



Online Resources and Links for the 3rd Edition Standards for Immune Effector Cells and Accreditation Manual

Resource Name/URL	Location Regular font indicates that the resource appears in the Accreditation Manual. Bold font indicates that the resource appears in both the Standards and the Accreditation Manual.
FACT Email	Contact Information
fact@factglobal.org	
JACIE Email	Contact Information
jacie@ebmt.org	
FACT Home Page	Contact Information, Introduction
http://www.factglobal.org	
JACIE Home Page	Contact Information, Introduction
https://www.ebmt.org/jacie	
Regulation (EC) No 765/2008	Introduction
http://data.europa.eu/eli/reg/2008/765/2021-07-16	
FACT Accredited Organizations	Introduction
https://accredited.factglobal.org	
JACIE Certified Centres	Introduction
https://www.ebmt.org/jacie-certified-centres	
21 USC 351: Adulterated drugs and devices	A4 (Good Manufacturing Practice)
https://uscode.house.gov/view.xhtml?req=(title:21%20section:351%20edition:prelim)	
ISBT 128 Standard Terminology for Medical Products of Human Origin (ST	
https://www.isbt128.org/standard-terminology	D7.1.1, D7.1.2, Appendix II
21 CFR Part 1271 Subpart D—Current Good Tissue Practice	B1.2, B4.4.2.5, C1.2, C4.4.2.5, D1.2,
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-L/part-1271/subpart-D	D4.4.2.5
Title 21 Food and Drugs	B1.2, C1.2
https://www.ecfr.gov/current/title-21	





Resource Name/URL	Regular font indicates that the resource appears in the Accreditation Manual.
Directive 2004/23/EC	fold font indicates that the resource appears in both the Standards and the Accreditation Manual. B1.2, B4.11.4.2, B8.2.1.4, B10.3.5, C1.2,
Directive 2004/25/2C	C4.11.4.2, D1.2, D3.1.1, D3.2.1,
http://data.europa.eu/eli/dir/2004/23/2009-08-07	D4.11.4.2, D13.3.5
Directive 2006/17/EC	B1.2, B4.11.4.2, C1.2, C4.11.4.2, C8.3.9,
http://data.europa.eu/eli/dir/2006/17/2012-12-17	D1.2, D4.11.4.2, D6.3.9, D11.1.1
Directive 2006/86/EC	B1.2, B4.11.4.2, B6.1, B7.4, B10.3.6,
http://data.compag.go./ali/dia/2006/06/2015 04/20	C1.2, C4.11.4.2, C10.2.2, C12.3.1, D1.2,
http://data.europa.eu/eli/dir/2006/86/2015-04-29	D4.11.4.2, D7.3.1.3, D7.4.3, D11.1.1
Directive 2001/83/EC	B1.2, C1.2, D1.1, D1.2
http://data.europa.eu/eli/dir/2001/83/2025-01-01	
Regulation (EC) No 1394/2007	B1.2, C1.2, D1.1, D1.2
http://data.europa.eu/eli/reg/2007/1394/2019-07-26	
Regulation (EU) 2024/1938	B1.2, C1.2, D1.2
http://data.europa.eu/eli/reg/2024/1938/2024-07-17	
Haute Autorité de Santé (HAS)	B1.9
https://www.has-sante.fr/	
Deutsche Krebsgesellschaft (DKG)	B1.9
https://www.krebsgesellschaft.de/	
Care Quality Commission (CQC)	B1.9
https://www.cqc.org.uk/	
Qualicor	B1.9
https://www.qualicor.eu/	
FACT Policies and Standard Operating Procedures (see Maintaining Accreditat	ion under Organizations, Policies) B2.1
https://www.factglobal.org/education-and-resources/general/fact-policies-and-standard-ope	erating-procedures/
Maintenance of Certification 2.0 — Strong Start, Continued Evolution	B3.1.1
https://www.nejm.org/doi/full/10.1056/NEJMp1409923	





