

International Compilation of Human Research Standards

2019 Edition

Compiled By:
Office for Human Research Protections
U.S. Department of Health and Human Services

PURPOSE

The International Compilation of Human Research Standards enumerates over 1,000 laws, regulations, and guidelines (collectively referred to as “standards”) that govern human subject protections in 131 countries, as well as standards from a number of international and regional organizations. First published in 2005, the Compilation was developed for use by researchers, IRBs/Research Ethics Committees, sponsors, and others who are involved in human subjects research protections around the world.

Content experts from around the world, listed at the end of the Compilation, provided updates (or confirmations of prior listings), which are reflected in the hundreds of changes entered into this Edition. One new country is featured in the 2019 Edition: El Salvador.

ORGANIZATION

The Table of Contents is on pages 3-4. For each country, the standards are categorized by row as:

1. General, i.e., applicable to most or all types of human subjects research
2. Drugs, Biologics, and Devices
3. Clinical Trial Registries
4. Research Injury
5. Social-Behavioral Research (also see Description and Analysis of Social-Behavioral Research Standards: <https://www.hhs.gov/ohrp/international/social-behavioral-research-standards/index.html>)
6. Privacy/Data Protection (also see Privacy International reports: <https://www.privacyinternational.org/reports>)
7. Human Biological Materials
8. Genetic (also see the HumGen International database: <http://www.humgen.umontreal.ca/int/>)
9. Embryos, Stem Cells, and Cloning

These nine categories often overlap, so it may be necessary to review the other standards to obtain an accurate understanding of the country’s requirements.

The information is then organized into four columns:

1. Key Organizations – include those groups that issue regulations or guidelines, or serve in a national oversight role for human subjects research.
2. Legislation – encompasses statutes, statutory instruments, and legislative decrees, as well as any pertinent constitutional provisions.
3. Regulations – refer to instruments that are created and issued in the name of governmental administrative bodies.
4. Guidelines – pertain to non-binding instruments.

The year of the document's most recent version (or date of initial approval, if never amended) is indicated in parenthesis when that information is available, unless the date is part of the document's title, e.g., Law No. 46/2018.

HOW TO ACCESS A DOCUMENT

Documents can be accessed in four possible ways:

1. Link to the web address (URL).
2. Search for a document at the website of the agency listed in the Key Organizations column.
3. Perform an Internet search on the document title.
4. Request a local research ethics committee to provide the document.

In many cases the documents are available in English. When the URL links to a non-English website or document, an online language translator usually can render an English version.

TOPICS NOT COVERED

In order to focus its scope, the International Compilation of Human Research Standards does not include standards from the state, provincial, or local levels. Nor does the Compilation cover:

1. Enabling legislation, i.e., laws that authorize an agency to promulgate human subjects standards, but do not direct the content of those regulations.
2. Laws, regulations, or guidelines that are disease-specific or focus on research integrity, clinical bioethics, product liability, clinical trial inspection procedures, intellectual property, good manufacturing practice, bioequivalence testing, or informed consent in clinical practice.
3. Ethics codes of academic, medical, or other professional organizations – see the Ethics Codes Collection: <http://ethics.iit.edu/ecodes/about>
4. Working papers, drafts, commentaries, or discussion papers.

NEW STANDARDS, UPDATES, AND BROKEN LINKS

To request inclusion of a new standard in the Compilation, or to report updates or broken links, contact Edward E. Bartlett, PhD, International Human Research Liaison, Office for Human Research Protections, U.S. Department of Health and Human Services: edward.bartlett@hhs.gov .

DISCLAIMER

Although this Compilation contains information of a legal nature, it has been developed for informational purposes only and does not constitute legal advice or opinions as to the current operative laws, regulations, or guidelines of any jurisdiction. In addition, because new standards are issued on a continuing basis, this Compilation is not an exhaustive source of all current applicable laws, regulations, and guidelines relating to human subject protections. While in-country persons have been requested to review listings to assure their accuracy and completeness, researchers and other individuals should check with local authorities and/or research ethics committees before commencing research activities.

TABLE OF CONTENTS

| | |
|--|-----------|
| INTERNATIONAL | 5 |
| NORTH AMERICA | 10 |
| Canada | 10 |
| United States | 12 |
| EUROPE | 20 |
| Regionwide | 20 |
| Armenia | 26 |
| Austria..... | 26 |
| Belarus | 28 |
| Belgium..... | 30 |
| Bosnia and Herzegovina | 32 |
| Bulgaria..... | 35 |
| Croatia..... | 38 |
| Cyprus..... | 40 |
| Czech Republic | 41 |
| Denmark..... | 42 |
| Estonia | 44 |
| Finland | 45 |
| France | 47 |
| Georgia..... | 49 |
| Germany..... | 50 |
| Greece | 54 |
| Hungary | 56 |
| Iceland..... | 59 |
| Ireland..... | 61 |
| Italy | 62 |
| Latvia | 64 |
| Lithuania | 66 |
| Luxembourg..... | 70 |
| Macedonia..... | 71 |
| Note: All websites and documents are available in Macedonian..... | 71 |
| Malta | 75 |
| Moldova | 76 |
| Montenegro..... | 78 |
| Netherlands | 80 |
| Norway..... | 81 |

| | |
|----------------------|-----|
| Poland | 84 |
| Portugal..... | 85 |
| Romania..... | 86 |
| Russia..... | 88 |
| San Marino..... | 90 |
| Serbia | 90 |
| Slovakia | 91 |
| Slovenia | 92 |
| Spain | 95 |
| Sweden..... | 99 |
| Switzerland | 101 |
| Ukraine..... | 105 |
| United Kingdom | 107 |

ASIA/PACIFIC

| | |
|----------------------------------|-----|
| Australia..... | 114 |
| Bangladesh..... | 117 |
| China, People’s Republic of..... | 117 |
| India | 120 |
| Indonesia | 123 |
| Japan | 124 |
| Kazakhstan..... | 127 |
| Korea..... | 127 |
| Kyrgyzstan | 130 |
| Malaysia..... | 132 |
| Myanmar | 132 |
| Nepal..... | 133 |
| New Zealand | 133 |
| Pakistan | 136 |
| Philippines | 136 |
| Singapore | 139 |
| Sri Lanka..... | 142 |
| Taiwan | 142 |
| Tajikistan | 145 |
| Thailand | 145 |
| Uzbekistan | 146 |
| Vietnam..... | 146 |

MIDDLE EAST/NORTH AFRICA.....

| | |
|--------------|-----|
| Egypt..... | 148 |
| Iran..... | 148 |
| Israel | 148 |
| Jordan..... | 149 |

| | |
|----------------------------|-----|
| Kuwait..... | 150 |
| Qatar | 150 |
| Saudi Arabia | 151 |
| Tunisia | 152 |
| Turkey..... | 152 |
| United Arab Emirates | 153 |

LATIN AMERICA AND THE CARIBBEAN.....

| | |
|--------------------------|-----|
| Regionwide | 154 |
| Argentina | 154 |
| Barbados | 155 |
| Bermuda..... | 155 |
| Bolivia..... | 156 |
| Brazil..... | 156 |
| Chile..... | 160 |
| Colombia..... | 162 |
| Costa Rica..... | 165 |
| Cuba..... | 165 |
| Dominica..... | 165 |
| Dominican Republic | 166 |
| Ecuador | 166 |
| El Salvador..... | 168 |
| Grenada..... | 168 |
| Guyana | 168 |
| Guatemala | 168 |
| Haiti | 169 |
| Honduras | 169 |
| Jamaica..... | 169 |
| México | 170 |
| Panamá..... | 171 |
| Perú | 172 |
| Saint Lucia | 172 |
| Trinidad and Tobago..... | 172 |
| Uruguay | 173 |
| Venezuela..... | 173 |

AFRICA

| | |
|--------------------|-----|
| Regionwide | 175 |
| Algeria | 175 |
| Benin..... | 175 |
| Botswana..... | 175 |
| Burkina Faso | 176 |
| Cameroon..... | 176 |

| | | | | | |
|------------------------------------|-----|--------------------|-----|------------------------------|------------|
| Congo, Democratic Republic of..... | 177 | Madagascar | 179 | South Africa | 183 |
| Côte-d'Ivoire | 177 | Malawi | 179 | Tanzania..... | 184 |
| Ethiopia..... | 177 | Mali..... | 181 | Uganda..... | 185 |
| Gambia..... | 177 | Mozambique | 181 | Zambia | 185 |
| Ghana | 177 | Nigeria | 181 | Zimbabwe | 186 |
| Guinea..... | 178 | Rwanda | 182 | ACKNOWLEDGEMENTS..... | 187 |
| Kenya..... | 178 | Senegal..... | 182 | | |
| Liberia..... | 179 | Sierra Leone | 182 | | |

Country

Key Organizations

Legislation

Regulations

Guidelines

| INTERNATIONAL | | | | |
|----------------------|--|--|--|--|
| <i>General</i> | Council for International Organizations of Medical Sciences (CIOMS): http://www.cioms.ch/ | | | International Ethical Guidelines for Health-Related Involving Humans (2016): https://cioms.ch/shop/product/international-ethical-guidelines-for-health-related-research-involving-humans/ |
| | International Committee of the Red Cross (ICRC): www.icrc.org | 1. Geneva Convention Relative to the Treatment of Prisoners of War, Articles 13 and 130 (1950): https://www.icrc.org/applic/ihl/ihl.nsf/7c4d08d9b287a42141256739003e636b/6fef854a3517b75ac125641e004a9e68 2. Additional Protocol I Relating to the Protection of Victims of International Armed Conflicts, Article 11 (1977): http://www.icrc.org/ihl.nsf/7c4d08d9b287a42141256739003e636b/f6c8b9fee14a77fdc125641e0052b079 | | |
| | Office of the United Nations High Commissioner for Human Rights (OHCHR): http://www.ohchr.org/english/ | International Covenant on Civil and Political Rights, Article 7 (1976): http://www.ohchr.org/en/professionalinterest/pages/ccpr.aspx | | |
| | TRUST Project: http://www.globalcodeofconduct.org | | | Global Code of Conduct for Research in Resource-Poor Settings (2018): http://www.globalcodeofconduct.org/wp-content/uploads/2018/05/Global-Code-of-Conduct-Brochure.pdf |
| | UNAIDS: http://www.unaids.org/ | | | 1. Good Participatory Practice: Guidelines for Biomedical HIV Prevention Trials (2011): http://www.unaids.org/sites/default/files/media_asset/JC1853_GPP_Guidelines_2011_en_0.pdf 2. Ethical Considerations in Biomedical HIV Prevention Trials (2012): http://www.unaids.org/en/media/unaids/contentassets/documents/unaidspublication/2012/jc1399_ethical_considerations_en.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--|---|-------------|---|---|
| <i>General</i> | United Nations Educational, Scientific, and Cultural Organization, Bioethics Program (UNESCO): https://en.unesco.org/ | | | Universal Declaration on Bioethics and Human Rights (2005): http://portal.unesco.org/en/ev.php-URL_ID=31058&URL_DO=DO_TOPIC&URL_SECTION=201.html |
| | World Health Organization: http://www.who.int/en/ | | | 1. Standards and Operational Guidance for Ethics Review of Health-Related Research with Human Participants (2011) 2. Ethical Issues in Patient Safety Research: Interpreting Existing Guidance (2013) 3. Managing Ethical Issues in Infectious Disease Outbreaks: Guidance Document (2016) 4. WHO Guidelines on Ethical Issues in Public Health Surveillance (2017) <i>Access:</i> http://www.who.int/ethics/publications/en/ |
| | World Medical Association: http://www.wma.net/e/ | | | Declaration of Helsinki (2013): https://www.wma.net/policies-post/wma-declaration-of-helsinki-ethical-principles-for-medical-research-involving-human-subjects/ |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | | | |
| | International Conference on Harmonization (ICH): http://www.ich.org/ | | | Integrated Addendum to ICH E6(R1): Guideline for Good Clinical Practice (2016): https://www.ich.org/fileadmin/Public_Web_Site/ICH_Products/Guidelines/Efficacy/E6/E6_R2_Step_4_2016_1109.pdf |
| | World Health Organization (WHO): http://www.who.int/en/ | | | 1. Handbook for Good Clinical Research Practice (GCP): Guidance for Implementation (2005): http://whqlibdoc.who.int/publications/2005/924159392X_eng.pdf 2. Operational Guidance: Information Needed to Support Clinical Trials of Herbal Products (2005) |
| | <i>Devices</i> | | | |
| International Medical Device Regulators Forum (IMDRF): http://www.imdrf.org/ | | | IMDRF: Statement Regarding Use of ISO 14155:2011 “Clinical Investigation of Medical Devices for Human Subjects- Good Clinical Practice” (2015): http://www.imdrf.org/docs/imdrf/final/proced | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|---|-------------|-------------|--|
| <i>Drugs, Biologics, and Devices</i> | | | | ural/imdrf-proc-150326-statement-iso141552011.pdf Archived Documents from the Global Harmonization Task Force (GHTF), replaced by the IMDRF in 2012: 1. Clinical Evaluation (2007) 2. Clinical Evidence – Key Definitions and Concepts (2007) 3. Post-Market Clinical Follow-Up Studies (2010) 4. Clinical Investigations (2010) 5. Reportable Events During Pre-Market Clinical Investigations (2012) 6. Clinical Evidence for IVD Medical Devices (2012) 7. Scientific Validity Determination and Performance Evaluation (2012) 8. Clinical Performance Studies for IVD Medical Devices (2012) Access: http://www.imdrf.org/ghf/ghf-archived-docs.asp |
| | International Standards Organization: http://www.iso.org/iso/home.html | | | Clinical Investigation of Medical Devices for Human Subjects -- Good Clinical Practice. Standard Number 14155:2011: http://www.iso.org/iso/iso_catalogue/catalogue_e_ics/catalogue_detail_ics.htm?csnumber=45557 |
| <i>Clinical Trials Registry</i> | World Health Organization – International Clinical Trials Registry Platform: http://www.who.int/ictrp/en/ | | | Resolution WHA 58.34 (2005): http://www.wpro.who.int/health_research/policy_documents/ministerial_summit_on_health_research_may2005.pdf?ua=1 |
| | World Medical Association: http://www.wma.net/e/ | | | Declaration of Helsinki, Article 35 (2013): http://www.wma.net/en/30publications/10policies/b3/index.html |
| | International Committee of Medical Journal Editors: http://www.icmje.org/ | | | Clinical Trial Registration: http://icmje.org/recommendations/browse/publishing-and-editorial-issues/clinical-trial-registration.html |
| <i>Research Injury</i> | World Medical Association: http://www.wma.net/e/ | | | Declaration of Helsinki, Paragraph 15 (2013): http://www.wma.net/en/30publications/10policies/b3/index.html |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|-----------------------------------|--|--------------------|--------------------|--|
| <i>Research Injury</i> | International Conference on Harmonization (ICH): http://www.ich.org/ | | | Integrated Addendum to ICH E6(R1): Guideline for Good Clinical Practice , Section 5.8 (2016): https://www.ich.org/fileadmin/Public_Web_Site/ICH_Products/Guidelines/Efficacy/E6/E6_R2_Step_4.pdf |
| | Council for International Organizations of Medical Sciences: http://www.cioms.ch/ | | | International Ethical Guidelines for Health-Related Involving Humans (2016), Guideline 14: https://cioms.ch/shop/product/international-ethical-guidelines-for-health-related-research-involving-humans/ |
| <i>Social-Behavioral Research</i> | UNESCO: http://www.unesco.org/ | | | Code of Conduct and Ethical Guidelines for Social Science Research: http://www.unesco.org/new/fileadmin/MULTIMEDIA/HQ/SHS/pdf/Soc_Sci_Code.pdf |
| <i>Privacy/Data Protection</i> | World Medical Association: http://www.wma.net/e/index.htm | | | 1. Declaration of Helsinki, Paragraph 24 (2013): http://www.wma.net/en/30publications/10policies/b3/index.html 2. Declaration of Taipei (2016): https://www.wma.net/policies-post/wma-declaration-of-taipei-on-ethical-considerations-regarding-health-databases-and-biobanks/ |
| <i>Human Biological Materials</i> | World Health Organization: http://www.who.int/en/ | | | 1. Guidelines for the Safe Transport of Infectious Substances and Diagnostic Specimens (1997): www.who.int/csr/emc97_3.pdf 2. Guideline for Obtaining Informed Consent for the Procurement and Use of Human Tissues, Cells, and Fluids in Research (2003): http://www.who.int/reproductivehealth/topics/ethics/human_tissue_use.pdf |
| | World Medical Association | | | Declaration of Taipei (2016): https://www.wma.net/policies-post/wma-declaration-of-taipei-on-ethical-considerations-regarding-health-databases-and-biobanks/ |
| | International Air Transport Association: http://www.iata.org/ | | | Infectious Substances and Diagnostic Specimens Shipping Guidelines (2005) |
| | International Society for Biological and Environmental Repositories: http://www.isber.org | | | Best Practices for Repositories I: Collection, Storage and Retrieval of Human Biological Materials for Research (2018): https://www.isber.org/page/BPR |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|--|-------------|-------------|---|
| <i>Genetic Research</i> | Human Genome Organization: http://www.hugo-international.org/ | | | 1. Statement on the Principled Conduct of Genetic Research (1996): http://www.eubios.info/HUGO.htm 2. Statement on DNA Sampling: Control and Access (1998): http://www.hugo-international.org/img/dna_1998.pdf 3. Statement on Gene Therapy Research (2001): http://www.hugo-international.org/img/gene_2001.pdf 4. Statement on Human Genomic Databases (2002): http://www.hugo-international.org/img/genomic_2002.pdf |
| | UNESCO Bioethics Program: http://portal.unesco.org/shs/en/ev.php-URL_ID=1372&URL_DO=DO_TOPIC&URL_SECTION=201.html | | | 1. Universal Declaration on the Human Genome and Human Rights Section 16 of III Programme for 1998-1999 (1997): http://unesdoc.unesco.org/images/0011/001102/110220e.pdf#page=47 2. International Declaration on Human Genetic Data: Section 22 of Major Programme III – Social and Human Sciences (2003): http://unesdoc.unesco.org/images/0013/001331/133171e.pdf#page=45 |
| <i>Embryos, Stem Cells, and Cloning</i> | International Society for Stem Cell Research: http://www.isscr.org/ | | | Guidelines for the Conduct of Human Embryonic Stem Cell Research (2006): http://www.isscr.org/docs/default-source/hesc-guidelines/issrhescguidelines2006.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--|--|-------------|---|--|
| NORTH AMERICA | | | | |
| Canada | | | | |
| Note: Several Canadian provinces and territories also have human subject research standards. | | | | |
| <i>General</i> | 1. Interagency Advisory Panel on Research Ethics (PRE): http://www.pre.ethics.gc.ca/eng/index 2. National Defence and the Canadian Armed Forces: http://www.forces.gc.ca/en/index.page 3. Correctional Service of Canada: http://www.csc-scc.gc.ca/index-eng.shtml | | | PRE: Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans, 2 nd Edition (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/Default/ National Defence and the Canadian Armed Forces: Research Involving Human Subjects (1998): http://www.forces.gc.ca/en/about-policies-standards-defence-admin-orders-directives-5000/5061-0.page Correctional Service of Canada: Commissioner's Directive - Research: DCOO9 (2017): http://www.csc-scc.gc.ca/acts-and-regulations/009-cd-en.shtml |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. Health Canada, Therapeutic Products Directorate: http://www.hc-sc.gc.ca/ahc-asc/branch-dirgen/hpfb-dgpsa/tpd-dpt/index-eng.php 2. Interagency Advisory Panel on Research Ethics (PRE): http://www.pre.ethics.gc.ca/eng/index | | 1. Regulations Amending the Food and Drug Regulations (1024 – Clinical Trials) (2001): http://www.hc-sc.gc.ca/dhp-mps/alt_formats/hpfb-dgpsa/pdf/compli-conform/1024-eng.pdf | Health Canada: Good Clinical Practice: Integrated Addendum to E6(R1) ICH Topic E6(R2) (2017) https://www.canada.ca/content/dam/hc-sc/migration/hc-sc/dhp-mps/alt_formats/pdf/prodpharma/applic-demande/guide-ld/ich/efficac/e6r2-step4-eng.pdf PRE: Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans, 2 nd Edition, Chapter 11: Clinical Trials (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/chapter11-chapitre11/ |
| | <i>Devices</i> Health Canada, Medical Devices: http://www.hc-sc.gc.ca/dhp-mps/md- | | Medical Devices Regulations (SOR/98-282) (1998): http://laws- | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--|--|---|--|--|
| <i>Drugs, Biologics, and Devices</i> | im/index-eng.php | | lois.justice.gc.ca/eng/regulations/sor-98-282/FullText.html | |
| <i>Clinical Trials Registry</i> | 1. Health Canada Clinical Trial Database: http://www.hc-sc.gc.ca/dhp-mpps/prodpharma/databasdonclin/index-eng.php 2. Interagency Advisory Panel on Research Ethics (PRE): http://www.pre.ethics.gc.ca/eng/index | | | PRE: Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans, Article 11.3 (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/chapter11-chapitre11/ |
| <i>Research Injury</i> | Interagency Advisory Panel on Research Ethics (PRE): http://www.pre.ethics.gc.ca/eng/index | | | Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans, 2 nd Edition, Article 3.2(j) (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/chapter3-chapitre3/ |
| <i>Social-Behavioral Research</i> | Interagency Advisory Panel on Research Ethics (PRE): http://www.pre.ethics.gc.ca/eng/index | | | Tri-Council Policy Statement, Ethical Conduct for Research Involving Humans. Exemptions (Chapter 2) and Qualitative Research (Chapter 10) (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/Default/ |
| <i>Privacy/Data Protection</i> Note: Each of the Canadian provinces and territories also has enacted privacy legislation. | 1. Office of the Privacy Commissioner of Canada (OPC): https://www.priv.gc.ca/en 2. Interagency Advisory Panel on Research Ethics (PRE): http://www.pre.ethics.gc.ca/eng/index 3. Canadian Institutes of Health Research (CIHR): http://www.cihr-irsc.gc.ca/e/193.html | 1. Privacy Act, Sections 7-8 (1983): http://laws-lois.justice.gc.ca/PDF/P-21.pdf 2. Personal Information Protection and Electronic Documents Act, Articles 5 and 7 (2001): http://laws-lois.justice.gc.ca/PDF/P-8.6.pdf | OPC: SOR/2001-6, SOR/2001-7, and SOR/2001-8 (September 29, 2014) | PRE: Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans, 2 nd Edition, Chapter 5: Privacy and Confidentiality (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/chapter5-chapitre5/ CIHR: CIHR Best Practices for Protecting Privacy in Health Research (2005): http://www.cihr-irsc.gc.ca/e/documents/et_pbp_nov05_sept2005_e.pdf |
| <i>Human Biological Materials</i> | Interagency Advisory Panel on Research Ethics (PRE): http://www.pre.ethics.gc.ca/eng/index | | | Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans, 2 nd Edition, Chapter 12: Human Biological Materials Including Materials Related to Human Reproduction (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/chapter12-chapitre12/ |
| <i>Genetic Research</i> | 1. Interagency Advisory Panel on Research Ethics (PRE): | | | PRE: Tri-Council Policy Statement: Ethical |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|---|---|---|---|
| <i>Genetic Research</i> | http://www.pre.ethics.gc.ca/eng/index 2. Canadian Biotechnology Advisory Committee (CBAC): http://www.hc-sc.gc.ca/sr-sr/biotech/role/strateg-eng.php 3. Health Canada, Biologics and Genetic Therapies Directorate: http://www.hc-sc.gc.ca/ahc-asc/branch-dirgen/hpfb-dgpsa/bgtd-dpbtg/index-eng.php | | | Conduct for Research Involving Humans, 2 nd Edition, Chapter 13: Human Genetic Research (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/chapter13-chapitre13/ |
| <i>Embryos, Stem Cells, and Cloning</i> | Interagency Advisory Panel on Research Ethics (PRE): http://www.pre.ethics.gc.ca/eng/index | Assisted Human Reproduction Act (2004): http://laws-lois.justice.gc.ca/eng/acts/A-13.4/ | Assisted Human Reproduction (Section 8 Consent) Regulations (2007): http://laws-lois.justice.gc.ca/eng/regulations/SOR-2007-137/index.html | PRE: Tri-Council Policy Statement: Ethical Conduct for Research Involving Humans, 2 nd Edition, Chapter 12, Sections E and F (2014): http://www.pre.ethics.gc.ca/eng/policy-politique/initiatives/tcps2-eptc2/chapter12-chapitre12/ |

