DISCLAIMERS

Products
The EUTC is a non-exhaustive list of (product codes for) substances of human origin which fall within the definition of either ‘tissue’ or ‘cells’ as defined in Directive 2004/23/EC. Without prejudice to the possibility for Member States to regulate such substances under the legislative frameworks for blood, organs, medicinal products (including advanced therapy medicinal products) or medical devices, substances in this list are regulated in at least one Member State under Directive 2004/23/EC and therefore subject to the EU requirements on the coding of human tissues and cells intended for human application.

Tissue Establishments
As defined in Directive 2006/86/EC as amended by Directive (EU) 2015/565, “EU Tissue Establishment Compendium” means the register of all tissue establishments which are authorised, licensed, designated or accredited by the Member States' competent authority or authorities and which contains the information about these tissue establishments as set out in Annex VIII to Directive 2006/86/EC as amended. Under Article 10b, point 2d of Directive 2006/86/EC as amended, Member States shall ensure that all competent authorities validate the data on the tissue establishments contained in the EU Tissue Establishment Compendium for their Member State and update this Compendium without undue delay. While the European Commission hosts the EU Coding Platform, the responsibility for ensuring that all data relating to tissue establishments within this Compendium is kept up-to-date and is correct, remains at all times with the Member State competent authorities.