Transfusion Medicine and Hemotherapy

Review Article

Transfus Med Hemother 2013;40:326–335 DOI: 10.1159/000354061 Received: May 23, 2013 Accepted: July 1, 2013 Published online: August 26, 2013

Platelet Lysate as Replacement for Fetal Bovine Serum in Mesenchymal Stromal Cell Cultures

Karen Bieback

Institute of Transfusion Medicine and Immunology, Medical Faculty Mannheim, Heidelberg University, German Red Cross Blood Service Baden-Württemberg – Hessen, Mannheim, Germany

Keywords

Platelet lysate · Platelet releasate · Mesenchymal stromal cells · Fetal bovine serum · Good manufacturing practice

Summary

Mesenchymal stromal cells (MSC) emerged as highly attractive in cell-based regenerative medicine. Initially thought to provide cells capable of differentiation towards mesenchymal cell types (osteoblasts, chondrocytes, adipocytes etc.), by and by potent immunoregulatory and pro-regenerative activities have been discovered, broadening the field of potential applications from bone and cartilage regeneration to wound healing and treatment of autoimmune diseases. Due to the limited frequency in most tissue sources, ex vivo expansion of MSC is required compliant with good manufacturing practice (GMP) guidelines to yield clinically relevant cell doses. Though, still most manufacturing protocols use fetal bovine serum (FBS) as cell culture supplement to isolate and to expand MSC. However, the high lot-to-lot variability as well as risk of contamination and immunization call for xenogenic-free culture conditions. In terms of standardization, chemically defined media appear as the ultimate achievement. Since these media need to maintain all key cellular and therapy-relevant features of MSC, the development of chemically defined media is still - albeit highly investigated - only in its beginning. The current alternatives to FBS rely on human blood-derived components: plasma, serum, umbilical cord blood serum, and platelet derivatives like platelet lysate. Focusing on quality aspects, the latter will be addressed within this review.

Platelet-Derived Factors for Cell Culture and Tissue Regeneration

Ex vivo/in vitro cell culture requires basal medium plus supplements providing growth factors, proteins, and enzymes to support attachment, growth, and proliferation. Fetal bovine serum (FBS) is commonly used to supplement cell culture media, because the fetal milieu is enriched in growth factors compared to the adult situation and poor in antibodies [1]. In contrast to plasma, serum contains a variety of growth factors, cytokines, and chemokines derived during blood coagulation and released by physiologically activated platelets [2]. Beside stop of bleeding, these factors mediate wound closure and healing. Studies in the 1980s defined growth-promoting effects of human platelet lysate (HPL) on various cell lines [3], tumor cells [4], and articular chondrocytes [5]. Especially the α-granule-derived factors such as platelet-derived growth factor (PDGF), transforming growth factor- β (TGF- β), insulinlike growth factor (IGF), vascular endothelial growth factor (VEGF), fibroblast growth factor (FGF-2/bFGF), hepatocyte growth factor (HGF), and epidermal growth factor (EGF) have been identified as cell mitogens with wound-healing capacity [6-8]. In conjunction with extracellular matrix molecules platelet factors, for instance TGF-\$1, provide osteoinductive capacity for osteoblasts [6] (fig. 1). To utilize these physiological functions, platelet-derived factors have been applied as therapeutic agents for wound healing and bone regeneration. To concentrate the factors, platelet-rich plasma (PRP) has been developed. This is derived by centrifugation of anticoagulated whole blood, yielding plasma enriched in platelets. Further concentration can be achieved by a second centrifugation step [6, 9]. Subsequent coagulation with calcium forms a fibrin gel, which, in conjunction with the platelet released growth factors, serves as a therapeutic agent in plastic surgery, orthopedic interventions, chronic wound healing,

KARGER

Fax +49 761 4 52 07 14 Information@Karger.com www.karger.com © 2013 S. Karger GmbH, Freiburg 1660-3796/13/0405-0326\$38.00/0

Accessible online at: www.karger.com/tmh PD Dr. Karen Bieback Institute of Transfusion Medicine and Immunology Medical Faculty Mannheim, Heidelberg University German Red Cross Blood Service Baden-Württemberg – Hessen Friedrich-Ebert Straße 107, 68167 Mannheim, Germany Karen.bieback@medma.uni-heidelberg.de



ophthalmology etc. [6, 9–13]. Despite the manifold interventions, evidence of effectiveness of PRP administration remains controversial: Sommeling et al. [10] reported a significant benefit in several indications, including wound healing as well as fat and bone grafting, whereas Martinez-Zapata et al. [13] found no evidence for a benefit of PRP in chronic wound treatment, similar to Sheth et al. [12], who state 'the current literature is complicated by a lack of standardization of study protocols, platelet separation techniques, and outcome measures'.

Certainly platelet-derived factors attracted attention as effective tool to supplement cell cultures replacing FBS. To improve safety for cell-based therapies Doucet et al. [14] initiated the use of HPL in supplementing mesenchymal stromal cell cultures.

Mesenchymal Stem/Stromal Cells

In the 1970s Friedenstein et al. [15, 16] described a population of non-hematopoietic progenitors isolated from the bone marrow. Although already described as being able to differentiate into a variety of mesodermal lineages such as bone, cartilage, fat, marrow stroma, tendon, muscle, dermis and connective tissues, further research rested until the 1990s [17, 18]. Only then, these non-hematopoietic cells were termed 'mesenchymal stem cells (MSC)', according to the hematopoietic stem cell (HSC) nomenclature [19]. Since later studies, however, failed to fulfill the self-renewal criterion of stem cells (1. self-renewal, 2. unspecialized, and 3. differentiation capacity to specialized cell types), it has been proposed to better name the cells 'mesenchymal stromal cells (MSC)' [20].

Of note, till now MSC are characterized as culture-adapted, ex vivo expanded cells. This population is still heterogeneous and contains progenitor cells at different maturation stages and also mature stromal cells [21]. In conjunction with the heterogeneity of cell preparations, inconsistencies in the use of different tissues as starting material and in isolation and cultivation protocols render comparability of results complicated. As an effort to standardize the term MSC, the International Society for Cell Therapy (ISCT) defined minimal criteria to be fulfilled [22]:

- adherence to cell culture plastic surfaces yielding cells of a fibroblastoid phenotype,
- expression of typical markers (CD105, CD73 and CD90) and lack of expression of CD45, CD34, CD14 (or CD11b), CD79α (or CD19) and HLA-DR surface molecules, and
- differentiation towards at least the three mesodermal chondrocyte, adipocyte and osteocyte lineages.

(for further overview see TRANSFUSION MEDICINE AND HEMO-THERAPY special issues Vol. 35, No. 3 and 4, 2008, and Vol. 37, No. 2, 2010 [23–41]).

Encouraged by this broad differentiation capacity, early clinical trials were initiated for a wide range of ailments [42, 43]. Based on the stromal origin within bone marrow, the stromal support capacity to facilitate HSC engraftment was assessed [44, 45]. Despite an unexpectedly low level of engraftment, a long-lasting therapeutic effect became apparent. To answer the question how MSC achieve this benefit without being actually present, MSC research returned 'back to bench' [46]. Now seminal studies identified strong immunomodulatory properties of MSC [47, 48]. In combination with their low immunogenicity based on the lack of HLA-DR and co-stimulatory molecules, MSC were rated well-suited for both autologous and allogeneic transplantation settings [49]. Continuatively the beneficial therapeutic effects could be attributed to the capacity of MSC to home to sites of inflammation and injury where MSC release a variety of pro-regenerative, anti-apoptotic and anti-fibrotic factors enabling endogenous repair processes [50].

