Requirements for human haematopoietic stem/progenitor cells

Xue Nan¹,² | Bowen Zhang¹ | Jie Hao³,⁴,⁵,⁶ | Wen Yue¹,² | Boqiang Fu²,⁷ | Mingyi Qu¹ | Yu Zhang³ | Haiyang Wang¹ | Fang Fang¹,²,⁸ | Jun Wei³,⁹ | Qiyuan Li³,¹⁰ | Shijun Hu³,¹¹ | Junying Yu³,¹² | Yingdai Gao¹³ | Qifa Liu¹⁴ | Jiani Cao³,⁴,⁵,⁶ | Lei Wang³,⁴,⁵,⁶ | Yaojin Peng³,⁴,⁵,⁶ | Huanxin Zhu³,⁴,⁵,⁶ | Lingmin Liang³,⁴,⁵,⁶ | Aijin Ma²,¹⁵ | Jiaxi Zhou³,¹³ | Tongbiao Zhao³,⁴,⁵,⁶ | Xuetao Pei¹,²

¹South China Research Center for Stem Cell & Regenerative Medicine, SCIB, Guangzhou, China
²South China Cell & Stem Cell Bank, SCIB, Guangzhou, China
³Chinese Society for Stem Cell Research, Shanghai, China
⁴National Stem Cell Resource Center, State Key Laboratory of Stem Cell and Reproductive Biology, Institute of Zoology, Institute for Stem Cell and Regeneration, Chinese Academy of Sciences, Beijing, China
⁵University of Chinese Academy of Sciences, Beijing, China
⁶Beijing Institute for Stem Cell and Regenerative Medicine, Beijing, China
⁷National Institute of Metrology, Beijing, China
⁸Department of State-owned Assets and Laboratory Management, Capital Medical University, Beijing, China
⁹Zephyrm Biotechnologies Co., Ltd, Beijing, China
¹⁰China National GeneBank, Shenzhen, China
¹¹Department of Cardiovascular Surgery of the First Affiliated Hospital & Institute for Cardiovascular Science, Collaborative Innovation Center of Hematology, State Key Laboratory of Radiation Medicine and Protection, Medical College, Soochow University, Suzhou, China
¹²Nuwacell Biotechnology Co., Ltd, Hefei, China
¹³Institute of Hematology and Blood Diseases Hospital, Chinese Academy of Medical Sciences & Peking Union Medical College, Tianjin, China
¹⁴Nanfang Hospital, Southern Medical University, Guangzhou, China
¹⁵Beijing Technology and Business University, Beijing, China

Correspondence
Aijin Ma, Jiaxi Zhou, Tongbiao Zhao,
Chinese Society for Stem Cell Research,
Shanghai, China.
Email: maaj@btbu.edu.cn (A. M.); zhoujx@ihcams.ac.cn (J. Z.); tzbzhao@ioz.ac.cn (T. Z.)
Xuetao Pei, South China Research Center for Stem Cell & Regenerative Medicine, SCIB, Guangzhou, China.
Email: peixt@bmi.ac.cn (X. P.)

Funding information
National Key R&D Program of China,
Grant/Award Number: 2017YFA0103104,

Abstract
‘Requirements for human haematopoietic stem/progenitor cells’ is the first set of guidelines on human haematopoietic stem/progenitor cells in China, jointly drafted and agreed upon by experts from the Chinese Society for Stem Cell Research. This standard specifies the technical requirements, inspection methods, inspection rules, instructions for usage, labelling requirements, packaging requirements, storage requirements and transportation requirements for human haematopoietic stem/progenitor cells, which is applicable to the quality control for human haematopoietic stem/progenitor cells. We hope that publication of these guidelines will promote...
1 | SCOPE

This document specifies the technical requirements, test methods, test regulations, instructions for use, labelling requirements, packaging requirements, storage requirements, transportation requirements and waste disposal requirements of human haematopoietic stem/progenitor cells. This document is applicable for the production and testing of human haematopoietic stem/progenitor cells. This document does not apply to haematopoietic stem cells involved in the Technical management specifications for haematopoietic stem cell transplantation and Cord blood haematopoietic stem cell bank management specifications (Trial).

