

TITLE

HIGH-SAFETY PROCESS FOR THE PREPARATION OF PURIFIED STEM
5 CELL FRACTIONS

STATE OF THE ART

Stem cells are gaining a growing interest in cosmetic and medical practice, in
10 connection with their capacity to generate new biological tissues applicable to
patients who, for various reasons, have lost these tissues or the capacity to
regenerate them.

In particular, adult stem cells, e.g. those from lipid or myeloid origin, have
attracted particular interest, due to their more controllable potency and also for
15 being exempt from the ethic restrictions applicable to embryonic ones.

A typical field of use for stem cells of lipid origin is that of tissue filling for
cosmetic purposes: here, the patient requiring treatment donates a part of his
own lipid tissue (lipoaspirate); this is processed by a laboratory recovering the
purified cell fraction, the latter being injected back into the patient in body areas
20 requiring filling. The cell fraction obtained is called Stromal Vascular Fraction
(SVF) and represents the total number of nucleated cells extracted from the
adipose tissue. Typically, 10 to 20% of these cells are stem cells, called also
mesenchymal stem cells. They are pluripotent and capable of repairing or
regenerating several types of human tissues. The use of autologous lipid tissue
25 avoids the risk of immune reactions and rejection of the tissue. Furthermore,
these extracted cells can then be stored frozen for long periods in liquid
nitrogen at disposal of the donor for further treatments including the use of stem
cells in human stem cell therapies.

Processes to prepare purified cells fractions of lipid tissue origin are known in the art.

5 These generally comprise the recovery of the original raw lipid material, the selective extraction of the stem cell component, the elimination of the exhausted lipid matrix, and the final washing, purification and preparation of the stem cells fraction in a convenient concentration and volume. These processes, involving the passage of the stem cells-containing material through a number of pipettes, tubes, beakers, etc. involve a considerable risk of bacterial contaminations and/or material loss: therefore they require very strict procedures as regards
10 sterility of the environment and of the different materials used, multiple washings to recover possibly adhered material, all adding up to the overall cost of the process. The multiple container transfers also increase the risk of inadvertent exchange of samples from different patients, exposing the final stem-cell receiver to the risk of non-autologous transfusion.

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The above procedure also requires a strict collaboration and understanding between the operator collecting the lipid raw material and the laboratory isolating the stem cells. Sometimes the raw material is sent in non-optimal containers (e.g. too large, not properly sealed, wrongly packaged, in not optimal
20 amounts, etc.): in all these cases, the laboratory is forced to work with a sub-optimal starting sample, which may affect the quality of the final product.

Aim of the invention is to provide an improved process to prepare stem cell fractions of lipid tissue origin, which is safer for the patient and more expedite for the process operator.

25 A further aim is to simplify / standardize the interface and cooperation between the operators collecting the lipid raw material, and those charged with the stem cell isolation.

A further aim is to reduce the number of steps and manipulations involved in processes for producing purified cell fractions.

A further aim is to make more rapid and effective those medical/cosmetic procedures involving the use of stem cells.

5 SUMMARY

The present invention relates to a process to obtain a stem cell fractions of lipid origin, essentially based on the steps of:

- 10 (a) collecting or receiving a sample of lipid tissue containing stem cells;
 - (b) washing the sample obtained in step (a) with a suitable aqueous buffer;
 - (c) incubating the sample obtained in step (b) with an enzyme capable to extract the
stem cells from the lipid tissue containing them;
 - 15 (d) recovering the aqueous phase from the product obtained in step (c);
 - (e) purifying the aqueous phase obtained in step (d);
 - (f) titrating the aqueous phase obtained in step (e) and optionally diluting it to obtain a final stem cell fraction with desired concentration and volume.
- 20 One essential feature of the invention consists in that the stem cells-containing material is treated within the same collecting device (herein referred as “single collecting device” or “SCD”), throughout at least the steps: (a), (b), and (c) of the process described herein. The use of the SCD avoids contact of the treated material with the external atmosphere, reduces to a minimum the risk of
- 25 contamination and the loss of active material linked to multiple container transfers, and simplifies the overall manipulations required to obtain purified stem cell fractions.

The SCD is a sterile container capable to draw and release a liquid; it is typically but not exclusively, a syringe. The SCD may have a large filling volume (e.g.20 to 100 mL), to allow a large scale harvesting of stem cells from the corresponding lipid material. It is preferably transparent or semi-transparent, with one mark indicating the optimal filling volume, and/or areas aimed at writing or labelling, to identify the sample source.