Thus based on their differentiation capacity, hematopoietic support as well as their immunomodulatory and pro-regenerative features, MSC are increasingly applied in cell-based therapy: currently 307 clinical trials are found searching for 'mesenchymal stem cells' and 32 for 'mesenchymal stromal cells' (*http://clinicaltrials.gov*; assessed June 2013). The high number of clinical trials raise an intense controversy regarding the pace of translation [51]. Despite fascinating in vitro results and promising preclinical data, clinical trials appears often rather weak, because, despite intense research, the mode of action of MSC remains elusive. Thus it is essential to carefully weigh the putative benefits with the risks.

Although, up to now, the majority of preclinical and clinical data fail to report severe adverse events after MSC application, suggesting that MSC can be applied safely [52], there have been reports of severe complications after stem cell infusions, including deaths [51, 53, 54]. In Korea, for instance, one patient died from pulmonary embolism after stem cell administration, and the International Cellular Medicine Society (ICMS) considered it 'likely to have been caused or triggered by the stem cell procedure' (www.cellmedicinesociety.org). Yet beyond these few cases possibly related to market-oriented companies that 'are setting up operations around the globe, and taking advantage of loopholes in other countries' regulations' [53], few adverse events have been reported. Calcifications were observed after transplantation of MSC or unfractionated bone marrow into infarcted hearts in a mouse model [55]. Tumor growth appeared facilitated after MSC infusion [56, 57]. Especially latent tumors, such as gliomas, sarcomas and melanomas, but also metastases became manifest after MSC infusion. This suggests that tumor surveillance may be impaired by MSC immunosuppressive activities or that direct integration into the tumor stroma and secretion of angiogenic factors may facilitate tumor growth [57]. It has been intensely debated whether or not the ex vivo expansion of MSC, which in most cases is needed to achieve clinically relevant cell numbers, provokes spontaneous transformations [58, 59]. The few cases reporting spontaneous transformation, however, had to retract their publications: 'spontaneous' transformation occurred due to laboratory cross-contamination with tumor cell lines. Thus yet consensus evolved that at least human MSC in general undergo replicative senescence instead of malignant transformation. Even documented events of aneuploidity, observed in few clinical-scale MSC preparations, in all cases caused progressive growth arrest and senescence [60]. Yet, by way of precaution, consensus on common standards and harmonized protocols need to be implemented to strengthen both efficacy and safety.

Towards Xenogenic-Free Manufacturing of Mesenchymal Stromal Cells

To ensure safety and efficacy, all steps within the MSC manufacturing process need to be standardized. Cellular quality and potency have to be reproducible. However, the manufacturing process for MSC is complex and composed of procurement, isolation, expansion, and quality control [37, 51, 61, 62]. Clinical success is linked to a sufficient number of vital and functional cells. Further, the cellular product shall bear no risks for infections, allergies, or malignancies. Thus within the manufacturing process, a number of considerations have to be taken into account [63]. Ancillary reagents pose a risk for the safety of the cellular product. FBS is typically used to isolate and expand MSC. It is a highly complex mixture of proteins and other factors and by nature ill-defined and variant from batch to batch [1]. Due to the high risk of contaminations (virus positivity reported to be as high as 20–50%), FBS is critically rated by the European Medicines Agency [64]. Because MSC internalize xenogenic proteins at high amounts, there is an additional risk of allergic reactions. FBS immunogenicity has already been demonstrated to compromise the therapeutic success [49, 65-67]. In view of these considerations xenogenic-free culture conditions appear desirable. Due to the numerous constituents of FBS, which positively (and negatively) affect adhesion (cell-cell and cell-matrix), mitosis, survival, apoptosis etc., a chemically defined medium needs an optimal composition of the few most essential factors to promote at least all key cellular features. By this it is hard to establish [68-70].

Although some regulatory agencies may tolerate xenogenic components, such as FBS, in phase I clinical trials, it is expected that later clinical trials including larger patient cohorts will require serum- or at least xenogenic-free cell preparations. Albeit washing procedures or sequential cultivation in human plasma or serum may help to reduce the content of xenogenic proteins, a residual risk is left over [65, 67].

To replace FBS, numerous studies now refer to 'humanized' culture conditions. 'Humanized' supplements include human serum (autologous or pooled allogeneic), cord blood serum as well as different platelet derivatives [69, 70]. Because these human blood components are in clinical use since years, can be derived from healthy blood donors and had been tested according to blood banking standards for infectious and immunological parameters, the potential risk is lowered. Similar to FBS, human blood component-derived supplements include a variety of essential factors capable of promoting cell growth. Human plasma, autologous and allogeneic serum as well as cord blood serum have been investigated as reviewed in [70]. Yet human platelet-derived factors emerge as the most intensely studied alternatives to FBS for MSC culture.

Platelet Releasate

As already stated, human platelets contain numerous factors to promote cell growth of cells and cell lines [3]. Early studies evaluated the MSC growth-promoting effects of platelet growth factors in PRP released by calcium and thrombin stimulation [71, 72]. Coagulated fibrin is subsequently removed by centrifugation and filtration. Both studies described accelerated expansion and migration but differed with respect to osteogenic differentiation potential. Thus platelet factors released by physiological stimuli might offer some advantages. Several substances have been shown to activate platelets, including thrombin, collagen, ADP/epinephrine and thrombin receptor-activating peptide (TRAP) [73]. Interestingly, own studies have shown that processed thrombin-activated platelet releasate in plasma (tPRP) and HPL promote different proliferation rates of bone marrow- and adipose tissue-derived MSC. Whereas HPL promoted a significant higher proliferation rate of bone marrow MSC than tPRP [74], adipose stromal cells exhibited, if at all, similar proliferative responses to HPL and tPRP [74, 75]. Differential proteomics of HPL and tPRP identified 20 differential proteins (Kinzebach et al. unpublished data). Identified proteins further denoted differences between bone marrow and adipose stromal cells: for example, fibrinogen significantly supported the expansion of adipose-derived stromal cells (ASC), and apolipoprotein A1 selectively reduced proliferation rate of bone marrow MSC.

Platelet Lysate

Platelet releasates contain only those factors released after platelet activation. Platelet lysates (HPL) in contrast contain all factors platelets are composed of. These can be easily derived by mechanical disruption of platelet concentrates via freezing and thawing. Subsequent centrifugation separates the platelet debris from the supernatant containing all bioactive platelet factors. Compared to chemical activation of platelets to gain the releasate, mechanical lysis to yield HPL appears preferable as being much easier, less time-consuming, and less cost-effective. It further avoids the use of additional substances such as thrombin which may cause side effects.

Platelet concentrates, used for HPL production, can be either frozen immediately after donation or used at the end of the shelf life (4–6 days after donation depending on the current local regulation) [76–78]. Thus those platelet concentrates not used for platelet transfusion can be allocated for HPL manufacturing, minimizing the decay of already donated units and consequently avoiding an additional donation from blood donors. Platelet concentrates can be then stored frozen, thawed, centrifuged, pooled, and sterile-filtered after quarantine storage (analogous to therapeutic fresh frozen plasma) to increase safety [77]. Final HPL can be stored at -20 °C for a prolonged time, maintaining a stable growth factor content [79].

Doucet et al. [14] have shown that PRP-derived HPL well supports MSC isolation and expansion whilst osteo-, adipoand chondrogenic differentiation propensity is retained compared to FBS. A variety of studies followed, exploiting HPL as GMP-compliant substitute for FBS in MSC expansion, supporting even significantly enhanced proliferation of MSC [73, 74, 80-85]. For large-scale MSC expansion 50-100 conventional tissue culture flasks may be needed. These are hard to handle and prone to contamination. Thus a few studies evaluated the isolation and expansion within a bioreactor using HPL [86, 87] in order to define the risks of MSC therapy. The majority of these studies report a lower size of MSC and accelerated proliferation compared to FBS. Furthermore, HPL increased not only size but also numbers of colonies [82, 88]. Importantly, only in FBS-supplemented media clonal chromosomal instabilities were monitored but not under the humanized culture conditions [59, 82, 89].

