Histological Evaluation of Decellularized Skeletal Muscle Tissue Using Two Different Decellularization Agents

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Summary: The aim of the present study was to determine effect of two decellularized agents, sodium dodecyl sulphate (SDS) and Triton X-100, to the skeletal muscle tissue. Final scaffold was evaluated by several histological techniques to analyse preservation of essential structures including collagen and elastic fibres, basement membranes, glycosaminoglycans and also to confirm elimination of nuclear and cytoplasmic components which are redundant in effectively prepared decellularized scaffolds. Comparison of tissue scaffolds processed with different detergents proved that SDS is superior to Triton X-100 as it can effectively decellularize muscle tissue.

Keywords: Decellularized scaffold; Histology; Skeletal muscle; SDS; Triton X-100

Introduction

A tissue loss in soft tissues caused by a trauma or a tumour requires a surgical reconstruction. Effective treatment of such defects can be prepared by tissue transplantation. Tissue engineering and regenerative medicine undergo significant development in the area of biomaterials and combination with stem cells offers accessible source for tissue reconstruction and transplantation (1–4). Biological cell-free material can serve as an optimal scaffold for wide applications as it does not evoke an immune response. Extracellular matrix (ECM) is a promising biomaterial alternative which provides a three-dimensional microenvironment niche for cells, which is highly specific for each type of tissue concerning the protein contents and complexity. There are several options how to substitute stromal components for scaffold production using: synthetic scaffold material, chemically cross-linked forms of ECM, purified ECM components or decellularization techniques. Successful decellularization produces biological material which combines advantages and properties of extracellular matrix as supportive microarchitecture, vascular network and participate in cell differentiation and proliferation (5, 6). Biological scaffolds composed of the extracellular matrix are commonly used in reconstructive surgical applications of many organs such as urinary bladder (7, 8), heart (9–11), lung (12, 13) or skin (14, 15).

This study compares decellularization effectiveness of two detergents, SDS (sodium dodecyl sulphate) and Triton X-100, which are both widely used as decellularization agents (16, 17). These agents supposed to maintain mechanical and structural integrity of the extracellular matrix with minimal damage of the scaffold, preserve vascular and neural networks and successfully remove nuclear and cytoplasmic components from ECM.

Material and Methods

All procedures were approved by the Ethical Committee supervising procedures on experimental animals at Charles University Medical Faculty in Hradec Králové.

Dissection of murine skeletal muscle tissue

A skeletal muscle tissue (approx. 1 × 0.5 × 0.5 cm pieces) was dissected from C57BL/6 mice and was immediately washed with PBS buffer and trimmed of a connective tissue and fat. The samples (n = 10) were fixed and processed for histological analysis.

Decellularization methods

We analysed two types of decellularization agents: 1% SDS (sodium dodecyl sulphate) and 1% Triton X-100 and samples were decellularized with the following protocol. Solutions were dissolved in distilled water.

A tissue was removed under aseptic conditions. Native samples were immediately processed after samples were delivered to the lab. The skeletal muscle was placed in 1% SDS or 1% Triton X-100 solution for 24 hours at room temperature under continuous shaking to remove cell components from the muscle tissue. The muscle sheets were
thoroughly washed with the fresh PBS buffer for 24 h at 4 °C to wash out cells and potential SDS or Triton X-100 residues. The samples were used on the same day when they were prepared. All steps were processed in centrifuge tube with approx. 40 ml of solution.

**Histological analysis**

The specimens (untreated and decellularized scaffolds) were fixed in 10% formalin, embedded in paraffin and sectioned into 5 µm slices. Sections were deparaffinised, rehydrated and washed with distilled water. For histological analysis, the sections were stained with haematoxylin eosin, Sirius red, Alcian blue, lamina externa was impregnated by Jones’ method, and elastic fibres were visualized with resorcin-fuchsin staining.

**Results**

**Morphological characterization of physiological skeletal muscle**

Native muscle samples were treated with two different decellularization agents and muscle morphology was examined with haematoxylin eosin staining compared to the untreated skeletal muscle sample. Figure 1A shows typical image of skeletal muscle represented by characteristic striated muscle fibres with nuclei localized beneath the sarcolemma revealing polygonal shape in a cross section.

**Morphology of decellularized skeletal muscle tissue treated with SDS or Triton X-100**

Haematoxylin eosin staining

SDS decellularization method resulted in a complete cell removal from the skeletal muscle tissue which was verified with standard haematoxylin-eosin staining. Tissue also revealed a general shrinkage. Microscopic architecture of the skeletal muscle tissue was preserved and typical polygonal shape of extracellular matrix, which surrounds empty spaces of striated skeletal fibres, was examined (Fig. 1B). Haematoxylin eosin staining revealed a complete removal of sarcoplasmic components which were washed out with PBS buffer. Three-dimensional architecture was preserved without nuclei remnants supported with DAPI staining showing no fluorescent signal (figure not shown).