2 | NORMATIVE REFERENCES

The following content constitutes indispensable articles of this standard through normative reference. For dated references, only the edition cited applies. For undated references, only the latest edition (including all amendments) applies. GB/T 6682 Water for analytical laboratory use—Specification and test methods WS 213 Diagnosis for hepatitis C WS 273 Diagnosis for syphilis WS 293 Diagnosis for HIV/AIDS T/CSCB 0001 General requirements for stem cells Pharmacopoeia of the People's Republic of China National Guide to Clinical Laboratory Procedures

3 | TERMS AND DEFINITIONS

For the purpose of this document, the terms and definitions in T/CSCB 0001, T/CSCB 0002 and following terms and definitions apply to this document.

3.1 | Human haematopoietic stem cells

Stem cells that have the ability to self-renew and differentiate into all mature blood cell types.

3.2 | Human haematopoietic progenitor cells

Progenitor cells that have the ability to differentiate into multiple or a particular lineage of blood cells.

3.3 | Haematopoietic colony

Cell clusters or colonies containing recognizable progeny generated from individual haematopoietic progenitor cells cultured in a semi-solid medium containing the appropriate cytokines.

4 | ABBREVIATIONS

BFU-E: burst-forming unit-erythroid CD: cluster of differentiation CFU-GEMM: colony-forming unit-granulocyte-erythroid-macrophage-megakaryocyte CFU-GM: colony-forming unit-granulocyte-macrophage EBV: Epstein-Barr virus HBV: hepatitis B virus HCMV: human cytomegalovirus HCV: hepatitis C virus HIV: human immunodeficiency virus HTLV: human T-cell lymphotropic virus NOD-SCID: non-obese diabetic-severe combined immunodeficient STR: short tandem repeat TP: treponema pallidum

5 | TECHNICAL REQUIREMENTS

5.1 | Source materials and ancillary materials

5.1.1 The requirements of T/CSCB 0001 shall be followed. 5.1.2 To ensure the safety of the donor and the donated cells, the process for donor evaluation and screening, cell collection, transportation and receipt shall be standardized. 5.1.3 The donor shall be screened for HIV, HBV, HCV, HTLV, EBV, HCMV and TP, and the results shall be documented.
5.2 | Primary quality attributes

5.2.1 | Cell morphology

Cells grown under suspended conditions shall be round and uniform in size. Cells shall be round, and exhibit high nuclear-to-cytoplasmic ratio, light blue cytoplasm and no granular structure in the cytoplasm after the Wright-Giemsa staining.

5.2.2 | Chromosome karyotype

The normal karyotype shall be 46, XY or 46, XX.

5.2.3 | Cell viability

Shall be ≥85% before cryopreservation and ≥70% after resuscitation.

5.2.4 | Cell markers

The expression of CD34 shall be ≥80% of the cell population.

5.2.5 | Colony-forming unit assays

Number of total colonies shall be ≥10 per $10^3$ cells, and number of CFU-GEMM colonies shall be ≥1 per $10^3$ cells.

5.2.6 | Haematopoietic reconstitution assays

Sixteen weeks post-transplantation into NOD-SCID Il2rgnull mice, human CD45+ cell reconstitution levels shall be ≥5%, and human CD45+CD19+ cells, CD45+CD3+ cells, CD45+CD33+ cells and CD45+CD235a+ cells shall be positive in total peripheral blood mononuclear cells of the recipient mice.

5.2.7 | Microorganisms

Fungi, bacteria, mycoplasma, HIV, HBV, HCV, HTLV, EBV, HCMV and TP shall be negative.

5.3 | Process control

5.3.1 The process of cell expansion, cryopreservation and resuscitation shall follow the requirements of T/CSCB 0001.

5.3.2 The identity of the cells shall match with that of donor cells by STR analysis.

6 | TEST METHODS

6.1 | Cell morphology

Observe the morphology of cells using a microscope. Cell staining shall be performed following the National Guide to Clinical Laboratory Procedures.

6.2 | Chromosome karyotype

The method in the Pharmacopoeia of the People's Republic of China shall be followed.

6.3 | Cell viability

The method in Appendix A shall be followed.

6.4 | Cell markers

The method in Appendix B shall be followed.

6.5 | Colony-forming unit assays

The method in Appendix C shall be followed.

6.6 | Haematopoietic reconstitution assays

The method in Appendix D shall be followed.

6.7 | Microorganisms

6.7.1 | Fungi

The method '1101 sterility test' in the Pharmacopoeia of the People's Republic of China shall be followed.

6.7.2 | Bacteria

The method '1101 sterility test' in the Pharmacopoeia of the People's Republic of China shall be followed.

