

Open Access

ORAL PRESENTATION

Cardio-vascular tissue processing

Ramadan Jashari

From 23rd World Congress of the World Society of Cardio-Thoracic Surgeons Split, Croatia. 12-15 September 2013

The Cardio-vascular Tissues aimed for clinical use are submitted to the European Regulation (Directives 2004/ 23EC, 2006/17EC and 2006/86EC) to assure high level of tissue quality and security. The European Directive was transposed into the Member state legislation and Standards, specific to their needs and possibilities.

The Cardio-vascular tissues are recovered in the operating theatre or another area which has a controlled environment conform to the GMP regulation. The aseptic infrastructure is mandatory for all steps of the tissue recovery, transport and processing.

Processing of cardio-vascular tissues has to be carried out in a class A laminar flow conditions, background class C, with fully controlled environment for particles and aerobe, anaerobe germs and fungi. The temperature and the air pressure need to be adapted to the GMP regulation, assuring the pressure-cascade with a higher level at processing area and lower pressure around. The substances that come in contact with tissues (Hanks, dimethyl-sulfoxide, Antibiotics) that are mandatory for assuring the quality of tissues, may not provoke any deterioration of the tissue, assuring that the cells and matrix remain intact at the end of processing. Morphology, measurements, functional test and cryopreservation and storage have to be strictly respected and notified for the implanting surgeon.

Regular environmental control is mandatory, in order to guarantee the "sterile conditions" in zone A, where the tissue is processed. The critical procedures have to be mentioned in the Quality Management System (QMS) and validated conform the type of the tissue. Any kind of non-conformity during the tissue processing must be notified and corrected.

The Tissue Establishment needs appropriate staff and equipment. The QMS has to be established and contain following elements: Quality Manager and a replacing person, a manual of quality which describes all the

Correspondence: r.jashari@ehb.org European Homograft Bank (EHB) International Association, Brussels, Belgium procedures and the way of managing the quality system, and the SOP (Standard Operating Procedures) which describe any kind of procedures regarding tissue processing. Of course, the appropriate equipment and material is needed. On top of all, a Medical Doctor has the entire responsibility of the Tissue Establishment.

Documenting of all steps of tissue processing from the recovery to the implantation of tissues is mandatory.

Published: 11 September 2013

doi:10.1186/1749-8090-8-S1-O104 Cite this article as: Jashari: Cardio-vascular tissue processing. Journal of Cardiothoracic Surgery 2013 8(Suppl 1):O104.

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

BioMed Central

Submit your manuscript at www.biomedcentral.com/submit



© 2013 Jashari; licensee BioMed Central Ltd. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/2.0), which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.