Although MSC are considered to escape allo-recognition, mitogens such as FGF and PDGF-BB as well as inflammatory cytokines have been documented to induce HLA-DR expression in MSC and by this the stimulation of CD4 T cells. Importantly HPL appears not to cause HLA-DR expression [90]. Regarding the maintenance of immunomodulatory properties, contradictory data exist. Flemming et al. [91] directly compared bone marrow-derived MSC cultivated in FBS or HPL regarding their immunosuppressive capacities. The inhibitory effect on T-cell proliferation was similar, likewise the activation of cytomegalovirus-specific T cells. Similar data were presented by Bernardo et al. [83]. In contrast, Abdelrazik et al. [92] defined an alteration in surface protein expression relevant for immunomodulation and adhesion after culture in HPL. This corresponded to a reduced inhibition of T and NK cell proliferation. These authors compared MSC expanded in three different batches of FBS (including two commercially available MSC growth media) with HPL supplementation.

Microarray analyses highlighted the down-regulation of several gene families that are included in differentiation/de-

velopment, cell adhesion / extracellular matrix-receptor interaction, TGF-ß signaling and thrombospondin-1-induced apoptosis. Gene clusters associated with cell cycle, DNA replication, and purine metabolism were up-regulated concomitant to the enhanced proliferation in HPL as well as in human serum [84, 93]. Albeit low, these changes in gene expression may cause differences in MSC therapeutic potential. Homing and engraftment are prerequisites for most therapeutic interventions. Changes in cell adhesion molecules thus may have relevant consequences. We detected that the reduced expression of integrin $\alpha 6$ (CD49f) in adipose tissue-derived stromal cells cultivated in a humanized medium correlated to decelerated integrin signaling and reduced adhesion to laminin [94]. We also observed reduced adhesion to endothelial cells. Finally fewer cells cultivated in human serum were detected in the lungs of mice infused with MSC in contrast to higher numbers of those cells cultivated in FBS. Whether this diminished entrapment is related to the lower size of cells, reduced interaction with extracellular matrix molecules, changed homing specificities, or other reasons is a matter of further analyses. Although observed for cells cultivated in human serum, similar ideas are discussed by Lucchini et al. [95]. Thus the choice of supplement may play a critical role when balancing risk and efficacy: reduced adhesion to the lung may lower the potential risk of pulmonary embolism. It may, however, also reduce the release of immunomodulatory TSG-6 (TNF-stimulated gene 6 protein) secreted of lung-adhered MSC to provide protection, e.g., in myocardial infarction [96].

Combining the growth-promoting effect of HPL on MSC, the osteoinductive properties in bone, and the scaffold properties of fibrin, HPL has been used to promote bone tissue engineering. Dozza et al. [97] for instance demonstrated that HPL-expanded MSC applied as collagen or fibrin construct to an uncemented hip prosthesis significantly promoted new bone formation compared to the prosthesis alone. Furthermore, there are indications that HPL may induce osteogenic differentiation without any further osteogenic stimuli. Only ceramics seeded with MSC grown in HPL were able of ectopic bone formation [88, 98]. Based on this differentiation potential, these cells have been used to treat patients in various orthopedic conditions. Centeno et al. [99] reported the results of 339 patients having received autologous bone marrow-derived MSC expanded in autologous HPL. Adverse event surveillance revealed a few cases most probably related to the reimplant procedure and three cases possibly related to stem cell applications. These few cases were either self-limited or cured by small therapeutic interventions. Importantly, no neoplastic transformations were observed at the stem cell injection site. Although in total two patients developed tumors, the neoplasm rate was similar to the control population. 53.1% of the patients reported symptom relief in the follow-up period of 11 months.

Lange et al. [100] observed decreased adipogenic differentiation potential of MSC expanded in HPL. Amongst others, the reduced expression of lipocalin-type prostaglandin D2 synthase was related to this effect. The authors conclude that HPL might offer an option to prevent unwanted adipogenic differentiation.

Quality Criteria for Platelet Lysate

Human supplements, likewise FBS, still need to be considered ill-defined, and they share some safety concerns. Generally platelet concentrates are manufactured and released for therapeutic purposes to be transfused to patients with severely reduced platelet numbers and/or impaired platelet function before they are converted for use as MSC supplement. Having been released as blood product, stringent blood donor eligibility criteria as well as sensitive viral NAT testing have already been fulfilled, ensuring safety of the starting material for HPL manufacturing [77, 78].

Autologous or Allogeneic

Platelet transfusion practices consider the blood groups and rhesus factors to avoid adverse events for the patients. For preparation of autologous MSC, autologous HPL may be considered. However, in severely sick patients a whole blood donation or apheresis may be risky. Due to the restricted volume of HPL from one patient the cell number achieved within the expansion steps might be restricted as well. Furthermore, variations between individual autologous donors hamper standardization and increase the need to quality control each individual HPL batch. Thus the majority of current protocols rely on pooling platelet concentrates from up to 50 healthy blood donors [85]. The cryopreservation step prior to further pooling allows quarantine storage of individual platelet concentrates. These can be released for clinical-grade HPL production once the donors have been retested negative for all infectious markers after a consecutive second donation [77]. The resulting large batch is easy to be quality controlled according to blood banking standards and for protein and growth factor content. For patients at risk, e.g. patients with known antibodies, blood group/Rhesus factor-matching allogeneic platelet preparations could be considered [78]. Interestingly, donor age appears to be of impact for HPL quality: comparing umbilical cord with adult PRP revealed a higher concentration of mitogenic growth factors in the cord blood-derived preparations [101]. Similar HPL from younger donors (<35 years) was more proliferative than that from older donors which increased the expression of senescence markers [102].

Thrombocyte Concentration

Crucial for manufacturing MSC is the concentration of thrombocytes which is directly related to the growth factor concentration. Lange et al. [84] evaluated different thrombocyte concentrations as 5% supplement in basal medium to evaluate the effect on MSC proliferation: 1.5, 1.0, 0.75, and 0.5×10^9 /ml. It became obvious that a platelet concentration below 1.5×10^9 /ml significantly reduced the pro-proliferative effect.

Shelf Life of Platelet Concentrates prior to HPL Production

To reduce the risk of transfusion-transmitted bacterial infections, in Germany the shelf life of platelet concentrates is 4 days. A variety of laboratories allocate the platelet concentrates at the end of this shelf life for HPL production. Bacterial screening complemented by subsequent donor testing is taken as safety measures. Fekete et al. [77] compared HPL prepared from platelet concentrates 2 days or 6 days after donation without any change in quality. Consequently, the platelet concentrate does not need to be discarded when not used for the initial intended therapeutic use but can be converted to HPL production after the shelf life has been exceeded.

Plasma or Thrombocyte Additive Solution

Although most critical, data regarding growth factor concentrations significantly differed in the various publications [79]. These discrepancies are predominantly due to different preparation methods to manufacture platelet concentrates, using either plasma or plasma additive solution. Plasma additive solution has been introduced to reduce the adverse effects of plasma [78]. Initially convinced that plasma components in conjunction with the platelet factors comprise the optimal MSC supplement, we for instance evaluated a pool of freshly prepared buffy coat-derived platelet concentrates prepared in human AB plasma of one of the blood donors [73, 74]. Later we and others tested outdated 'routine' platelet concentrates. As pool of 8-50 donors, no divergence from freshly prepared platelet lysate was obvious, suggesting that outdated platelet concentrates - instead of autoclaving - can be cryopreserved to manufacture HPL [77, 85]. For instance the group of Dirk Strunk filed an US-patent entitled 'Plasmafree platelet lysate for use as a supplement in cell cultures and for the preparation of cell therapeutics' (Pub. No.: US 2009/0305401 A1; Dec 10, 2009).

