Abstract

Different types of biological grafts are useful for the replacement of infected prostheses in vascular surgery. In cases, when autologous grafts or homografts are not available, xenografts can be an alternative choice. To reduce immunogenicity, decellularization of the xenogeneic graft is necessary. Our goal was to determine the long-lasting stability of our decellularized porcine vascular grafts, in the mirror of differential scanning calorimetry (DSC) and routine histological examination. In the first experiment, light and electron microscopic confirmation of adequate decellularization of our used protocol was assessed. In the second series, DSC measurements were performed both on native and on treated samples; thereafter, histological evaluation of the analyzed samples was performed. Light and electron microscopy evaluation showed complete removal of cells and preserved connective tissue fiber structure after decellularization in the first series. In the second part of our investigation, DSC measurements demonstrated reversible change in the structure of elastic fibers and irreversible change in the structure of collagen fiber structure supported by histological findings. Our examination strengthens the earlier observations that decellularized biological grafts can tend to late aneurysmatic degeneration, due to their weakened fiber structure, besides the known immunological circumstances.

Keywords

Decellularization · Aorta · Biological graft · Calorimetry

Introduction

Currently, bypass surgery is still a frequent revascularization method in vascular surgery. The application of autologous graft remained the gold standard in lower-extremity reconstruction; however, in aortoiliac position, prosthetic grafts, particularly Dacron prostheses are the most commonly utilized synthetic vessels. Long-lasting stability, easy handling and unlimited availability are advantageous, but in case of graft infection, treatment and replacement are considered the most difficult challenges in vascular surgery, including high morbidity and mortality rate. Limb salvage and survival of the patient can be effectively achieved by graft replacement. Among other options (autologous deep vein, homograft, impregnated graft), the usage of biological xenogeneic graft can be an alternative solution.

After doubtful early attempts [1, 2], the attention has pointed toward the development of biological xenografts with adequate biomechanical properties, high resistance against infections and unlimited availability [3–5]. By decellularization of the vessel, the immunogenicity can be reduced, preventing the implanted graft from deterioration [3–10]. Several efficient decellularization methods were previously published testing the mechanical properties of treated vessels with different methodologies [3, 4, 6–10]; however, mechanical stability of cell-free grafts based on differential scanning calorimetry (DSC) in consideration of histological evaluation was first assessed in our study.

Our aim was to analyze the effects of decellularization on the connective tissue fiber structure of the treated porcine aortic wall with calorimetric and histological examinations, thus concluding on the long-term outcome after implantation.
Materials and methods

Surgical preparation

The present study conforms to the Guide for the Care and Use of Laboratory Animals published by the US National Institute of Health (NIH Publication No. 85-23, revised 1996) and was approved by the local institutional Committee on Animal Research, University of Pécs (BA02/2000-29/2001).

Infrarenal aortic segments were freshly excised under sterile surgical conditions from euthanized Yorkshire pigs (mass from 22 to 27 kg) of other ongoing experiments, declared free of clinically evident diseases. Following separation of the adhesive tissue, a macroscopic examination was performed to exclude any pathology, and longitudinal aortic segments (about 10 mm in length and 6 mm in width) were next washed in sterile phosphate-buffered saline (PBS) to remove residual traces of blood.

Decellularization protocol

Aortic segments were decellularized through continuous shaking in 1% sodium dodecyl sulfate (SDS; Sigma-Aldrich Chemie GmbH, Steinheim, Germany), 0.05% sodium azide (NaN₃; Alfa Aesar GmbH & Co, Karlsruhe, Germany) and phosphate-buffered saline (PBS; Sigma-Aldrich Chemie GmbH, Steinheim, Germany), at room temperature, for 48 h. The solution was exchanged in every 6 h. At the end of the decellularization protocol, the aortic segments were washed using PBS for 12 h to effectively remove residual detergents and cell debris, then stored in Medium 199 with Earle’s salts (Biowest, Nuaillé, France) and maintained at 4 °C, until measurements [9, 10].

Light microscopy evaluation

Samples were fixed for 24 h in 10% neutral buffered formalin, in support of routine histological examination. Longitudinal and cross-sectional tissue blocks were embedded using paraffin, and 2–4-μm-thick sections were sliced. Sections were stained applying hematoxylin–eosin (HE) staining to evaluate the effectiveness of decellularization. Besides this, Masson trichrome (MT) and orcein (O) histochemistry was used to examine the changes in the structure of the connective tissue fibers, using protocol recommended by the manufacturers. HE-, MT- and orcein-stained histological sections of decellularized and native samples before and after DSC

Fig. 1 Hematoxylin–eosin (HE), Masson trichrome (MT) and orcein (O) staining of native (N) and decellularized (D) aortic wall
were compared and analyzed using Pannoramic Viewer soft-
ware (3DHistech Ltd.) and were evaluated by two independ-
ent pathologists. Six samples \( (n = 6) \) were assessed in each

group.