Resource Name/URL Bold font	Location Regular font indicates that the resource appears in the Accreditation Manual. indicates that the resource appears in both the Standards and the Accreditation Manual.
Boarded to Death — Why Maintenance of Certification Is Bad for Doctors and Patient	B3.1.1
https://www.nejm.org/doi/full/10.1056/NEJMp1407422	
General Medical Council – Specialist or GP applications	B3.1.1, B3.2.2
https://www.gmc-uk.org/registration-and-licensing/our-registers/a-guide-to-our-registers/specialist-	and-gp-application-types
Educational Activities Form (under Forms on the FACT Immune Effector Cell Standard	s webpage) B3.1.5, B3.4.2, B3.10.2, C3.1.3.3, D3.1.4
https://factglobal.org/standards/immune-effector-standards/	
ASTCT – The Learning Center	B3.1.5, C3.1.3.3, D3.1.4
https://learn.astct.org/	
ESH – eLearning	B3.1.5, B3.3.4, B3.5.2, C3.1.3.3, D3.1.4
https://elearning.esh.org/esh	
EBMT – e-Learning	B3.1.5, B3.3.4, B3.5.2, C3.1.3.3, D3.1.4
https://www.ebmt.org/learning-projects/e-learning	
BSBMTCT – e-Learning	B3.3.4, B3.5.2
https://bsbmtct.org/e-learning/	
AABB – Cellular Therapy: A Handbook, 2 nd Edition	B3.3.4.13, C9.2
https://www.aabb.org/aabb-store/product/cellular-therapy-a-handbook-2nd-editionprint-17073790	
NIH Chronic Graft-Versus-Host Disease Consensus Conference 2025 Update	B3.3.5.6
https://pubmed.ncbi.nlm.nih.gov/40409691/	
Consensus Conference on Clinical Practice in Chronic GVHD: Second-Line Treatment	of Chronic Graft-versus-Host Disease B3.3.6.5, C9.11.2
https://pubmed.ncbi.nlm.nih.gov/20685255/	
Extracorporeal photopheresis for the treatment of acute and chronic graft-versus-hosp practice recommendations from an Italian Society of Hemapheresis and Cell Manip Bone Marrow Transplantation (GITMO) consensus process	
https://pubmed.ncbi.nlm.nih.gov/23305044/	





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·	ource appears in both the Standards and the Accreditation Manual.
Extracorporeal photopheresis for treatment of adults and children with acute GVHD: UK consensus staten	
published literature	
https://pubmed.ncbi.nlm.nih.gov/24887389/	
Diagnosis and management of chronic graft-versus-host disease	B3.3.6.5, C9.11.2
https://pubmed.ncbi.nlm.nih.gov/22533811/	
Diagnosis and management of acute graft-versus-host disease	B3.3.6.5, C9.11.2
https://pubmed.ncbi.nlm.nih.gov/22533831/	
Guidelines on the Use of Therapeutic Apheresis in Clinical Practice - Evidence-Based Approach from the of the American Society for Apheresis: The Ninth Special Issue	Writing Committee B3.3.6.5, C9.11.2
https://pubmed.ncbi.nlm.nih.gov/37017433/	
ASTCT Special Interest Groups	B3.5.2, B3.7.4
https://www.astct.org/Membership/Special-Interest-Groups	
ONCC – Blood & Marrow Transplant Certified Nurse (BMTCN®)	B3.6.1
https://www.oncc.org/blood-marrow-transplant-certified-nurse-bmtcn	
The EBMT Textbook for Nurses, EBMT Online Learning Course for Nurses	B3.6.2.7
https://www.ebmt.org/ebmt-jacie-books/ebmt-textbook-nurses	
American Nurses Association – Nurse Staffing	B3.6.4
https://www.nursingworld.org/practice-policy/nurse-staffing/	
Press Ganey National Database of Nursing Quality Indicators (NDNQI)	B3.6.4
https://info.pressganey.com/press-ganey-blog-healthcare-experience-insights/your-comprehensive-guide-to-the-press-ganey	-national-database-of-
nursing-quality-indicators-ndnqi	
Conditioning Chemotherapy Dose Adjustment in Obese Patients: A Review and Position Statement by the	e American Society B3.7.2.5
for Blood and Marrow Transplantation Practice Guideline Committee	
https://www.astctjournal.org/article/S1083-8791(14)00050-0/fulltext	