United States

All of the following departments and agencies subscribe to subpart A, often referred to as the Common Rule (last updated in 2018), and codified in the relevant section of the Code of Federal Regulations: <http://www.hhs.gov/ohrp/humansubjects/guidance/45cfr46.html>. Some departments and agencies subscribe to additional subparts, such as:

- Subpart B: Additional Protections for Pregnant Women, Human Fetuses, and Neonates (2001)
- Subpart C: Additional Protections Pertaining to Biomedical and Behavioral Research Involving Prisoners as Subjects (1978)
- Subpart D: Additional Protections for Children Involved as Subjects in Research (1991)
- Subpart E: Institutional Review Board Registration Requirements (2009)

| | | | | |
|----------------|---|---|--|---|
| <i>General</i> | Agency for International Development: www.usaid.gov/ | | 22 CFR 225, Subpart A | Protection of Human Subjects in Research Supported by USAID: A Mandatory Reference for ADS Chapter 200 (2015): https://www.usaid.gov/sites/default/files/documents/1870/200.pdf |
| | Central Intelligence Agency: https://www.cia.gov/index.html | | Executive Order 12333, Subparts A, B, C, and D | |
| | Consumer Product Safety Commission: www.cpsc.gov/ | | 16 CFR 1028, Subpart A | |
| | Department of Agriculture: www.usda.gov/wps/portal/usdahome/ | | 1. 7 CFR 1c, Subpart A 2. 45 CFR 46, Subparts B, C, and D | Protection of Human Subjects (2011): https://www.afm.ars.usda.gov/media/10444/pp605-1.pdf |
| | Department of Commerce, National Institute of Standards and Technology: www.commerce.gov/ | | 15 CFR 27, Subpart A | |
| | Department of Defense, Human and Animal RDT&E Protection Programs: http://www.acq.osd.mil/rd/hptb/programs | United States Code Title 10, Section 980: Limitation on Use of Humans as Experimental | 1. 32 CFR 219, Subpart A 2. DoDI 3216.02 (2011): http://www.dtic.mil/whs/directives/corres/pdf/321602p.pdf | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|----------------|--|--|---|--|
| <i>General</i> | /regulatory/ | Subjects | | |
| | Department of Education: www.ed.gov/ | 1. Protection of Pupil Rights Amendment (1974) 2. Family Educational Rights and Privacy Act (1974) | 1. 34 CFR 97 subparts A (1991) and D (1997) 2. 34 CFR 98 (1984) 3. 34 CFR 99 (2000) 4. 34 CFR 350.4(c) (1991) 5. 34 CFR 356.3(c) (1991) | |
| | Department of Energy: http://science.energy.gov/ber/human-subjects/ | | 1. 10 CFR 745 (1991), Subpart A 2. DOE Order 443.1B 3. DOE Order 481.1 | |
| | Department of Health and Human Services, Office for Human Research Protections: www.hhs.gov/ohrp/ | Public Health Service Act (1993): http://history.nih.gov/research/downloads/PL103-43.pdf | 45 CFR 46, Subparts A, B, C, D, and E: http://www.hhs.gov/ohrp/humansubjects/guidance/45cfr46.html | Various: http://www.hhs.gov/ohrp/regulations-and-policy/ |
| | Department of Health and Human Services, Food and Drug Administration: https://www.fda.gov/ | | <i>FDA is not a Common Rule agency. For studies funded by FDA:</i> 45 CFR 46, Subparts A, D, and E: http://www.hhs.gov/ohrp/humansubjects/guidance/45cfr46.html | |
| | Department of Health and Human Services, National Institutes of Health: https://www.nih.gov/ | | | NIH Single IRB Policy (2016): https://grants.nih.gov/policy/clinical-trials/single-irb-policy-multi-site-research.htm |
| | Department of Homeland Security: www.dhs.gov/ | Public Law 108-458, Section 8306 | 1. 45 CFR 46, Subparts A-D 2. DHS Directive 026-04, Human Subjects Research (2007): https://www.dhs.gov/xlibrary/assets/f oia/mgmt-directive-026-04-protection-of-human-subjects.pdf | |
| | Department of Housing and Urban Development: www.hud.gov/ | | 24 CFR 60.101, which cites 45 CFR part 46, subpart A. | |
| | 1. Department of Justice Office of Justice Programs: http://ojp.gov/ 2. Bureau of Prisons: www.bop.gov | | 1. 28 CFR 22 Privacy Regulation (1976): http://www.ecfr.gov/cgi-bin/text-idx?c=ecfr&tpl=/ecfrbrowse/Title28/28cfr22_main_02.tpl 2. 42 U.S.C. § 3789g Confidentiality of Information (1984) http://www.gpo.gov/fdsys/pkg/USCODE-2010-title42-chap46-subchapVIII-sec3789g.htm | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|----------------|--|-------------|---|---|
| <i>General</i> | | | 3. 28 CFR 46 (1991), Subpart A: http://www.ecfr.gov/cgi-bin/text-idx?c=ecfr&tpl=/ecfrbrowse/Title28/28cfr46_main_02.tpl | |
| | Department of Labor: https://www.dol.gov/ | | 29 CFR 21 | |
| | Department of Transportation: www.dot.gov/ | | 49 CFR 11, Subpart A | |
| | Department of Veterans Affairs: 1. Office of Research Oversight (ORO): http://www1.va.gov/oro/ 2. Office of Research and Development: www.research.va.gov | | 1. 38 FR 16 (1991), Subpart A 2. 38 CFR 17.85 (1998) | Various: https://www.research.va.gov/resources/policies/human_research.cfm |
| | Environmental Protection Agency, Program in Human Research Ethics: https://www.epa.gov/osa/basic-information-about-human-subjects-research-0 | | 40 CFR 26 1. Subpart A: Basic EPA Policy for Protection of Subjects in Human Research Conducted or Supported by EPA (Common Rule) 2. Subpart B: Prohibition of Research Conducted or Supported by EPA Involving Intentional Exposure of Human Subjects who are Children or Pregnant or Nursing Women (2006) 3. Subpart C: Observational Research: Additional Protections for Pregnant Women and Fetuses Involved as Subjects in Observational Research Conducted or Supported by EPA (2006) 4. Subpart D: Observational Research: Additional Protections for Children Involved as Subjects in Observational Research Conducted or Supported by EPA (2006) 5. Subpart K: Basic Ethical Requirements for Third-Party Human Research for Pesticides Involving Intentional Exposure of Non-pregnant, Non-nursing | 1. Scientific and Ethical Approaches for Observational Exposure Studies (2008): http://www.epa.gov/nerl/sots/SEAOES_doc20080707.pdf 2. EPA Order 1000.17A: Policy and Procedures on Protection of Human Subjects in EPA Conducted or Supported Research (2016) https://www.epa.gov/osa/epa-order-100017-policy-and-procedures-protection-human-research-subjects-epa-conducted-or |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | | | Adults (2013) 6. Subpart L: Prohibition of Third-Party Research Involving Intentional Exposure to a Pesticide of Human Subjects who are Children or Pregnant or Nursing Women (2013) 7. Subpart M: Requirements for Submission of Information on the Ethical Conduct of Completed Human Research (2013) 8. Subpart O: Administrative Actions for Noncompliance (2013) 9. Subpart P: Review of Proposed and Completed Human Research (2013) 10. Subpart Q: Standards for Assessing Whether To Rely on the Results of Human Research in EPA Actions (2013) | |
| | National Aeronautics and Space Administration: www.nasa.gov/ | | 14 CFR 1230, Subpart A | |
| | National Science Foundation: www.nsf.gov/ | | 45 CFR 690, Subpart A | |
| | Social Security Administration: http://www.ssa.gov/ | | 45 CFR 46, Subpart A: http://www.hhs.gov/ohrp/humansubjects/guidance/45cfr46.html | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs and Biologics</i> Food and Drug Administration: http://www.fda.gov/Drugs/default.htm and https://www.fda.gov/BiologicsBloodVaccines/default.htm | 1. Food, Drug, and Cosmetic Act, 21 USC Sections 355 and 371 (2012): http://uscode.house.gov/browse/prelim@title21&edition=prelim 2. Public Health Service Act, 42 USC Section 262 (1998): https://www.fda.gov/RegulatoryInformation/LawsEnforcedbyFDA/ucm149278.htm 3. 21 st Century Cures Act, Section 3024 (2016): https://www.govinfo.gov/content/pkg/PLAW-114publ255/pdf/PLAW-114publ255.pdf | 1. 21 CFR 50 (Informed Consent) 2. 21 CFR 312 (Investigational New Drug Application) 3. 21 CFR 56 (Institutional Review Boards) 4. 21 CFR 314 (Applications for Approval to Market a New Drug) 5. 21 CFR 54 (Financial Disclosure by Clinical Investigators) 6. 21 CFR 320 (Bioavailability and Bioequivalence Requirements) | General: Good Clinical Practice and Human Subject Protections in FDA-Regulated Clinical Trials: http://www.fda.gov/ScienceResearch/SpecialTopics/RunningClinicalTrials/default.htm Drugs: http://www.fda.gov/Drugs/GuidanceComplianceRegulatoryInformation/Guidances/default.htm Biologics: https://www.fda.gov/BiologicsBloodVaccines/GuidanceComplianceRegulatoryInformation/default.htm |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|---|--|---|---|
| <i>Drugs, Biologics, and Devices</i> | <p><i>Devices</i></p> <p>Food and Drug Administration, Center for Devices and Radiological Health: http://www.fda.gov/MedicalDevices/default.htm</p> | <p>1. Food, Drug, and Cosmetic Act, 21 USC Section 360 (2012): https://www.fda.gov/RegulatoryInformation/LawsEnforcedbyFDA/FederalFoodDrugandCosmeticActFDCAct/FDCActChapterVDrugsandDevices/default.htm#Part_A</p> <p>2. 21st Century Cures Act, Section 3024 (2016): https://www.gpo.gov/fdsys/pkg/PLAW-114publ255/pdf/PLAW-114publ255.pdf</p> | <p>1. 21 CFR 50 (Informed Consent)</p> <p>2. 21 CFR 56 (Institutional Review Boards)</p> <p>3. 21 CFR 807, Subpart E</p> <p>4. 21 CFR 812 (Investigational Device Exemptions)</p> <p>5. 21 CFR 814 (Pre-market Approval of Medical Devices)</p> <p>6. 21 CFR 54 (Financial Disclosure by Clinical Investigators)</p> | <p>Good Clinical Practice and Human Subject Protections in FDA-Regulated Clinical Trials: http://www.fda.gov/ScienceResearch/SpecialTopics/RunningClinicalTrials/default.htm</p> <p>Other: http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/default.htm</p> |
| <i>Clinical Trials Registry</i> | <p>Food and Drug Administration: http://www.fda.gov/Drugs/default.htm</p> | <p>1. Food and Drug Administration Modernization Act, Section 113 (1997): https://www.fda.gov/RegulatoryInformation/LawsEnforcedbyFDA/SignificantAmendments-to-the-FDCA/FDAMA/default.htm</p> <p>2. Food and Drug Administration Amendments Act, Section 801 (2007): https://www.fda.gov/RegulatoryInformation/LawsEnforcedbyFDA/SignificantAmendments-to-the-FDCA/FoodandDrugAdministrationAmendmentsActof2007/default.htm</p> | | |
| | <p>National Institutes of Health ClinicalTrials.gov: https://www.clinicaltrials.gov/ct2/home</p> | | <p>1. Clinical Trials Regulation and Results Information Submission, 42 CFR 11 (2016): https://www.federalregister.gov/documents/2016/09/21/2016-22129/clinical-trials-registration-and-results-information-submission</p> <p>2. NIH Policy on the Dissemination of NIH-Funded Clinical Trial Information (2016): https://www.federalregister.gov/documents/2016/09/21/2016-22379/dissemination-of-nih-funded-clinical-trial-information</p> | <p>FAQs on ClinicalTrials.gov: https://www.clinicaltrials.gov/ct2/manage-recs/faq</p> |
| | <p>Department of Veterans Affairs: 1. Office of Research Oversight (ORO): http://www1.va.gov/oro/ 2. Office of Research and</p> | | | <p>FAQ: http://www.research.va.gov/resources/ORD_Admin/clinical_trials/registration-faq.pdf</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|-----------------------------------|---|---|--|---|
| <i>Clinical Trials Registry</i> | Development: www.research.va.gov | | | |
| <i>Research Injury</i> | | | Sections 116(a)(6) and (7) of the Common Rule. | |
| | Department of Defense, Regulatory Affairs: http://www.acq.osd.mil/rd/hptb/programs/regulatory/ | | DoDI 3216.02 (2011): http://www.dtic.mil/whs/directives/colres/pdf/321602p.pdf | |
| | Department of Veterans Affairs: 1. Office of Research Oversight (ORO): www1.va.gov/oro/ 2. Office of Research and Development: www.research.va.gov | 38 CFR 17.85: Treatment of Research-Related Injuries to Human Subjects: https://www.gpo.gov/fdsys/pkg/CFR-2013-title38-vol1/pdf/CFR-2013-title38-vol1-sec17-85.pdf | Handbook 1200.5, Appendix F, Paragraph 2a(11) | |
| <i>Social-Behavioral Research</i> | All Common Rule agencies. | | Predominantly Exempt categories 1, 2, and 3, and Expedited categories 6 and 7. | |
| | National Science Foundation: https://www.nsf.gov/ | | | Frequently Asked Questions and Vignettes: https://www.nsf.gov/bfa/dias/policy/hsfaqs.jsp |
| <i>Privacy/Data Protection</i> | All Common Rule agencies | | Common Rule 111(a)(7) (2018) | |
| | Department of Health and Human Services (DHHS): 1. National Institutes of Health (NIH): http://privacyruleandresearch.nih.gov/ 2. Agency for Healthcare Research and Quality (AHRQ): https://www.ahrq.gov/ 3. Office for Civil Rights (OCR): http://www.hhs.gov/ocr/hipaa/ | 1. Privacy Act, 5 U.S.C. § 552a (1974): http://www.justice.gov/opcl/privacyact1974.htm 2. Health Insurance Portability and Accountability Act (1996): https://www.gpo.gov/fdsys/pkg/PLAW-104publ191/content-detail.html 3. Confidential Information Protection and Statistical Efficiency Act (CIPSEA) (2002): http://www.eia.gov/cipsea/cipsea.pdf 4. Health Information Technology for Economic and Clinical Health (HITECH) Act (2009): https://www.gpo.gov/fdsys/pkg/PLAW-111publ5/pdf/PLAW-111publ5.pdf | 1. HIPAA Privacy Rule, 45 CFR parts 160 and 164, Subparts A and C (2002): http://www.hhs.gov/ocr/privacy/hipaa/administrative/privacyrule/index.html 2. HIPAA Security Rule, 45 CFR parts 160, 162, and 164 (2009): http://www.hhs.gov/ocr/privacy/hipaa/administrative/securityrule/index.html 3. HIPAA Breach Notification Rule, 45 CFR § 164.400-414: http://www.hhs.gov/ocr/privacy/hipaa/administrative/breachnotificationrule/index.html | NIH: 1. NIH Policy on Certificates of Confidentiality (2017): https://humansubjects.nih.gov/coc/index 2. Various: http://privacyruleandresearch.nih.gov/ AHRQ: Confidentiality in AHRQ-Supported Research (2018): https://grants.nih.gov/grants/guide/notice-files/NOT-HS-18-012.html OCR: Various: https://www.hhs.gov/hipaa/for-professionals/special-topics/research/index.html and https://www.hhs.gov/hipaa/for-professionals/faq/research-uses-and-disclosures |
| | Department of Homeland Security: www.dhs.gov/ | Public Law 107-347: https://www.gpo.gov/fdsys/pkg/PLAW-107publ347/pdf/PLAW-107publ347.pdf | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|---|---|---|--|
| <i>Privacy/Data Protection</i> | Social Security Administration: http://www.ssa.gov/ | Privacy Act (1974): http://www.hhs.gov/foia/privacy/index.html | | |
| <i>Human Biological Materials</i> | All Common Rule agencies | | Numerous provisions in the Common Rule (2018) | |
| | Department of Health and Human Services, Office for Human Research Protections (OHRP): http://www.hhs.gov/ohrp/ | | | 1. Issues to Consider in the Research Use of Stored Data or Tissues (1997) 2. Guidance on Research Involving Coded Private Information or Biological Specimens (2008) |
| | Food and Drug Administration: a. Office of In Vitro Diagnostic Device Evaluation and Safety: http://www.fda.gov/MedicalDevices/ProductsandMedicalProcedures/InVitroDiagnostics/default.htm b. Center for Biologics Research and Evaluation (CBER): - Office of Cellular, Tissue and Gene Therapies - Office of Blood Research and Review: http://www.fda.gov/BiologicsBloodVaccines/default.htm | | | 1. Guidance on Informed Consent for In Vitro Diagnostic Device Studies Using Leftover Human Specimens That are Not Individually Identifiable (2006): http://www.fda.gov/MedicalDevices/DeviceRegulationandGuidance/Documents/ucm078384.htm 2. In Vitro Diagnostic (IVD) Device Studies - Frequently Asked Questions (2010) http://www.fda.gov/downloads/MedicalDevices/DeviceRegulationandGuidance/GuidanceDocuments/UCM071230.pdf 3. CBER-Specific: Various: http://www.fda.gov/BiologicsBloodVaccines/GuidanceComplianceRegulatoryInformation/OtherRecommendationsforManufacturers/ucm094338.htm |
| <i>Genetic Research</i> | All Common Rule agencies | | Common Rule 116(c)(9) (2018) | |
| | 1. DHHS Office for Human Research Protections (OHRP): http://www.hhs.gov/ohrp/ 2. DHHS National Institutes of Health, Office of Science Policy, Biosafety, Biosecurity, and Emerging Biotechnology Policy Division: https://osp.od.nih.gov/biosafety-biosecurity-and-emerging-biotechnology/ | 1. Research on Transplantation of Fetal Tissue, Public Law 103-43 (1993): http://www.hhs.gov/ohrp/regulations-and-policy/guidance/public-law-103-43/index.html 2. Genetic Information Nondiscrimination Act (2008): https://www.gpo.gov/fdsys/pkg/PLAW-110publ233/content-detail.html | | OHRP: Guidance on the Genetic Information Nondiscrimination Act: Implications for Investigators and Institutional Review Boards (2009): http://www.hhs.gov/ohrp/regulations-and-policy/guidance/guidance-on-genetic-information-nondiscrimination-act/index.html NIH: NIH Guidelines for Research Involving Recombinant or Synthetic Nucleic Acid Molecules (2016): https://osp.od.nih.gov/wp-content/uploads/2013/06/NIH_Guidelines.pdf |
| <i>Embryos, Stem Cells, and Cloning</i> | Food and Drug Administration, Center for Biologics Evaluation and | | | Application of Current Statutory Authorities to Human Somatic Cell |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | Research: http://www.fda.gov/BiologicsBloodVaccines/default.htm | | | Therapy Products and Gene Therapy Products. October 14, 1993. 58 FR 53248: http://www.fda.gov/downloads/BiologicsBloodVaccines/SafetyAvailability/UCM148113.pdf |
| | National Academy of Sciences (NAS): http://www.nationalacademies.org/nrc/ | | | 1. Guidelines for Human Embryonic Stem Cell Research (2005): http://www.nap.edu/catalog.php?record_id=11278 2. 2008 Amendments to the National Academies' Guidelines for Human Embryonic Stem Cell Research: http://books.nap.edu/catalog.php?record_id=12260 3. 2010 Final Report of the National Academies Human Embryonic Stem Cell Research Advisory Committee and 2010 Amendments to the National Academies Guidelines for Human Embryonic Stem Cell Research: http://www.nap.edu/catalog.php?record_id=12923 |
| | National Institutes of Health: http://stemcells.nih.gov/ | Research on Transplantation of Fetal Tissue. Public Law 103-43 (1993): https://history.nih.gov/research/downloads/PL103-43.pdf | | 1. Removing Barriers to Responsible Scientific Research Involving Human Stem Cells, Executive Order 13505 (2009): https://www.gpo.gov/fdsys/pkg/DCPD-200900136/pdf/DCPD-200900136.pdf 2. NIH Guidelines on Human Stem Cell Research (2009): http://stemcells.nih.gov/policy/2009-guidelines.htm 3. NIH Human Embryonic Stem Cell Registry (2016): https://grants.nih.gov/stem_cells/registry/current.htm Access: http://stemcells.nih.gov/ |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|---|--|-------------|--|
| EUROPE | | | | |
| Regionwide | | | | |
| <i>General</i> | European Commission: 1. European Group on Ethics in Science and New Technologies (EGE): https://ec.europa.eu/research/ege/index.cfm 2. Directorate-General for Research and Innovation: http://ec.europa.eu/research/participants/docs/h2020-funding-guide/cross-cutting-issues/ethics_en.htm | | | EGE: 1. Ethical Aspects of Clinical Research in Developing Countries (2003): http://ec.europa.eu/bepa/european-group-ethics/docs/avis17_en.pdf 2. Horizon 2020: How to Complete your Ethics Self –Assessment (2015): http://ec.europa.eu/research/participants/portal/doc/call/h2020/h2020-msca-if-2015/1645175-h2020_-_guidance_ethics_self_assess_en.pdf |
| | Council of Europe, Bioethics Unit: http://www.coe.int/bioethics | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 15-18, ETS No. 164 (1997): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=164&CM=7&DF=9/15/2008&CL=ENG 2. Additional Protocol to the Convention on Human Rights and Biomedicine concerning Biomedical Research, CETS No. 195 (2005): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=195&CM=1&DF=10/24/2007&CL=ENG | | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> European Commission: DG SANTE: Directorate-General for Health and Food Safety: http://ec.europa.eu/health/index_en.htm | 1. Directive 2001/20/EC on the Approximation of the Laws, Regulations and Administrative Provisions of the Member States Relating to the Implementation of Good Clinical Practice in the Conduct of Clinical Trials on Medicinal Products for Human Use: http://ec.europa.eu/health/files/eudralex/vol-1/dir_2001_20/dir_2001_20_en.pdf 2. Directive 2005/28/EC Laying | | EudraLex Volume 10: Clinical Trials: http://ec.europa.eu/health/documents/eudralex/vol-10/ |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|--|---|-------------|---|
| <i>Drugs, Biologics, and Devices</i> | | <p>Down Principles and Detailed Guidelines for Good Clinical Practice as Regards Investigational Medicinal Products for Human Use, as Well as the Requirements for Authorization of the Manufacturing or Importation of Such Products: http://ec.europa.eu/health/files/eudralex/vol-1/dir_2005_28/dir_2005_28_en.pdf</p> <p>3. Regulation No. 536/2014 of the European Parliament and of the Council on Clinical Trials on Medicinal Products for Human Use, Repealing Directive 2001/20/EC: http://ec.europa.eu/health/files/eudralex/vol-1/reg_2014_536/reg_2014_536_en.pdf</p> <p>4. Commission Implementing Regulation (EU) 2017/556 of 24 March 2017 on the Detailed Arrangements for the Good Clinical Practice Inspection Procedures Pursuant to Regulation (EU) No 536/2014 of the European Parliament and Council: https://eur-lex.europa.eu/legal-content/EN/TXT/PDF/?uri=CELEX:32017R0556&from=EN</p> | | |
| | <p>European Medicines Agency: http://www.ema.europa.eu/</p> | | | <p>1. Note for Guidance on Good Clinical Practice (CPMP/ICH/135/95) (1997): https://ec.europa.eu/health/sites/health/files/files/eudralex/vol-10/3cc1aen_en.pdf</p> <p>2. Reflection Paper on Ethical and GCP Aspects of Clinical Trials of Medicinal Products for Human Use Conducted Outside of the EU/EEA and Submitted in Marketing Authorization Applications to the EU Regulatory Authorities (2012): http://www.ema.europa.eu/docs/en_GB/docum</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | | ent library/Regulatory and procedural guideline/2012/04/WC500125437.pdf 3. Guideline for Good Clinical Practice E6(R2) (2016): http://www.ema.europa.eu/docs/en_GB/document_library/Scientific_guideline/2009/09/WC50002874.pdf |
| | <i>Devices</i> DG GROWTH: Internal Market, Industry, Entrepreneurship, SMEs: https://ec.europa.eu/growth/sectors/medical-devices_en | 1. Directive 93/42/EEC Concerning Medical Devices: http://eur-lex.europa.eu/LexUriServ/LexUriServ.do?uri=CONSLEG:1993L0042:20071011:en:PDF 2. Directive 98/79/EC on in vitro Diagnostic Medical Devices (IVD): https://ec.europa.eu/growth/single-market/european-standards/harmonised-standards/iv-diagnostic-medical-devices_en 3. Directive 2007/47/EC of the European Parliament and of the Council of 5 September 2007 Amending Council Directive 90/385/EEC on Approximation of the Laws of the Member States Relating to Active Implantable Medical Devices: http://eur-lex.europa.eu/legal-content/EN/TXT/PDF/?uri=CELEX:32007L0047&from=EN | | Various: http://ec.europa.eu/consumers/sectors/medical-devices/documents/guidelines/index_en.htm |
| <i>Clinical Trials Registry</i> | EU Clinical Trials Register: https://www.clinicaltrialsregister.eu/ | | | FAQs: https://www.clinicaltrialsregister.eu/doc/EU_CTR_FAQ.pdf |
| <i>Research Injury</i> | European Commission: DG SANTE: Directorate-General for Health and Food Safety: http://ec.europa.eu/health/index_en.htm | 1. Clinical Trials Directive 2001/20/EC: http://ec.europa.eu/health/human-use/clinical-trials/directive/index_en.htm 2. Regulation No. 536/2014 of the European Parliament and of the Council on Clinical Trials on Medicinal Products for Human Use, Repealing Directive 2001/20/EC: | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------|---|--|-------------|---|
| <i>Research Injury</i> | | http://ec.europa.eu/health/files/eudralex/vol-1/reg_2014_536/reg_2014_536_en.pdf | | |
| | Council of Europe, Bioethics Unit: http://www.coe.int/bioethics | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Article 24, ETS No. 164 (1997): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=164&CM=7&DF=9/15/2008&CL=ENG 2. Additional Protocol to the Convention on Human Rights and Biomedicine concerning Biomedical Research, Article 13, CETS No. 195 (2005): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=195&CM=1&DF=10/24/2007&CL=ENG | | |
| <i>Privacy/Data Protection</i> | European Data Protection Board (EDPB): https://edpb.europa.eu/ | Regulation (EU) 2016/679 of the European Parliament and of the Council of 27 April 2016 on the Protection of Natural Persons with Regard to the Processing of Personal Data and on the Free Movement of Such Data, and Repealing Directive 95/46/EC (General Data Protection Regulation): http://eur-lex.europa.eu/legal-content/EN/TXT/PDF/?uri=CELEX:32016R0679&from=EN | | EDPB: 1. Transfers of Personal Data to Third Countries: Applying Articles 25 and 26 of the EU Data Protection Directive (1998): Consent (2018): http://ec.europa.eu/justice/article-29/documentation/opinion-recommendation/files/1998/wp12_en.pdf 2. Adequacy Referential (2018): http://ec.europa.eu/newsroom/article29/document.cfm?action=display&doc_id=49724 |
| | European Medicines Agency (EMA): http://www.ema.europa.eu/ | | | 1. External Guidance on the Implementation of the European Medicines Agency Policy on the Publication of Clinical Data for Medicinal Products for Human Use (2016): http://www.ema.europa.eu/docs/en_GB/document_library/Regulatory_and_procedural_guideline/2016/12/WC500218567.pdf 2. European Medicines Agency policy on publication of clinical data for medicinal products for human use https://www.ema.europa.eu/documents/other/e |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Privacy/Data Protection</i> | <p>Council of Europe: Data Protection and Cybercrime Division: http://www.coe.int/t/dghl/standardsetting/dataprotection/default_EN.asp</p> | <p>Convention for the Protection of Individuals with Regard to Automatic Processing of Personal Data (1981): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=108&CL=ENG</p> | | <p>uropean-medicines-agency-policy-publication-clinical-data-medicinal-products-human-use_en.pdf 3. Questions and Answers on the European Medicines Agency Policy on Publication of Clinical Data for Medicinal Products for Human Use (2015): http://www.ema.europa.eu/docs/en_GB/document_library/Report/2014/10/WC500174378.pdf 4. External Guidance on the Implementation of the European Medicines Agency Policy on the Publication of Clinical Data for Medicinal Products for Human Use (2016): https://www.ema.europa.eu/documents/regulatory-procedural-guideline/external-guidance-implementation-european-medicines-agency-policy-publication-clinical-data_en-1.pdf</p> <p>1. Recommendation No. R (97) 5 on the Protection of Medical Data (1997): https://wcd.coe.int/ViewDoc.jsp?id=571075&Site=CM&BackColorInternet=C3C3C3&BackColorIntranet=EDB021&BackColorLogged=F5D383 2. Article 29 Working Party Documentation: http://ec.europa.eu/justice/data-protection/article-29/index_en.htm</p> |