Native skeletal muscle tissue treated with SDS revealed translucent composition; Triton X-100 did not have the same effect on the muscle tissue and remnants of unaffected tissue in the final scaffold could be observed.

Standard haematoxylin eosin staining proved presumption of inefficient decellularization based on Triton X-100 due to presence of some nuclei. Skeletal muscle fibres with occasional presence of striation were observed and presence of several nuclei underneath the sarcolemma (Fig. 1C) could

![Fig. 1: Histological appearance after haematoxylin eosin staining of native muscle tissue (A) and muscle tissue treated with SDS (B) and Triton X-100 (C). Cross section of untreated muscle has typical architecture of skeletal muscle tissue while SDS-treated muscle shows disappearance of all cytoplasmic and nuclear components with preservation of ECM. Longitudinal section of a sample treated with Triton X-100 reveals presence of striated muscle fibres with preserved nuclei (arrows). Scale bar represents 50 µm.](image-url)
be detected. Decellularization was inefficient, because ECM components were disrupted and debris surrounded striated muscle fibres DAPI staining showed positive fluorescent signal because of nuclei presence (figure not shown).

**Sirius Red**

Collagen structure is highly desirable to preserve after decellularization mainly for its structural function to sustain a shape of the scaffold. Sirius staining proved preservation of collagen in scaffold treated by SDS without any loss of collagen density or visible disruption of collagen fibres (Fig. 2A). Figure 2B showed another example of ineffective decellularization with Triton X-100; collagen fibres were heavily damaged and presence of striated muscle fibres was observed. Deterioration of collagen fibres caused lack of cohesion in scaffold and resulted in deformation of scaffold.

**Alcian blue staining**

Alcian blue is a polyvalent basic dye for visualization of glycosaminoglycans (GAG) as a part of the ground substance which plays an important role in water retention and growth factor adhesion. SDS treatment preserved GAGs (Fig. 3A), but light blue staining probably illustrated a partial loss of these macromolecules. Nevertheless, glycosaminoglycans were preserved and intact. Decellularization by Triton X-100 preserved GAGs as well and a density of blue staining showed higher amount of GAGs in samples treated with Triton X-100 than in scaffolds decellularized by SDS. Glycosaminoglycans, presented in Triton X-100 scaffold, showed disruption and fragmentation.

![Fig. 2](image1.png)

*Fig. 2:* Visualization of collagen fibres by Sirius Red staining in the skeletal muscle tissue decellularized by SDS (A) and Triton X-100 (B). Collagen fibres are stained deeply red as they are major components of ECM; high density of red staining in case of SDS treated tissue was observed. Triton X-100 decellularization preserved collagen fibres as well, but they lacked intactness and they were significantly reduced. Scale bar represents 50 µm.

![Fig. 3](image2.png)

*Fig. 3:* Alcian blue staining illustrated presence of glycosaminoglycans in scaffold treated with SDS (A) or Triton X-100 (B). Scaffold prepared by SDS decellularization method demonstrates absence of striated muscle fibres and a typical honey-comb shape was preserved which is visualised by surrounding ECM, particularly blue coloured glycosaminoglycans. Triton treated samples also contained preserved glycosaminoglycans, but these scaffolds were unefficiently decellularized because of striated muscle fibres presence (pink stained structures with nuclei localized on the periphery, not shown in figure). Scale bar represents 50 µm.
**Jones’ impregnation method**

Preservation of the lamina externa was determined with impregnation technique by Jones. The lamina externa is an essential component for cell attachment and tissue regeneration so it is crucial to preserve this structure. The lamina externa contains silver-reactive sites which surround adipocytes, peripheral nerve fibres and first of all muscle fibres. Decellularization had no severe effect upon lamina externa structure and both samples preserved this structure stained in black colour. Figure 4 shows impregnation of basement membrane which belongs to muscle fibres or to blood capillaries.

**Resorcin-fuchsin staining**

Elastic fibres are arranged in a branching pattern to form a three dimensional network and they are localized closely to collagen fibres to limit their distensibility of the tissue and prevent tearing from excessive stretching. No distortion or fragmentation was observed in case of a scaffold treated by SDS agens (Fig. 5A); elastic fibres were stained slightly lighter than in case of Triton X-100 decellularization. Triton X-100 scaffold stained with resorcin-fuchsin showed disruption of elastic fibres (Fig. 5B).