Buffy Coat- or Apheresis-Derived Platelet Concentrates

Within their study Fekete et al. [77] furthermore compared GMP-grade HPL obtained from pooled whole blood-derived buffy coats with those from apheresis-derived platelet concentrates. There were no significant differences regarding the cytokine content (bFGF, sCD40L, PDGF-AA, PDGF-AB/BB, sVCAM-1, sICAM-1, RANTES, TGF- β 1) fitting to the similar support of MSC proliferation.

Specific Cytokines

As already mentioned, HPL contains a plethora of cytokines, chemokines, and soluble adhesion molecules which have been intensely studied by proteomic approaches [8, 82, 103]. However, concentrations between the groups differ significantly [77, 79]. And those growth factors essential for MSC isolation and expansion have not been defined yet. It needs to be established whether the composition and concentration of platelet-derived factors can serve as quality control parameters for HPL.

FGF2/bFGF, PDGF, EGF, IGF, and TGF, among others, have been studied but failed to support MSC growth when supplemented solitary or in combination to serum-free medium [104]. Yet added to FBS-supplemented media, these growth factors were capable of fostering proliferation and differentiation. This effect, however, may not necessarily hold true in 'humanized' culture systems. Similar to our own yet unpublished data, bFGF appears to support expansion of FBS-supplemented bone marrow MSC cultures, but fails to do so in HPL-supplemented systems [105]. Neutralization of PDGF-AB/BB, TGF-β1 and FGF significantly diminished the proliferative effect of HPL, with the strongest effects seen by neutralizing FGF alone or in combination with PDGF-BB [77]. Although these factors appear to be essential, it was not possible to induce proliferation by a cocktail of these growth factors in serum-free medium. Providing extracellular matrix molecules as attachment factors in addition likewise failed to mimic HPL. Obviously, additional components are necessary to fully support MSC proliferation.

Pathogen Reduction Strategies

Combining a number of advantages, human supplements still pose the risk of transferring infectious agents. Hemovigilance plus quarantine storage can in part overcome the risk of the diagnostic window [77]. However, there is still a residual risk due to pathogens which are currently not routinely tested (especially viruses). Then, being stored at room temperature, platelet concentrates bear the risk of undetected bacterial contamination. Thus various pathogen reduction or inactivation strategies became investigated for erythrocyte and platelet concentrates [106]. Pathogen inactivation protocols based on photochemical treatment have been established and pathogen-inactivated human serum showed no differences regarding the quality of MSC compared to control serum [107]. A virally inactivated HPL was introduced by Shih et al. [108]. Here the HPL was treated by solvent/detergent, then extracted by soybean oil, and further purified by C18 chromatography and sterile filtration. Via a semiquantitative human cytokine antibody array cross-reacting with some bovine proteins, the growth factor cocktail was compared. 22 cytokines showed a higher concentration in the virally inactivated HPL than in FBS, and only two cytokines (angiopoietin-2 and bFGF) were found at lower concentrations. Virally inactivated HPL induced massive proliferation compared to FBS. The typical MSC characteristics, phenotype, immune phenotype, and differentiation were maintained, indicating the feasibility of this approach.

Clinical Trials with Mesenchymal Stromal Cells Expanded in Human Supplements

A few studies already applied 'humanized' culture conditions to expand MSC for clinical trials. The study presented by von Bonin et al. [109] evaluated bone marrow MSC expanded in human HPL in patients with refractory graft-versus-host disease (GvHD). Two out of 13 patients treated in total benefitted from the treatment. After a second dose, 5 out of 11 patients responded with a mitigation of their symptoms. Similar findings were obtained in a study where 11 pediatric patients suffering from acute or chronic GvHD have been treated with MSC expanded in HPL. Some patients received up to 5 MSC infusions with no acute and late side effects. A complete response was observed in 23.8% and a partial remission reported in 47.6% of patients [95]. In contrast, a higher rate of complete response has been reported analyzing MSC expanded in FBS. Here a complete response in 30 from a total of 55 patients has been obtained [110]. Further, 9 patients profited with an improvement of GvHD symptoms. It is discussed whether differences in the mode of MSC expansion may cause different migration towards chemotactic stimuli and by this favor one or the other organ affected by GvHD. By this, the discussion fits to the described changes in gene expression related to adhesion and homing.

Conclusion

For standardized MSC manufactured on a routine basis, chemically defined media approved for GMP and clinical use are regarded as ultimate endpoint. Until achieving this, objective supplements derived from human blood products (serum or platelet lysate) emerged as reasonable alternatives to FBS. Rauch et al. [79] defined quality criteria for HPL: a 10- to 20-fold enrichment of α -granule factors compared to serum and a reduced overall protein content, including immunoglobulins and albumin achieved by washing.

Any change of culture conditions can be of major impact on cellular quality. Switching from FBS to human supplements induces measurable changes. Although main characteristics of MSC, as defined by the ISCT, appear retained, continuative studies indicated that the choice of supplement for instance affected gene and protein expression. Adhesion as well as homing are changed by humanized culture systems compared to FBS. It is thus recommended to study the cellular effects in detail before – simply acting on the assumption that FBS and HPL have similar effects – changing culture conditions between preclinical and clinical trials. This is, however, complicated by the fact that the mode of action has not yet been defined comprehensively for FBS-cultivated cells. Thus the effects of a changed culture condition cannot be compared. These effects are expected to be even more drastic when developing chemically defined media, composed of only few factors and lacking the manifold other undefined factors present in HPL or serum. Otherwise, identifying the impact of defined media components may allow generating designed MSC for specific therapeutic applications, similar to the differentiation media already applied for targeting differentiation of MSC.

Of note, culture medium is only one building block in the complex structure of the GMP-compliant MSC manufacturing process. The establishment of standardized manufacturing protocols, quality control parameters, and assays is of utmost importance. This becomes even more important given the rapid pace in which clinical trials are initiated, often based on only weak scientific evidence. Despite the intensified research work in the translational field, only agreement on standardized protocols in conjunction with a strictly regulated environment performed by experts in the field of GMP manufacturing is expected to enable meaningful comparative multicenter studies assessing feasibility, safety, and efficacy.

The fact that the transition to clinical trials employing MSC is evolving rapidly, even without knowing the mode of action, complicates this demand. To not slow down clinical MSC evolution, an intense interaction between basic and clinical research is demanded to deepen our knowledge and to develop safe but efficacious MSC-based novel therapies.

Acknowledgement

This work was supported by research funds of the German Federal Ministry of Education and Research (START-MSC: 01GN0531 and 01GN0939) and a project commissioned by the European Community (CASCADE: FP7–223236).

Disclosure Statement

There is no conflict of interest.

References

- 1 Gstraunthaler G: Alternatives to the use of fetal bovine serum: serum-free cell culture. Altex-Altern Tierexp 2003;20:275–281.
- 2 Nurden AT: Platelets, inflammation and tissue regeneration. Thromb Haemost 2011;105(suppl 1): S13–33.
- 3 Eastment CT, Sirbasku DA: Human platelet lysate contains growth factor activities for established cell lines derived from various tissues of several species. In vitro 1980;16:694–705.
- 4 Hara Y, Steiner M, Baldini MG: Platelets as a source of growth-promoting factor(s) for tumor cells. Cancer Res 1980;40:1212–1216.
- 5 Choi YC, Morris GM, Sokoloff L: Effect of platelet lysate on growth and sulfated glycosaminoglycan synthesis in articular chondrocyte cultures. Arthritis Rheum 1980;23:220–224.
- 6 Dugrillon A, Eichler H, Kern S, Kluter H: Autologous concentrated platelet-rich plasma (cprp) for local application in bone regeneration. Int J Oral Maxillofac Surg 2002;31:615–619.
- 7 Rendu F, Brohard-Bohn B: The platelet release reaction: Granules' constituents, secretion and functions. Platelets 2001;12:261–273.