| <i>Human Biological Samples</i> | <p>European Commission: European Group on Ethics in Science and New Technologies: http://ec.europa.eu/research/ege/index.cfm</p> <p>Council of Europe, Bioethics Unit: http://www.coe.int/bioethics</p> | <p>Directive 2004/23/EC on Setting Standards of Quality and Safety for the Donation, Procurement, Testing, Processing, Preservation, Storage, and Distribution of Human Tissues and Cells: http://eur-lex.europa.eu/LexUriServ/LexUriServ.do?uri=CELEX:32004L0023:EN:HTML</p> <p>Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 21-22, ETS No. 164 (1997): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=164&CM=7&DF=9/15/2008&CL=ENG</p> | | <p>Recommendation Rec (2016) 6 of the Committee of Ministers to Member States on Research on Biological Materials of Human Origin: https://search.coe.int/cm/Pages/result_details.aspx?ObjectId=090000168064e8ff</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | European Medicines Agency: http://www.ema.europa.eu/ | Regulation (EC) No. 1394/2007 on Advanced Therapy Medicinal Products and Amending Directive 2001/83/EC and Regulation (EC) No. 726/2004: http://ec.europa.eu/health/files/eudra_lex/vol-1/reg_2007_1394/reg_2007_1394_en.pdf | | |
| | Council of Europe, Bioethics Unit: http://www.coe.int/bioethics | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 12-14, ETS No. 164 (1997): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=164&CM=7&DF=9/15/2008&CL=ENG 2. Additional Protocol to the Convention on Human Rights and Biomedicine Concerning Biomedical Research, CETS No. 195 (2005): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=195&CM=1&DF=10/24/2007&CL=ENG | | 1. Recommendation No. R (92) on Genetic Testing and Screening for Health Care Purposes (1992): http://wcd.coe.int/ViewDoc.jsp?id=612007&Site=CM&BackColorInternet=9999CC&BackColorIntranet=FFBB55&BackColorLogged=FFAC75 2. Recommendation Rec (2006)4 of the Committee of Ministers to Member States on Research on Biological Materials of Human Origin (2016): https://search.coe.int/cm/Pages/result_details.aspx?ObjectId=090000168064e8ff 3. Recommendation Rec(2016)8 of the Committee of Ministers to Member States on the Processing of Personal Health-Related Data for Insurance Purposes, Including Data Resulting from Genetic Tests (2016): https://search.coe.int/cm/Pages/result_details.aspx?ObjectId=09000016806b2c5f |
| <i>Embryos, Stem Cells, and Cloning</i> | European Commission: European Group on Ethics in Science and New Technologies: http://ec.europa.eu/research/ege/index.cfm | 1. Statements by the Commission Re: Article 6 (2006): http://www.uv.es/operuv/docs_7pm/FP7ECStatementsComm_Ethical.pdf 2. Statement of the Commission Related to Research Activities Involving Human Embryonic Stem Cells (2013): http://eur-lex.europa.eu/LexUriServ/LexUriServ.do?uri=OJ:C:2013:373:0012:0015:EN:PDF | | 1. Commission Staff Working Paper Report on Human Embryonic Stem Cell Research (2003): https://ec.europa.eu/research/press/2003/pdf/sec2003-441report_en.pdf 2. Opinion No. 22 - The Ethics Review of hESC FP7 Research Projects (2007): http://bookshop.europa.eu/ga/recommendations-on-the-ethical-review-of-hesc-fp7-research-projects-pbKAAJ07022/downloads/KA-AJ-07-022-EN-C/KAAJ07022ENC_002.pdf;pgid=y8dIS7GUWMdSR0EAlMEUUsWb0000dz-kvfzb;sid=lexx3tq0IOFxyokBvtfvebiRJj93DZfXP54=?FileName=KAAJ07022ENC_002.pdf&SKU=KAAJ07022ENC_PDF&CatalogueNumber=KA-AJ-07-022-EN-C |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | Council of Europe, Bioethics Unit: http://www.coe.int/bioethics | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 18, ETS No. 164 (1997): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=164&CM=7&DF=9/15/2008&CL=ENG 2. Additional Protocol on Prohibition of Human Cloning, ETS No. 168 (1998): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=168&CM=7&DF=9/15/2008&CL=ENG | | Statement on Genome Editing Technologies by the Committee on Bioethics (2015): https://rm.coe.int/168049034a |
| Armenia | | | | |
| For an overview of human subject protections in Armenia, see “Ethical Review of Biomedical Research in the CIS Countries,” Chapter 3, Section 1: http://www.unesco.org/new/fileadmin/MULTIMEDIA/FIELD/Moscow/pdf/ethical_review_cis_book_kubar_english.pdf | | | | |
| Note: All websites and documents are in Armenian. | | | | |
| <i>Drugs, Biologics, and Devices</i> | 1. Drug and Medical Technology Agency: http://www.pharm.am/ 2. Ethics Committee of the Ministry of Health | 1. Law of the Republic of Armenia of May 4, 1996: About Medical Aid, The Maintenance of the Population, Article 21: http://www.arlis.am/DocumentView.aspx?DocID=71619 2. Resolution of the Government of Armenia of January 24, 2002: Procedure for Clinical Trials of New Medications in Armenia: http://www.arlis.am/DocumentView.aspx?docID=9154 | | |
| <i>Human Biological Materials</i> | Ethical Committee of the National Center for AIDS Prevention: http://www.armajids.am/main/index.php?lng=1 | RA Law on Prevention of Disease Caused by HIV (2012): http://www.arlis.am/DocumentView.aspx?DocID=78616 | | Declaration of Helsinki: Ethical Principles for Medical Research Involving Human Subjects (2013) |
| Austria | | | | |
| <i>General</i> | 1. Ministry of Health: http://www.bmg.gv.at 2. Forum of Austrian Ethics Committees: http://www.ethikkommissionen.at 3. Bioethics Commission: http://www.bundeskanzleramt.at/site/3575/default.aspx | 1. University Act (2011): http://www.ris.bka.gv.at/Dokumente/ErV/ERV_2002_1_120/ERV_2002_1_120.pdf 2. Hospitals Act (2014): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=10010285&ShowPrintPreview=True | Regulation on Leading Ethics Committees (2004): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=20003352&ShowPrintPreview=True | Bioethics Commission: 1. Codification of Legislation on Medical Research (2011) 2. Research on Persons without the Capacity to Consent (2013) Access: http://www.bundeskanzleramt.at/site/4070/default.aspx |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. Ministry of Health: http://www.bmg.gv.at 2. Austrian Agency for Health and Food Safety: http://www.ages.at/ages/en/ages-austrian-agency-for-health-and-food-safety/ 3. Austrian Federal Office for Safety in Health Care: http://www.basg.gv.at/en/basg-austrian-federal-office-for-safety-in-health-care/ | Austrian Drug Law (2013): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=10010441&ShowPrintPreview=True | | Various: http://www.basg.at/arsneimittel/vor-der-zulassung/klinische-pruefungen/ |
| | <i>Devices</i> 1. Ministry of Health: http://www.bmg.gv.at 2. Austrian Agency for Health and Food Safety: http://www.ages.at/ages/en/ages-austrian-agency-for-health-and-food-safety/ 3. Austrian Federal Office for Safety in Health Care: http://www.basg.gv.at/en/basg-austrian-federal-office-for-safety-in-health-care/ | Medical Devices Act (2014): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=10011003 | | Various: http://www.basg.at/medizinprodukte/formulare/klinische-pruefung/ |
| <i>Research Injury</i> | 1. Austrian Agency for Health and Food Safety: http://www.ages.at/ages/en/ages-austrian-agency-for-health-and-food-safety/ 2. Austrian Federal Office for Safety in Health Care: http://www.basg.at/en/austrian-federal-office-for-safety-in-health-care/ | 1. Austrian Drug Law, Article 32 (2013): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=10010441&ShowPrintPreview=True 2. Austrian Medical Devices Law, Article 47 (2017): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=10011003&ShowPrintPreview=True | | |
| <i>Privacy/Data Protection</i> Note: The Austrian states also have privacy/data protection laws | Austrian Data Protection Authority: https://www.dsb.gv.at/DesktopDefault.aspx?alias=dsken | 1. Data Protection Act No. 165/1999: https://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=10001597 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | | |
| <i>Human Biological Materials</i> | 1. Ministry of Health: http://www.bmg.gv.at 2. Bioethics Commission: http://www.bundeskanzleramt.at/site/3575 | 1. Law on Safety of Blood (2009): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen | Regulation on Tissue Banks (2014): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen& | Bioethics Commission: 1. Opinion of the Bioethics Commission at the Federal Chancellery: Biobanks for Medical Research (2007) |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | /default.aspx | &Gesetzesnummer=10011145&ShowPrintPreview=True 2. Law on Quality and Safety of Human Tissue and Cells (2013): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=20005698&ShowPrintPreview=True | Gesetzesnummer=20005848&ShowPrintPreview=True | 2. Biobanks for Medical Research - Amendments to the Bioethics Commission Report of May 2007 (2011) <i>Access:</i> http://www.bundeskanzleramt.at/site/4070/default.aspx |
| <i>Genetic Research</i> | 1. Ministry of Health: http://www.bmg.gv.at 2. Bioethics Commission: http://www.bundeskanzleramt.at/site/3575/default.aspx | Gene Technology Act (2012): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=10010826&ShowPrintPreview=True | | |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Ministry of Health: http://www.bmg.gv.at 2. Bioethics Commission: http://www.bundeskanzleramt.at/site/3575/default.aspx | Reproductive Medicine Act (2010): http://www.ris.bka.gv.at/GeltendeFassung.wxe?Abfrage=Bundesnormen&Gesetzesnummer=10003046&ShowPrintPreview=True | | Bioethics Commission: 1. Stem Cell Research in the Context of the EU's Sixth Framework Programme Research (2002) 2. Research on Human Embryonic Stem Cells (2009): http://www.bundeskanzleramt.at/DocView.axd?CobId=34240 |
| Belarus | | | | |
| For an overview of human subject protections in Belarus, see "Ethical Review of Biomedical Research in the CIS Countries," Chapter 3, Section 3: http://www.unesco.org/new/fileadmin/MULTIMEDIA/FIELD/Moscow/pdf/ethical_review_cis_book_kubar_english.pdf | | | | |
| <i>General</i> | 1. Ministry of Health (MOH): http://minzdrav.by/en/ 2. National Bioethics Committee | 1. Constitution of the Republic of Belarus, Article 25 (2004): http://www.pravo.by/WEBNPA/text.asp?RN=v19402875 2. Law on Health Care System, Articles 40, 46 (2010): http://pravo.by/webnpa/text.asp?RN=v19302435 | MOH: 1. Ordinance No. 274 on Establishing the National Bioethics Committee (2006) 2. Decree No. No. 55 on Ethics Committees (2008) (Russian): http://www.levonevski.net/pravo/norm2009/num05/d05639.html | MOH: 1. Code of Medical Ethics (1999): http://www.levonevski.net/pravo/norm2009/num37/d37726.html 2. Guidelines for Ethics Committees on Standard Operational Proceedings (No. 55-0004, 2000): http://www.levonevski.net/pravo/norm2009/num35/d35896/index.html 3. Methodological Guidelines of Health Ministry (2000) |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. Ministry of Health (MOH): http://minzdrav.by/en/ 2. State Pharmacological Committee 3. Centre for Expertise and Testing in Health Care: http://rceeth.by/ | 1. Law on Drugs, Articles 15,16 (2009): http://pravo.by/webnpa/text.asp?RN=h10600161 2. Law on Health Care System, Article 40 (2010): http://pravo.by/webnpa/text.asp?RN=v19302435 | MOH: 1. Ordinance No. 254 on Clinical Drug Trials and Good Clinical Practice (1999): http://www.levonevski.net/pravo/norm2009/num36/d36922/index.html 2. Ordinance No. 161 on Accreditation of Health Care Institutions and Attestation of | MOH: Instruction on Accreditation of Health Care Institutions and Attestation of Specialists Involved in Conducting Clinical Trials of Drugs and Medical Devices (No. 55-0504, 2004) (Russian): http://www.levonevski.net/pravo/norm2009/num24/d24926.html |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | Specialists Involved in Conducting Clinical Trials of Drugs and Medical devices (1999): http://www.levonevski.net/pravo/norm2009/num37/d37336.html 3. Decree No. 55 on Ethics Committees (2008): http://www.levonevski.net/pravo/norm2009/num05/d05639.html 4. Decree No. 50 on Certain Aspects of Clinical Drug Trials (2009) | |
| | <i>Devices</i> | | | |
| | 1. Ministry of Health (MOH): http://minzdrav.by/en/ 2. Centre for Expertise and Testing in Health Care: http://rceth.by/ | Law on Health Care System, Article 40 (2010): http://pravo.by/webnpa/text.asp?RN=v19302435 | MOH: 1. Ordinance No. 161 on Accreditation of Health Care Institutions and Attestation of Specialists Involved in Conducting Clinical Trials of Drugs and Medical devices (1999): http://www.levonevski.net/pravo/norm2009/num37/d37336.html 2. Decree No. 216 on Certain Aspects of Clinical Trials of Medical Devices (2008) (Russian): http://86.57.250.247/data/pravo/ipb_prikazmz/N216_2008.htm | MOH: Instruction on Accreditation of Health Care Institutions and Attestation of Specialists Involved in Conducting Clinical Trials of Drugs and Medical Devices (No. 55-0504, 2004): http://www.levonevski.net/pravo/norm2009/num24/d24926.html |
| <i>Privacy/Data Protection</i> | 1. Ministry of Health: http://minzdrav.by/en/ 2. National Bioethics Committee | 1. Constitution of the Republic of Belarus, Article 28 (2004): http://www.pravo.by/WEBNPA/text.asp?RN=v19402875 2. Law on Health Care System, Article 46 (2010): http://pravo.by/webnpa/text.asp?RN=v19302435 | | |
| <i>Human Biological Materials</i> | 1. Ministry of Health (MOH): http://minzdrav.by/en/ 2. National Bioethics Committee 3. State Service of Forensic Medicine (SSFM) | Law on Health Care System, Articles 40 and 46 (2010): http://pravo.by/webnpa/text.asp?RN=v19302435 | MOH: Ordinance No. 111 on Further Development of National Pathology Service (1993): http://86.57.250.247/data/pravo/ipb_prikaznew/N111_1993(1994).doc | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | | | SSFM: Ordinance No. 38-c on Rules for Conducting Morphological Examinations (1999) | |
| Belgium | | | | |
| <i>General</i> | Belgium Advisory Committee on Bioethics (BACB): https://www.health.belgium.be/en/belgian-advisory-committee-bioethics | Law Relating to Experimentation on Humans (2004): http://www.erasme.ulb.ac.be/page.asp?id=11365&langue=EN | | BACB: 1. Opinion No. 13: Regarding Experimentation on Man (2001) 2. Opinion No. 31: Regarding Experimentation Involving Pregnant and Breastfeeding Women (2004) 3. Opinion 36: Ethical Evaluation of Research in Certain Areas of the Humanities (French and Dutch) (2006) 4. Opinion No. 40: Scope of the Law Relating to Experimentation on Humans (French and Dutch) (2007) 5. Opinion No. 51: Publication of the Results of Human Experimentation (2012) 6. Opinion No. 69: Experiments and Other Scientific Research Involving Inmates (2017) <i>Access:</i> http://www.health.belgium.be/en/belgian-advisory-committee-bioethics |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | Medicines Directorate-General: http://www.health.belgium.be/eportal | | <ol style="list-style-type: none"> 1. Royal Decree of September 27, 1994 2. Royal Decree of June 30, 2004 Determining the Implementation Measures of the Law 3. Royal Decree of June 30, 2004 Modifying the Royal Decree of June 6, 1960 4. Royal Decree of July 15, 2004 Determining Payments for Ethical Opinions or Authorization for the Conduct of a Clinical Trial or Experiment 5. Application of the Law of May 7, 2004 Relating to Experiments on Human Volunteers who Participate in Phase I Trials (2004) 6. Explanations Concerning the Submission of a Request for an Ethical Opinion or Authorization for the Conduct of a Clinical Trial (2004) | |
| <i>Research Injury</i> | | Law Relating to Experimentation on Humans, Chapter XVII (Responsibility and Insurance) Article 29 (2004) | | |
| <i>Privacy/Data Protection</i> | Commission for the Protection of Privacy: http://www.privacycommission.be/ | <ol style="list-style-type: none"> 1. Privacy Act (1992): http://www.privacycommission.be/en/privacy-act 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | Decree of February 13, 2001 Implementing the Law of December 8, 1999: http://www.privacycommission.be/sites/privacycommission/files/documents/Royal_Decree_2001.pdf | <ol style="list-style-type: none"> 1. Legal Basis (2018): https://www.autoriteprotectiondonnees.be/fondement-legal-pour-le-traitement-de-donnees-a-caractere-personnel 2. Consent (2018): https://www.autoriteprotectiondonnees.be/consentement 3. International Data Transfer (2018): https://www.autoriteprotectiondonnees.be/international-0 |
| <i>Human Biological Materials</i> | <ol style="list-style-type: none"> 1. Belgian Advisory Committee on Bioethics: http://www.health.belgium.be/en 2. Superior Health Council (CSS): http://www.health.belgium.be/eportal/Aboutus/relatedinstitutions/SuperiorHealthCouncil/index.htm | <ol style="list-style-type: none"> 1. Royal Decree (1987) Regarding the Expression of Consent for the Removal of Organs and Tissues on Living Donors 2. Royal Decree (1997) Regarding the Removal and | | <p>Belgian Advisory Committee on Bioethics:</p> <ol style="list-style-type: none"> 1. Opinion No. 52: Use of Human Tissues and Cells in Reproductive Medicine (2012) 2. Opinion No. 54: Post Mortem Removal of Human Body Material for Human |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | 3. Federal Public Service: www.health.fgov.be | Allocation of Organs of Human Origin 3. Act on the Removal and Transplantation of Organs (2006) 4. 2007 Amendment | | Medical Applications or for Scientific Research Purposes (2012) <i>Access:</i> http://www.health.belgium.be/en/belgian-advisory-committee-bioethics <i>CSS:</i> Various: http://www.health.belgium.be/eportal/Aboutus/relatedinstitutions/SuperiorHealthCouncil/domains/cellstissuesorgans/index.htm#.Viovr88XQOU |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Belgian Advisory Committee on Bioethics: https://www.health.belgium.be/en/belgian-advisory-committee-bioethics 2. Federal Commission for Medical and Scientific Research on Embryos in Vitro: http://health.belgium.be/eportal/Healthcare/Consultativebodies/Commissions/Embryoinvitro/19076630?ie2Term=research&ie2section=83 | 1. Royal Decree Fixing the Criteria for the Program Applicable to the Care Programs ‘Reproductive Medicine’ (15/02/1999) 2. Act on Research on Embryos in Vitro (2003): http://www.eshre.com/ESHRE/English/Legal-Matters-and-Guidelines/Legal-documentation/Belgium/page.aspx/164 3. Law on Medically Assisted Reproduction and the Destination of Supernumerary Embryos and Gametes (2007): http://www.staatsbladclip.be/lois/2007/07/17/loi-2007023090.html | | Belgian Advisory Committee on Bioethics: 1. Opinion No. 33: Somatic and Germinal Line Gene Modification (2005) 2. Opinion No. 52: Use of Human Tissues and Cells in Reproductive Medicine (2012) <i>Access:</i> http://www.health.belgium.be/en/belgian-advisory-committee-bioethics |
| Bosnia and Herzegovina | | | | |
| Note: All websites and documents are in Bosnian. | | | | |
| <i>General</i> | | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 15-18, ETS No. 164 (2007): 2. Additional Protocol Concerning Biomedical Research, CETS No. 195 (2007) 3. Law on Health Protection, MoH Republic of Srpska (2015): http://www.vladars.net/sr-SP-Cyrl/Vlada/Ministarstva/MZSZ/Doc | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | | <p>uments/Zakon%20o%20zdravstvenoj%20zastiti%20sa%20izmjenama%20106-99%20%2044-15.pdf</p> <p>4. Law on Health Protection, MoH Federation of Bosnia and Herzegovina, No 46/10: http://www.fmoh.gov.ba/index.php/zakoni-i-strategije/zakoni/zakon-o-zdravstvenoj-zastiti</p> | | |
| <i>Drugs, Biologics, and Devices</i> | <p><i>Federation of Bosnia and Herzegovina</i></p> <p>1. Ministry of Health: http://www.fmoh.gov.ba/</p> <p>2. Medicines and Medical Devices Agency of Bosnia and Herzegovina: http://www.almbih.gov.ba/</p> | <p>1. Law on Drugs No. 58/08: http://www.almbih.gov.ba/doc/regulative/medicinal_products_and_medical_devices_act.pdf</p> <p>2. Law on Changes and Amendments of the Law on Drugs No. 29/05: http://www.almbih.gov.ba/doc/regulative/fbih/Zakon_o_lijekovima-sluzbene_novine_FBiH_broj_29-05.pdf</p> <p>3. Law on Drugs Federation of Bosnia and Herzegovina, No 109/2012: http://www.fmoh.gov.ba/index.php/zakoni-i-strategije/zakoni/zakon-o-lijekovima-fbih</p> | <p>1. Regulation about Clinical testing of IMP and Medical Devices (2010): http://www.almbih.gov.ba/doc/regulative/pravilnik_klinicka_bos.pdf</p> <p>2. Regulation about Medical Devices (2010): http://www.almbih.gov.ba/doc/regulative/pravilnik_ms_bos.pdf</p> <p>3. Standards of GCP in Conducting CTs (2012): http://www.almbih.gov.ba/doc/regulative/Smjernice_dobre_klinicke_prakse-bo.pdf</p> <p>4. Instructions on Manner of Reporting on Safety in the Framework of Clinical Trials (2016): http://www.almbih.gov.ba/doc/upustva-vodici/uputstvo_o_nacinu_izvjestavanja_o_sigurnosti.pdf</p> | |
| | <p><i>Republic of Srpska</i></p> <p>1. Ministry of Health and Social Welfare (Bosnian): http://www.vladars.net/sr-SP-Cyrl/Vlada/Ministarstva/MZSZ/OMin/Pages/Splash.aspx</p> <p>2. Medicines and Medical Devices Agency of Bosnia and Herzegovina: http://www.almbih.gov.ba/</p> | <p>1. Law on Drugs No. 58/08: http://www.almbih.gov.ba/doc/regulative/medicinal_products_and_medical_devices_act.pdf</p> <p>2. Law on Changes and Amendments of Law on Drugs No. 34/08: http://www.almbih.gov.ba/doc/regulative/rs/ID_Zakona_o_lijekovima_34_08.pdf</p> | <p>1. Regulation about Clinical testing of IMP and Medical Devices (2010): http://www.almbih.gov.ba/doc/regulative/pravilnik_klinicka_bos.pdf</p> <p>2. Regulation about Medical Devices (2010): http://www.almbih.gov.ba/doc/regulative/pravilnik_ms_bos.pdf</p> <p>3. Standards of GCP in Conducting CTs (2012): http://www.almbih.gov.ba/doc/regulative/Smjernice_dobre_klinicke_prakse-bo.pdf</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | se-bo.pdf 4. Instructions on Manner of Reporting on Safety in the Framework of Clinical Trials (2016): http://www.almbih.gov.ba/doc/upustva-vodici/uputstvo_o_nacinu_izvjestavnja_o_sigurnosti.pdf | |
| <i>Research Injury</i> | <i>Federation of Bosnia and Herzegovina</i> | | | |
| | Medicines and Medical Devices Agency of Bosnia and Herzegovina: http://www.almbih.gov.ba/ | 1. Medicinal Products and Medicinal Devices Act, Articles 52 and 116 (2008): http://www.almbih.gov.ba/doc/regulative/medicinal_products_and_medical_devices_act.pdf 2. Law on Health Insurance of the Federation of Bosnia and Herzegovina, Official Gazette No. 46/10 | Regulation about Clinical Testing of IMP and Medical Devices, 4/10: http://www.almbih.gov.ba/doc/regulative/pravilnik_klinicka_bos.pdf | |
| <i>Research Injury</i> | <i>Republic of Srpska</i> | | | |
| | Ministry of Health and Social Welfare (Bosnian): http://www.vladars.net/sr-SP-Cyril/Vlada/Ministarstva/MZSZ/OMin/Pages/Splash.aspx | 1. Medicinal Products and Medicinal Devices Act, Article 52 and 116 2. Law on Health Insurance of the Republic of Srpska, Official Gazette Republic of Srpska No. 106/09: http://www.farmaceutska-komora.org/images/stories/5Zakon_o_zdravstvenoj_zastiti.pdf | Regulation about Clinical Testing of IMP and Medical Devices, 4/10: http://www.almbih.gov.ba/doc/regulative/pravilnik_klinicka_bos.pdf | |
| <i>Privacy/Data Protection</i> | Personal Data Protection Agency of Bosnia and Herzegovina: http://www.azlp.gov.ba/Default.aspx?langTag=en-US&template_id=147&pageIndex=1 | 1. Law on the Protection of Personal Data in Bosnia and Herzegovina (2005): http://www.azlp.gov.ba/propisi/Default.aspx?id=5&langTag=en-US&pageIndex=1 2. Law on Amendments to the Law on the Protection of Personal Data in Bosnia and Herzegovina, Official Gazette of Bosnia and Herzegovina No. 76/11 (2011): http://www.azlp.gov.ba/Default.aspx?langTag=en-US&template_id=147&pageIndex=1 | Regulation on the Manner of Keeping the Records of Personal Data Filing Systems and the Pertinent Records Form (2009): http://www.azlp.gov.ba/propisi/default.aspx?id=1321&langTag=bs-BA | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--|---|--|-------------|--|
| <i>Embryos, Stem Cells and Cloning</i> | <p><i>Federation of Bosnia and Herzegovina</i></p> <p>Ministry of Health: http://www.fmoh.gov.ba/</p> | <p>1. Law on Transplantation of Organs and Tissues, Official Gazette of Bosnia and Herzegovina No. 75/09: http://www.fmoh.gov.ba/index.php/zakoni-i-strategije/zakoni/zakon-o-transplantaciji-organa-i-tkiva-u-svrhu-lijecenja</p> <p>2. Law on Blood and Blood Products, Official Gazette of Bosnia and Herzegovina No. 09/10: http://www.fbihvlada.gov.ba/bosanski/zakoni/2010/zakoni/8bos.htm</p> | | |
| | <p><i>Republic of Srpska</i></p> <p>Ministry of Health and Social Welfare (Bosnian): http://www.vladars.net/sr-SP-Cyrl/Vlada/Ministarstva/MZSZ/OMin/Pages/Splash.aspx</p> | <p>1. Law on Transplantation of Organs (2010): http://www.vladars.net/sr-SP-Cyrl/Vlada/Ministarstva/MZSZ/Documents/Zakon%20o%20transplantaciji%20judskih%20organa.pdf</p> <p>2. Law on Transplantation of Human Tissues and Cells (2010): http://www.vladars.net/sr-SP-Cyrl/Vlada/Ministarstva/MZSZ/Documents/Zakon%20o%20transplantaciji%20judskih%20tkiva%20i%20celija.pdf</p> | | <p>Rulebook about Testing Procedure for Donor of Transplant Organs in Terms of Diseases Which can be Transmitted by Transplantation (2010): http://www.vladars.net/sr-SP-Cyrl/Vlada/Ministarstva/MZSZ/Documents/%d0%9f%d1%80%d0%b0%d0%b2%d0%b8%d0%bb%d0%bd%d0%b8%d0%ba %d0%be %d0%ba%d1%80%d0%b8%d1%82%d0%b5%d1%80%d0%b8%d1%98%d1%83%d0%bc%d0%b8%d0%bc%d0%b0 %d0%b7%d0%b0 %d1%82%d0%b5%d1%81%d1%82%d0%b8%d1%80%d0%b0%d1%9a%d0%b5 %d0%b4%d0%b0%d0%b2%d0%b0%d0%bb%d0%b0%d1%86%d0%b0 %d1%99%d1%83%d0%b4%d1%81%d0%ba%d0%b8%d1%85 %d0%be%d1%80%d0%b3%d0%b0%d0%bd%d0%b0_64_10.pdf</p> |
| Bulgaria | <p>Ministry of Healthcare: http://www.mh.government.bg/</p> | <p>1. Constitution of the Republic of Bulgaria, Amendment SG. 18/25, Article 29 (2015): http://www.parliament.bg/bg/const</p> <p>2. Oviedo Convention on Human Rights and Biomedicine (2003)</p> <p>3. Law Ratifying the Additional Protocol on Biomedical Research (2006): https://www.mh.government.bg/me</p> | | |
| <i>General</i> | | | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|---|---|--|---|