**Discussion**

The main goal of this study was to clarify efficiency of two decellularization agents, SDS and Triton X-100, which
Detergents are effective in cell removal from the scaffold cell membrane causing cell lysis by osmotic gradient. Both of these agents disrupt was based on studies (18–20) which used these two agents to prepare decellularized scaffolds. Both of these agents disrupt the cell membrane causing cell lysis by osmotic gradient. Detergents are effective in a cell removal from in the scaffold which can have a crucial role in an immune response, but they also can affect negatively structure of the scaffold; for instance, Triton X-100 can affect a decrease of GAG and damage ultrastructure of tissue or organ (21, 22). SDS also removes effectively cell nuclei and cytoplasmic components, especially from dense tissues, but they tend to damage collagen and decrease GAG content (23, 24), so they have very similar adverse consequence on the extracellular matrix. As both these agents are considered to be effective decellularization reagents, we would like to know which one is more effective in decellularizing the skeletal muscle tissue and for that reason we choose the same conditions in decellularization process (including length of decellularization, material and technique to apply the agent).

After decellularization, the skeletal muscle tissue appeared almost translucent in SDS protocol; Triton protocol did not affect a muscle so well and we could observe cytoplasmic debris in the tissue. Overall, our study proved higher efficiency of SDS agent to decellularize the skeletal muscle tissue which was supported by several histological findings. Haematoxylin eosin, as a standard method for the first tissue inspection, showed persistence of striated muscle fibres and nuclei after treatment with Triton X-100 which is according to criteria stated by Gilbert and Crapo (25) (lack of visible nuclear and cytoplasmic material in tissue section stained with haematoxylin and eosin staining) considered as ineffective decellularization. Elimination of cells from tissue or organ is crucial for subsequent implantation of the scaffold into living organism and significantly decreases a risk of rejection.

The extracellular matrix is a complex of structural and functional proteins and each type of tissue has specific composition of these proteins (26). The most abundant ECM protein is collagen which is present in several motifs as collagen type IV in basal membrane of vascular structures, collagen type I in collagen fibres, collagen type VII as an anchoring fibrils. Preservation of a different type of collagen was evaluated by two methods as Sirius Red and Jones’ impregnation. Sirius Red is specific staining method to distinguish collagen fibres, especially it allows determining difference between collagen type I and III. This type of staining revealed no changes in collagen distribution after SDS cell extraction, either fragmentation or distortion. These results were not observed in Triton X-100 treatment; collagen fibres were unevenly distributed in ECM and were heavily damaged which could cause incoherence and mechanical endurance of tissue or organ. Triton protocol also decreased collagen density as reported in the study by Graus et al. (27). Elastin fibres accompany collagen fibres and are arranged in a branching pattern forming a three dimensional network. These fibres play an important role in distensibility of the tissue and can prevent tearing from stretching. SDS and Triton X-100 treatment of the skeletal muscle tissue preserved elastin fibres, but Triton X-100 damaged fibres morphology and altered molecular structure of elastin resulting in uneven distribution of elastic fibres. SDS decellularization affects elastin fibres in content which was decreased. Laminin, collagen type IV and anchoring fibrils are the most prominent structures in basement membrane and they specify its ultrastructure and function as an attachment site for cells, filtration, and regulation and also provide scaffold during regeneration (28). Decellularized samples processed from both decellularized agents revealed intactness of basement membrane as visualized by Jones’ impregnation.

The ability of preserving glycosaminoglycans was also compared in both skeletal muscle tissues treated by SDS and Triton X-100. SDS decellularization protocol successfully retains these heteropolysaccharides which are bound to proteoglycans and serve as binding sites for growth factors, cytokines and contribute to water retention due to their negative charge (29). Triton X-100 treatment results in significant elimination of GAGs from the skeletal muscle tissue as analysed by Alcian blue staining. Reduced content of GAGs in the scaffold may decrease mechanical resistance because of GAGs are responsible for gel consistency which can resist deformations during tissue function (24, 28). Decellularization with SDS as a main agent was more successful in term of GAGs preservation, but amount of these macromolecules was also decreased. These results are consistent with other studies (30, 21) using SDS as a main decellularization agent.

**Conclusion**

ECM alteration or damage can be caused during decellularization by several factors including type of used detergent, concentration of detergent, duration of decellularization process, presence of protease inhibitor etc. (31). Differences between various decellularization agents are based upon their function and these variations determine discrepancies in effect on the extracellular matrix. Our study proves that ionic detergent SDS is sufficient agent for cells removal in the skeletal muscle tissue and preservation of the extracellular matrix microarchitecture providing microenvironment suitable for cell attachment and growth.
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Abbreviation

ECM extracellular matrix
DAPI 4',6-diamidino-2-phenylindole
GAG glycosaminoglycans
PBS phosphate buffered saline
SDS sodium dodecyl sulphate

References


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