As will be evident from the description, the SCD fulfils the functions of a collecting device (like a pipette), a phase separator, and process reactor, depending on the particular process step involved; all these functions are thus advantageously performed without transferring the sample from one container to another, and/or contacting it with the external atmosphere.

The SCD is essentially used throughout steps (a),(b) and (c) of the process: however, if desired, the SCD can be maintained also throughout one or more of steps (d),(e),(f), with the same functions and advantages described above.

Based upon the above premises, the process of the invention comprises the following steps:

- (a) collecting or receiving, in a SCD, a sample of lipid tissue containing the stem cells;
- (b) in said SCD, washing the sample of step (a) with a suitable aqueous buffer;
- (c) in said SCD, incubating the sample of step (b) with an enzyme capable to digest the lipid tissue and extract therefrom the stem cells ;
- (d) recovering the aqueous phase from the product of step (c);
- (e) purifying the aqueous phase obtained in step (d);
- (f) titrating the aqueous phase obtained in step (e) and, if necessary, diluting it to a final stem cell fraction with desired concentration and volume.

DETAILED DESCRIPTION OF THE INVENTION

Step (a):

In this step, a sample of lipid tissue containing stem cells is drawn from a
5 suitable source into the SCD.

Lipoaspirates are typical lipid tissue samples. The sample must be liquid or at least fluid for the purpose of the present process; insufficiently fluid materials can be rendered such by further homogenisation and/or addition of liquid media, e.g. buffered solutions.

10 In step (a) the SCD is put in contact with the lipid tissue sample and is operated to draw a suitable volume thereof; drawing is halted before filling completely the available volume of the SCD, thus allowing a further drawing capability (typically one half of the SCD volume) for washing buffers and other reagents as described next.

15 The expression “collecting or receiving” in step (a) accounts for the fact that this step may be performed by an institution/operator being the same or different from the one performing the other steps (b)-(f): in the first option, in step (a) the operator “collects” the lipid sample and processes it directly as per steps (b)-(f); in the second case, the operator “receives” the lipid sample, collected by
20 someone else, and processes it per steps (b)-(f); typically, step (a) can be performed by a hospital or an aesthetic centre; steps(b)-(f) are performed by a laboratory specialized in stem cells processing.

In this second option, the present process is particularly advantageous in that it removes a primary cause for contamination occurring in known processes
25 where the lipoaspirate is collected into a first container, stored, sent to an external laboratory and then transferred into a suitable reactor: all these transfers/manipulations involved a contact of the sample with the external atmosphere, with the connected risk of contamination, along with an inevitable percent of product loss. Such disadvantages and risks are now minimized by
30 the present process.

The use of the SCD is further advantageous in that it provides the initial operator, i.e. the one collecting the lipid sample, with a standardized container, suitably adapted for the further processing from the point of view of filling volume, void volume, air-tightness, packaging material, etc.

- 5 The following steps can be performed immediately after step (a); alternatively the partially filled SCD is stored for a certain time, at conditions maintaining the viability of the stem cells, until the time of further processing as per steps (b)-(f).

Step (b).

- 10 In this step, the drawn lipid sample from step (a) is washed inside the SCD, with an aqueous buffer solution.

To do so, the partially filled SCD from step (a) is operated to draw a volume of a buffer solution, which mixes with the lipid tissue sample present inside the SCD; homogeneous mixing of the two phases can be facilitated e.g. by applying
15 vibrations /shaking to the SCD. The used buffer solution is a stem cell-compatible one, typically a PBS buffer supplemented with a calcium and/or magnesium salts useful as enzyme nutrients; the volume ratio of buffer to lipid tissue is e.g. from about 0.5:1.5 to about 1.5:0.5; preferably it is about 1:1.

After said mixing/homogenizing, the SCD is kept still until the two phases (lipid
20 and aqueous) separate. Then the SCD is then operated to eject the aqueous phase while retaining the lipid phase: in particular, when the SCD is a syringe, this can be done by orienting it in downward (needle-down) position: this causes the lipid phase to move in the upper section of the syringe, distal from the needle, while the aqueous phase aggregates into the opposite section, proximal
25 to the needle: in this position, a pressure on the syringe plunger causes the aqueous phase to be ejected from the needle; the pressure is suitably maintained until the water/lipid interface reaches the needle: at this point the aqueous phase (to be discarded) is substantially eliminated, and the syringe contains only the lipid phase, upon which the next step is to be performed.

- 30 The above-described washing step can be repeated more times, e.g. one or two, until the desired degree of washing of the lipid phase is reached.