| <i>General</i> | | dia/filer_public/c6/12/c612c85a-519f-4a22-b390-5fad298ce80b/zakon-ratifitsirane-protokol-konventsiya-zashhita-pravata-na_choveka_29-08-2006.pdf 4. Medicinal Products in Human Medicine Act (2017): http://www.bda.bg/images/stories/documents/regulations/zakoni/ZLPHM_28122017.pdf 5. Healthcare Act, Articles 197-206 (2018): http://www.mh.government.bg/media/filer_public/2018/02/27/zakon-zazdraveto.pdf | | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | 1. Ministry of Healthcare (MOH): http://www.mh.government.bg/ 2. Bulgarian Drug Agency (BDA): http://www.bda.bg/en/ | Medicinal Products in Human Medicine Act, Chapter 4 (2018): https://www.lex.bg/laws/ldoc/2135549536 | Regulation No. 31 from 12 August 2007 for Determining the Principles of Good Clinical Practice (2012): http://www.bda.bg/images/stories/documents/regulations/naredbi/20180320_Naredda_31.pdf |
| | <i>Devices</i> | Bulgarian Drug Agency (BDA): http://www.bda.bg/en/ | Medical Devices Act (2016): http://www.bda.bg/images/stories/documents/legal_acts/ZMI_en_20160308.pdf | Ordinance No. 10 of 2008 on the Documents Required from the Principal/Coordinating Investigator or Sponsor for Obtaining an Ethics Committee Statement and on the Procedure for Safety Monitoring of Medical Devices During Clinical Investigations and Assessment of the Clinical Data Collected During such Investigations (2010): http://www.bda.bg/images/stories/documents/legal_acts/Ordinance_Clinical_investigations_MD_EN.pdf |
| | | | | Various: http://www.bda.bg/en/114-information-for-companies-section/medical-devices-category |
| <i>Research Injury</i> | Bulgarian Drug Agency (BDA): http://www.bda.bg/en/ | Medicinal Products in Human Medicine Act, Chapter 4, Articles 91 and 92 (2016): http://www.bda.bg/images/stories/documents/legal_acts/ZLPHM_en.pdf | Regulation 31 from 12 August 2007 for Determining the Principles of Good Clinical Practice, Section 5.8 (2012) (Bulgarian): http://www.bda.bg/images/stories/doc | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|--|---|---|--|
| <i>Research Injury</i> | | | uments/regulations/naredbi/20180320_Naredba_31.pdf | |
| <i>Privacy/Data Protection</i> | <p>1. Bulgarian Commission for Personal Data Protection: https://www.cdpd.bg/en/index.php?p=rubric&aid=2</p> <p>2. Ombudsman: www.ombudsman.bg</p> | <p>1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj</p> <p>2. Law for Protection of Personal Data (2018): https://www.cdpd.bg/en/index.php?p=element&aid=373</p> | | <p>1. General (2018): https://www.cdpd.bg/index.php?p=element&aid=1163</p> <p>2. Research (2018): https://www.cdpd.bg/en/index.php?p=element&aid=1162</p> <p>3. Consent (2018): https://www.cdpd.bg/en/index.php?p=element&aid=1162</p> |
| <i>Human Biological Materials:</i> | <p>1. Executive Agency for Transplantation: http://www2.bgtransplant.bg/bg</p> <p>2. Council of Ministers, Ethics Committee for Transplantation</p> | <p>1. Law Ratifying the Additional Protocol to the Convention on Human Rights and Biomedicine concerning Transplantation of Organs and Tissues of Human Origin (2006): https://www.mh.government.bg/media/filer_public/c6/12/c612c85a-519f-4a22-b390-5fad298ce80b/zakon-ratifikatsiya-protokol-konventsia-zashhita-pravata-na-choveka-29-08-2006.pdf</p> <p>2. Law on Transplantation of Organs, Tissues, and Cells (2013): http://bgtransplant.bg/iat/docs/Zakoni_ZTOTK.doc</p> | <p>Regulation No. 13 of 4 April 2007 for the Terms and Conditions of Informing Bulgarian Citizens on the Activities regarding the Transplantation of Organs, Tissues and Cells: http://www2.bgtransplant.bg/sites/default/files/docs/naredbi/Naredba_no13_ot_04_april_2007_g.rtf</p> | |
| <i>Embryos, Stem Cells, and Cloning</i> | <p>Ministry of Healthcare: http://www.mh.government.bg/</p> | <p>1. Law Ratifying the Additional Protocol to the Convention on Human Rights and Biomedicine concerning Transplantation of Organs and Tissues of Human Origin (2006): https://www.mh.government.bg/media/filer_public/c6/12/c612c85a-519f-4a22-b390-5fad298ce80b/zakon-ratifikatsiya-protokol-konventsia-zashhita-pravata-na-choveka-29-08-2006.pdf</p> <p>2. Law on Transplantation of Organs, Tissues, and Cells (2013): http://bgtransplant.bg/iat/docs/Zakoni_ZTOTK.doc</p> | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|--|--|---|--|
| Croatia | | | | |
| Note: All websites and documents are in Croatian. | | | | |
| <i>General</i> | Central Ethics Committee: http://www.halmed.hr/en/O-HALMED-u/Sredisnje-eticko-povjerenstvo-SEP/ | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 15-18, ETS No. 164 (1997): http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=164&CM=7&DF=9/15/2008&CL=ENG 2. Patient Protection Act, Article 20 (2008): http://www.zakon.hr/z/255/Zakon-o-za%C5%A1titi-prava-pacijenata | | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | 1. Ministry of Health: https://zdravlje.gov.hr/ 2. Agency for Medicinal Products and Medical Devices: http://www.halmed.hr/ | 1. Medicinal Product Act (2013): http://narodne-novine.nn.hr/clanci/sluzbeni/2013_06_76_1522.html 2. Rule Book on Amendments to Medicinal Product Act (2014): http://narodne-novine.nn.hr/clanci/sluzbeni/2014_07_90_1809.html | 1. Ordinance on Clinical Trials and Good Clinical Practice (2015): http://narodne-novine.nn.hr/clanci/sluzbeni/2015_03_25_534.html 2. Ordinance on Amendments to the Ordinance on Clinical Trials and Good Clinical Practice (2015): https://narodne-novine.nn.hr/clanci/sluzbeni/2014_07_90_1809.html |
| | <i>Devices</i> | 1. Ministry of Health: https://zdravlje.gov.hr/ 2. Agency for Medicinal Products and Medical Devices: http://www.halmed.hr/ | Medical Devices Act (2013): http://narodne-novine.nn.hr/clanci/sluzbeni/2013_06_76_1521.html | |
| | <i>Research Injury</i> | 1. Agency for Medicinal Products and Medical Devices of Croatia: http://www.halmed.hr/ 2. Ministry of Health: https://zdravlje.gov.hr/ 3. Croatian Health Insurance Fund: http://www.hzzo.hr/en/ | 1. Law on Mandatory Health Insurance (2013): http://www.hzzo.hr/wp-content/uploads/2013/10/ZOZO_PR_OCISCENI_TEKSTv2.pdf?6d8ad4 2. Medicinal Product Act (2013): http://narodne-novine.nn.hr/clanci/sluzbeni/2013_06_76_1522.html 3. Rule Book on Amendments to Medicinal Product Act (2014): http://narodne-novine.nn.hr/clanci/sluzbeni/2014_07_90_1809.html | Ordinance on Clinical Trials and Good Clinical Practice, Articles 11 and 16, Act 5.8., 6.8. and 8.2.5 (2015): http://narodne-novine.nn.hr/clanci/sluzbeni/2015_03_25_534.html |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|---|---|--|---|
| <i>Privacy/Data Protection</i> | Croatian Personal Data Protection Agency: http://www.azop.hr/ | <p>7 90 1809.html</p> <p>1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj</p> <p>2. Implementation Act of the General Data Protection Act (NN 42/18) (2018): https://narodne-novine.nn.hr/clanci/sluzbeni/2018_05_42_805.html</p> | | General (2018): http://azop.hr/info-servis/detaljnije/smjernice |
| <i>Human Biological Materials</i> | Ministry of Health: https://zdravlje.gov.hr/ | <p>1. Law about Blood and Blood Products (2006): http://narodne-novine.nn.hr/clanci/sluzbeni/2006_07_79_1916.html</p> <p>2. Rule Book on Amendments to Law about Blood and Blood Products (2011): http://narodne-novine.nn.hr/clanci/sluzbeni/2011_11_124_2476.html</p> <p>3. Law on the Implementation of Human Tissues and Cells (2012): http://narodne-novine.nn.hr/clanci/sluzbeni/2012_12_144_3070.html</p> <p>4. Law on Transplantation of Human Organs for the Purpose of Treatment (2012): http://narodne-novine.nn.hr/clanci/sluzbeni/2012_12_144_3071.html</p> | Ordinance on the Conditions of Space, Professional Workers, Medical-Technical Equipment and Quality Assurance for Collection, Retrieval, Testing, Processing, Preservation, Storage, and Allocation of Human Tissues and Cells (2013): http://www.propisi.hr/print.php?id=9354 | |
| <i>Embryos, Stem Cells, and Cloning</i> | Ministry of Health: https://zdravlje.gov.hr/ | <p>1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (2003): http://zakon.poslovna.hr/public/Konvencija-o-zastiti-ljudskih-prava-i-dostojanstva-ljudskog-bica-u-pogledu-primjene-biologije-i-medicine-u-vezi-presadivanja-organa-i-tkiva-ljudskog-porijekla/243337/zakoni.aspx</p> <p>2. Medical Fertilization Act,</p> | Ordinance on the Conditions of Space, Professional Workers, Medical-Technical Equipment and Quality Assurance for Collection, Retrieval, Testing, Processing, Preservation, Storage and Allocation of Human Tissues and Cells (2013): http://www.propisi.hr/print.php?id=9354 | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|---|--|-------------|------------|
| <i>Embryos, Stem Cells, and Cloning</i> | | Article 32: (2012): http://www.hzzo-net.hr/dload/zakoni/20_01.pdf 3. Law on the Implementation of Human Tissues and Cells (2012): http://narodne-novine.nn.hr/clanci/sluzbeni/2012_12_144_3070.html | | |
| Cyprus | | | | |
| <i>General</i> | | 1. Law No. 31 (III)/2001: Oviedo Convention on Human Rights and Biomedicine 2. The Safeguarding and Protection of Patients' Rights Law (2004): http://www.bioethics.gov.cy/Moh/cnbc/cnbc.nsf/All/745717D26F068582C2257CCA003B350F/\$file/Patients%20Rights%20Law-English%20translation.pdf | | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health, Pharmaceutical Services: http://www.moh.gov.cy/Moh/phs/phs.nsf/dmlindex_en/dmlindex_en?opendocument 2. Ministry of Health, National Bioethics Committee: http://www.bioethics.gov.cy/moh/cnbc/cnbc.nsf/index_en/index_en?OpenDocument | Law for Good Clinical Practice (2004): http://www.moh.gov.cy/Moh/phs/phs.nsf/All/9C064264122B82BEC22572FA003433A5/\$file/%CE%9A.%CE%94.%CE%A0.%20452%20CF%84%CE%BF%CF%85%202004.pdf?OpenElement | | |
| <i>Research Injury</i> | Ministry of Health, Pharmaceutical Services: http://www.moh.gov.cy/moh/moh.nsf/index_en/index_en?OpenDocument | Legislation Concerning Medicinal Products of Human Use (Good Clinical Practice) No. 452/2004 Article 11 (8): http://www.moh.gov.cy/Moh/phs/phs.nsf/All/9C064264122B82BEC22572FA003433A5/\$file/%CE%9A.%CE%94.%CE%A0.%20452%20CF%84%CE%BF%CF%85%202004.pdf?OpenElement | | |
| <i>Privacy/Data Protection</i> | Commissioner's Office for the Protection of Personal Data: http://www.dataprotection.gov.cy/dataprotection/dataprotection.nsf/index_en/index_en?opendocument | 1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 2. Protection of Natural Persons Against the Processing of Personal Data and the Free Circulation of such Data Act of | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines | |
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| <i>Privacy/Data Protection</i> | | 2018 (Law 125 (I)): http://www.dataprotection.gov.cy/dataprotection/dataprotection.nsf/all/DE97F6F59835A03AC22582DD003D895E/\$file/%CE%9D%CF%8C%CE%BC%CE%BF%CF%82%20125(%CE%99) 2018.pdf?openelement | | | |
| <i>Embryos, Stem Cells, and Cloning</i> | | Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (2002) | | | |
| Czech Republic | | | | | |
| <i>General</i> | Ministry of Health, Central Ethics Committee: http://www.mzcr.cz | 1. Oviedo Convention on Human Rights and Biomedicine (2001) 2. Act No. 130/2002 Collection on Research and Development Support, as Amended 3. Act No. 372/2011 on Healthcare Services, As Amended (2018) 4. Act. No. 373/2011 on Specific Healthcare Services, As Amended (2018) | | | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | 1. Ministry of Health (MOH): http://www.mzcr.cz 2. State Institute for Drug Control (SUKL): http://www.sukl.cz/index.php?lchan=1&led=1 | Act No. 378/2007 Collection on Pharmaceuticals, As Amended (2017) | MOH: Decree No. 226/2008 on Good Clinical Practices and on Detailed Conditions for Evaluation of Pharmaceutical Products | SUKL: Various: http://www.sukl.cz/medicinal-products-clinical-trials-guidelines-1 |
| | <i>Devices</i> | State Institute for Drug Control (SUKL): http://www.sukl.cz/index.php?lchan=1&led=1 | 1. Act No 268/2014 Coll., on Medical Devices and on Amendment to Act. 634/2004 Coll., on Administrative Fees, As Amended (2017) 2. Decree No 62/2015 Coll. Implementing Certain Provisions of the Act on Medical Devices | Various: http://www.sukl.cz/medical-devices?highlightWords=501%2F2000 | Various: http://www.sukl.cz/medical-devices-guidelines |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|---|--|---|--|
| <i>Research Injury</i> | | <p>1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Article 24, ETS No. 164 (2001)</p> <p>2. Law No. 89/2012 Coll. Civil Code: http://www.czechlegislation.com/en/89-2012-sb</p> | | |
| <i>Privacy/Data Protection</i> | <p>Office for Personal Data Protection: https://www.uoou.cz/en/</p> | <p>1. Act No. 101/2000 Coll., On Protection of Personal Data and Amending Certain Laws, As Amended (2015): http://www.uoou.cz/uoou.aspx?menu=4&submenu=5</p> <p>2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj</p> | <p>1. EU General Data Protection Regulation (2016)</p> <p>2. Position No. 3/2004 Personal Data Processing in the Context of Clinical Testing of Drugs and Other Medical Substances</p> | <p>1. General (2018): https://www.uoou.cz/gdpr-strucne/ds-4843/p1=4843</p> <p>2. International Data Transfer (2018): https://www.uoou.cz/en/vismo/zobraz_dok.asp?id_org=200156&id_ktg=1165&p1=1165</p> |
| <i>Embryos, Stem Cells, and Cloning</i> | <p>1. Ministry of Education, Youth, and Sport: http://www.msmt.cz/index.php?lchan=1&lred=1</p> <p>2. Research and Development Council, Bioethical Commission: http://www.vyzkum.cz/FrontClanek.aspx?idsekce=15908</p> | <p>Act of 26 April 2006 on Research on Human Embryonic Stem Cells No. 227/2006 Sb. (Coll.)</p> | | |
| Denmark | | | | |
| <i>General</i> | <p>National Committee on Health Research Ethics (NVK): http://www.nvk.dk/english</p> | <p>Act No. 1083 on Research Ethics Review of Health Research Projects (2017): https://www.retsinformation.dk/Forms/R0710.aspx?id=192671</p> <p>2013 version (English): http://www.nvk.dk/english/act-on-research</p> | <p>Executive Order No. 1464 on Information and Consent at Inclusion of Trial Subjects in Biomedical Research Projects (2018) (Danish): https://www.retsinformation.dk/Forms/R0710.aspx?id=201254</p> | <p>Guidelines about Notification (Checklist) (2017): http://www.nvk.dk/forsker/forskervejledning</p> |
| <i>Drugs, Biologics, and Devices</i> | <p>Danish Medicines Agency: https://laegemiddelstyrelsen.dk/en/</p> | <p>1. Act No. 506 on Medicinal Products (2013): https://www.retsinformation.dk/forms/r0710.aspx?id=146586</p> <p>2. Act No. 620 on Clinical trials on Medical Products No. 620 (2016): https://www.retsinformation.dk/Forms/r0710.aspx?id=180117</p> | <p>1. Executive Order No. 295 on Clinical Trials of Medicinal Products on Humans (2006): https://www.retsinformation.dk/Forms/R0710.aspx?id=9891</p> <p>2. Executive Order No. 1464 on Information and Consent at Inclusion of Trial Subjects in Biomedical Research Projects (2016): https://www.retsinformation.dk/Forms/R0710.aspx?id=185233</p> | <p>Guidelines for Applications for Authorisation of Clinical Trials of Medical Products in Humans (2017): https://laegemiddelstyrelsen.dk/en/licensing/clinical-trials/trials-in-humans/guideline-for-applications-for-authorisation-of-clinical-trials-of-medicinal-products-in-humans/</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|---|--|--|---|
| <i>Research Injury</i> | Patient Compensation Association: http://pebl.dk/en.aspx | 1. Liability for Damages Act (2007): http://pebl.dk/Patientskader/Love-og-regler/Lov-om-klage-og-erstatningsadgang/Behandlingsskader 2. Act No. 1022 on the Right to Complain and Receive Compensation within the Health Service (2017): https://www.retsinformation.dk/Forms/R0710.aspx?id=192623 | | |
| <i>Privacy/Data Protection</i> | Danish Data Protection Agency (DPA): https://www.datatilsynet.dk/english/the-danish-data-protection-agency/introduction-to-the-danish-data-protection-agency/ | 1. Act No. 429 on Processing of Personal Data (2007): https://www.datatilsynet.dk/english/the-act-on-processing-of-personal-data/read-the-act-on-processing-of-personal-data/compiled-version-of-the-act-on-processing-of-personal-data/ 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | Executive Order No. 1188 on Health Law, Chapter 9 (2016): https://www.retsinformation.dk/forms/r0710.aspx?id=183932#idc101cee1-c9c0-4880-ac97-586a56134f56 | |
| <i>Human Biological Materials</i> | National Committee on Health Research Ethics (NVK): http://www.nvk.dk/english | 1. Act No. 1083 on Research Ethics Review of Health Research Projects (2017): https://www.retsinformation.dk/Forms/R0710.aspx?id=192671 2. Data Protection Act 2018: https://www.datatilsynet.dk/media/6894/danish-data-protection-act.pdf | Executive Order No. 1188 on Health Law (2018): https://www.retsinformation.dk/Forms/R0710.aspx?id=199871 | Guidelines on the Use of Biological Material in Health Research Projects (2017): http://www.nvk.dk/emner/biobanker/vejledning-om-bio-mat |
| <i>Genetic Research</i> | National Committee on Health Research Ethics (NVK): http://www.nvk.dk/english | Act No. 1083 on Research Ethics Review of Health Research Projects (2017): https://www.retsinformation.dk/Forms/R0710.aspx?id=192671 2013 version (English): http://www.nvk.dk/english/act-on-research | | Guidelines on Health Research Projects Involving Genome Research (2018): http://www.nvk.dk/emner/genomer/vejledning-om-genomer |
| <i>Embryos, Stem Cells, and Cloning</i> | Danish Council of Ethics: http://www.etiskraad.dk/english | Act No. 440 on Danish Council of Ethics (2004): https://www.retsinformation.dk/forms/r0710.aspx?id=9909 | Executive Order No. 93 on Medically Assisted Procreation (2015): https://www.retsinformation.dk/forms/r0710.aspx?id=167647 | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|--|--|--|---|
| Estonia | | | | |
| <i>General</i> | Estonian Council on Bioethics: http://www.eetikakeskus.ut.ee/en | 1. Oviedo Convention on Human Rights and Biomedicine (2002) 2. Constitution of the Republic of Estonia, Paragraph 18 (2016): https://www.riigiteataja.ee/en/eli/521052015001/consolide | | Code of Ethics of Estonian Scientists: http://www.akadeemia.ee/repository/File/ALUSDOKUD/Code-ethics.pdf |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs:</i> 1. State Agency of Medicines: http://www.sam.ee/en/clinical-trials-medicinal-products-estonia 2. Minister of Social Affairs (MSA): https://www.sm.ee/en | Medicinal Products Act, Chapter 5 (2015): https://www.riigiteataja.ee/en/eli/ee/525112013005/consolide/current | MSA: 1. 1 RTL 2005, 22, 298: Rules of Procedure of Medical Ethics Committee for Clinical Trials, a List of Data to be Submitted for Obtaining Approval, Procedure for Adoption of Resolutions and Format of Application for Obtaining Approval (2005): https://www.riigiteataja.ee/en/eli/502052017001/consolide 2. Conditions and Procedure for Conducting Clinical Trials of Medicinal Products. Regulation No. 23 (2005): https://www.riigiteataja.ee/en/eli/502052017002/consolide | |
| | <i>Devices:</i> Estonian Health Board: http://www.terviseamet.ee/en/medical-devices.html | Medical Devices Act (2004): https://www.riigiteataja.ee/en/eli/ee/509012015001/consolide/current | Regulation No 86: 2010 of the Minister of Social Affairs on the Conditions and Procedures for the Clinical Investigation of Medical Devices | |
| <i>Research Injury</i> | 1. Minister of Social Affairs (MSA): https://www.sm.ee/en 2. Estonian Health Insurance Fund: https://www.haigekassa.ee/en | Medicinal Products Act, Section 90: https://www.riigiteataja.ee/en/eli/ee/525112013005/consolide/current | Conditions and Procedure for Conducting Clinical Trials of Medicinal Products. Regulation No. 23 of the Minister of Social Affairs of (2005): https://www.riigiteataja.ee/en/eli/502052017002/consolide | |
| <i>Privacy/Data Protection</i> | Estonian Data Protection Inspectorate: http://www.aki.ee/en/inspectorate | 1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 2. Personal Data Protection Act (2016): https://www.riigiteataja.ee/en/eli/ee/512112013011/consolide/current | | 1. Research (2018) 2. International Data Transfer (2018): http://www.aki.ee/en/guidelines/transfer-personal-data-foreign-country |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | | Human Genes Research Act (RT I 2000, 104, 685) (2014): https://www.riigiteataja.ee/en/eli/ee/518062014005/consolide | | |
| <i>Embryos, Stem Cells, and Cloning</i> | | 1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (2002) (Estonian): https://www.riigiteataja.ee/akt/78569 2. Artificial Insemination and Embryo Protection Act, RT I 1997, 51, 824 (2011): https://www.riigiteataja.ee/en/eli/ee/530102013057/consolide/current | | |
| Finland | | | | |
| <i>General</i> | 1. Ministry of Social Affairs and Health (MSAH): http://www.stm.fi/en/frontpage 2. National Committee on Medical Research Ethics (TUKIJA): http://www.tukija.fi/en 3. Finnish Advisory Board on Research Integrity (TENK): http://www.tenk.fi/en | Medical Research Act No. 488/1999 (Amended 295/2004, 794/2010, and 143/2015): http://www.finlex.fi/en/laki/kaannokset/1999/en19990488 | Decree on Medical Research Nos. 986/1999, 313/2004, and 65/2016: http://www.finlex.fi/en/laki/kaannokset/1999/en19990986.pdf | TUKIJA: 1. Report on Children in Medical Research (2003) 2. Operating Procedures of the National Committee on Medical Research Ethics (2017) <i>Access:</i> http://tukija.fi/en/publications1 |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. Finnish Medicines Agency (FIMEA): http://www.fimea.fi/frontpage 2. Ministry of Social Affairs and Health (MSAH): http://stm.fi/en/frontpage 3. National Committee on Medical Research Ethics (TUKIJA): http://www.tukija.fi/en | 1. Medicines Act No. 395/1987: http://www.finlex.fi/fi/laki/smur/1987/19870395 2. Medical Research Act No. 488/1999 (Amended 295/2004, 794/2010, and 143/2015): http://www.finlex.fi/en/laki/kaannokset/1999/en19990488 | 1. Decree on Clinical Trials on Medicinal Products No. 841/2010 2. Other Decrees: http://www.finlex.fi/fi/laki/smur/1987/19870395#nojalla FIMEA: Administrative Regulation on Clinical Trials on Medicinal Products in Human Subjects No. 2/2012: http://www.fimea.fi/download/22302_Maarays_2-2012_kliiniset_laaketutkimukset.pdf | TUKIJA: Templates for Clinical Trial Information Leaflet and Consent Form (2016) <i>Access:</i> http://tukija.fi/en/publications1 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | <p><i>Devices</i></p> <p>National Supervisory Authority for Welfare and Health (VALVIRA): http://www.valvira.fi/en/licensing/medical_devices</p> | <p>Medical Devices Act No. 629/2010 (Finnish): http://www.finlex.fi/fi/laki/kokoelma/2010/20100085.pdf</p> <p>EU Regulations: Medical Device Regulation 2017/745: http://eur-lex.europa.eu/legal-content/EN/TXT/PDF/?uri=CELEX:32017R0745&from=EN In Vitro Diagnostic Medical Devices Regulation 2017/746: http://eur-lex.europa.eu/legal-content/EN/TXT/PDF/?uri=CELEX:32017R0746&from=EN</p> | <p>1. Decree (Decision) on Clinical Investigations (2010): http://www.finlex.fi/data/normit/39644-maarays_3_2010_kliininen_laitetutkimus.pdf</p> <p>2. Various: http://www.valvira.fi/en/licensing/medical_devices/legislation</p> | |
| <i>Research Injury</i> | <p>1. Finnish Patient Insurance Centre: http://www.potilasvakuutuskeskus.fi/www/page/pvk_www_2181</p> <p>2. Pharmaceutical Injuries Insurance http://www.laakevahinko.fi/in-english/</p> | <p>Patient Injuries Act No. 585/1986: http://www.finlex.fi/fi/laki/ajantasa/1986/19860585</p> | | <p>Pharmaceutical Injuries Insurance: General Terms and Conditions (2017): http://www.laakevahinko.fi/in-english/terms-and-conditions/</p> |
| <i>Social-Behavioral Research</i> | <p>Finnish Advisory Board on Research Integrity (TENK): http://www.tenk.fi/en/</p> | | | <p>Ethical Principles of Research in the Humanities and Social and Behavioural Sciences and Proposals for Ethical Review (2009): http://www.tenk.fi/sites/tenk.fi/files/ethicalprinciples.pdf</p> |
| <i>Privacy/Data Protection</i> | <p>Office of the Data Protection Ombudsman: http://www.tietosuoja.fi/1560.htm</p> | <p>1. Personal Data Act No. 523/1999: http://www.finlex.fi/fi/laki/ajantasa/1999/19990523</p> <p>2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj</p> | | |
| <i>Human Biological Materials</i> | <p>National Supervisory Authority for Welfare and Health (Valvira): http://www.valvira.fi/web/en</p> | <p>1. Act on the Medical Use of Human Organs, Tissues, and Cells No. 101/2001 (Finnish and Swedish): http://www.finlex.fi/fi/laki/ajantasa/2001/20010101</p> <p>2. Law on Biobanks, No 688/2012 (Finnish and Swedish): http://www.finlex.fi/fi/laki/ajantasa/2012/20120688</p> | <p>1. Decree on Consent for Biobank No. 643/2013: http://www.finlex.fi/fi/laki/alkup/2013/20130643</p> <p>2. Decree on information on Biobank No. 649/2013: http://www.finlex.fi/fi/laki/alkup/2013/20130649</p> <p>3. Government Decree on Medical Use of Human Organs, Tissues, and Cells No. 594/2007</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | | | 4. Ministry Decree on Medical Use of Human Organs, Tissues, and Cells No. 1302/2007 | |
| <i>Genetic Research</i> | National Committee on Medical Research Ethics (TUKIJA): http://www.tukija.fi/en | Medical Research Act No. 488/1999 (Amended 295/2004, 794/2010, and 143/2015): http://www.finlex.fi/en/laki/kaannokset/1999/en19990488 | | |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. National Supervisory Authority for Welfare and Health: http://www.valvira.fi/web/en 2. National Committee on Medical Research Ethics (TUKIJA) http://www.tukija.fi/en 3. Finnish Advisory Board on Research Integrity (TENK): http://www.tenk.fi/en/ 4. National Advisory Board on Social Welfare and Health Care Ethics (ETENE): http://www.etene.fi/en | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Additional Protocol on Prohibition of Human Cloning (2002) 2. Medical Research Act No. 488/1999 (amended 295/2004, 749/2010, and 143/2015): http://www.finlex.fi/en/laki/kaannokset/1999/en19990488 3. Act on Assisted Fertility Treatments No. 1237/2006: http://www.finlex.fi/fi/laki/ajantasa/2006/20061237 4. Criminal Code of Finland (39/1889), Chapter 22, Section 4: https://www.finlex.fi/en/laki/kaannokset/1889/en18890039.pdf | | TUKIJA: Report on Stem Cells, Cloning, and Research (2005): http://tukija.fi/documents/1481661/1546647/2005cells.pdf/c14b7dd0-11b4-428d-bdae-539566ade614 |

