI [107–109]. Matrix metalloproteinase (MMP) inhibitors have been utilized to inhibit these cleavages [110, 111], with an MMP8-specific inhibitor, MMP8-I, superior to a pan-MMP inhibitor, GM6001, in suppressing this effect [70]. We developed KP-457, which prevents the cleavage activity of ADAM17 and ADAM10, thus retaining the expression of GPIbα and GPVI on imMKCL-derived platelets [112].

The role of hydrodynamic forces in platelet generation from mature megakaryocytes and the development of bioreactors

Based on the live imaging analysis of proplatelet extension and platelet release in mouse bone marrow, Junt et al. concluded that shear stress caused by blood flow is the primary biophysical determinant for platelet release from megakaryocytes in vivo [47]. In fact, a simple study using flow chambers coated with vWF or extracellular matrix (ECM) showed that a high shear rate accelerated the in vitro platelet production from megakaryocytes derived from cord blood or bone marrow HSCs [113]. Accordingly, many groups have attempted to mimic the bone marrow environment for megakaryocytes in vitro through systems that are designed to have megakaryocytes extending proplatelets into artificial flow streams (Table 3).

Sullenbarger et al. created a 3D system in which cord blood cells were cultured in polyester fabric or hydrogel scaffolds [114]. In the material was TPO and/or fibronecin, and MK differentiating medium and gasses were continuously perfused. Compared to 2D systems, platelets were produced for a longer period of time and at a higher yield. We developed a microfluidic chip system in which PSC-derived megakaryocytes are trapped in slits [115].

### Table 3 Proposed systems for platelet bioreactors

| Reference          | Design                        | Platform         | Extracellular component                  | Endothelial cell |
|--------------------|-------------------------------|------------------|------------------------------------------|------------------|
| Sullenbarger et al. [114] | Continuously perfused 3D compartment | Hydrogel         | Fibronectin                               | None             |
| Thon et al. [117]      | Microfluidic chip with pores   | Silicon-based organic polymer | Fibrinogen, Fibronectin, Matrigel         | HUVEC cells      |
| Di Buduo et al. [118] | 3D vascular microtubes         | Vascular tube in silk protein sponge | Collagen, Fibronectin, Fibrinectin, Laminin | Primary endothelial cells |
| Blin et al. [119]       | Microfluidic chip with pillars | Silicon-based organic polymer | vWF                                       | None             |
| Avanzi et al. [120]     | Pseudo-3D membrane            | Nanofiber membrane       | -                                         | None             |
| Martinez et al. [116]   | Microfluidic chip with slits   | Silicon-based organic polymer | -                                         | None             |
| Ito et al. [73]         | Vertical-type shaker           | Tank               | (Endogenously included)                   | None             |
| Fujiiyama et al. [121]  | Closed bone culture circuit    | Isolated porcine thigh bone | (Endogenously included)                   |                  |

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Once trapped, the cells extend proplatelets from the pore to incur shear flow. Later, Martinez et al. designed slit-type bioreactors with optimal flow patterns and uniform shear stress based on extensive computational fluid dynamics analysis [116]. Thon et al. designed a microfluidic chip-based bioreactor that incorporates chemical and physical components of the bone marrow such as stiffness, the ECM, pore size, endothelial cells and shear stress flow [117]. Additionally, their bioreactor is capable of live-cell microscopy to observe the PSC-derived megakaryocytes extending proplatelets from the pores. To faithfully recapitulate the bone marrow niche environment, Di Buduo et al. created an artificial bone marrow niche system made from vascular tubes in natural silk protein sponges harboring cytokines, ECM components and endothelial-derived proteins [118]. Blin et al. built a different-type of a microfluidic device in which multiple vWF-coated micropillars stand within a wider channel to capture megakaryocytes. The cells extend proplatelets and release platelets depending on the controlled shear rates [119]. Avanzi et al. designed a bioreactor with a nanofiber membrane, through its loose texture megakaryocytes extend proplatelets into. They used this bioreactor to assemble a stepwise platelet production line from cord blood cell-derived CD34+ cells [120]. Fujiyama et al. used isolated porcine thighbone to create a closed culture circuit in a xenogeneic bone environment [121]. However, despite the many types of bioreactors, the yield of platelets is estimated to be no more than 20–30 per megakaryocyte. This number falls far short of the 800–2000 platelets per megakaryocyte in vivo [53, 54], and the quality of the produced platelets are inferior.