Step (c)

In this step the washed lipid phase from step (b) is incubated in the SCD with an enzyme capable to extract the stem cells from the lipid material containing
5 them.

To perform this step, the SCD is operated to further draw an aliquot of a liquid medium containing said enzyme. The enzyme is typically a collagenase, The liquid medium is typically a buffer, preferably a PBS buffer optimized for enzyme activity, in particular supplemented with calcium and/or magnesium salts; the
10 liquid medium has a known enzymatic titre, allowing the operator to draw a desired and reproducible amount of the enzyme.

The thus filled SCD, optionally inserted within a sealed envelope, is then placed into an incubator, typically a temperature-controlled oven provided with an oscillating tray. Prior to incubation, the SCD is preferably agitated to
15 homogenize the content; mixing is then continued within the incubator, by the oscillation movement of the tray.

The incubator may be operated under the following non-exhaustive conditions: incubation time 20-80 minutes, preferably 30-60 minutes, most preferably 45 minutes; temperature of 30-45 °C, preferably at 37°C; agitation: 1-5 rpm,
20 preferably 2 or 3 rpm.

Step (d)

In this step the enzyme reaction is blocked, the lipid phase is eliminated, and the aqueous phase (containing the stem cells, liberated by the enzyme) is
25 recovered for further processing per steps (e)-(f).

To perform this step, the SCD is removed from the incubator and extracted from the (optionally used) envelope; the incubated suspension is mixed with an aliquot of an enzyme-inactivating solution. A suitable mode of mixing the two liquids consist in drawing the inactivating solution into the SCD, agitating the

SCD to obtain complete homogenization, keeping the SCD still until the lipid and aqueous phases separate, and recovering the aqueous phase.

Recovery of the aqueous phase can be done by ejecting it from the SCD (ejection mode), or, alternatively, by retaining it into the SCD (retention mode).

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The “ejection mode” can be performed by orienting the syringe it in the downward (needle-down) position: this causes the lipid phase to move in the upper section of the syringe, distal from the needle, while the aqueous phase aggregates into the opposite section, proximal to the needle: in this position, a pressure on the syringe plunger causes the aqueous phase to be ejected from the needle; the pressure is suitably maintained until the water/lipid interface reaches the needle: at this point the aqueous phase (to be collected for further processing per steps (e)-(f)) is substantially ejected from the syringe; the latter contains the stem cell-depleted lipid phase which can now be ejected separately and eliminated.

Alternatively to the ejection mode, the “retention mode” can be performed by orienting the syringe in the upward (needle-up) position: this causes the lipid phase to move in the section of the syringe, proximal to the needle, while the aqueous phase aggregates into the opposite section, distal from the needle: in this position, a pressure on the syringe piston causes the lipid phase to be ejected from the needle; suitable means can be used to avoid the dispersion of the liquid ejected from the up-oriented syringe: for example, prior to ejection, the needle may be inserted through the rubber stopper of a flask, into which the ejected liquid is then collected. The pressure is suitably maintained until the water/lipid interface reaches the needle: at this point the lipid phase (to be discarded) is substantially ejected from the syringe; the latter contains the aqueous phase meant for further processing per steps (e)-(f).

At the end of step (d) the aqueous phase is recovered from the SCD and treated separately, unless the SCD has a shape and consistence allowing it to be centrifuged: in this case, the subsequent process steps can also be

performed into the SCD, adding further protection/simplification to the overall process.

5 The emptied SCD, if not adapted for centrifugation, is preferably washed one or more times with an appropriate solution (preferably the inactivating solution described above) to recover possible stem cells adhering to its surfaces, and all the resulting aqueous phases are pooled for the further processing according to steps (e)-(f))

Step (e)

10 In this step, the (pooled) aqueous phases from step (d) are purified from possible soluble/insoluble impurities derived from the original lipid sample. Purification is generally obtained by centrifugation, elimination of the supernatant, re-suspension of the pellet, filtration. After eliminating the supernatant, the re-suspension of the pellet can be performed by using a
15 suitable buffer (e.g. a PBS buffer) or the inactivating solution described above.

The above centrifugation and re-suspension can be repeated one or more times to increase purification of the particulate (stem cell) fraction from water soluble impurities.

20 The (finally) re-suspended pellet is then filtered one or more times, to eliminate particulate impurities being oversized with respect to the stem cell fraction. To do so, the suspended pellet is filtered through a membrane with an appropriate pore size, e.g. 80-120 μm , preferably about 100 μm , retaining the oversized particulate material, and allowing the (lower sized) stem cells to pass in the filtrate. The resulting liquid can be filtered again with progressively finer filters,
25 e.g. 60-80 μm , preferably about 40 μm , to allow a finer elimination of the oversized particulate. The finally filtered liquid, containing the purified stem cells is further processed per step (f).