- 8 Maynard DM, Heijnen HFG, Horne MK, White JG, Gahl WA: Proteomic analysis of platelet alphagranules using mass spectrometry. J Thromb Haemost 2007;5:1945–1955.
- 9 Jonas JB, Dugrillon A, Kluter H, Kamppeter B: Subconjunctival injection of autologous platelet concentrate in the treatment of overfiltrating bleb. J Glaucoma 2003;12:57–58.
- 10 Sommeling CE, Heyneman A, Hoeksema H, Verbelen J, Stillaert FB, Monstrey S: The use of plateletrich plasma in plastic surgery: A systematic review. J Plast Reconstr Aesthet Surg 2013;66:301–311.
- 11 Vavken P, Sadoghi P, Murray MM: The effect of platelet concentrates on graft maturation and graftbone interface healing in anterior cruciate ligament reconstruction in human patients: a systematic review of controlled trials. Arthroscopy 2011;27: 1573–1583.
- 12 Sheth U, Simunovic N, Klein G, Fu F, Einhorn TA, Schemitsch E, Ayeni OR, Bhandari M: Efficacy of autologous platelet-rich plasma use for orthopaedic indications: A meta-analysis. The Journal Bone Joint Surg 2012;94:298–307.
- 13 Martinez-Zapata MJ, Marti-Carvajal AJ, Sola I, Exposito JA, Bolibar I, Rodriguez L, Garcia J: Autologous platelet-rich plasma for treating chronic wounds. Cochrane Database Syst Rev 2012;10: CD006899.
- 14 Doucet C, Ernou I, Zhang Y, Llense JR, Begot L, Holy X, Lataillade JJ: Platelet lysates promote mesenchymal stem cell expansion: a safety substitute for animal serum in cell-based therapy applications. J Cell Physiol 2005;205:228–236.
- 15 Friedenstein AJ, Deriglasova UF, Kulagina NN, Panasuk AF, Rudakowa SF, Luria EA, Ruadkow IA: Precursors for fibroblasts in different populations of hematopoietic cells as detected by the in vitro colony assay method. Exp Hematol 1974;2:83–92.
- 16 Friedenstein AJ, Petrakova KV, Kurolesova AI, Frolova GP: Heterotopic of bone marrow. Analysis of precursor cells for osteogenic and hematopoietic tissues. Transplantation 1968;6:230–247.
- 17 Pittenger MF, Mackay AM, Beck SC, Jaiswal RK, Douglas R, Mosca JD, Moorman MA, Simonetti DW, Craig S, Marshak DR: Multilineage potential of adult human mesenchymal stem cells. Science 1999;284:143–147.
- 18 Caplan AI: The mesengenic process. Clin Plast Surg 1994;21:429–435.
- 19 Caplan AI: Mesenchymal stem cells. J Orthop Res 1991;9:641–650.
- 20 Horwitz EM, Le Blanc K, Dominici M, Mueller I, Slaper-Cortenbach I, Marini FC, Deans RJ, Krause DS, Keating A: Clarification of the nomenclature for MSC: The International Society for Cellular Therapy position statement. Cytotherapy 2005;7:393–395.
- 21 Pevsner-Fischer M, Levin S, Zipori D: The origins of mesenchymal stromal cell heterogeneity. Stem Cell Rev 2011;7:560–568.
- 22 Dominici M, Le Blanc K, Mueller I, Slaper-Cortenbach I, Marini F, Krause D, Deans R, Keating A, Prockop D, Horwitz E: Minimal criteria for defining multipotent mesenchymal stromal cells. The International Society for Cellular Therapy position statement. Cytotherapy 2006;8:315–317.
- 23 Schafer R, Northoff H: Characteristics of mesenchymal stem cells – new stars in regenerative medicine or unrecognized old fellows in autologous regeneration? Transfus Med Hemother 2008;35: 154–159.
- 24 Bourin P, Gadelorge M, Peyrafitte JA, Fleury-Cappellesso S, Gomez M, Rage C, Sensebe L: Mesenchymal progenitor cells: tissue origin, isolation and culture. Transfus Med Hemother 2008;35:160–167.

- 25 Bieback K: Basic biology of mesenchymal stem cells. Transfus Med Hemother 2008;35:151–152.
- 26 Rojewski MT, Weber BM, Schrezenmeier H: Phenotypic characterization of mesenchymal stem cells from various tissues. Transfus Med Hemother 2008;35:168–184.
- 27 Wagner W, Saffrich R, Ho AD: The stromal activity of mesenchymal stromal cells. Transfus Med Hemother 2008;35:185–193.
- 28 Bifari F, Lisi V, Mimiola E, Pasini A, Krampera M: Immune modulation by mesenchymal stem cells. Transfus Med Hemother 2008;35:194–204.
- 29 Collas P, Noer A, Sorensen AL: Epigenetic basis for the differentiation potential of mesenchymal and embryonic stem cells. Transfus Med Hemother 2008;35:205–215.
- 30 Frith J, Genever P: Transcriptional control of mesenchymal stem cell differentiation. Transfus Med Hemother 2008;35:216–227.
- 31 Gimble JM, Guilak F, Nuttall ME, Sathishkumar S, Vidal M, Bunnell BA: In vitro differentiation potential of mesenchymal stem cells. Transfus Med Hemother 2008;35:228–238.
- 32 Quinn C, Flake AW: In vivo differentiation potential of mesenchymal stem cells: Prenatal and postnatal model systems. Transfus Med Hemother 2008;35:239–247.
- 33 Ratajczak MZ, Zuba-Surma EK, Wojakowski W, Ratajczak J, Kucia M: Bone marrow – home of versatile stem cells. Transfus Med Hemother 2008;35: 248–259.
- 34 Tonn T, Barz D: Msc a multipotent stromal cell in search of clinical application. Transfus Med Hemother 2008;35:269–270.
- 35 Klingemann H, Matzilevich D, Marchand J: Mesenchymal stem cells – sources and clinical applications. Transfus Med Hemother 2008;35:272–277.
- 36 Brooke G, Rossetti T, Ilic N, Murray P, Hancock S, Pelekanos R, Atkinson K: Points to consider in designing mesenchymal stem cell-based clinical trials. Transfus Med Hemother 2008;35:279–285.
- 37 Bieback K, Schallmoser K, Kluter H, Strunk D: Clinical protocols for the isolation and expansion of mesenchymal stromal cells. Transfus Med Hemother 2008;35:286–294.
- 38 Slaper-Cortenbach IC: Current regulations for the production of multipotent mesenchymal stromal cells for clinical application. Transfus Med Hemother 2008;35:295–298.
- 39 Lepperdinger G, Brunauer R, Gassner R, Jamnig A, Kloss F, Laschober GT: Changes of the functional capacity of mesenchymal stem cells due to aging or age-associated disease – implications for clinical applications and donor recruitment. Transfus Med Hemother 2008;35:299–305.
- 40 Henschler R, Deak E, Seifried E: Homing of mesenchymal stem cells. Transfus Med Hemother 2008;35:306–312.
- 41 Lindner U, Kramer J, Rohwedel J, Schlenke P: Mesenchymal stem or stromal cells: toward a better understanding of their biology? Transfus Med Hemother 2010;37:75–83.
- 42 Bernardo ME, Pagliara D, Locatelli F: Mesenchymal stromal cell therapy: a revolution in regenerative medicine? Bone Marrow Transplant 2012;47: 164–171.
- 43 Horwitz EM, Prockop DJ, Fitzpatrick LA, Koo WW, Gordon PL, Neel M, Sussman M, Orchard P, Marx JC, Pyeritz RE, Brenner MK: Transplantability and therapeutic effects of bone marrow-derived mesenchymal cells in children with osteogenesis imperfecta. Nat Med 1999;5:309–313.