6.7.3 | Mycoplasma

The method '3301 sterility test' in the Pharmacopoeia of the People's Republic of China shall be followed.
6.7.4 | HIV

The nucleic acid test method in WS 293 shall be followed.

6.7.5 | HBV

The nucleic acid test method in the National Guide to Clinical Laboratory Procedures shall be followed.

6.7.6 | HCV

The nucleic acid test method in WS 213 shall be followed.

6.7.7 | HTLV

The nucleic acid test method in WS 273 shall be followed.

6.7.8 | EBV

The nucleic acid test method in the National Guide to Clinical Laboratory Procedures shall be followed.

6.7.9 | HCMV

The nucleic acid test method in WS 213 shall be followed.

7.2 | Quality inspection and release

Each batch of cell preparation shall be subject to the quality inspection before release. The quality inspection items shall include all the attributes specified in 5.2. The inspection reports shall be attached.

7.3 | Review inspection

Review inspection shall be performed by professional cytological testing institutions/laboratories as necessary.

7.4 | Decision rules

Products that pass all requirements in 5.2 for the quality inspection and quality review inspection are considered to be qualified. Products that do not meet these criteria should be considered unqualified.

8 | INSTRUCTION FOR USAGE

The instructions for usage shall include, but not limited to:

- Product name;
- Passage number;
- Cell numbers;
- Production date;
- Lot number;
- Production organization;
- Storage conditions;
- Shipping conditions;
- Operation manual;
- Execution standard number;
- Manufacturing address;
- Contact information;
- Postal code;
- Matters that need attention.

Note: Upon user’s requirement, endotoxin test results can be provided.

9 | LABELS

The label shall include, but not limited to:
10 | PACKAGE, STORAGE AND TRANSPORTATION

10.1 | Package

The appropriate materials and containers shall be selected to ensure maintenance of the primary quality attributes of human hematopoietic stem/progenitor cells.

10.2 | Storage

10.2.1 T/CSCB 0001 shall be followed.

10.2.2 Productions should be stored at a temperature below -130°C.

10.3 | Transportation

10.3.1 T/CSCB 0001 shall be followed.

10.3.2 Cryopreserved cell products shall be transported in dry ice or at temperature below -130°C. Non-cryopreserved cell products shall be transported at temperature between 2 and 8°C.

11 | WASTE DISPOSALS

Waste that arises during human hematopoietic stem/progenitor cell production and detection shall be disposed according to the regulations in T/CSCB 0001.

ACKNOWLEDGEMENT

This work was supported by grants from the National Key R&D Program of China 2017YFA0103104, 2018YFA0108400 and 2018YFE0204400; the Strategic Priority Research Program of the Chinese Academy of Sciences XDA16040501, XDA16040502 and XDA16040504; and Guangzhou Health Care and Cooperative Innovation Major Project 201803040005.

ORCID

Jie Hao https://orcid.org/0000-0002-7739-1850
Shijun Hu https://orcid.org/0000-0002-0068-8429
Yaojin Peng https://orcid.org/0000-0002-3909-6041
Tongbiao Zhao https://orcid.org/0000-0003-1429-5818

How to cite this article: Nan X, Zhang B, Hao J, et al. Requirements for human hematopoietic stem/progenitor cells. Cell Prolif. 2022;55:e13152. doi:10.1111/cpr.13152
APPENDIX A

(NORMATIVE)
Cell Viability Test and Cell Counting

A.1 | Instruments
A.1.1 Microscope
A.1.2 Haemocytometer (XB-K-25)

A.2 | Reagents
Unless otherwise specified, all reagents shall be of analytical purity. The water used in all tests shall be deionized.
A.2.1 Phosphate-buffered saline: pH 7.4.
A.2.2 Trypan blue solution: concentrated stock solution is diluted to 0.4% (W/V) with phosphate-buffered saline (A.2.1) for use.