Step (f)

In this step, the purified liquid from step (e) is titrated and then diluted to obtain a final stem cell fraction with desired concentration and volume.

5 Titration can be done by withdrawing a precise volume of the liquid of step (e) (e.g. 50 μ L) and subjecting it to a stem cell count by means of a suitable counting apparatus, typically a FACS with optimized gates to obtain Adipose-Derived Mesenchymal Stem Cells counts; alternatively, the stem cell content can be assessed indirectly by means of other instruments e.g. a haemocytometer: the latter counts the total number of nucleated cells which, at
10 this stage of the process, are found to be stem cells by 10-20%. Preferably, two or more readings are taken and averaged, for a higher precision.

Based upon its known titre, the liquid from step (e) can, if necessary, be diluted to an appropriate concentration. Dilution can be performed by using a suitable buffer (e.g. a PBS buffer) or the inactivating solution described above. The final
15 concentration value is chosen in function of the desired level of potency of the final stem cells fraction; useful.

The final stem cell fraction can then be packaged in a suitable container (e.g. mini-syringe) as a unit with an appropriate volume, e.g. 1 mL; the final volume is chosen to be compatible and handy with the site of administration (e.g. wrinkle
20 filling, tissue reconstruction, etc.) of the final stem cell fraction.

The final stem cell fraction is preferably used as soon as possible or, alternatively, it is stored in suitable conditions of sterility and temperature, until the time of use.

It can be used for any application in which stem cells of lipid origin are useful.
25 Non limitative examples are in the field of aesthetic or reconstructive treatments, in particular tissue filling, wound healing, tissue or organ reconstruction. The so obtained stem cell fraction and its medical uses form part of the present invention.

A suitable, non-exhaustive procedure in accordance with the present invention is described as follows.

5 CLAIMS

1. Processes to prepare stem cells fractions of lipid tissue origin, comprising the steps of:

- 10 (a) collecting or receiving a sample of lipid tissue containing the stem cells;
(b) washing the sample of step (a) with a suitable aqueous buffer;
(c) incubating the sample of step (b) with an enzyme capable to digest the lipid tissue and extract therefrom the stem cells;
(d) recovering the aqueous phase from the product of step (c);
15 (e) purifying the aqueous phase obtained in step (d);
(f) titrating the aqueous phase obtained in step (e) and, if necessary, diluting it to obtain a final stem cell fraction with desired concentration and volume

wherein the stem cells-containing material is treated within a single collecting
20 device (SCD) throughout at least the steps: (a), (b), and (c), said SCD performing the functions of collecting means, phase separator and process reactor.

2. Process according to claim 1, wherein the SCD has a filling volume
25 comprised between 20 and 100 mL).

3. Process according to claims 1-2, wherein the SCD is a syringe, optionally provided with one or more marks to indicate the optimal filling volume(s), and/or with areas for writing or labelling.
- 5 4. Process according to claims 1-3, wherein the lipid material is a lipoaspirate.
5. Process according to claims 1-4, wherein the step (a) and (b-d) respectively, are performed by two different operators at locations remote from each other.
- 10 6. Process according to claims 1-5, wherein in step (b) the buffer is a PBS buffer supplemented with enzyme nutrients.
7. Process according to claims 1-6, using a syringe as SCD, in which the steps (b) and/or (c) include orienting the syringe downwards (needle down) or
15 upwards (needle up), followed by ejecting the phase proximal to the needle.
8. Process according to claims 1-7, wherein in step (c) the enzyme is a liberase, and the incubation is performed for 20 to 80 minutes, at a temperature from 30 to 45°C.
- 20
9. Process according to claims 1-8, wherein in step (d), the incubated mixture is mixed with an albumin-containing solution, then the lipid phase is discarded and the aqueous phase is recovered for the subsequent process steps.
- 25 10. Process according to claims 1-9, where the stem cells-containing material is further maintained within the said SCD during one or more of the steps (d)-(e).

11. Process according to claims 1-10, wherein purification in step (e) is performed by centrifugation(s) and/or filtration(s).

12. Process according to claims 1-11, wherein the final stem cell fraction is formulated as one or more units of 1-5 ml, with a total nucleated cell concentration comprised between 10^8 to 10^4 .

13. A stem cell fraction obtained by the process of claims 1-12.

14. A stem cell fraction of claim 13, for use as an ingredient for tissue filling, wound healing, tissue or organ reconstruction.