| France | | | | |
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| <i>General</i> | 1. Ministry of Social affairs and Health: http://www.sante.gouv.fr/ 2. National Consultative Bioethics Committee for Health and Life Sciences (CCNE): http://www.ccne-ethique.fr/en 3. National Commission for Informatics and Freedoms (CNIL): http://www.cnil.fr/english/the-cnil/ | 1. Law No. 2012-300 of 5 March 2012 on Research Involving Human Persons: https://www.legifrance.gouv.fr/affichTexte.do?cidTexte=JORFTEXT00025441587 2. Law No. 2011-814 of 7 July 2011 on Bioethics | Public Health Code Articles R1121-1 and subsequent sections: http://legifrance.gouv.fr/ | CCNE: Various: http://www.ccne-ethique.fr/en/type_publication/avis |
| <i>Drugs, Biologics, and Devices</i> | 1. National Consultative Bioethics Committee for Health and Life Sciences (CCNE): http://www.ccne-ethique.fr 2. National Health Products Safety Agency (ANSM): http://ansm.sante.fr/ | Medications for Human Use, Articles L5121-11, L5124-1, and L5126-1 (2004): http://www.legifrance.gouv.fr/ | Decision on Good Clinical Practices: http://www.legifrance.gouv.fr/affichTexte.do?cidTexte=JORFTEXT00000819256 | CCNE: Phase I Trials in Cancer (2002) Access: http://www.ccne-ethique.fr/en/type_publication/avis |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|-----------------------------------|---|---|---|--|
| <i>Social-Behavioral Research</i> | National Consultative Ethics Committee | | | Opinion on the Ethics of Research in the Sciences of Human Behavior No. 38 (1993): http://www.ccne-ethique.fr/en/publications/opinion-ethics-research-sciences-human-behaviour#.WNkybNfyEY |
| <i>Privacy/Data Protection</i> | <ol style="list-style-type: none"> National Commission of Information and Liberty (CNIL): http://www.cnil.fr/english/ National Consultative Bioethics Committee for Health and Life Sciences (CCNE): http://www.ccne-ethique.fr | <ol style="list-style-type: none"> Law 2004-801 of August 6, 2004 Modifying Law 78-17 of January 6, 1978 Relating to the Protection of Data Subjects as Regards the Processing of Personal Data Law No. 2016-1321 of 7 October 2016 for a Numeric Republic: https://www.legifrance.gouv.fr/affichLoiPubliee.do?idDocument=JORFDOLE000031589829&type=general&legislature=14 EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj Data Protection Act (2018): https://www.legifrance.gouv.fr/affichLoiPreparation.do;jsessionid=AD5660270AD9F70B94275AC823321680.tplgfr22s_3?idDocument=JORFDOLE000036195293&type=contenu&id=2&typeLoi=proj&legislature=15 | CNIL: Decree No. 2005-1309 of 20 October 2005 Enacted for the Application of Act No. 78-17 of 6 January 1978 on Data Processing, Files and Individual Liberties (Amended by Decree 2007-451 of 25 March 2007): http://www.cnil.fr/fileadmin/documents/en/Decree%202005-1309.pdf | CNIL: <ol style="list-style-type: none"> Health Research: CNIL Adopts New Simplification Measures (2018): https://www.cnil.fr/fr/recherches-dans-le-domaine-de-la-sante-la-cnil-adopte-de-nouvelles-mesures-de-simplification Health Research with Consent (2018): https://www.cnil.fr/fr/declaration/mr-001-recherches-dans-le-domaine-de-la-sante-avec-recueil-du-consentement Health Research without Consent (2018): https://www.cnil.fr/fr/declaration/mr-003-recherches-dans-le-domaine-de-la-sante-sans-recueil-du-consentement Practical Guide on the Protection of Personal Data: What Framework Applies to Research? (2018): https://www.cnil.fr/sites/default/files/atoms/files/guide-cnom-cnil.pdf CCNE: <ol style="list-style-type: none"> Ethical Questions Arising from the Transmission of Scientific Information Concerning Research in Biology and Medicine (1995) Biometrics, Identifying Data and Human Rights (2007) Access: http://www.ccne-ethique.fr/en/type_publication/avis |
| <i>Human Biological Materials</i> | <ol style="list-style-type: none"> National Consultative Bioethics Committee for Health and Life Sciences (CCNE): http://www.ccne-ethique.fr Ministry of Higher Education, Research, and Innovation: http://www.enseignementsup-recherche.gouv.fr/ | <ol style="list-style-type: none"> Donation and Use of the Components and Products of the Human Body, Articles L1211-1 to L1274-3 (2004): http://www.legifrance.gouv.fr/ Public Health Code Articles L1241-1 and following sections: (2010): http://www.legifrance.gouv.fr/initR | | CCNE: <ol style="list-style-type: none"> Umbilical Cord Blood Banks for Autologous Use for Research (2002) Ethical Issues Raised by Collections of Biological Material and Associated Information Data: “Biobanks,” “Biolibraries” (2003) |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | | echCodeArticle.do | | Access: http://www.ccne-ethique.fr/en/type_publication/avis |
| <i>Genetic Research</i> | 1. National Consultative Bioethics Committee for Health and Life Sciences (CCNE): http://www.ccne-ethique.fr 2. Biomedicine Agency: http://www.enseignementsup-recherche.gouv.fr/ | Civil Code Articles 16-10 to 16-13: http://www.legifrance.gouv.fr/affichCode.do;jsessionid=D2DE023194483D3384DE19DE8959BDDA.tpdjo17v_3?idSectionTA=LEGISCTA000006136513&cidTexte=LEGITEXT000006070721&dateTexte=20131006 | | 1. Ethical Issues in Connection with the Development of Foetal Genetic Testing on Maternal Blood (2013) 2. Ethical Reflection on Developments in Genetic Testing in Connection with Very High Throughput Human DNA Sequencing (2016) Access: http://www.ccne-ethique.fr/en/type_publication/avis |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. National Consultative Bioethics Committee for Health and Life Sciences (CCNE): http://www.ccne-ethique.fr 2. Biomedicine Agency: http://www.enseignementsup-recherche.gouv.fr/ | Law No. 2013-715 of 6th August 2013: http://www.legifrance.gouv.fr/affichTexte.do?cidTexte=JORFTEXT000027811435&dateTexte=&categorieLien=id | Decree No. 2015-155 of 11 February, 2015: Public Health Code on Research on Embryos Article R2151-1 and Following Sections: http://legifrance.gouv.fr/affichCode.do?idArticle=LEGIARTI000030233469&idSectionTA=LEGISCTA000006190409&cidTexte=LEGITEXT000006072665&dateTexte=20151015 | 1. Commercialization of Human Stem Cells and Other Cell Lines (2006) 2. Opinion on the Ethical Reflection Concerning Research on Human Embryonic Cells and on Human Embryos in Vitro (2010) Access: http://www.ccne-ethique.fr/en/type_publication/avis |
| Georgia | | | | |
| For an overview of human subject protections in Georgia, see “Ethical Review of Biomedical Research in the CIS Countries,” Chapter 3, Section 4: http://www.unesco.org/new/fileadmin/MULTIMEDIA/FIELD/Moscow/pdf/ethical_review_cis_book_kubar_english.pdf | | | | |
| <i>General</i> | Bioethics and Health Law Studies Society: http://www.patientsrights.ge/index.php?page=385&lang=geo | 1. Oviedo Convention on Human Rights and Biomedicine ETS No.164 (2001) 2. Additional Protocol to the Convention’s on Human Rights and Biomedicine, concerning Biomedical Research, ETS No. 195 (2010) 3. Law on Health Care, Chapter XIX (2017): https://matsne.gov.ge/en/document/view/29980?publication=37 | | |
| <i>Drugs, Biologics, and Devices</i> | State Regulation Agency for Medical Activities (LEPL) of the Ministry of Labor, Health, and Social Affairs: http://rama.moh.gov.ge/ | Law on Medicines and Pharmaceutical Activities No. 659 and 1586 (2015): https://matsne.gov.ge/en/document/view/29836?impose=translateEn | Regulation about the Rules and Conditions of Issuing of the Approval of Clinical Trials Approved #176 (2005): http://rama.moh.gov.ge/res/docs/20160809105943176.pdf | Order of Health Minister about Implementation of “ICH: E6 Good Clinical Practice: Consolidated Guidance” (1996) including WMA: Declaration of Helsinki (2013): http://rama.moh.gov.ge/res/docs/9539N233.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|--|--|-------------|--|
| <i>Research Injury</i> | | Convention on Human Rights and Biomedicine (Convention of Oviedo), Article 24, ETS No. 164 (2001) | | |
| <i>Privacy/Data Protection</i> | Office of the Personal Data Protection Inspector: https://personaldata.ge/en/home | 1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 2. Law on Data Protection (2018): https://matsne.gov.ge/en/document/view/1561437?publication=15 | | |
| <i>Embryos, Stem Cells, and Cloning</i> | | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Additional Protocol on Prohibition of Human Cloning ETS No. 168 (2001) 2. Law on Health Care, Article 142 (2017): https://matsne.gov.ge/en/document/view/29980?publication=37 | | |
| Germany | | | | |
| <i>General</i> | 1. German Medical Association (BÄK): http://www.bundesaerztekammer.de/weitere-sprachen/english/german-medical-association/ 2. Central Ethics Committee of the German Medical Association (ZEKO): http://www.zentrale-ethikkommission.de/ 3. Permanent Working Party of Research Ethics Committees in Germany: http://www.ak-med-ethik-komm.de/ 4. German Ethics Council: https://www.ethikrat.org/en/ 5. Federal Ministry of Health (BMG): http://www.bundesgesundheitsministerium.de/en/en.html 6. German Research Foundation (DFG), Permanent Senate Commission on Key Questions in Clinical Research (SCCR): http://www.dfg.de/en/dfg_profile/statutory_bodies/senate/clinical_research/index.ht | | | BÄK: (Model) Professional Code for Physicians in Germany, Article 15 (2018): http://www.bundesaerztekammer.de/fileadmin/user_upload/downloads/pdf-Ordner/MBO/MBO-AE_EN_2018.pdf |

