

Week 7 - Required and Additional reading



This document was produced under the EU4Health Programme under a service contract with the European Health and Digital Executive Agency (HaDEA) acting under the mandate from the European Commission. The information and views set out in this document are those of the author(s) and do not necessarily reflect the official opinion of the Commission/Executive Agency. The Commission/Executive Agency do not guarantee the accuracy of the data included in this document. Neither the Commission/Executive Agency nor any person acting on the Commission's/Executive Agency's behalf may be held responsible for the use which may be made of the information contained therein.



Week 7 - SoHO VIGILANCE AND BIOVIGILANCE

22/01/24 - 26/01/24

Tutors and External experts involved: A.P. Barreiros, F. Bariani

		REQUIRED READING	SELECTED PARTS			
EU	LEGAL ACTS					
BLO	BLOOD					
1.	DIRECTIVE 2002/98/EC	Setting standards of quality and safety for the collection, testing, processing, storage and distribution of human blood and blood components and amending Directive 2001/83/EC	Art. 3 (l) Chapter V Haemovigilance Chapter VII Data Protection			
2.	DIRECTIVE 2005/61/EC	Implementing Directive 2002/98/EC of the European Parliament and of the Council as regards traceability requirements and notification of serious adverse reactions and events	all			
TIS	TISSUE					
3.	DIRECTIVE 2004/23/EC	Setting standards of quality and safety for the donation, procurement, testing, processing, preservation, storage and distribution of human tissues and cells	all			
4.	DIRECTIVE 2006/86/EC	Implementing Directive 2004/23/EC of the European Parliament and of the Council as regards traceability requirements, notification of serious adverse reactions and events and certain technical requirements for the coding, processing, preservation, storage and distribution of human tissues and cells	all			
OR	ORGANS					
5.	DIRECTIVE 2010/53/EU	Standards of quality and safety of human organs intended for transplantation	Art. 3 (n) (o) (s) Art. 4 Framework for quality and safety Art. 10 Traceability Art. 11 Reporting system and management concerning serious adverse events and reactions			





			Annex Part B			
6.	DIRECTIVE 2012/25/EU	Commission Implementing Directive 2012/25/EU of 9 October 2012 laying down information procedures for the exchange, between Member States, of human organs intended for transplantation Text with EEA relevance	all			
GL	GUIDELINE DOCUMENTS					
7.	EDQM Guide	EDQM Guide to the preparation, use and quality assurance of blood component 21st Edition	2.3.4. Interventions and treatment Chapter 10 Haemovigilance 11.12. Traceability and haemovigilance			
8.	EDQM Guide	EDQM Guide to the quality and safety of tissues and cells for human application 5th Edition	Chapter 16 Traceability Chapter 17 Vigilance			

		SELECTED PARTS					
GL	GUIDELINE DOCUMENTS						
	Deceased donation article	The critical pathway for deceased donation: reportable uniformity in the approach to deceased donation.					
		$Transpl\ Int.\ 2011\ Apr; 24(4): 373-8.\ doi:\ 10.1111/j.1432-2277.2011.01243.x.\ (Domínguez-Gil\ B,\ Delmonicons)$	All				
		FL, Shaheen FA, Matesanz R, O'Connor K et al.)					
2.	TRANSPOSE D5.2	Report and a scientific paper on donor eligibility criteria	All				
3.	TRANSPOSE D5.3	Criteria for the selection and protection of donors	All				
4.	VOX SANGUINIS 2020	Putting the spotlight on donation-related risks and donor safety - are we succeeding in protecting donors?	All				
5.	VOX SANGUINIS 2022	Blood donor eligibility criteria for medical conditions: A BEST collaborative study	All				
6.	EDQM Guide	EDQM Guide to the quality and safety of tissues and cells for human application 5th Edition	Appendices 28, 29, 30, 31				





EU	EU LEGAL ACTS				
1.	DIRECTIVE 2005/62/EC	Implementing Directive 2002/98/EC of the European Parliament and of the Council as regards Community standards and specifications relating to a quality system for blood establishments	Art. 1 (g) Annex - 9. Non-Conformance		
2.	DIRECTIVE 2016/1214/EC	Amending Directive 2005/62/EC as regards quality system standards and specifications for blood establishments	Art. 1		
3.	DIRECTIVE 2006/17/EC	Implementing Directive 2004/23/EC of the European Parliament and of the Council as regards certain technical requirements for the donation, procurement and testing of human tissues and cells including ANNEX I, ANNEX II	Art. 1 (g) Traceability Art. 2.11. Annex IV 1.4.4.		
4.	Commission Decision of 3.8.2010	Establishing TC guidelines concerning the conditions of inspections and control measures, and on the training and qualification of officials	all		
5.	Commission Decision of 3.7.2015	Establishing a model for agreements between the Commission and relevant organisations on the provision of product codes for use in the Single European Code	all		
6.	EC REPORT on standards of quality and safety of human organs	Report from the Commission to the European Parliament, the Council, the European Economic and Social Committee and the Committee of the Regions on the implementation of Directive 2010/53/EU of the European Parliament and of the Council of 7 July 2010 on standards of quality and safety of human organs intended for transplantation	Part 4		