- 44 Koc ON, Gerson SL, Cooper BW, Dyhouse SM, Haynesworth SE, Caplan AI, Lazarus HM: Rapid hematopoietic recovery after coinfusion of autologous-blood stem cells and culture-expanded marrow mesenchymal stem cells in advanced breast cancer patients receiving high-dose chemotherapy. J Clin Oncol 2000;18:307–316.
- 45 Le Blanc K, Samuelsson H, Gustafsson B, Remberger M, Sundberg B, Arvidson J, Ljungman P, Lonnies H, Nava S, Ringden O: Transplantation of mesenchymal stem cells to enhance engraftment of hematopoietic stem cells. Leukemia 2007;21:1733– 1738.
- 46 Prockop DJ: 'Stemness' does not explain the repair of many tissues by mesenchymal stem/multipotent stromal cells (MSCS). Clin Pharmacol Ther 2007; 82:241–243.
- 47 Le Blanc K, Rasmusson I, Sundberg B, Gotherstrom C, Hassan M, Uzunel M, Ringden O: Treatment of severe acute graft-versus-host disease with third party haploidentical mesenchymal stem cells. Lancet 2004;363:1439–1441.
- 48 Le Blanc K, Tammik L, Sundberg B, Haynesworth SE, Ringden O: Mesenchymal stem cells inhibit and stimulate mixed lymphocyte cultures and mitogenic responses independently of the major histocompatibility complex. Scand J Immunol 2003; 57:11–20.
- 49 Sundin M, Ringden O, Sundberg B, Nava S, Gotherstrom C, Le Blanc K: No alloantibodies against mesenchymal stromal cells, but presence of antifetal calf serum antibodies, after transplantation in allogeneic hematopoietic stem cell recipients. Haematologica 2007;92:1208–1215.
- 50 Caplan AI, Correa D: The MSC: an injury drugstore. Cell Stem Cell 2011;9:11–15.
- 51 Bianco P, Barker R, Brustle O, Cattaneo E, Clevers H, Daley GQ, De Luca M, Goldstein L, Lindvall O, Mummery C, Robey PG, Sattler de Sousa EBC, Smith A: Regulation of stem cell therapies under attack in europe: for whom the bell tolls. EMBO J 2013;32:1489–1495.
- 52 Sensebe L, Krampera M, Schrezenmeier H, Bourin P, Giordano R: Mesenchymal stem cells for clinical application. Vox Sang 2010;98:93–107.
- 53 Cyranoski D: Korean deaths spark inquiry. Nature 2010;468:485.
- 54 Tuffs A: Stem cell treatment in germany is under scrutiny after child's death. Br Med J 2010;341
- 55 Breitbach M, Bostani T, Roell W, Xia Y, Dewald O, Nygren JM, Fries JW, Tiemann K, Bohlen H, Hescheler J, Welz A, Bloch W, Jacobsen SE, Fleischmann BK: Potential risks of bone marrow cell transplantation into infarcted hearts. Blood 2007;110:1362–1369.
- 56 Ramasamy R, Lam EW, Soeiro I, Tisato V, Bonnet D, Dazzi F: Mesenchymal stem cells inhibit proliferation and apoptosis of tumor cells: impact on in vivo tumor growth. Leukemia 2007;21:304–310.
- 57 Klopp AH, Gupta A, Spaeth E, Andreeff M, Marini F 3rd: Concise review: dissecting a discrepancy in the literature: do mesenchymal stem cells support or suppress tumor growth? Stem Cells 2011;29:11–19.
- 58 Lepperdinger G, Brunauer R, Jamnig A, Laschober G, Kassem M: Controversial issue: is it safe to employ mesenchymal stem cells in cellbased therapies? Exp Gerontol 2008;43:1018–1023.
- 59 Prockop DJ, Brenner M, Fibbe WE, Horwitz E, Le Blanc K, Phinney DG, Simmons PJ, Sensebe L, Keating A: Defining the risks of mesenchymal stromal cell therapy. Cytotherapy 2010;12:576–578.

- 60 Tarte K, Gaillard J, Lataillade JJ, Fouillard L, Becker M, Mossafa H, Tchirkov A, Rouard H, Henry C, Splingard M, Dulong J, Monnier D, Gourmelon P, Gorin NC, Sensebe L: Clinicalgrade production of human mesenchymal stromal cells: occurrence of aneuploidy without transformation. Blood 2010;115:1549–1553.
- 61 Bieback K, Kinzebach S, Karagianni M: Translating research into clinical scale manufacturing of mesenchymal stromal cells. Stem Cells Int 2011; 2010:193519.
- 62 Sensebe L: Clinical grade production of mesenchymal stem cells. Biomed Mater Eng 2008;18 (1 suppl):S3-10.
- 63 EMEA, Committee for Advanced Therapies (CAT): Reflection Paper on Stem Cell-Based Medicinal Products. EMA/CAT/571134, 2011. www.ema.europa.eu/docs/en_GB/document_library/ Scientific_guideline/2011/02/WC500101692.pdf.
- 64 EMEA, Committee for Proprietary Medicinal Products (CPMP): Note for Guidance on the Use of Bovine Serum in the Manufacture of Human Medicinal Products. EMEA CPMP/BWP/1793/02, 2003. www.ema.europa.eu/docs/en_GB/document_ library/Scientific_guideline/2009/09/WC500003675. pdf.
- 65 Heiskanen A, Satomaa T, Tiitinen S, Laitinen A, Mannelin S, Impola U, Mikkola M, Olsson C, Miller-Podraza H, Blomqvist M, Olonen A, Salo H, Lehenkari P, Tuuri T, Otonkoski T, Natunen J, Saarinen J, Laine J: N-glycolylneuraminic acid xenoantigen contamination of human embryonic and mesenchymal stem cells is substantially reversible. Stem Cells 2007;25:197–202.
- 66 Horwitz EM, Prockop DJ, Gordon PL, Koo WW, Fitzpatrick LA, Neel MD, McCarville ME, Orchard PJ, Pyeritz RE, Brenner MK: Clinical responses to bone marrow transplantation in children with severe osteogenesis imperfecta. Blood 2001; 97:1227–1231.
- 67 Spees JL, Gregory CA, Singh H, Tucker HA, Peister A, Lynch PJ, Hsu SC, Smith J, Prockop DJ: Internalized antigens must be removed to prepare hypoimmunogenic mesenchymal stem cells for cell and gene therapy. Mol Ther 2004;9:747–756.
- 68 Mannello F, Tonti GA: Concise review: no breakthroughs for human mesenchymal and embryonic stem cell culture: conditioned medium, feeder layer, or feeder-free; medium with fetal calf serum, human serum, or enriched plasma; serum-free, serum replacement nonconditioned medium, or ad hoc formula? All that glitters is not gold! Stem Cells 2007;25:1603–1609.
- 69 Tekkatte C, Gunasingh GP, Cherian KM, Sankaranarayanan K: 'Humanized' stem cell culture techniques: the animal serum controversy. Stem Cells Int 2011;2011:504723.
- 70 Kinzebach S, Bieback K: Expansion of mesenchymal stem/stromal cells under xenogenic-free culture conditions. Adv Biochem Eng Biotechnol 2013;129:33–57.
- 71 Kilian O, Flesch I, Wenisch S, Taborski B, Jork A, Schnettler R, Jonuleit T: Effects of platelet growth factors on human mesenchymal stem cells and human endothelial cells in vitro. Eur J Med Res 2004;9:337–344.
- 72 Gruber R, Karreth F, Kandler B, Fuerst G, Rot A, Fischer MB, Watzek G: Platelet-released supernatants increase migration and proliferation, and decrease osteogenic differentiation of bone marrowderived mesenchymal progenitor cells under in vitro conditions. Platelets 2004;15:29–35.