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| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. Federal Institute for Drugs and Medical Devices (BfArM): http://www.bfarm.de/EN/Home/home_no_de.html 2. Paul-Ehrlich-Institut (PEI): http://www.pei.de/EN/home/node.html4 3. Federal Ministry of Health (BMG): http://www.bundesgesundheitsministerium.de/en/en.html | Medicinal Products Act, Sixth Chapter (2017): http://www.gesetze-im-internet.de/amg_1976/ <i>2016 English version, without amendments:</i> Medicinal Products Act, Sixth Chapter (2016): http://www.gesetze-im-internet.de/englisch_amg/englisch_amg.html#p0925 | 1. Promulgation on the Principles of the Conduct of Clinical Trials of Drugs According to the Rules (1987) 2. Second Promulgation on the Clinical Trial of Drugs in Human (1997) 3. Regulation on the Application of Good Clinical Practice in the Conduct of Clinical Trials of Medicinal Products for Human Use (2012): http://www.gesetze-im-internet.de/gcp-v/ | BfArM and PEI: Third Notification on the Clinical Trials of Medicinal Products for Humans (2006): http://www.pei.de/SharedDocs/Downloads/EN/pu/clinical-trials/3rd-notification-clinical-trials-2006-08-10.pdf?__blob=publicationFile&v=1 |
| | <i>Devices</i> 1. Federal Institute for Drugs and Medical Devices (BfArM): http://www.bfarm.de/EN/Home/home_no_de.html 2. Paul-Ehrlich-Institut (PEI): http://www.pei.de/EN/home/node.html4 3. Federal Ministry of Health (BMG): http://www.bundesgesundheitsministerium.de/en/en.html | Act on Medical Devices, Fourth Chapter (2017): http://www.gesetze-im-internet.de/mpg/ | Regulation on Clinical Trials of Medical Devices (2014): http://www.gesetze-im-internet.de/mpkpv/ | |
| <i>Clinical Trials Registry</i> | German Clinical Trials Register (DRKS): https://www.drks.de/drks_web/setLocale_EN.do | | | FAQs: https://www.drks.de/drks_web/navigate.do?navigationId=faq&messageEN=FAQ |
| <i>Research Injury</i> | | 1. Medicinal Products Act, Section 40(3) (2016): http://www.gesetze-im-internet.de/englisch_amg/englisch_amg.html#p0926 2. Act on Medical Devices, Section 20(3) (2017): http://www.gesetze-im-internet.de/mpg/_20.html | | |
| <i>Privacy/Data Protection</i> Note: The 16 German states also have data protection | 1. Federal Commissioner for Data Protection and Freedom of Information: https://www.bfdi.bund.de/ 2. Datenschutzkonferenz (DSK): https://www.datenschutzkonferenz-online.de/ | 1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 2. Federal Data Protection Act (BDSG) (2017): https://www.gesetze-im-internet.de/englisch_bdsg/index.htm | | DSK: Short Paper No. 4: Data Transmission to Third Countries: https://www.bfdi.bund.de/SharedDocs/Downloads/DE/Datenschutz/Kurzpapier_Drittlaender.pdf?__blob=publicationFile&v=3 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| laws: http://www.datenschutzbayern.de/infoquel/ds-inst/deutschland.html | | 1 | | |
| <i>Human Biological Materials</i> | | 1. Act of Quality and Security of Human Tissue and Cells (2007): http://www.gesetze-im-internet.de/gewebeg/ 2. Transfusion Law (2017): http://www.gesetze-im-internet.de/tfg/ 3. Transplantation Law (2017): http://www.gesetze-im-internet.de/tpg/ | | |
| | German Ethics Council: https://www.ethikrat.org/en/ | | | Opinion on Human Biobanks for Research (2010): https://www.ethikrat.org/fileadmin/Publikationen/Stellungnahmen/englisch/DER_StnBiob_Engl_Online_mitKennwort.pdf |
| | Central Ethics Committee of the German Medical Association (ZEKO) (German): http://www.zentrale-ethikkommission.de/ | | | 1. Opinion on the (Re)Use of Human Body Material for Medical Research Purposes (2003): http://www.zentrale-ethikkommission.de/fileadmin/user_upload/downloads/pdf-Ordner/Zeko/Koerpermat-1.pdf 2. First Addendum: The (Re)Use of Human Body Material of Deceased Persons for Medical Research Purposes (2003): http://www.zentrale-ethikkommission.de/fileadmin/user_upload/downloads/pdf-Ordner/Zeko/Erste_Ergaenzung_Koerpermaterialien.pdf |
| | German Society of Surgery (DGCH): http://www.dgch.de/index.php?id=118 | | | DGCH Guidelines on Good Professional Practice (GPP) for the Procurement of Human Tissue and Cells for Drug Production: http://www.dgch.de/fileadmin/media/pdf/servicemeldungen/069_Gewebegesetz_GFP-Leitfaden_der_DGCH_fuer_die_Gewinnung_menschlicher_Gewebe.pdf |
| <i>Genetic Research</i> | | 1. Embryo Protection Act (2011): http://www.gesetze-im-internet.de/eschg/ | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | | 2. Genetic Engineering Act (2017): http://www.gesetze-im-internet.de/gentg/ | | |
| | German Society of Human Genetics (GfH): http://www.gfhev.de/en/gfh/ | | | 1. DNA Banking and Personal Data in Biomedical Research: Technical, Social, and Ethical Questions (2004): http://www.medgenetik.de/sonderdruck/en/DNA%20Banking_engl_060605.pdf 2. Position Paper of the German Society of Human Genetics (2007): http://www.medgenetik.de/sonderdruck/2007_gfh_positionspapier.pdf |
| | German Research Foundation (DFG), Permanent Senate Commission on Genetic Research: http://www.dfg.de/en/dfg_profile/statutory_bodies/senate/genetic_research/index.html | | | Statements: http://www.dfg.de/en/dfg_profile/statutory_bodies/senate/genetic_research/publications/index.html |
| <i>Embryos, Stem Cells, and Cloning</i> | Federal Ministry of Education and Research (BMBF): http://www.bmbf.de/en/index.php | 1. Embryo Protection Act (2011): http://www.gesetze-im-internet.de/eschg/ 2. Stem Cell Act (2017): http://www.gesetze-im-internet.de/stzg/ | Regulation on the Central Ethics Committee for Stem Cell Research and the Competent Authority Pursuant to the Stem Cell Act (2017): http://www.gesetze-im-internet.de/zesv/ | |
| | German Ethics Council: https://www.ethikrat.org/en/ | | | 1. The Import of Human Embryonic Stem Cells (2001): https://www.ethikrat.org/fileadmin/Publikation/en/Stellungnahmen/Archiv/Stellungnahme_Stammzellimport.pdf 2. Cloning for Reproductive Purposes and Cloning for the Purposes of Biomedical Research (2004): https://www.ethikrat.org/fileadmin/Publikation/en/Stellungnahmen/Archiv/Stellungnahme_Klonen.pdf 3. Should the Stem Cell Law be Amended? (2007): https://www.ethikrat.org/fileadmin/Publikation/en/Stellungnahmen/Archiv/Stn_Stammzellgesetz.pdf 4. Human-Animal Mixtures in Research (2011): https://www.ethikrat.org/fileadmin/Publikation/en/Stellungnahmen/englisch/opinion-human- |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | | | | animal-mixtures-in-research.pdf 5. Stem Cell Research - New Challenges for the Ban on Cloning and Treatment of Artificially Created Germ Cells? (2014): https://www.ethikrat.org/fileadmin/Publikation/en/Ad-hoc-Empfehlungen/englisch/recommendation-stem-cell-research.pdf 6. Germline Intervention in the Human Embryo (2017): http://www.zentrale-ethikkommission.de/fileadmin/user_upload/downloads/pdf-Ordner/Zeko/Stammzell.pdf |
| | Central Ethics Committee of the German Medical Association (ZEKO): http://www.zentrale-ethikkommission.de/ | | | Opinion on Stem Cell Research (2002): http://www.zentrale-ethikkommission.de/downloads/Stammzell.pdf |
| | German Research Foundation (DFG): http://www.dfg.de/en/ | | | Opinion on Stem Cell Research (2006): http://www.dfg.de/download/pdf/dfg_magazin/forschungspolitik/stammzellforschung/stammzellforschung_deutschland_lang_0610.pdf |
| | Central Ethics Committee for Stem Cell Research (ZES): http://www.rki.de/EN/Content/Institute/Committees/StemCell/StemCell_content.html | | | |
| Greece | | | | |
| <i>General</i> | National Bioethics Commission (NBC): http://www.bioethics.gr/ | | | 1. Research Ethics for Biological Sciences (2008): http://www.bioethics.gr/index.php/en/gnomes/86-research-ethics-in-biological-sciences 2. A Guide for Research Ethics Committees for Biological Research (2008): http://www.bioethics.gr/images/pdf/ENGLISH/OPINIONS_REPORTS/guide.pdf 3. Conflict of Interest in Biomedical Research (2014): http://www.bioethics.gr/images/pdf/EKDOSEIS/OPINIONS_AND_REPORTS_2008-2013_EN.pdf 4. Incidental Findings in Research and Clinical Practice (2015): http://www.bioethics.gr/index.php/en/gnomes/983-incident- findings-in-research-and- |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|--|---|--|--|
| <i>General</i> | | | | clinical-practice |
| <i>Drugs, Biologics, and Devices</i> | <p>1. National Organization for Medicines (NOM): http://www.eof.gr/web/guest/home, then click on “EN” in upper left hand section for English</p> <p>2. National Bioethics Commission (NBC): http://www.bioethics.gr/index.php?category_id=3</p> | <p>1. Act 2619/98: Ratification of the Convention for the Protection of Human Rights and Dignity of the Human Being with Regard to the Application of Biology and Medicine (1998)</p> <p>2. Act 3418/2005 Code on Medical Ethics</p> | <p>1. Ministerial Decision ΔΥΤ3 89292/2003: Harmonization of the Greek Legislation with EU Legislation, According to the Directive 2001/20/EC</p> <p>2. Ministerial Decision ΔΥΤ 3 α/79602/2007: Harmonization of the Greek Legislation with EU Legislation, according to the Directive 2005/28/EC</p> | <p>NBC:</p> <p>1. Recommendation on Clinical Trials: http://www.bioethics.gr/images/pdf/ENGLISH/OPINIONS_REPORTS/recom_clinical_trials_en.pdf</p> <p>2. Control of Non-Invasive Clinical Trials for Drugs (2013): http://www.bioethics.gr/index.php/en/gnomes/532-control-of-non-invasive-clinical-trials-for-drugs</p> |
| <i>Research Injury</i> | <p>National Bioethics Commission (NBC): http://www.bioethics.gr/index.php?category_id=3</p> | <p>1. Act 2619/98: Ratification of the Convention for the Protection of Human Rights and Dignity of the Human Being with Regard to the Application of Biology and Medicine (1998)</p> <p>2. Act 3418/2005 Code on Medical Ethics</p> | <p>1. Ministerial Decision ΔΥΤ3 89292/2003: Harmonization of the Greek Legislation with EU Legislation, According to the Directive 2001/20/EC</p> <p>2. Ministerial Decision ΔΥΤ 3 α/79602/2007 Harmonization of the Greek Legislation with EU Legislation, According to the Directive 2005/28/EC:</p> | |
| <i>Privacy/Data Protection</i> | <p>Hellenic Data Protection Authority: http://www.dpa.gr/</p> | <p>1. Greek Constitution 1975/1986/2001 Article 9.1</p> <p>2. Act 2619/98 (Biomedicine Convention of the Council of Europe) (1998)</p> <p>3. Act 2472/1997 on the Protection of Individuals with Regard to the Processing of Personal Data (As Amended by Laws 2819/2000 and 2915/2000): http://www.dpa.gr/portal/page?_page_id=33.19052&_dad=portal&_schema=PORTAL</p> <p>4. Act 3418/2005 Code on Medical Ethics</p> <p>5. EU General Data Protection Regulation (2016): https://www.lawspot.gr/nomikes-pleirotories/nomothesia/genikos-kanonismos-gia-tin-prostasia-dedomenon?lspt_context=gdpr</p> | | |
| <i>Genetic Research</i> | <p>National Bioethics Commission (NBC):</p> | <p>1. Greek Constitution 1975/1986/2001, Article 5.5</p> | | <p>1. Recommendation on Banks of Biological Material of Human Origin</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | http://www.bioethics.gr/index.php?category_id=3 | <p>2. Act 2619/98: Ratification of the Convention for the Protection of Human Rights and Dignity of the Human Being with Regard to the Application of Biology and Medicine (1998)</p> <p>3. Act 2472/1997 on the Protection of Individuals with Regard to the Processing of Personal Data (As Amended by Laws 2819/2000 and 2915/2000): http://www.dpa.gr/pls/portal/docs/PAGE/APDPX/ENGLISH_INDEX/LEGAL%20FRAMEWORK/LAW%202472-97-NOV2013-EN.PDF</p> <p>4. Act 3418/2005 Code on Medical Ethics</p> | | <p>(Biobanks) in Biomedical Research: http://www.bioethics.gr/images/pdf/ENGLISH/OPINIONS_REPORTS/biobanks_recom_eng.pdf</p> <p>2. Recommendation on the Collection and Use of Genetic Data: http://www.bioethics.gr/images/pdf/ENGLISH/OPINIONS_REPORTS/recom_genetic_data_eng.pdf</p> <p>3. Opinion on Prenatal and Pre-Implantation Diagnosis and Embryo Treatment: http://www.bioethics.gr/images/pdf/ENGLISH/OPINIONS_REPORTS/1_pd_pg_d_opin_eng2.pdf</p> <p>4. Opinion on Direct-To-Consumer Genetic Testing (2012): http://www.bioethics.gr/index.php/en/gnomes/91-direct-to-consumer-dtc-genetic-testing</p> <p>5. Opinion on Incidental Findings in Research and Clinical Practice (2015): http://www.bioethics.gr/images/pdf/GNOMES/OPINION_Incidental_Findings_FINAL_.pdf</p> <p>6. Opinion on Advances in Human Genome Editing (2016): http://www.bioethics.gr/images/pdf/GNOMES/OPINION_gene%20editing_Final_EN.pdf</p> |
| <i>Embryos, Stem Cells, and Cloning</i> | <p>1. National Bioethics Commission (NBC): http://www.bioethics.gr/index.php?category_id=3</p> <p>2. National Authority for Medically Assisted Reproduction</p> | <p>1. Act 2619/98: Ratification of the Convention for the Protection of Human Rights and Dignity of the Human Being with Regard to the Application of Biology and Medicine (1998)</p> <p>2. Civil Code (Act 3089/2002, Medically Assisted Reproduction)</p> <p>3. Act 3305/2005 Application of Medically Assisted Reproduction</p> | | <p>NBC:</p> <p>1. Recommendation on the Use of Stem Cells in Biomedicine and Clinical Medicine</p> <p>2. Recommendation on Human Reproductive Cloning</p> <p>3. Opinion on Prenatal and Pre-implantation Diagnosis and Embryo Treatment</p> <p>Access: http://www.bioethics.gr/index.php/gnomes</p> |
| <p>Hungary Note: All webpages and documents are in Hungarian.</p> | | | | |
| <i>General</i> | <p>1. Ministry of Human Capacities (EMMI): http://www.kormany.hu/hu/emberi-</p> | <p>1. Fundamental Law of Hungary, Updated with the Fifth Amendment (2016), Articles II-</p> | <p>1. Decree 23/2002 (V. 9.) of the Minister of Health on Biomedical Research on Human Beings:</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|---|--|---|------------|
| <i>General</i> | eroforrasok-miniszteriuma 2. Medical Research Council, Research Ethics Committees (KFEB, TUKEB, HRB): https://ett.aek.hu/en/secretariat/ | III: http://njt.hu/cgi_bin/njt_doc.cgi?docid=140968.322953 2. Act CLIV of 1997 on Health Care, Chapters VIII and IX: http://njt.hu/cgi_bin/njt_doc.cgi?docid=30903.339193 3. Act VI. of 2002 on the Promulgation of the Oviedo Convention on Human Rights and Biomedicine: http://njt.hu/cgi_bin/njt_doc.cgi?docid=64201.264663 4. Act LXXXI of 2006 on the Promulgation of the Additional Protocol to the Convention on Human Rights and Biomedicine, Concerning Biomedical Research 5. Act C of 2012 on the Criminal Code, Chapter XVI Medical Procedures and Criminal Offenses Against the Order of Research, Sections 168-175 | http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0200023.EUM&celpara=#xcelparam 2. Decree 35/2005 (VIII.26.) of the Minister of Health on the Clinical Trials of Investigational Medicinal Products for Human Use and on the Application of Good Clinical Practice: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0500035.EUM 3. Decree No. 235/2009 (X.20.) from the Hungarian Government on the Regulations of Giving Permission for Human Medical Experiments, for Clinical Studies of Experimental Medicinal Products, and for the Clinical Studies of the Medical Devices: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0900235.KOR&celpara=#xcelparam | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. National Institute of Pharmacy and Nutrition: http://www.ogvei.gov.hu 2. Medical Research Council, Ethics Clinical Pharmacology Ethics Committee (KFEB): https://ett.aek.hu/kfeb/ | <i>Clinical Trials:</i> Act XCV of 2005 on Medicinal Products for Human Use, Section 3: http://net.jogtar.hu/jr/gen/getdoc.cgi?docid=a0500095.tv&dbnum=62 <i>Non-Interventional Trials:</i> Act CLIV of 1997 on Health Care, Chapter VIII, Section 164/A: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=99700154.TV | <i>Clinical Trials:</i> Decree 35/2005 (VIII. 26) of the Minister of Health on the Clinical Trial and Application of Correct Clinical Practices of Investigational Medicinal Products Intended for Use in Humans: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0500035.EUM&celpara=#xcelparam <i>Non-Interventional Trials:</i> Decree 23/2002. (V. 9) of the Minister of Health on Biomedical Research on Human Beings: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0200023.EUM&celpara=#xcelparam | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|--|---|---|---|
| <i>Drugs, Biologics, and Devices</i> | <p><i>Devices</i></p> <p>1. Authority for Medical Devices, National Healthcare Service System: http://www.enkk.hu/index.php/hun/</p> <p>2. Medical Research Council, Ethics Committee for Clinical Pharmacology: https://ett.aeek.hu/kfeb/</p> | <p>Act CLIV of 1997 on Health Care, Chapter VIII, Section 159: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=99700154.TV</p> | <p><i>Clinical Trials:</i></p> <p>Decree 4/2009. (III. 17.) of the Minister of Health on Medical Devices: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0900004.EUM&celpara=#xcelparam</p> <p><i>Non-Interventional Trials:</i></p> <p>1. Decree 23/2002. (V. 9.) of the Minister of Health on Biomedical Research on Human Beings http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0200023.EUM&celpara=#xcelparam</p> <p>2. Government Decree 235/2009. (X.20.) on the Regulations of Giving Permission for Human Medical Experiments, for Clinical Studies of Experimental Medicinal Products and for the Clinical Studies of the Medical Devices: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0900235.KOR&celpara=#xcelparam</p> <p>3. Government Decree 27/2015 (II.25.) About the National Health Care Service System: http://njt.hu/cgi_bin/njt_doc.cgi?docid=174246.343548</p> | |
| <i>Research Injury</i> | <p>National Institute of Pharmacy and Nutrition: http://www.ogyei.gov.hu</p> | <p>Act XCV of 2005 on Medicinal Products for Human Use, Section 3, Paragraph 5: http://net.jogtar.hu/jr/gen/getdoc.cgi?docid=a0500095.tv&dbnum=62</p> | | |
| <i>Privacy/Data Protection</i> | <p>Hungarian National Authority for Data Protection and Freedom of Information: http://www.naih.hu/general-information.html</p> | <p>1. Act XLVII of 1997 on the Handling of Medical and Other Related Data: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=99700047.TV&celpara=#xcelparam</p> <p>2. Act CXII of 2011 on Right of Informational Self-Determination and Freedom of</p> | | <p>Preparing to Apply the Privacy Policy in 12 Steps: Guidance for Data Controllers and Data Processors (2018): http://www.naih.hu/felkeszueles-az-adatvedelmi-rendelet-alkalmazasara.html</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|---|---|--|---|
| <i>Privacy/Data Protection</i> | | Information: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A1100112.TV&celpara=#xcelparam 3. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | | |
| <i>Human Biological Materials</i> | Ministry of Human Capacities (EMMI): http://www.kormany.hu/hu/emberi-eroforrasok-miniszteriuma | Act LXXX of 2006 on the Promulgation of the Additional Protocol to the Convention on Human Rights and Biomedicine, Concerning Transplantation of Organs and Tissues of Human Origin: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0600080.TV&celpara=#xcelparam | Decree 18/1998 (XII 27) EüM on Implementing Act CLIV of 1997 on Health Care as Regards Transplantation and Storage of Organs and Tissues and Certain Histopathology Examinations: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=99800018.EUM&celpara=#xcelparam | |
| <i>Genetic Research</i> | 1. Ministry of Human Capacities (EMMI): http://www.kormany.hu/hu/emberi-eroforrasok-miniszteriuma 2. Medical Research Council, Committee for Human Reproduction (HRB): https://ett.aEEK.hu/hrb/ | Act XXI of 2008 on the Rules of Protection of Human Genetic Data, of Human Genetic Examinations and Research and of the Operation of Biobanks: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0800021.TV&celpara=#xcelparam | | Decree 60/2003. (X. 20.) of the Minister of Health, Social and Family Affairs on the Minimum Professional Requirements Necessary for Providing Health Services: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0300060.ESC&celpara=#xcelparam |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Ministry of Human Capacities (EMMI): http://www.kormany.hu/hu/emberi-eroforrasok-miniszteriuma 2. Medical Research Council, Research Ethics Committees (KFEB, TUKEB, HRB): https://ett.aEEK.hu/hrb/ | 1. Act CLIV of 1997 on Health Care, Chapter IX 2. Act VI of 2002 on the Promulgation of the Convention on Human Rights and Medicine and the Additional Protocol on Cloning: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=A0200006.TV&celpara=#xcelparam | Decree 30/1998. (VI. 24.) of the Minister of Welfare on Regulations on Specific Procedures for Human Reproduction: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=99800030.NM&celpara=#xcelparam | Decree 18/1998. (XII. 27.) of the Minister of Health on Implementing Act CLIV of 1997 on Health Care Regarding Transplantation and Storage of Organs and Tissues and Certain Histopathology Examinations: http://net.jogtar.hu/jr/gen/hjegy_doc.cgi?docid=99800018.EUM&celpara=#xcelparam |
| Iceland | | | | |
| <i>General</i> | 1. Ministry of Welfare (MOW): http://eng.velferdarraduneyti.is/ 2. National Bioethics Committee (NBC): http://www.vsn.is/en | 1. Act on Scientific Research in the Health Sector No. 44/2014: https://www.government.is/media/velferdarraduneyti-media/media/acrobat-enskar-sidur/Health-Sector-Research-Act-No-44-2014.pdf 3. Oviedo Convention on Human Rights and Biomedicine (2004) | Regulation on the Structure of Research Projects in the Health Sector, Including Research Protocol, Internal Monitoring, and the Responsibilities of the Principal Investigator No. 520/2018: https://www.reglugerd.is/reglugerdir/efrir-raduneytum/velferdarraduneyti/nr/21073 | NBC: 1. Vulnerable Groups Including Children: http://www.vsn.is/en/content/vulnerable-groups-including-children 2. Informed Consent: http://www.vsn.is/en/content/informed-consent 3. Withdrawal of Consent: http://www.vsn.is/en/content/withdrawal-consent 4. Duty to Report Unexpected Events: http://www.vsn.is/en/content/duty-report- |