A.3 | Procedure
A.3.1 Preparing single-cell suspension
Collect the cells to be tested and resuspend the cells in phosphate-buffered saline (A.2.1).
A.3.2 Trypan blue staining
Dilute the cell suspension (A.3.1) in trypan blue solution (A.2.2) at 1:1, and mix well.
A.3.3 Cell counting
Put coverslips on each chamber of a clean haemocytometer (A.1.2). Transfer 10 μL trypan blue/cell suspension (A.3.2) to the edge of the coverslip, allowing the cell suspension to fully fill the chambers under the coverslip without over- or underfill. Repeat with the second chamber. Let stand for 30 seconds, and count all cells (stained and unstained) in the stained cells in each chamber under microscope (A.1.1).
Repeat steps A.3.2 to A.3.3 once.
A.3.4 Calculating the viability

A.4 | Calculation
The cell viability can be calculated with the following formula (A.1):
\[ S = (M - D) / M \times 100\% \] (A.1)

Note:
S—cell viability
M—total cell number
D—stained cell number
Calculate the average viability of two repeats. This result is the average cell viability.

A.5 | Precision
Under the same condition, the absolute deviation of the two repeats should not exceed 10% of the arithmetic mean.

APPENDIX B

(NORMATIVE)
Cell markers and flow cytometry analysis

B.1 | INSTRUMENTS
B.1.1 Flow cytometer
B.1.2 Horizontal centrifuge
B.1.3 Electronic balance

B.2 | REAGENTS
All reagents in this method are of analytical purity. Unless otherwise specified, the water used in all tests is the level 1 water as specified in GB/T 6682.
B.2.1 Phosphate-buffered saline (PBS): pH 7.4.
B.2.2 Bovine serum albumin (BSA): purity ≥98%
B.2.3 Anti-human CD34 antibody and isotype control.
B.2.4 Prepare solutions needed for flow cytometry assay using electronic balance (B.1.3): washing buffer, fixation buffer, blocking/permeabilization buffer and antibody dilution buffer.

B.3 | PROCEDURE
B.3.1 Sample preparation
Harvest single cells by centrifugation at 300 g for 5 min. Discard the supernatant. Wash the cell samples with an appropriate volume of wash solution, then collect samples by centrifuging at 300 g for 5 min, discard the supernatant and repeat 2 times.
B.3.2 Antibody incubation
Divide the cell suspension into two equal aliquots for isotype control and test. Dilute and stain the cells with anti-human CD34 antibody (labelled as sample group) and isotype control (labelled as negative control group) separately according to their instructions/manuals. Wash the cells with washing buffer twice, collect samples by centrifuging at 300 g for 5 min, and discard the supernatant.
B.3.3 Flow cytometry analysis
Resuspend the cells with the washing solution, and then transfer the cell suspension to the tube through a 40-μm filter, and test sample on the flow cytometer according to the application manual.
B.3.4 Gating strategies
First, exclude events of debris, dead cells and untargeted cell populations by drawing a gate (gate 1) according to estimated cell size (FSC) and granularity (SSC). Next, by comparing the test and isotype control, place the gate for positive staining population (gate 2) to exclude the cells not labelled with fluorescent antibodies. Isotype antibodies should be used as negative control.

B.4 | RESULT ANALYSIS
The results of flow cytometry analysis are analysed comprehensively with appropriate software following its user manual.

APPENDIX C

(NORMATIVE)
Colonies-forming unit assays

C.1 | Instruments
C.1.1 Microscope
C.1.2 Haemocytometer
C.2 | Reagents
C.2.1 Medium for colony-forming unit assays: Iscove’s Modified Dulbecco’s Medium (IMDM), methylcellulose, bovine serum albumin, foetal bovine serum, β-mercaptoethanol, L-glutamine, supplemented with stem cell factor (SCF), granulocyte-macrophage colony-stimulating factor (GM-CSF), interleukin-3 (IL-3), erythropoietin (EPO) and other cytokines. Store the medium at −20°C and avoid light. The medium shall be thawed overnight at 4°C before use and shall be used within 1 week.

C.2.2 IMDM: Store at 4°C and avoid light.
C.2.3 Acetic acid with methylene blue: 3%.
C.2.4 Trypan blue solution: 0.4%.

C.3 | Procedure
C.3.1 Nucleated cell counting
Collect the cells and resuspend the cells in IMDM. Dilute the cell suspension in 3% acetic acid with methylene blue at 1:50, and mix well. Draw up 10 μl of the diluted sample and examine the density of the nucleated cells according to the method in Appendix A. Determine the concentration of the nucleated cells in suspension using the following calculations: the nucleated cells per ml = average count per square × 50 (dilution factor) × 10⁶.

C.3.2 Viable cell counting
Draw up 10 μl of the cell suspension, and determine the concentration of the viable cells in suspension using the haemocytometer (C.1.2) and microscope (C.1.1), according to the method in Appendix A.