Resource Name/URL Regular font indicates that the resource appears in both to	Location source appears in the Accreditation Manual. he Standards and the Accreditation Manual.
The Hematopoietic Cell Transplant Pharmacist: Roles, Responsibilities, and Recommendations from the ASBMT Pharmacy Special Interest Group	B3.7.2.5
https://pubmed.ncbi.nlm.nih.gov/29292057/	
Consensus recommendations for the role and competencies of the EBMT clinical pharmacist and clinical pharmacologist involved in hematopoietic stem cell transplantation	B3.7.2.5
https://pubmed.ncbi.nlm.nih.gov/31101890/	
EBMT – Pharmacist Committee	B3.7.4
https://www.ebmt.org/pharmacist-committee	
UK BMT Pharmacists Group (training passport)	B3.7.4
https://www.bopa.org.uk/membergroups/specialist-advisory-groups/bmt-group/	
American Society for Quality (ASQ) – Certification Pathway Tool	B3.9.2, C3.3.2, D3.3.2
https://www.asq.org/cert	
National Association for Healthcare Quality (NAHQ)	B3.9.2, C3.3.2, D3.3.2
https://nahq.org/credentials/	
FACT Quality Management Resource Center (see FACT Quality Handbook, FACT Quality Management Series Webinars)	B3.9.2, B4.8, B4.14, B4.15, B5.1, C3.3.2,
https://www.factglobal.org/education-and-resources/general/quality-management-resource-center/	C4.8, C4.14, C4.15, C5.1, D3.3.2, D4.8, D4.14, D4.15, D5.1
EBMT – The JACIE Guide	B3.9.2, B4.8, B4.14, B4.15, B5.1, C3.3.2,
https://www.ebmt.org/education/jacie-guide	C4.8, C4.14, C4.15, C5.1, D3.3.2, D4.8,
	D4.14, D4.15, D5.1 B3.10.2
FACT Data Management Resource Center	B3. 10.2
https://www.factglobal.org/education-and-resources/general/data-management-resource-center	D2 10 2
The EBMT Registry	B3.10.2
https://www.ebmt.org/registry/ebmt-registry	201110
Anthony Nolan – Psychological Assessment Guidance in HSCT (adults)	B3.11.1.2
https://www.anthonynolan.org/clinicians-researchers-hub/healthcare-professionals/patient-services/latest-clinical-guidelines	





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Psychological Consequences of Hematopoietic Stem Cell Transplant		B3.11.1.2
https://pmc.ncbi.nlm.nih.gov/articles/PMC3105969/		
HTA Guide to Quality and Safety Assurance for Tissues and Cells for Patien	t Treatment	B4.4, C4.4, D4.4
https://www.hta.gov.uk/guidance-professionals/regulated-sectors/human-application/html	a-guide-quality-and-safety-assurance	
ISO 9001:2015		B4.4.2.5, C4.4.2.5, D4.4.2.5
https://www.iso.org/standard/62085.html		
21 CFR PART 211—CURRENT GOOD MANUFACTURING PRACTICE FOR F	INISHED PHARMACEUTICALS	B4.4.2.5, C4.4.2.5, D4.4.2.5
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-C/part-211		
EDQM – Guide to the quality and safety of tissues and cells for human app	lication	B4.4.2.5, C4.4.2.5, D4.4.2.5
https://freepub.edqm.eu/publications/AUTOPUB_17/detail		
CIBMTR Data Collection Forms (Cellular Therapy Essential Data [CTED] for	rms)	B4.8.5.1, B9.2
https://cibmtr.org/CIBMTR/Data-Operations/Data-Collection-Forms		
EBMT Registry Data Collection Forms		B4.8.5.1, B9.2
https://www.ebmt.org/registry/ebmt-data-collection		
21 CFR PART 1271—HUMAN CELLS, TISSUES, AND CELLULAR AND TISSUE	E-BASED PRODUCTS B	4.10.6, C1.2.1, C4.10.4, D1.2.1,
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-L/part-1271		D4.10.7
E2E Pharmacovigilance Planning		B4.11.4.2, C4.11.4.2
https://www.fda.gov/regulatory-information/search-fda-guidance-documents/e2e-pharm	acovigilance-planning	
Good Pharmacovigilance Practices and Pharmacoepidemiologic Assessmen	ıt .	B4.11.4.2, C4.11.4.2
https://www.fda.gov/regulatory-information/search-fda-guidance-documents/good-pharm	macovigilance-practices-and-pharmacoepidemiologic-	
<u>assessment</u>		
The science and practice of current environmental risk assessment for gene	e therapy: a review	B5.1.16
https://www.isct-cytotherapy.org/article/S1465-3249(24)00677-7/abstract		
Preparing for the Unthinkable: Emergency Preparedness for the Hematopol	ietic Cell Transplant Program	B5.1.17, C5.1.20, D5.1.19
https://pmc.ncbi.nlm.nih.gov/articles/PMC7129195/		