**Transmission electron microscopy (TEM) assessment**

Samples were prepared for TEM examination by fixation in
2% formaldehyde- and 2.5% glutaraldehyde-containing
solution, for 12 h, at 4 °C. Following fixation, 50–60-nm-
thick sections were cut, and examination was performed
using a JEOL JEM-1200EX II transmission electron micro-
scope, at 80 kV.

**DSC measurements**

The thermal denaturation of the samples was investigated
with a SETARAM Micro DSC-II calorimeter. Each analysis

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**Fig. 2** TEM images of native (N) and decellularized (D) aortic walls

**Fig. 3** HE staining of native (N) and decellularized (D) aortic walls after DSC analysis

**Fig. 4** Orcein staining of native (N) and decellularized (D) aortic walls after DSC analysis
was performed within the range of 0 and 100 °C with a heating rate of 0.3 K min⁻¹. Conventional Hastelloy batch vessels (Vₘₐₓ = 1 mL) were applied for the experiment with an average sample mass of ~150 mg (range 110–200 mg). Normal saline solution was used as a reference. The reference and sample vessels were equilibrated with a precision of ±0.1 mg, so heat capacity correction between the sample and reference was not needed. Routine baseline correction is performed by a second scan of the denatured samples. In recent experiments, our samples showed the presence of partially reversible compounds, so we used the appropriate part of the Origin 6.0 software to smooth the experimental curves and correct the baseline. The melting temperature of the samples (Tₘ) was defined as the peak of the heat transition curves (at this point the 50% of sample turns from native into denatured state). The calorimetric enthalpy change (ΔH(cal)) was calculated from the area under the heat absorption curve with the SETARAM two-point setting software. The sample size in grams was used to normalize the ΔH(cal) values (in J g⁻¹ unit).

**Results and discussion**

Nowadays, the utilization of biological xenografts (graft originates from another species) shows an increasing number, as a result of the growing demands in vascular surgery, decisively in the treatment of septic vascular prostheses [11–14]. However, frequent, late aneurysm formation due to their degeneration is the most unfavorable consequence of their use. A well-known reason behind it is the immune-mediated graft rejection [6, 10, 15]. Several different types of mechanical tests were previously performed; nevertheless, none of them could provide information on, whether there was any association between late graft degeneration and decellularization treatment. DSC analysis can reveal structural alteration of macromolecules, before the defect of tissues becomes evident [16]. Up to the present day, we found just a few publications in the literature used DSC analysis

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**Fig. 5** MT staining of native (N) and decellularized (D) aortic walls after DSC analysis.

**Fig. 6** a The denaturation and renaturation heat flows of untreated aorta to illustrate the main thermal processes in the first heating–cooling cycle (after smoothing and baseline correction using Origin 6). b The denaturation and renaturation heat flows of untreated aorta in the second heating–cooling cycle focusing on the low-temperature range (after smoothing and baseline correction using Origin 6).
for characterization of decellularized aortic wall [17, 18].
We have planned and performed histological evaluation and
DSC analysis of the decellularized porcine aortic wall, to
assess the structural changes following treatment, which can
affect the long-term mechanical stability.

**Light microscopy evaluation of native and decellularized aortic wall**

In the first series, decellularized samples were compared
with native aortic sections as controls, to confirm the
effectiveness of decellularization protocol. Histological
findings with HE staining demonstrated complete cell
removal in treated samples, and there were not visible,
blue-stained cell nuclei on the slices of decellularized
aortic wall. MT and orcein histochemistry confirmed pre-
served structure of connective tissue fibers (collagen and
elastic fibers) after decellularization. Compared with the
untreated aortic wall, on cross-sectional slices of decellu-
larized aorta only the two layers, tunica media and adven-
titia, were identifiable, and decreased wall thickness was
noticeable in case of each treated sample. These findings
were proved consistently uniform throughout all investi-
gated samples (Fig. 1).

**Transmission electron microscopy assessment**

TEM images confirmed the absence of cells and cell nuclei
among the preserved, regular-structured collagen fibers
(Fig. 2).