| Country | Key Organizations | Legislation | Regulations | Guidelines | |
|--------------------------------------|---|---|--|---|---|
| <i>General</i> | | | | unexpected-events 5. Advertising to Recruit Participants: http://www.vsn.is/en/content/advertising-recruit-participants | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | 1. Icelandic Medicines Agency (MCA): http://www.ima.is/ 2. National Bioethics Committee (NBC): www.visindasidanefnd.is | Medicinal Products Act No. 93/1994 (2013): http://eng.velferdarraduneyti.is/acts-of-Parliament/nr/20128 | MCA: Regulation on Clinical Trials of Medicinal Products in Humans No. 443/2004 (2010): http://eng.velferdarraduneyti.is/media/acrobat-enskar_sidur/Medicinal-Products-Act-NoMedicinal-Products-Act-No-93-1994-as-amended.pdf | NBC: Various: http://www.vsn.is/en/content/clinical-trials |
| | <i>Devices</i> | Ministry of Welfare: http://eng.velferdarraduneyti.is/ | Act on Medical Devices No 16/2001 (2011): https://www.government.is/media/velferdarraduneyti-media/media/acrobat-enskar_sidur/Medicinal-Products-Act-NoMedicinal-Products-Act-No-93-1994-as-amended.pdf | 1. Regulation on Medical Devices No. 934/2010 (2010): http://eng.velferdarraduneyti.is/media/acrobat-enskar_sidur/16012012_Act-on-Medical-Devices-No-16-2001-as-amended.pdf 2. Regulation on Active Implantable Medical Devices No. 320/2011: http://www.stjornartidindi.is/Advert.aspx?ID=c50d676c-4651-46c2-83b5-ad946f3deaaa 3. Regulation on In Vitro Diagnostic Medical Devices No. 936/2011: http://stjornartidindi.is/Advert.aspx?ID=f20b3e4e-ab25-44d3-8e32-e5f42a7b02f0 | |
| <i>Research Injury</i> | Icelandic Health Insurance Agency (MCA): http://www.sjukra.is/english | 1. Act on Patient Insurance No. 111/2000 (2011): https://www.government.is/media/velferdarraduneyti-media/media/acrobat-enskar_sidur/Act-on-Patient-Insurance-as-amended.pdf 2. Act on Health Insurance No. 112/2008 (2012): https://www.government.is/media/velferdarraduneyti-media/media/acrobat-enskar_sidur/Act-on-Health-Insurance-No-112-2008-16.pdf | Regulation on Clinical Trials of Medicinal Products in Humans No 443/2004 (2010): https://www.government.is/media/velferdarraduneyti-media/media/Reglugerdir-enska/Regulation-on-clinical-trials-of-medicinal-products-in-humans-no-443-2004-as-amended.pdf | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|--|--|--|--|
| <i>Privacy/Data Protection</i> | Data Protection Authority: http://www.personuvernd.is/information-in-english/ | 1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 2. Act No. 90/2018 on Data Protection and the Processing of Personal Data: https://www.althingi.is/altext/148/s/1296.html | | |
| <i>Human Biological Materials</i> | 1. Ministry of Welfare: http://eng.velferdarraduneyti.is/ 2. National Bioethics Committee (NBC): www.visindasidanefnd.is/en | Biobanks Act No. 110/2000 (2015): https://www.government.is/media/velferdarraduneyti-media/media/acrobat-enskar_sidur/Biobanks-Act-as-amended-2015.pdf | Regulations on the Keeping and Utilization of Biological Samples in Biobanks No. 1146/2010: https://www.reglugerd.is/reglugerdir/efrir-raduneytum/heilbrigdisraduneyti/nr/16910 | NBC: 1. Access to and Utilisation of Health Data and Bio-Samples: http://www.vsn.is/en/content/access-and-utilisation-health-data-and-bio-samples 2. Biobanks: http://www.vsn.is/en/content/biobanks |
| <i>Embryos, Stem Cells, and Cloning</i> | | 1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (2004) 2. Act on Artificial Fertilisation and Use of Human Gametes and Embryos for Stem-Cell Research, No. 55/1996 (2010): http://eng.velferdarraduneyti.is/media/acrobat-enskar_sidur/Act_No_55_1996_on_Artificial_Fertilisation_etc_as_amended.pdf | Regulation on Artificial Fertilization No. 144/2009: https://www.reglugerd.is/reglugerdir/efrir-raduneytum/heilbrigdis/nr/10797 | |
| Ireland | | | | |
| See this summary on Clinical Trials Involving Medical Products: http://health.gov.ie/blog/policy/clinical-trials-involving-medicinal-products/ | | | | |
| <i>General</i> | Department of Health: http://health.gov.ie/ | | | 1. Operational Procedures for Research Ethics Committees: Guidance 2004: http://health.gov.ie/wp-content/uploads/2014/07/Operational_Procedures1.pdf 2. Health Service Executive National Consent Policy, Part 3: http://www.hse.ie/eng/about/Who/qualityandpatientsafety/National_Consent_Policy/ |
| <i>Drugs, Biologics, and Devices</i> | 1. Department of Health: http://health.gov.ie/ 2. Health Products and Regulatory | European Communities (Clinical Trials on Medicinal Products for Human Use) | European Communities (Clinical Trials on Medicinal Products for Human Use) Regulations 2004 | Various: https://www.hpra.ie/homepage/site-tools/search?query=clinical%20trials |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | Authority: https://www.hpra.ie/ | Amendment 2004 (S.I. No. 190 of 2004): http://www.irishstatutebook.ie/eli/2004/si/878/made/en/print | (S.I. No. 190 of 2004): http://www.irishstatutebook.ie/eli/2004/si/190/made/en/html | |
| <i>Research Injury</i> | Health Products and Regulatory Authority: https://www.hpra.ie/ | | European Communities (Clinical Trials on Medicinal Products for Human Use) Regulations 2004, Section 13(6)(k) and Schedule 1, Part 2, Paragraph 4 (S.I. No. 190 of 2004): http://www.irishstatutebook.ie/eli/2004/si/190/made/en/html | |
| <i>Privacy/Data Protection</i> | Data Protection Commissioner (DPC): http://www.dataprotection.ie/docs/Home/4.htm Health Research Board (HRB): http://www.hrb.ie/ | 1. Data Protection Act (1988), as Amended (2003): http://www.irishstatutebook.ie/2003/en/act/pub/0006/index.html 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 3. Data Protection Act 2018: https://www.oireachtas.ie/en/bills/bills/2018/10/ | Data Protection Act 2018 (Section 36(2)) (Health Research) Regulations 2018: http://www.hrb.ie/funding/gdpr-guidance-for-researchers/gdpr-and-health-research/health-research-regulations-2018/ | DPC: 1. 12 Steps to Being Prepared (2018): http://gdprandyou.ie/organisations/ 2. Transfers Abroad (2018): https://www.dataprotection.ie/docs/Transfers-Abroad/y/37.htm HRB: Health Research Regulations 2018 FAQ: http://www.hrb.ie/funding/gdpr-guidance-for-researchers/general-gdpr-faq/ |
| <i>Human Biological Materials</i> | Health Products and Regulatory Authority: https://www.hpra.ie/ | | | Human Biological Material: Recommendations for Collection, Use, and Storage in Research (2005): http://health.gov.ie/wp-content/uploads/2014/07/Human_Biological_Material1.pdf |
| <i>Genetic Research</i> | Health Products and Regulatory Authority: https://www.hpra.ie/ | | | Guidelines for Pharmacogenetic Research (2006): http://lenus.ie/hse/bitstream/10147/96983/1/Pharmacogenetic06.pdf |
| Italy | | | | |
| <i>General</i> | 1. National Bioethics Committee (CNB): http://www.governo.it/bioetica/eng/index.html 2. National Monitoring Center for Clinical Trials (OSS): http://oss-sper-clin.agenziafarmaco.it/ | | OSS: Ministerial Decree of 12 May 2006: Terms of Reference for the Establishment and the Functioning of Ethics Committees | CNB: Various: http://www.governo.it/bioetica/eng/opinions.html |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | | | |
| | 1. National Monitoring Center for Clinical Trials: | 1. Decree of the President of the Republic: Regulations to | 1. Ministerial Decree of 21 December 2007: Directions for | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | http://www.agenziafarmaco.com/en/content/national-monitoring-centre-clinical-trials 2. Italian Medicines Agency: http://www.agenziafarmaco.it/ 3. Ministry of Health (MOH): http://www.ministerosalute.it | Simplify the Procedures and to Verify and Check New Systems and Experimental Therapeutic Protocols (September 21, 2001) (Italian) 2. Legislative Decree No. 211: Transposition of Directive 2001/20/EC Relating to the Implementation of Good Clinical Practice in the Conduct of Clinical Trials on Medicinal Products for Clinical Use (2003) 3. Legislative Decree No. 200: Transposition of Directive 2005/28 EC Laying down Principles and Detailed Guidelines as Regards Investigational Medical Products for Human Use, as Well as the Requirements for Authorizing of Manufacturing or Importing of such Products (2007) (Italian): http://www.aifa.gov.it/allegati/dlgs_200-6nov2007.pdf | Submitting the Request for Authorisation of a Clinical Trial on a Medicinal Product for Human Use to the Competent Authority, for Communicating Substantial Amendments, for Declaring the End of the Trial and for the Request of an Opinion to the Ethics Committee 2. Ministerial Decree of 31 March 2008: Definition of the Minimum Requirements that Contract Research Organisations (CROs) Shall Satisfy in Order to Work within Clinical Trials on Medicinal Products | |
| | <i>Devices</i> | Ministry of Health, Directorate General for Medicines and Medical Devices: http://www.ministerosalute.it | | Ministerial Decree 2 of August 2005: Procedures for the Presentation of Documentation to Notify about Clinical Investigations with Medical Devices |
| <i>Research Injury</i> | Ministry of Labour and Social Policy: www.lavoro.gov.it | | Ministerial Decree 14 of July 2009: Minimum Requirements for Insurance Policies Which Safeguard Participants to Clinical Trials of Medicinal Products | |
| <i>Privacy/Data Protection</i> | Italian Data Protection Independent Authority: http://www.garanteprivacy.it/garante/navi/g/jsp/index.jsp?solotesto=N | 1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 2. Italian Personal Data Protection Code, Legislative Decree No. 196 of June 30, 2003: http://www.garanteprivacy.it/garant | 1. Administrative Decree: Electronic Data Transmission Pertaining to Clinical Medical Experimentation (May 25, 2000) 2. Regulation for the Implementation of Articles No. 20 and 21 of the Legislative | General Principles of Processing Personal Data (2018): https://www.garanteprivacy.it/home/doveri#2 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Privacy/Data Protection</i> | | e/navig/jsp/index.jsp?folderpath=Normativa%2FItaliana%2FII+Codice+in+materia+di+protezione+dei+dati+personali | Decree No. 196 of June 30, 2003 3. Ministerial Decree No. 277 (2007) | |
| <i>Genetic Research</i> | 1. Istituto Superiore di Sanita (ISS): http://www.iss.it/chis/?lang=2 2. Italian Society of Human Genetics (SIGU): http://www.sigu.net/ | | | ISS: Guidelines for Phase I Clinical Trials with Investigational Medicinal Products Employed in Gene Somatic Therapy (2004): http://www.iss.it/binary/publ/publi/0478.1106653420.pdf SIGU: Various: http://www.sigu.net/show/documenti/5/1/linee%20guida |
| <i>Embryos, Stem Cells, and Cloning</i> | | Regulation of Medically Assisted Reproduction, Law No. 40, Article 13 (2004) | | |