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Impact of Severe Weather Conditions on Biological Products	B5.1.17, C5.1.20, D5.1.19
$\underline{\text{https://www.fda.gov/vaccines-blood-biologics/safety-availability-biologics/impact-severe-weather-conditions-biological-products}}$	
The Art of Writing and Implementing Standard Operating Procedures (SOPs) for Laboratories in Low-Resource Settings: Review of Guidelines and Best Practices	B5.3, C5.3, D5.3
https://journals.plos.org/plosntds/article?id=10.1371/journal.pntd.0005053	
Adolescents and Young Adults with Cancer	B5.3.6
https://www.cancer.gov/types/aya	
Integrating Geriatric Assessment into Cancer Care: A Conversation with Dr. Supriya Mohile	B5.3.6
https://www.cancer.gov/news-events/cancer-currents-blog/2018/geriatric-assessment-cancer-care-mohile	
CLSI Home Page	B5.3.10
https://clsi.org/	
Family donor care management: principles and recommendations	B6.3.1.2, B6.4.3, C6.3.1.2, C6.4.1
https://pubmed.ncbi.nlm.nih.gov/20023708/	
WMDA S(P)EAR alert: August 2011	B6.3.2.1, C6.3.10, C6.3.11
https://wmda.info/wp-content/uploads/2020/08/20110824-CLWG-SEAR-Alert-August-2011.pdf	
Granulocyte colony-stimulating factor (G-CSF) administration in individuals with sickle cell disease: time for a moratorium?	B6.3.6.2, C6.3.6.1
https://pubmed.ncbi.nlm.nih.gov/19513902/	
Granulocyte colony-stimulating factor-based stem cell mobilization in patients with sickle cell disease	B6.3.6.2, C6.3.6.1
https://pubmed.ncbi.nlm.nih.gov/18489998/	
Mobilization, collection, and processing of peripheral blood stem cells in individuals with sickle cell trait	B6.3.6.2, C6.3.6.1
https://pubmed.ncbi.nlm.nih.gov/11806986/	
WHO guiding principles on human cell, tissue and organ transplantation	B6.3.12.1, C6.3.12.1
https://iris.who.int/handle/10665/341814	





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Allogeneic hematopoietic stem cell donation-standardized assessment of donor outcome data: a consensus statement from the Worldwide Network for Blood and Marrow Transplantation (WBMT) https://pubmed.ncbi.nlm.nih.gov/22773129/	
NMDP Home Page https://www.nmdp.org/	B6.4.2
Anthony Nolan Home Page https://www.anthonynolan.org/	B6.4.2
HTA BMT code of practice (see Code G: Donation of allogeneic bone marrow and peripheral blood stem cells for transplantation) https://www.hta.gov.uk/guidance-professionals/codes-practice-standards-and-legislation/codes-practice	B6.4.3, C6.4.1
Eligibility Determination for Donors of Human Cells, Tissues, and Cellular and Tissue-Based Products (2007) https://www.fda.gov/regulatory-information/search-fda-guidance-documents/eligibility-determination-donors-human-cells-tissues-and-cellular-and-tissue-based-products	B6.4.4.1, B6.4.9, B6.4.17, C6.1, C6.4.7, C7.4.5
Recommendations for Determining Eligibility of Donors of Human Cells, Tissues, and Cellular and Tissue-Based Products (HCT/Ps) (2025 draft)	B6.4.4.1, B6.4.9, B6.4.17, C6.1, C6.4.2.1, C6.4.7, C7.4.5
https://www.fda.gov/regulatory-information/search-fda-guidance-documents/recommendations-determining-eligibility-donors-human-cells-tissuesand-cellular-and-tissue-based	<u>is</u>
21 CFR 1271.80(d)(2) https://www.ecfr.gov/current/title-21/chapter-l/subchapter-L/part-1271/subpart-C/section-1271.80#p-1271.80(d)(2)	B6.4.4.1, C6.4.2.1
Chapter 7: haemodilution, transfusion and donor testing (Guidance from the UK's Dept. of Health & Social Care) https://www.gov.uk/government/publications/guidance-on-the-microbiological-safety-of-human-organs-tissues-and-cells-used-in-transplantation/chapter-7-haemodilution-transfusion-and-donor-testing	B6.4.4.1, C6.4.2.1
FACT – Hematopoietic Progenitor Cell, Apheresis and Marrow Donor History Questionnaire https://factglobal.org/education-and-resources/general/applicant-education-and-resources/resources/donor-history-questionnaires-hpc-apheresis-and-hpc-marrow/	B6.4.8.8
21 CFR 1271.75 How do I screen a donor?	B6.4.8.8
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-L/part-1271/subpart-C/section-1271.75	