To readdress the hydrodynamic factors involved in platelet biogenesis, we performed in vivo live imaging analysis of mouse bone marrow megakaryocytes with flow dynamic visualization and observed that megakaryocytes actively generating platelets were exposed to turbulent flow [73]. We then tested a vertical reciprocal-motion stirrer, in which turbulent flow is generated, to cultivate imMKCLs at the maturation stage and succeeded in dramatically improving the yield and quality of iPSC-PLTs [73]. Simulation analysis identified shear stress and turbulent energy as key parameters of the turbulent flow conditions. By adjusting these two parameters, the same turbulent flow conditions were applied to an 8 L-scale bioreactor, which enabled the production of approximately 10^11 competent iPSC-PLTs, half the number needed for a single transfusion [73]. Interestingly, the megakaryocytes in the new bioreactor released soluble factors that autonomously promoted the iPSC-PLT production [73]. Two of these, IGFBP2 and MIF, promoted the release of ECM proteins from the megakaryocytes. This release could create a scaffold-like condition for the megakaryocytes. Another soluble factor, NRDC, promoted the cleavage of platelet precursors from elongated proplatelets.

Alternatively, instead of pursuing bioreactors to produce platelets ex vivo, the intravenous infusion of megakaryocytes differentiated from cord blood HSCs has been proposed [122, 123]. Animal models have shown that the infused megakaryocytes can reach the lungs, where they are trapped by the capillary bed to produce platelets as endogenous megakaryocytes have been observed to do [50]. The potential disadvantages of this approach, however, are the delayed circulation of platelets compared with the direct infusion of platelets and the unknown safety and immunogenicity profiles of the infused nucleated megakaryocytes. Furthermore, it is difficult to continuously supply cells from the same source, because cord blood HSCs cannot be expanded like PSCs. However, this approach omits the necessity of producing platelets ex vivo, and the lung-generated platelets may be more physiological in nature.

**Platelet purification and packaging**

To prepare iPSC-PLTs into a transfusable format, the produced platelets must be washed, purified and resuspended in storage solution with appropriate conditions. Based on differences in size and specific gravity, filtrations and continuous centrifugation can be applied for the washing, purification and concentration before resuspension [73, 124]. Bicarbonate Ringer’s solution added with ACD-A solution (BRS-A) is used for the washed platelet products [125] and could also be used as a resuspension solution for ex vivo platelets. Other solutions such as PAS solutions recently tested for blood products may also be applied [126]. Given the bacteria-free production procedure for iPSC-PLTs, the development of solutions that retain platelet function could contribute to significantly extending the shelf life of iPSC-PLTs, which is currently only 4 or 5 days, and reduce the difficulty in balancing the supply and demand.

**Preclinical studies of the safety and efficacy of iPSC-PLTs**

**General overview and in vitro studies**

Before clinical trials take place, rigorous preclinical tests to assess the safety and efficacy are required from regulation authorities (Fig. 4). A master cell bank, i.e. cell stock, that will be used as the original source for each batch of platelet production, whether cord blood HSCs, PSCs or megakaryocyte lines, will need to be thoroughly tested for viruses and any other infectious pathogens. A platelet production process in compliance with GMP grade from a certified pathogen-free master cell bank will assure products devoid of infection. Safety of the final platelet product needs to be tested in animals. The toxicity and mutagenicity of any additives
used in the production process need to be tested in animals and in vitro unless previously proven safe or the final concentration of the additives are below the level determined by the International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use (ICH).

A safety parameter always tested in regenerative medicine is tumorigenicity, which applies to iPSC-PLTs as well (Fig. 4). Platelets themselves are anucleate cells, which means they cannot proliferate. However, the batch may include contaminating nucleated cells. iPSCs form teratoma upon transfer in vivo [2–4], and expandable megakaryocytes may also proliferate after infusion [71, 72]. Chances are extremely low that PSCs remain in the long cultivation for megakaryocyte differentiation and maturation, but confirmation assays are still required. On the other hand, depending on the method, megakaryocytes with proliferation potential could substantially remain in the final platelet suspension [73]. These nucleated cells may also accumulate oncogenic mutations during the culture [127, 128]. Membrane filtration and centrifugation can reduce the number of nucleated cells, but not completely. To eliminate the risk of tumorigenicity, irradiation similar to that used to prevent GVHD after transfusion with current blood donation products [129] could be performed and should not affect the function of the platelets. Even if these procedures are taken, however, the risk of tumorigenicity still needs to be confirmed through in vitro culture and in vivo administration tests.

Platelets can be activated through various mechanisms, and thus are easy to deteriorate or aggregate by stimulation including low temperatures and physical pressures [126]. Therefore, it is crucial to evaluate the functional viability of the produced platelets (Fig. 4). Various in vitro measures include flow cytometry analysis of the CD42b and CD62P expressions, Annexin V binding, and electron microscopy observation.

Platelets can be activated through various mechanisms, and thus are easy to deteriorate or aggregate by stimulation including low temperatures and physical pressures [126]. Therefore, it is crucial to evaluate the functional viability of the produced platelets (Fig. 4). Various in vitro measures include flow cytometry analysis of the CD42b and CD62P expressions, Annexin V binding, and electron microscopy observation.