**Histological analysis after thermal denaturation**

In the second series, we compared the native, untreated
and the decellularized aortic sections before and after DSC
measurements. The routine, HE-stained sections showed
minor alteration in the structure of connective tissue fibers
after DSC analysis, and this change was more significant in
the surface near the luminal, where some fragmentation of
fibers was detectable (Fig. 3).

Collagen and elastic fibers were stained selectively with
specific immunohistochemistry to better illustrate the struc-
tural changes of the different types of fibers. Orcein staining
showed no detectable alteration in the elastic fiber structure
after DSC measurement, the run of the fibers is regular, and
there was no significant difference compared with the native
aorta (Fig. 4).

In case of MT staining, which is specific for collagen,
there are some irregular, more undulatory, slightly frag-
mented collagen fibers, especially near the luminal sur-
face, which suggests mild damage in their structure fol-
lowing decellularization treatment (Fig. 5).

**Thermal denaturation measurements**

Aortic segments underwent two heating–cooling cycles. In
the case of the first step, we have checked the reversibility
of thermal transitions under cooling, and the second one is
generally used to perform instrument baseline correction.
The samples were denoted as untreated and decellularized.
Figure 6a shows a typical first heating–cooling cycle of
untreated aorta wall.

| Table 1 | The average characteristic thermal parameters of the denatur-
| | nation processes: ΔT°C stands for the temperature range of transition,
| | Tm°C is the denaturation temperature at the maximum of the given
| | heat flow and ΔH/J g−1 is the calorimetric enthalpy normalized on the
| | wet mass of the sample |
| **Native samples** | **High-temperature range** |
| Low-temperature range | ΔT°C | Tm°C | ΔH/J g−1 | ΔT°C | Tm°C | Tm2°C | ΔH/J g−1 |
| n = 6 |  |  |  |  |  |  |  |
| First heating | 15 | 30.8 | 5.4 | 24 | 53.4 | 59.6 | 25 |
| Cooling | 16 | 23.5 | −1.3 |  |
| Second heating | 16 | 29.2 | 7.1 |  |
| Cooling | 16 | 22.5 | −1.3 |  |
| Treated samples | **Low-temperature range** | **High-temperature range** |
| n = 8 | ΔT°C | Tm°C | ΔH/J g−1 | ΔT°C | Tm°C | Tm2°C | ΔH/J g−1 | ΔT°C | Tm°C | Tm2°C | ΔH/J g−1 |
| First heating | 16 | 7.4 | 2.95 | 16 | 37.8 | 2.14 | 33 | 47 | 56 | 6.8 |
| Cooling | 14 | 21.5 | −1.18 |  |
| Second heating | 13 | 9.1 | 1.68 | 19 | 31.2 | 1.93 |  |
| Cooling | 15 | 22.5 | −2.8 |  |
During heating, two major denaturation processes are going on: Very probably around 30 °C is the thermal transition of elastin [19] and ~ 60 °C the collagen denaturation [20–24]. Some small pretransition can be seen ~ 20 and 48 as well as at 54 °C. These two later peaks (which give only small enthalpy contribution) should have a myosin and actin origin on the basis of measurements performed on muscle protein solutions and skeletal muscle fibers [25–31]. In the literature, we have not found similar experiments performed on smooth muscles. The cooling process showed a near 100% renaturation in enthalpy of the denaturation peak at ~ 30 °C as an exothermic peak of 25°C. The main denaturation transition (~ 60 °C) disappeared during cooling, indicating that the change in collagen structure during denaturation is irreversible (Table 1).

The second heating–cooling cycle exhibited surprising result (Fig. 6b). The low-temperature endotherm and exotherm remained in the same temperature range and \( T_{m} \) with practically unchanged calorimetric enthalpies, while the higher transitions disappeared.

During the first heating–cooling cycle of the treated aorta wall, a low endotherm appeared around 10 °C, which was detectable many times in case of cross-striated muscle fibers (unpublished data), and we could not attach it to any structural element (Fig. 7a).

The elastin compound enthalpy contribution ~ 30 °C decreased, and the main transition became wider. The so-called \( T_{1/2} \) half width (the temperature range where the heat flow is half of its maximum) extremely increased, informing about the structural loosening compared with the untreated samples, as a consequence of removing most of the muscle proteins during the treatment. It means that the strength of cooperativity depends on the protein collagen matrix interaction. It can be the sign of the decreased elastic capability of aorta wall. In case of the cooling phase, only one dominating sharp exothermic peak was detected, referring basically to the elastin refolding.