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Testing Human Cells, Tissues, and Cellular and Tissue-Based Product (HCT/	P) Donors for Relevant Communicable Disease	B6.4.8.8
Agents and Diseases		
https://www.fda.gov/vaccines-blood-biologics/safety-availability-biologics/testing-human-cdonors-relevant-communicable	cells-tissues-and-cellular-and-tissue-based-product-hctp-	
21 CFR 1271.55 What records must accompany an HCT/P after the donor-e records must I retain?	ligibility determination is complete; and what	B6.4.17, C6.1, C6.4.7, C7.4.5
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-L/part-1271/subpart-C/section	n-1271.55	
Circular of Information for the Use of Cellular Therapy Products, June 2024		B7.6.5, C7.4.4 , D7.4.2, D7.4.4 ,
https://factglobal.org/education-and-resources/general/applicant-education-and-resource	es/resources/	D11.2.4.3
The why, what, and how of the new FACT standards for immune effector cel	lls	B7.8
https://jitc.biomedcentral.com/articles/10.1186/s40425-017-0239-0		
Production of chimeric antigen receptor (CAR) T cells		B7.8
https://www.nature.com/nprot/posters		
ASTCT Consensus Grading for Cytokine Release Syndrome and Neurologic	Toxicity Associated with Immune Effector Cells	B7.8
https://pubmed.ncbi.nlm.nih.gov/30592986/		
CDC – Adult Immunization Schedule Notes		B7.11.1
https://www.cdc.gov/vaccines/hcp/imz-schedules/adult-notes.html		
Physical, psychological, and social sequelae following hematopoietic stem c	ell transplantation: a review of the literature	B7.11.2
https://onlinelibrary.wiley.com/doi/10.1002/pon.1399		
Psychosocial aspects of hematopoietic stem cell transplantation		B7.11.2
https://pmc.ncbi.nlm.nih.gov/articles/PMC8290998/		
Commission Directive (EU) 2015/565		B8.2.1.4
http://data.europa.eu/eli/dir/2015/565/oj		
21 CFR PART 54—FINANCIAL DISCLOSURE BY CLINICAL INVESTIGATORS		B8.4
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-A/part-54		