Animal models for in vivo studies of efficacy

For infused platelets to be effective in vivo, they need to circulate well and show hemostasis function at the bleeding site but at the same time not form unnecessary thrombi in the lungs or other organs. In humans, platelets are considered to have up 10 days of life, after which they are trapped and destroyed by splenic macrophages. However,
in thrombocytopenia patients, the transfused platelets have a shorter lifetime, sometimes 3 days or less according to radiolabeling studies [132].

In vitro functionality tests are ultimately not sufficient to assess the in vivo functionality of platelets, therefore, to test the circulation and hemostasis function for the preclinical studies, animal models are adopted [Fig. 4]. Using such animal models, the function of the platelets of interest could be compared with donor-derived product platelets. Immunodeficient mice have been used to assess circulation by flow cytometry and hemostasis through tail bleeding [73, 133, 134]. Radiation or treatment with anti-mouse CD41 antibodies is used to prepare thrombocytopenic conditions. In addition, mice with laser-induced vascular injury have been used to observe the incorporation of infused platelets by fluorescence intravital microscopy, [68, 72].

In addition to mice, rat, guinea pigs and dogs are also used for in vivo evaluations, but rabbit models are most commonly used, owing partly to the ease of assessment through the ear veins [135, 136]. Rabbits are pretreated with ethyl palmitate to suppress clearance by the reticuloendothelial system. Cytotoxic drugs such as busulfan are used to create thrombocytopenic conditions. Our collaborator found splenectomy improves the circulation [137]. Notably, rabbit and rat kidney or liver injury models have been used to assess hemostasis in massive visceral hemorrhaging for various platelet substitutes [130].

As mentioned above, iPSC-PLTs have a fetal/neonatal nature, which may give them longer survival, but also hyporeactivity compared with adult-type platelets [44, 45]. However, we observed iPSC-PLTs and blood donor-derived platelets have generally the same level of in vitro and in vivo functionality [73]. iPSC-PLTs are initially larger but become smaller after the transfusion and show a temporary increase in circulation number. We speculate that the younger, larger platelets become fragmented in the lung upon the circulation.

**Applying iPSC-PLTs to alloimmune platelet transfusion refractoriness**

**Autologous product as a complete match source**

As alloantigens, platelets express ABO antigens, human leukocyte antigen class I (HLA-I) and human platelet antigen (HPA). The production of alloantibodies against these antigens causes platelet transfusion refractoriness (allo-PTR), which appears as a poor increase of post-transfusion platelet count in transfused patients [14]. The contribution of ABO antigens is considered to play only a minor role in the platelet transfusion settings but may affect significantly platelets that highly express them [138, 139]. Alloantibodies against non-self HLA-I and HPA are produced through pregnancy or previous platelet transfusions or allotransplantations. Measures such as the leukoreduction of blood components or the use of single donor-derived products rather than multiple donor-derived pooled products have lowered the incidence of allo-PTR from about 15% to about 5% of patients receiving transfusions [14].

HLA-I is the major cause of allo-PTR, while HPA causes other forms of alloimmune responses, including post-transfusion purpura (PTP), in which the post-transfusion platelet count somehow becomes even lower several days after the transfusion, and fetal and neonatal alloimmune thrombocytopenia (FNAIT), in which platelets are cleared from infants by maternal anti-HPA alloantibodies [14, 140]. In patients with allo-PTP, only platelet products that match these alloantigens are effective, but such products may not be available for rare HLA-I or HPA or in emergencies.

iPSC-PLTs can cope with allo-PTP in several ways (Fig. 5). First, autologous iPSC-PLTs can be produced by establishing iPSCs from allo-PTP patients themselves. In 2019, we started a clinical trial of autologous iPSC-PLTs for an aplastic anemia patient with allo-PTP who had no matching blood donor in Japan [141]. Even from aplastic anemia patients, skin fibroblasts or blood lymphocytes can be used to establish iPSCs and produce iPSC-PLTs. This approach is also ethically highly compatible since it should not elicit an alloimmune response or blood-borne infections [142]. The limitation of autologous iPSC-PLTs, however, is that the iPSC production process is time-consuming, labor-intensive, and too costly for every patient. In addition, the production efficiency and quality of the platelets could vary depending on the source individual and iPSC line. Therefore, good autologous iPSC-PLTs cannot always be assured in a timely manner.