- 73 Kocaoemer A, Kern S, Klüter H, Bieback K: Human AB serum and thrombin-activated platelet-rich plasma are suitable alternatives to fetal calf serum for the expansion of mesenchymal stem cells from adipose tissue. Stem Cells 2007;25:1270–1278.
- 74 Bieback K, Hecker A, Kocaomer A, Lannert H, Schallmoser K, Strunk D, Kluter H: Human alternatives to fetal bovine serum for the expansion of mesenchymal stromal cells from bone marrow. Stem Cells 2009;27:2331–2341.
- 75 Muller AM, Davenport M, Verrier S, Droeser R, Alini M, Bocelli-Tyndall C, Schaefer DJ, Martin I, Scherberich A: Platelet lysate as a serum substitute for 2d static and 3d perfusion culture of stromal vascular fraction cells from human adipose tissue. Tissue Eng Part A 2009;15:869–875.
- 76 Bernardo ME, Cometa AM, Pagliara D, Vinti L, Rossi F, Cristantielli R, Palumbo G, Locatelli F: Ex vivo expansion of mesenchymal stromal cells. Best Pract Res Clin Haematol 2011;24:73–81.
- 77 Fekete N, Gadelorge M, Furst D, Maurer C, Dausend J, Fleury-Cappellesso S, Mailander V, Lotfi R, Ignatius A, Sensebe L, Bourin P, Schrezenmeier H, Rojewski MT: Platelet lysate from whole blood-derived pooled platelet concentrates and apheresis-derived platelet concentrates for the isolation and expansion of human bone marrow mesenchymal stromal cells: Production process, content and identification of active components. Cytotherapy 2012;14:540–554.
- 78 Schrezenmeier H, Seifried E: Buffy-coat-derived pooled platelet concentrates and apheresis platelet concentrates: which product type should be preferred? Vox Sang 2010;99:1–15.
- 79 Rauch C, Feifel E, Amann EM, Spotl HP, Schennach H, Pfaller W, Gstraunthaler G: Alternatives to the use of fetal bovine serum: human platelet lysates as a serum substitute in cell culture media. Altex-Altern Anim Ex 2011;28:305–316.
- 80 Avanzini MA, Bernardo ME, Cometa AM, Perotti C, Zaffaroni N, Novara F, Visai L, Moretta A, Del Fante C, Villa R, Ball LM, Fibbe WE, Maccario R, Locatelli F: Generation of mesenchymal stromal cells in the presence of platelet lysate: A phenotypic and functional comparison of umbilical cord blood- and bone marrow-derived progenitors. Haematologica 2009;94:1649–1660.
- 81 Schallmoser K, Bartmann C, Rohde E, Reinisch A, Kashofer K, Stadelmeyer E, Drexler C, Lanzer G, Linkesch W, Strunk D: Human platelet lysate can replace fetal bovine serum for clinical-scale expansion of functional mesenchymal stromal cells. Transfusion 2007;47:1436–1446.
- 82 Crespo-Diaz R, Behfar A, Butler GW, Padley DJ, Sarr MG, Bartunek J, Dietz AB, Terzic A: Platelet lysate consisting of a natural repair proteome supports human mesenchymal stem cell proliferation and chromosomal stability. Cell Transplant 2011; 20:797–811.
- 83 Bernardo ME, Avanzini MA, Perotti C, Cometa AM, Moretta A, Lenta E, Del Fante C, Novara F, de Silvestri A, Amendola G, Zuffardi O, Maccario R, Locatelli F: Optimization of in vitro expansion of human multipotent mesenchymal stromal cells for cell-therapy approaches: further insights in the search for a fetal calf serum substitute. J Cell Physiol 2007;211:121–130.
- 84 Lange C, Cakiroglu F, Spiess AN, Cappallo-Obermann H, Dierlamm J, Zander AR: Accelerated and safe expansion of human mesenchymal stromal cells in animal serum-free medium for transplantation and regenerative medicine. J Cell Physiol 2007;213:18–26.

- 85 Schallmoser K, Strunk D: Preparation of pooled human platelet lysate (phpl) as an efficient supplement for animal serum-free human stem cell cultures. J Vis Exp 2009;(32).pii: 1523. doi: 10.3791/ 1523.
- 86 Rojewski MT, Fekete N, Baila S, Nguyen K, Furst D, Antwiler D, Dausend J, Kreja L, Ignatius A, Sensebe L, Schrezenmeier H: GMP-compliant isolation and expansion of bone marrow-derived MSCs in the closed, automated device quantum cell expansion system. Cell Transplant 2012; http:// dx.doi.org/10.3727/096368912X657990.
- 87 Nold P, Brendel C, Neubauer A, Bein G, Hackstein H: Good manufacturing practice-compliant animal-free expansion of human bone marrow derived mesenchymal stroma cells in a closed hollow-fiber-based bioreactor. Biochem Biophys Res Commun 2013;430:325–330.
- 88 Salvade A, Della Mina P, Gaddi D, Gatto F, Villa A, Bigoni M, Perseghin P, Serafini M, Zatti G, Biondi A, Biagi E: Characterization of platelet lysate cultured mesenchymal stromal cells and their potential use in tissue-engineered osteogenic devices for the treatment of bone defects. Tissue Eng Part C Methods 2010;16:201–214.
- 89 Dahl JA, Duggal S, Coulston N, Millar D, Melki J, Shahdadfar A, Brinchmann JE, Collas P: Genetic and epigenetic instability of human bone marrow mesenchymal stem cells expanded in autologous serum or fetal bovine serum. Int J Dev Biol 2008; 52:1033–1042.
- 90 Bocelli-Tyndall C, Zajac P, Di Maggio N, Trella E, Benvenuto F, Iezzi G, Scherberich A, Barbero A, Schaeren S, Pistoia V, Spagnoli G, Vukcevic M, Martin I, Tyndall A: Fibroblast growth factor 2 and platelet-derived growth factor, but not platelet lysate, induce proliferation-dependent, functional class ii major histocompatibility complex antigen in human mesenchymal stem cells. Arthritis Rheum 2010;62:3815–3825.
- 91 Flemming A, Schallmoser K, Strunk D, Stolk M, Volk HD, Seifert M: Immunomodulative efficacy of bone marrow-derived mesenchymal stem cells cultured in human platelet lysate. J Clin Immunol 2011;31:1143–1156.
- 92 Abdelrazik H, Spaggiari GM, Chiossone L, Moretta L: Mesenchymal stem cells expanded in human platelet lysate display a decreased inhibitory capacity on T- and NK-cell proliferation and function. Eur J Immunol 2011;41:3281–3290.
- 93 Bieback K, Ha VA, Hecker A, Grassl M, Kinzebach S, Solz H, Sticht C, Kluter H, Bugert P: Altered gene expression in human adipose stem cells cultured with fetal bovine serum compared to human supplements. Tissue Eng Part A 2010;16: 3467–3484.
- 94 Dreher L, Elvers-Hornung S, Brinkmann I, Huck V, Henschler R, Gloe T, Kluter H, Bieback K: Cultivation in human serum reduces adipose tissuederived mesenchymal stromal cell adhesion to laminin and endothelium and reduces capillary entrapment. Stem Cells Dev 2013;22:791–803.
- 95 Lucchini G, Introna M, Dander E, Rovelli A, Balduzzi A, Bonanomi S, Salvade A, Capelli C, Belotti D, Gaipa G, Perseghin P, Vinci P, Lanino E, Chiusolo P, Orofino MG, Marktel S, Golay J, Rambaldi A, Biondi A, D'Amico G, Biagi E: Platelet-lysateexpanded mesenchymal stromal cells as a salvage therapy for severe resistant graft-versus-host disease in a pediatric population. Biol Blood Marrow Transplant 2010;16:1293–1301.