Under the second heating–cooling cycle of treated sample (Fig. 7b), during heating the low-temperature melting position has shown about 2 °C shift to lower value, and the elastin melting became higher in calorimetric enthalpy. The cooling cycle exhibited a “fusion” exotherm of elastic compounds during refolding.

**Conclusions**

Earlier attempts demonstrated adequate biomechanical properties and suture retention of cell-free vascular grafts [3, 10]. However, these investigations did not provide eligible information about the microstructural changes in the connective tissue fibers that may influence the mechanical stability and long-term behavior of these decellularized vessels. For this purpose, DSC analysis was performed.

Our results showed that after decellularization of the aortic tissue, minor structural changes develop in the scaffold, which can decrease the mechanical stability, and consequently the long-term durability of the treated xenograft. Decellularization process causes mild connective tissue fiber damage: irreversible structural change in collagen fibers and reversible alteration in the structure of elastin fibers, which leads to weakened scaffold strength and reduced elasticity. These factors, together with the decreased vessel wall thickness, predispose for the development of late degenerative complications and consecutive aneurysm formation.
This work demonstrated that for the above-mentioned late consequences of cell-free grafts’ utilization not only the well-known graft versus host immune reaction, but also the decellularization treatment of the vessel is responsible. Therefore, in cases, when biological xenografts’ application is necessary for human application, postoperative close follow-up is mandatory.

For the future, DSC examination can be used to investigate the effects of different types of decellularization methods on the vessels’ connective tissue fiber structure.

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References

1. Nemes A, Acsády G, Fraelw F, Lichten H, Monos E, Oertl R, Somogyi E, Sótonyi P. Application of a vascular graft material (Solcograft-P) in experimental surgery. Biomaterials. 1985;6(5):303–11.

2. Schröder A, Imig H, Peiper U, Neidel J, Petereit A. Results of conventional prostheses. PLoS ONE. 2014;9(7):e103588.

3. MintáI T, Pataczai B, Wiegand N, Kereskai L, Váncsodi J, Lőrinczy D. The effect of deep-freezing on the structure of patellar and Achilles tendon allografts used for ACL reconstruction. J Therm Anal Calorim. 2017;127:1171–5.

4. Willett TL, Labow RS, Lee JM. Increased proteolysis of collagen in an in vitro tensile overload tendon model. Ann Biomed Eng. 2007;35(15):1514–9.

5. Lőrinczy D, Könczöl F, Gaszner B, Belágyi J. Structural stability of actin as studied by DSC and EPR. Thermochim Acta. 1997;296:18–161.

6. Willett TL, Labow RS, Lee JM. Thermal stability of collagen in an in vitro tensile overload tendon model. J Orthop Res. 2008;26(12):1605–10.

7. Willett TL, Labow RS, Lee JM. Thermal stability of collagen in an in vitro tensile overload tendon model. J Bone Joint Soc Br. 2009;91:217–20.

8. Wiegand N, Vándhy L, Lőrinczy D. Differential scanning calorimetric examination of ruptured lower limb tendons in human. J Therm Anal Calorim. 2010;101:487–92.

9. Lőrinczy D, Belágyi J. Effects of nucleotide on skeletal muscle myosin unfolding in myofibrils by DSC. Biochem Biophys Res Commun. 1995;217:592–8.

10. Lőrinczy D, Belágyi J. Internal motions in the catalytic domain of myosin heads in muscle fibres. Effects of ADP and ATP. Biochem Biophys Res Commun. 1996;219:936–40.

11. Lőrinczy D, Belágyi J. Comparative study of myosins in solutions and supramolecular complexes. Effect of nucleotides. Thermochim Acta. 1997;296:18–161.

12. Lőrinczy D, Könczöl F, Gaszner B, Belágyi J. Structural stability of actin as studied by DSC and EPR. Thermochim Acta. 1998:322:95–100.

13. Lőrinczy D, Belágyi J. Nucleotide binding induces global and local structural changes of myosin head in muscle fibres. Eur J Biochem. 2001;268:5970–6.
30. Dergez T, Lőrinczy D, Könczöl F, Farkas N, Belagyi J. Differential scanning calorimetry study of glycerinated rabbit psoas muscle fibres in intermediate state of ATP hydrolysis. BMC Struct Biol. 2007;7:41–50.

31. Lőrinczy D, Vértes ZS, Könczöl F, Belagyi J. Thermal transitions of actin. J Therm Anal Calorim. 2009;95:713–9.

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