**Allogeneic HLA-homozygous haplotype products as a stocked source**

To overcome the limitations in the cost and availability of autologous iPSC-PLTs, premanufactured off-the-shelf allogeneic iPSC-PLTs from megakaryocyte cell lines that were confirmed to expand and produce platelets with high efficiency could be applied (Fig. 5). Allogeneic preparations are subject to strategies that overcome allogeneic immune responses. As for the ABO antigen, O-type donors could be selected. For HLA class I, good donors for the iPSCs are people with homozygous HLA. HLA genes are located as a cluster within chromosome 6p21, and two sets of HLA haplotypes exist in each person. HLA-homozygous grafts are compatible with a wide range of recipients because matching just one of the two haplotypes of the recipient evades rejection. In fact, blood donors with homozygous HLA are often registered and called upon when there is a demand for
Generation and manipulation of human iPSC-derived platelets.

HLA-compatible platelets. Based on this wide compatibility of HLA-homologous grafts, iPSCs from HLA-homozygous subjects or cord blood are being stocked for regenerative medicine use [143, 144]. These iPSCs can be used to produce iPSC-PLTs. It is estimated that iPSC lines expressing the most frequent HLA-I would cover 50%, 80% and 90% of the Japanese population, respectively [144].

**HLA-deficient products as universal products**

With regards to HLA, platelets nullified of HLA class I could become a universal product [70, 145–148], and the mass production of one line would significantly lower the cost compared with the preparation of a product library (Fig. 5). HLA-I can be either depleted (knockout; KO) using CRISPR/Cas9 or other gene-editing technologies or downregulated (knockdown; KD) using RNA silencing technologies. One common KO and KD target has been beta2-microglobulin (B2M), a molecule composing the HLA-I heterodimer complex and required for the cell surface expression of all HLA-I family molecules [70, 145–148].

Another target would be HLA-ABC, because antigens of anti-HLA-I antibodies in allo-PTR are mostly HLA-A and HLA-B, with rare cases of HLA-C [149].

A general concern in HLA-I deficient products is that the total elimination of HLA-I could elicit rejection by natural killer (NK) cells because HLA-I molecules are inhibitory ligands of killer immunoglobulin-like receptors (KIRs) and CD94/NKG2 on NK cells [150, 151]. As means to avoid rejection by NK cells, KD, which can retain a low but not null expression level, is suggested. Other proposals include overexpressing HLA-E [152] or retaining the expression of HLA-C [153]. Interestingly, we found that B2M-KO iPSC-PLTs, which are completely devoid of HLA-I expression, do not elicit an NK cell response in vitro [147]. To confirm the circulation in vivo in the presence of NK cells, we developed humanized immune system (HIS) mice, which were reconstituted with a sufficient number of human NK cells to reject B2M-knockout iPSC-derived hematopoietic cells. B2M-KO iPSC-PLTs circulated comparably with wild-type iPSC-PLTs in the NK cell reconstituted HIS mice and also circulated in the presence of anti-HLA-I antibody [147]. While this observation removes the concern of NK cells rejecting HLA-KO platelets, the reason for the low immunogenicity of platelets remains unknown.

Low immunogenic O-type HLA-depleted iPSC-PLTs are suitable as a common platform for products with further genetic modifications or drug loading to endow an alternative or a new phenotype (Fig. 5). As for HPA incompatibility, the deletion strategy used for HLA-I cannot be applied, because HPA is a group of molecules essential for platelet function. Given that allo-PTR due to HPA frequently
involves HLA-I as well, it would be appropriate to convert HPA on HLA-depleted iPSC-PLTs to the desired HPA type by gene modification [154].

Meanwhile, platelets are known to have functions besides hemostasis, such as contributing to the maintenance of vascular integrity, vascular and lung development in newborns, tissue regeneration, anti-bacterial immunity and anti-malarial immunity, and are also involved in the pathogenesis of rheumatoid arthritis and cancer [155]. Taking these into account, platelets have been studied as carriers to incorporate coagulation factors [156–158], anti-cancer drugs [159, 160], and oncolytic virus [161]. Platelets enable these factors to circulate in an encapsulated form and deliver to the sites they are attracted such as bleeding and cancer [162]. These substances could be loaded onto HLA-KO iPSC-PLTs through a defined process including genetic manipulation [163], providing a novel drug delivery system of standardized quality.

Conclusion

The discovery of iPSCs has brought new possibilities in regenerative medicine, but to reach clinical application, many qualitative and quantitative issues needed to be addressed. Platelets derived from iPSCs have reached a clinical trial owing to the development of expandable megakaryocyte lines, hydrodynamically designed bioreactors and clinical trial owing to the development of expandable megakaryocyte lines, hydrodynamically designed bioreactors and novel drugs. Given that HLA-null universal iPSC-PLTs are suitable as a mass commercial product, transfusion medicine could evolve to a new generation, assuring a ubiquitous and safe supply of platelet products. Furthermore, iPSC-PLTs are amenable to other genetic manipulations, which may bring forth novel therapeutic measures.

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