- 96 Lee RH, Pulin AA, Seo MJ, Kota DJ, Ylostalo J, Larson BL, Semprun-Prieto L, Delafontaine P, Prockop DJ: Intravenous hMSCs improve myocardial infarction in mice because cells embolized in lung are activated to secrete the anti-inflammatory protein TSG-6. Cell Stem Cell 2009;5:54–63.
- 97 Dozza B, Di Bella C, Lucarelli E, Giavaresi G, Fini M, Tazzari PL, Giannini S, Donati D: Mesenchymal stem cells and platelet lysate in fibrin or collagen scaffold promote non-cemented hip prosthesis integration. J Orthop Res 2011;29:961– 968.
- 98 Chevallier N, Anagnostou F, Zilber S, Bodivit G, Maurin S, Barrault A, Bierling P, Hernigou P, Layrolle P, Rouard H: Osteoblastic differentiation of human mesenchymal stem cells with platelet lysate. Biomaterials 2010;31:270–278.
- 99 Centeno CJ, Schultz JR, Cheever M, Freeman M, Faulkner S, Robinson B, Hanson R: Safety and complications reporting update on the re-implantation of culture-expanded mesenchymal stem cells using autologous platelet lysate technique. Curr Stem Cell Res Ther 2011;6:368–378.
- 100 Lange C, Brunswig-Spickenheier B, Eissing L, Scheja L: Platelet lysate suppresses the expression of lipocalin-type prostaglandin d2 synthase that positively controls adipogenic differentiation of human mesenchymal stromal cells. Exp Cell Res 2012;318:2284–2296.

- 101 Murphy MB, Blashki D, Buchanan RM, Yazdi IK, Ferrari M, Simmons PJ, Tasciotti E: Adult and umbilical cord blood-derived platelet-rich plasma for mesenchymal stem cell proliferation, chemotaxis, and cryo-preservation. Biomaterials 2012;33:5308–5316.
- 102 Lohmann M, Walenda G, Hemeda H, Joussen S, Drescher W, Jockenhoevel S, Hutschenreuter G, Zenke M, Wagner W: Donor age of human platelet lysate affects proliferation and differentiation of mesenchymal stem cells. PLoS One 2012;7: e37839.
- 103 Senzel L, Gnatenko DV, Bahou WF: The platelet proteome. Curr Opin Hematol 2009;16:329–333.
- 104 Ng F, Boucher S, Koh S, Sastry KS, Chase L, Lakshmipathy U, Choong C, Yang Z, Vemuri MC, Rao MS, Tanavde V: PDGF, TGF-beta, and FGF signaling is important for differentiation and growth of mesenchymal stem cells (MSCs): Transcriptional profiling can identify markers and signaling pathways important in differentiation of MSCs into adipogenic, chondrogenic, and osteogenic lineages. Blood 2008;112:295–307.
- 105 Perez-Ilzarbe M, Diez-Campelo M, Aranda P, Tabera S, Lopez T, del Canizo C, Merino J, Moreno C, Andreu EJ, Prosper F, Perez-Simon JA: Comparison of ex vivo expansion culture conditions of mesenchymal stem cells for human cell therapy. Transfusion 2009;49:1901–1910.

- 106 Janetzko K, Bugert P: Pathogen reduction in blood products: what's behind these techniques? Transfus Med Hemother 2011;38:5–6.
- 107 Stahle M, Carlsson B, Le Blanc K, Korsgren O, Knutson F: Photochemical pathogen inactivation of human serum enables its large-scale application in clinical cell transplantation. Vox Sang 2010;98:e364–365.
- 108 Shih DT, Chen JC, Chen WY, Kuo YP, Su CY, Burnouf T: Expansion of adipose tissue mesenchymal stromal progenitors in serum-free medium supplemented with virally inactivated allogeneic human platelet lysate. Transfusion 2011; 51:770–778.
- 109 Von Bonin M, Stolzel F, Goedecke A, Richter K, Wuschek N, Holig K, Platzbecker U, Illmer T, Schaich M, Schetelig J, Kiani A, Ordemann R, Ehninger G, Schmitz M, Bornhauser M: Treatment of refractory acute GVHD with third-party MSC expanded in platelet lysate-containing medium. Bone Marrow Transplant 2009;43:245–251.
- 110 Le Blanc K, Frassoni F, Ball L, Locatelli F, Roelofs H, Lewis I, Lanino E, Sundberg B, Bernardo ME, Remberger M, Dini G, Egeler RM, Bacigalupo A, Fibbe W, Ringden O: Mesenchymal stem cells for treatment of steroid-resistant, severe, acute graft-versus-host disease: a phase II study. Lancet 2008;371:1579–1586.

Imprint

ISSN Print Edition: 1660–3796 **ISSN Online Edition:** 1660–3818

Journal Homepage: http://www.karger.com/tmh

Publication Data: Volume 40, 2013 of 'TRANSFUSION MEDICINE AND HEMOTHERAPY' appears with 6 issues.

Copyright: © 2013 by S. Karger Verlag für Medizin und Naturwissenschaften GmbH, Freiburg (Germany). All rights reserved. No part of the journal may be reproduced in any form without the written permission of the publisher. This includes digitalisation and any further electronic computing, like saving, copying, printing or electronic transmission of digitalized material from this journal (online or offline). Authorization to photocopy items for internal or personal use of specific clients is granted by Karger.

Disclaimer: The statements and data contained in this publication are solely those of the individual authors and contributors and not of the publisher and the editor(s). The appearance of advertisements in the journal is not a warranty, endorsement, or approval of the products or services advertised or of their effectiveness, quality or safety. The publisher and the editor(s) disclaim responsibility for any injury to persons or property resulting form any ideas, methods, instructions or products referred to in the content or advertisements.

Distribution and Subscription: Karger offers three types of subscription: Print Only, Online Only and the combined Print + Online. The basic annual subscription rate is the same for all three delivery forms; however, a fee for the combined print and online subscription is levied, and there is a postage and handling charge for Print Only and Print + Online. Subscriptions run for a full calendar year. Prices are given per volume.

Print subscription: EUR 173.– + postage and handling. Online subscription: EUR 173.–. Combined (print + online) subscription: EUR 223.– + postage and handling. For customers in Germany: Please turn to your bookshop or to S. Karger Verlag für Medizin und Naturwissenschaften GmbH Wilhelmstr. 20A, 79098 Freiburg (Germany) Tel. +49 761 45 20 70, Fax +49 761 45 20 714 E-mail information@karger.com

For customers in all other countries: Please contact your bookshop or S. Karger AG Allschwilerstr. 10, 4009 Basel (Switzerland) Tel. +41 61 3 06 11 11, Fax +41 61 3 06 12 34 E-mail karger@karger.com

Advertising: Correspondence should be addressed to the publisher. S. Karger Verlag für Medizin und Naturwissenschaften GmbH Attn. Ellen Zimmermann (Head of Marketing) E-mail e.zimmermann@karger.com

Price list No. 27 of January 1, 2013 is effective.

V.i.S.d.P. (Person responsible according to the German Press Law): Sibylle Gross

Type setting and printing: Kraft Druck GmbH, 76275 Ettlingen, Germany.

Bibliographic Services Biological Abstracts Current Contents/Clinical Medicine Excerpta Medica/EMBASE Medical Documentation Service Reference Update Research Alert Science Citation Index SCISEARCH Database PubMed Central

KARGER

© 2013 S. Karger GmbH, Freiburg

Fax +49 761 4 52 07 14 Information@Karger.com www.karger.com

Accessible online at: www.karger.com/tmh