C.3.3 Cell plating
Adjust the concentration of cell suspension to 1 × 10⁴ cells per ml. Drop up 0.5 ml of the cell suspension and dilute cells to 5 ml of the medium for colony-forming unit assays. Vortex the tube to mix the contents thoroughly. Let stand for several minutes to allow the bubbles to rise to the top. Draw up the media containing cells into the syringe with blunt-end needle, and dispense a volume of 1.1 ml into each 35-mm dish. Rotate the dish to allow the medium to attach to the wall of the dish on all sides.

C.3.4 Colony formation
Incubate at 37°C, in 5% CO₂ with ≥95% humidity for 14–16 days.

C.4 | Analysis of results
Count and evaluate the colonies including BFU-E, CFU-G/M/GM and CFU-GEMM using the microscope and the gridded scoring dish.

APPENDIX D
(NORMATIVE)
In vivo haematopoietic reconstruction transplantation experiment

D.1 | Instruments
D.1.1 Microscope
D.1.2 Centrifuge
D.1.3 Flow cytometer
D.1.4 Haemocytometer

D.2 | Reagents
D.2.1 Phosphate-buffered saline (PBS): pH 7.4.
D.2.2 Disodium ethylenediaminetetraacetic acid solution: 1%
D.2.3 Red blood cell lysis buffer.
D.2.4 Antibodies and isotype controls. For antibody storage conditions, please refer to product instructions.

D.3 | Procedure
D.3.1 Sample preparation
Harvest single cells by centrifugation (D.1.2) at 300 g for 5 min. Discard the supernatant. Collect the cells and resuspend the cells. Determine the concentration of the viable cells in suspension, according to the method in Appendix A. Adjust the concentration of cell suspension to 5 × 10⁶ cells per ml.

D.3.2 Cell transplantation
5.0 × 10⁶ cells were injected into sublethally irradiated (200-300 cGy) NOD-SCID Il2rg−/− mice at 6 to 8 weeks of age via the tail vein, setting up a blank control group that was injected with an equal volume of phosphate buffer to the cell suspension. The number of mice per group was ≥8.

D.3.3 Peripheral blood cell collection and antibody labelling
After 16 weeks of cell injection, collect the anticoagulated peripheral blood of transplanted mice, lyse the red blood cells, and label the human CD45 antibody, CD19 antibody, CD3 antibody, CD33 antibody, CD235a antibody or the corresponding isotype control, according to the method in Appendix A. After cell incubation and washing, perform flow cytometry (D.1.3) to analyse the percentage of human CD45⁺ cells, human CD45⁺ CD19⁺ cells, human CD45⁺ CD33⁺ cells, human CD45⁺ CD33⁺ cells and human CD45⁺CD235a⁺ cells.

D.3.4 Result analysis
Using flow cytometry software to analyse the results of flow cytometry, the percentage of human CD45⁺ cells in the PBMCs is not less than 5%.

Unlabelled PBSCs from transplanted mice were used as negative control 1, and antibody-labelled PBMCs from transplanted mice were used as biological control 2.

Firstly, according to the FSC and SSC, the target cell group 1 was gated, and the dead cells, platelets, red blood cells and cell fragments were excluded. According to the fluorescence intensity of negative control 1, the positive gate was delimited, and the positive cell group 2 was gated based on group 1.

Following the above gate setting principles, human CD45⁺ CD19⁺ cells, human CD45⁺ CD33⁺ cells, human CD45⁺ CD33⁺ cells and human CD45⁺ CD235a⁺ cells were detected. The tested positive rate minus the positive rate of control 2 is the actual positive rate of the tube. If the value is greater than 1%, it is considered positive.

Bus et molenesis magnis aut remquea ulluptae velit elis sinh vel eosio. Abo. Se cum dem volupta cum voloreer cusum venturi aliquum tiatur autatatenda conseque conseceae eni dolorectotat et idem. Magnatem rero tet, te simusdum, sumquat iortantia quatur aut et maxim fugitius dolore, ornit parum inits idusape rspanienit mi, qui doloreque volessenet ad quodi autectus sit que expilbus voluptum rerovit eic tet hilt, vollore doloro sunt qui dolestiurem recta que molupta testem fugia duci conoectum quis est, officiu ndessum hic torior-est, sit arum volph net isstitiam quam, quatatia auda voluptae la sit.