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Adoptive Cellular Therapies (ACT) Stakeholder's Council	B9.2
https://cibmtr.org/CIBMTR/About/Administrative-Committees/Adoptive-Cellular-Therapies-ACT-Council	
Part 11, Electronic Records; Electronic Signatures - Scope and Application	B10.4.1, C12.6.1, D13.4.1
https://www.fda.gov/regulatory-information/search-fda-guidance-documents/part-11-electronic-records-electronic-signatures-scope-and-application	
MoReg®	B10.4.1, C12.6.1, D13.4.1
https://www.moreq.info/	
NMDP – Policies & Protocols (see Data Use and Processing Policies under Transplant centers, Additional Resources)	B10.5.2
https://network.nmdp.org/policies-protocols	
21 CFR PART 207—REQUIREMENTS FOR FOREIGN AND DOMESTIC ESTABLISHMENT REGISTRATION AND LISTING FOR HUMAN DRUGS, INCLUDING DRUGS THAT ARE REGULATED UNDER A BIOLOGICS LICENSE APPLICATION, AND ANIMAL DRUGS, AND THE NATIONAL DRUG CODE	C1.2.1, D1.2.1
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-C/part-207	
21 CFR PART 807—ESTABLISHMENT REGISTRATION AND DEVICE LISTING FOR MANUFACTURERS AND INITIAL IMPORTERS OF DEVICES	C1.2.1, D1.2.1
https://www.ecfr.gov/current/title-21/chapter-I/subchapter-H/part-807	
Tissue Establishment Registration	C1.2.1, D1.2.1
https://www.fda.gov/vaccines-blood-biologics/biologics-establishment-registration/tissue-establishment-registration	
21 CFR 211.42(b)	C2.1.3, D2.1.3
https://www.ecfr.gov/current/title-21/part-211/section-211.42#p-211.42(b)	
CAR-T Cell Therapies From the Transfusion Medicine Perspective	C5.1.6
https://pubmed.ncbi.nlm.nih.gov/27067907/	
NICE – Guidance on the use of ultrasound locating devices for placing central venous catheters http://guidance.nice.org.uk/TA49	C6.3.11





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Practice Guidelines for Central Venous Access 2020: An Updated Report	by the American Society of Anesthesiologists Task C6.3.11
Force on Central Venous Access	
https://journals.lww.com/anesthesiology/fulltext/2020/01000/practice_guidelines_for_	central venous access.9.aspx
ICCBBA Home Page	C7.1.1, C7.1.2, D7.1.1, D7.1.2
https://www.isbt128.org/	
ISO 3166	C7.1.1, D7.1.1
https://www.iso.org/iso-3166-country-codes.html	
Eurocode-IBLS Home Page	C7.1.1, C7.1.2, D7.1.1, D7.1.2
https://www.eurocode.org/	
ICCBBA ST-018 (ISBT 128 Standard Labeling of Collection Products for C	Cellular Therapy Manufacturing) C7.1.3, C7.3.2.1, D7.1.3, D7.3.2.1
https://www.isbt128.org/ST-018	
Recognition and Use of a Standard for Uniform Blood and Blood Compo	nent Container Labels C7.2.11, D7.2.11
https://www.fda.gov/regulatory-information/search-fda-guidance-documents/recognit	ion-and-use-standard-uniform-blood-and-blood-component-
<u>container-labels</u>	
FACT ISBT 128 Hybrid (Split) Label Webinar	C7.3.2.1, D7.3.2.1
https://learn.factglobal.org/2024-on-demand-webinar-isbt128-hybrid-split-label	
ST-004 ISBT 128 Standard Labeling of Cellular Therapy Products	C7.4.1, D7.4.1
https://www.isbt128.org/ST-004	
European Commission	C8.3.9
https://commission.europa.eu/index_en	
21 CFR 1271.200 Equipment	C8.4.2.3
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-L/part-1271/subpart-D/sec	ction-1271.200
21 CFR Part 211 Subpart D—Equipment	C8.4.2.3
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-C/part-211/subpart-D	
21 CFR 1271.270(a) Records. General.	C9.10.1, D8.9.2
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-L/part-1271/subpart-D/sec	ction-1271.270





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Regulation (EU) 2016/679	C12.4
http://data.europa.eu/eli/reg/2016/679/2016-05-04	
Directive 2001/20/EC	D1.1
http://data.europa.eu/eli/dir/2001/20/2022-01-01	
21 CFR PART 210—CURRENT GOOD MANUFACTURING PRACTICE IN MANUFACTURING, PROCESSING, PACKING, OR HOLDING OF DRUGS; GENERAL	D1.2, D2.5
https://www.ecfr.gov/current/title-21/chapter-l/subchapter-C/part-210	
21 CFR PART 211—CURRENT GOOD MANUFACTURING PRACTICE FOR FINISHED PHARMACEUTICALS	D1.2, D2.5
https://www.ecfr.gov/current/title-21/chapter-I/subchapter-C/part-211	
21 CFR 211.52 Washing and toilet facilities.	D2.1.2
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