| Latvia | | | | |
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| <i>General</i> | Central Medical Ethics Committee | | Statutes of Central Medical Ethics Committees (1998): http://likumi.lv/doc.php?id=46597 | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. State Agency of Medicines: http://www.zva.gov.lv/?setlang=en&large 2. Central Medical Ethics Committee | Law on Pharmacy, Section 26 (2013): https://likumi.lv/ta/en/en/id/203008-law-on-the-rights-of-patients | Cabinet Regulation No. 289: Regulations Regarding the Procedures for Conduct of Clinical Trials and Non-interventional Trials of Medicinal Products, Labelling of Investigational Medicinal Products and the Procedures for Assessment of Conformity of Clinical Trial of Medicinal Products with the Requirements of Good Clinical Practice: https://likumi.lv/ta/en/en/id/207398-regulations-regarding-the-procedures-for-conduct-of-clinical-trials-and-non-interventional-trials-of-medicinal-products-labelling-of-investigational-medicinal-products-and-the-procedures-for-assessment-of-conformity-of-clinical-trial-of- | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <p><i>Drugs, Biologics, and Devices</i></p> | | | <p>medicinal-products-with-the-requirements-of-good-clinical-practice</p> | |
| | <p><i>Devices</i></p> <p>State Agency of Medicines: http://www.zva.gov.lv/?setlang=en&large =</p> | <p>Medical Treatment Law, Section 34 (2014): https://likumi.lv/ta/en/en/id/44108-medical-treatment-law</p> | <p>Cabinet Regulation No. 891: Procedures for the Clinical Trial of Medical Devices Intended for Human Use (2010): https://likumi.lv/ta/en/en/id/218764-procedures-for-the-clinical-trial-of-medical-devices-intended-for-human-use</p> | |
| <p><i>Research Injury</i></p> | <p>State Agency of Medicines: http://www.zva.gov.lv/?setlang=en&large =</p> | | <p><i>Drugs:</i></p> <p>Cabinet Regulation No. 289: Regulations Regarding the Procedures for Conduct of Clinical Trials and Non-interventional Trials of Medicinal Products, Labelling of Investigational Medicinal Products and the Procedures for Assessment of Conformity of Clinical Trial of Medicinal Products with the Requirements of Good Clinical Practice, Sections 22, 31.6, 54.10, 55.9, and 61.14 (2010): https://likumi.lv/ta/en/en/id/207398-regulations-regarding-the-procedures-for-conduct-of-clinical-trials-and-non-interventional-trials-of-medicinal-products-labelling-of-investigational-medicinal-products-and-the-procedures-for-assessment-of-conformity-of-clinical-trial-of-medicinal-products-with-the-requirements-of-good-clinical-practice</p> <p><i>Devices:</i></p> <p>Cabinet Regulation No. 891: Procedures for the Clinical Trial of Medical Devices Intended for Human Use, Sections 42.7 and 62.5 (2010): https://likumi.lv/ta/en/en/id/218764-</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | | | procedures-for-the-clinical-trial-of-medical-devices-intended-for-human-use | |
| <i>Privacy/Data Protection</i> | 1. Data State Inspectorate: http://www.dvi.gov.lv/en/ 2. Central Medical Ethics Committee | 1. Personal Data Processing Law (2014): https://likumi.lv/ta/id/300099-fizisko-personu-datu-apstrades-likums 2. Law on the Rights of Patients, Section 10 (2013): https://likumi.lv/ta/en/en/id/203008-law-on-the-rights-of-patients 3. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | Cabinet Regulation No. 446: Procedures for Using Patient Data in a Specific Research Study (2015): https://likumi.lv/ta/en/en/id/275747-procedures-for-using-the-patient-data-in-a-specific-research | |
| <i>Human Biological Materials</i> | Central Medical Ethics Committee | Law on the Protection of Dead Human Beings and Use of Human Organs and Tissue (2008): https://likumi.lv/ta/en/en/id/62843-on-the-protection-of-the-body-of-deceased-human-beings-and-the-use-of-human-tissues-and-organs-in-medicine | Cabinet Regulation No. 1176 (2013) Procedures for Use of Human Tissues and Cells: http://likumi.lv/ta/id/261810-cilveka-audu-un-sunu-izmantosanas-kartiba | |
| <i>Genetic Research</i> | 1. Ministry of Health: http://www.vm.gov.lv/en/ 2. Data State Inspectorate: http://www.dvi.gov.lv/en/ 3. Central Medical Ethics Committee | 1. Human Genome Research Law (2005): https://likumi.lv/ta/en/en/id/64093-human-genome-research-law 2. Law on the Development and Use of the National DNA Database (2006): https://likumi.lv/ta/en/en/id/90819-law-on-development-and-use-of-the-national-dna-database | Regulation of the Cabinet of Ministers: “Procedures for Genetic Research” (2004): http://likumi.lv/doc.php?id=92330 | |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Ministry of Health: http://www.vm.gov.lv/en/ 2. Central Medical Ethics Committee | Sexual and Reproductive Health Law, Sections 15-20 (2004): https://likumi.lv/ta/en/en/id/58982-sexual-and-reproductive-health-law | Cabinet Regulation No. 1176 (2013) Procedures for Use of Human Tissues and Cells: http://likumi.lv/ta/id/261810-cilveka-audu-un-sunu-izmantosanas-kartiba | |
| Lithuania | | | | |
| Note: All websites and documents are in Lithuanian. | | | | |
| <i>General</i> | Ministry of Health (MOH): http://www.sam.lt/go.php/lit/IMG | 1. Oviedo Convention on Human Rights and Biomedicine (2002): http://conventions.coe.int/treaty/en/treaties/html/164.htm | 1. V-405, Decree on the Procedure for Keeping a Record of Biomedical Research, Collecting, Storage, and Providing Information on | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | | <p>2. Law on Ethics of Biomedical Research (2016): https://www.e-tar.lt/portal/lt/legalAct/TAR.234B15954C2F/wKarWpLPIL</p> <p>3. Changes of Law on Ethics of Biomedical Research No. 536/2014 (2017): https://www.e-tar.lt/portal/lt/legalAct/43db5e50d05f11e7910a89ac20768b0f</p> | <p>Biomedical Research (2010): https://e-seimas.lrs.lt/portal/legalAct/lt/TAD/TAIS.372121</p> <p>2. Government of the Republic of Lithuania: Decree No. 1458 on State Fees (2017): https://www.e-tar.lt/portal/lt/legalAct/TAR.E3A145C8DD49/adJtSaHbRM</p> <p>3. V-15, Decree on the Procedure for Calculating and Paying Compensation for the Expenses Incurred Due to Participation in Biomedical Research and the Time Spent (2018): https://www.e-tar.lt/portal/lt/legalAct/2a0242a0b5fe11e5a6588fb85a3cc84b/ILdhwknYPP</p> <p>4.V-28, Decree on the Detailed Requirements for the Content of a Person's Consent to Participate in Biomedical Research and for the Information about the Biomedical Research as well as a Procedure for Giving and Withdrawing the Consent (2016): https://www.e-tar.lt/portal/lt/legalAct/0f2f1b70b9db11e5a6588fb85a3cc84b</p> <p>5. V-1483, Decree on the List of Interventional Methods of Biomedical Research Causing a Slightly Detrimental and Temporary Impact on the Subject's Health (2014): http://www3.lrs.lt/pls/inter3/dokpaies.ka.showdoc_1?p_id=1002481&p_tr2=2</p> <p>6. V-235/A1-83, Decree on the Procedure for a Minor's Participation in Biomedical Research (2018): https://e-seimas.lrs.lt/portal/legalAct/lt/TAD/8</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | Lithuanian Bioethics Committee (LBEC): http://bioetika.sam.lt/index.php?1608991497 | | 28d53e095ef11e4b92e9028929aad91/QTkdEmVfmU 1. V-28, Decree on the Procedure to Conduct Biomedical Research on Medical Documents, No. V-28 (2011): https://www.e-tar.lt/portal/lt/legalAct/352d55b0c44111e583a295d9366c7ab3/Maiuzzfyns 2. V-7, Decree on the Sample Form of the Biomedical Research Protocol, Summary of the Protocol and the CV of Investigator (2017): https://www.e-tar.lt/portal/lt/legalAct/3790a050be7e11e5a6588fb85a3cc84b 3. V-24, Decree on the Procedure for Submission of the Documents to the Lithuanian Bioethics Committee to Issue Favorable Opinion to Conduct a Clinical Trial on Medicinal Products or Approval to Conduct Biomedical Research by the Sponsor of the Clinical Trial on Medicinal Product or Other Type of Biomedical Research (2016): https://www.e-tar.lt/portal/lt/legalAct/27a3460090f011e4bb408baba2bddd3/UqgJXDRUqi 4. V-4, Decree on the Request to Issue Approval to Conduct Biomedical Research, the Application Form and the Biomedical Research Ethical Assessment Form (2016): https://www.e-tar.lt/portal/lt/legalAct/TAR.EF5F8A | Guidelines for Patient Information Sheet and Informed Consent Form, Adopted by the Group of Experts on Biomedical Research of the LBEC (2017): http://bioetika.sam.lt/index.php?3396441505 |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. Ministry of Health (MOH): http://www.sam.lt/go.php/lit/IMG 2. State Medicines Control Agency (SMCA): | 1. Law on Ethics of Biomedical Research (2016): https://www.e-tar.lt/portal/lt/legalAct/TAR.234B15954C2F/wKarWpLPIL | 1. Decree No. 320 on the Rules of Good Clinical Practice (2004): https://www.e-tar.lt/portal/lt/legalAct/TAR.EF5F8A | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | http://www.vvkt.lt/lit/English | 2. Law on Pharmacy (2017): https://www.e-tar.lt/portal/lt/legalAct/TAR.FF33B3BF23DD/gRoLvrgCbW 3. Changes of Law on Ethics of Biomedical Research (2017): https://www.e-tar.lt/portal/lt/legalAct/43db5e50d05f11e7910a89ac20768b0f | 32B830/WkRbILGNxF 2. Corrections of GCP Terminology in Lithuanian (2006) https://www.e-tar.lt/portal/lt/legalAct/TAR.1C6613E02B96 3. Decree No. 435 on the Procedure for Issuing a Favorable Opinion to Conduct Clinical Trials on Medicinal Product, Approval for Clinical Trials on Medicinal Product, and Conducting and Controlling Clinical Trials (2017): https://e-seimas.lrs.lt/portal/legalAct/lt/TAD/TAIS.277308/QPLLKpOUKw | |
| | Lithuanian Bioethics Committee (LBEC): http://bioetika.sam.lt/index.php?1608991497 | | Decree No. V-6 on the Sample Form of the Request to Issue Favorable Opinion to Conduct Clinical Trial on Medicinal Product Form and the Ethical Assessment Form (2016): https://www.e-tar.lt/portal/lt/legalAct/b65b5ca0c44011e583a295d9366c7ab3/qcrDrSCSCJ | Guidelines to Advertise Clinical Trials, Adopted by the Group of Experts on Biomedical Research of the LBEC (2018): http://bioetika.sam.lt/index.php?3396441505 |
| | <i>Devices</i> | | | |
| | Ministry of Health (MOH): http://www.sam.lt/go.php/lit/IMG | | Decree No. V-2 on the Procedure to Issue Approvals to Conduct Biomedical Research (2017): https://www.e-tar.lt/portal/lt/legalAct/TAR.47B235393D3A/zpczrvbOOR | |
| | State Health Care Accreditation Agency Under the Ministry of Health (SHCA): http://www.vaspvt.gov.lt/en | 1. Law on Ethics of Biomedical Research (2016): https://www.e-tar.lt/portal/lt/legalAct/TAR.234B15954C2F/dReKXfNOaQ 2. Changes of Law on Ethics of Biomedical Research (2017): https://www.e-tar.lt/portal/lt/legalAct/43db5e50d05f11e7910a89ac20768b0f | | |
| <i>Research Injury</i> | Ministry of Health (MOH): http://www.sam.lt/go.php/lit/IMG | 1. Law on Ethics of Biomedical Research (2016): https://www.e-tar.lt/portal/lt/legalAct/TAR.234B15954C2F/wKarWpLPIL 2. Changes of Law on Ethics of | MOH: Decree No. 745 on the Rules of Compulsory Civil Liability Insurance for the Principal Investigator and the Sponsor | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | | Biomedical Research (2017): https://www.e-tar.lt/portal/lt/legalAct/43db5e50d05f11e7910a89ac20768b0f | (2016): https://www.e-tar.lt/portal/lt/legalAct/c86cf490b3be11e598c4c7724bda031b/1aIhDiebov | |
| <i>Privacy/Data Protection</i> | State Data Protection Inspectorate: https://www.ada.lt/go.php/eng | 1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 2. Law on Legal Protection of Personal Data (2018): https://www.e-tar.lt/portal/lt/legalAct/TAR.5368B592234C/VCRurdZydD | | |
| <i>Human Biological Materials</i> | Ministry of Health (MOH): http://www.sam.lt/go.php/lit/IMG | 1. Law on Ethics of Biomedical Research (2016): https://www.e-tar.lt/portal/lt/legalAct/TAR.234B15954C2F/wKarWpLPIL 2. Changes of Law on Ethics of Biomedical Research (2017): https://www.e-tar.lt/portal/lt/legalAct/43db5e50d05f11e7910a89ac20768b0f | | |
| <i>Embryos, Stem Cells, and Cloning</i> | Ministry of Health (MOH): http://www.sam.lt/go.php/lit/IMG | 1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (2002): http://www.coe.int/en/web/conventions/full-list/-/conventions/treaty/168 2. Law on Ethics of Biomedical Research (2016): https://www.e-tar.lt/portal/lt/legalAct/TAR.234B15954C2F/wKarWpLPIL 3. Changes of Law on Ethics of Biomedical Research (2017): https://www.e-tar.lt/portal/lt/legalAct/43db5e50d05f11e7910a89ac20768b0f | 1. Decree No. V-660 on the Procedure to Issue Authorization for the Transit of Tissues of Human Embryonic Tissue, Embryonic Stem Cells and their Lines, Fetal Tissue, and Fetal Stem Cells throughout the Territory of the Republic of Lithuania (2007): https://www.e-tar.lt/portal/lt/legalAct/TAR.8A75E79827FD 2. Decree No. V-659 on the Procedure for Importing of the Stem Cells Taken from the Umbilical Cord or Placenta after the Birth of a Child and the Samples Taken for Genetic Research into the Territory of the Republic of Lithuania and Exporting Therefrom (2017): https://www.e-tar.lt/portal/lt/legalAct/TAR.E2473B1958CA/gEtbNSRzcc | |

Luxembourg

Note: All websites and documents are available in French.

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | National Ethics Consultative Commission: http://www.cne.public.lu/fr/commission/statut.html | | | Various: http://www.cne.public.lu/fr/publications/avis.html |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health: http://www.ms.public.lu and http://www.sante.lu 2. National Research Ethics Committee: http://www.cner.lu 3. Division of Pharmacy and Medicines of the Ministry of Health: http://www.sante.public.lu/fr/politique-sante/ministere-sante/direction-sante/div-pharmacie-medicaments/index.html | Hospitals Act of 1998, Article 25 (2010): http://www.legilux.public.lu/leg/a/archives/1998/0078/a078.pdf | Grand-Ducal Decree of May 30, 2005 on the Conduct of Clinical Trials on Medicinal Products for Human Use: http://www.legilux.public.lu/leg/a/archives/2005/0084/2005A15161.html | |
| <i>Privacy/Data Protection</i> | National Commission for Data Protection: http://www.cnpd.public.lu/fr/index.html | 1. Law of August 2, 2002 on the Protection of Persons with Regard to the Processing of Personal Data as amended by a law of July 27, 2007: http://www.cnpd.public.lu/fr/legislation/droit-lux/doc_loi02082002_en.pdf 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | Grand-Ducal Decree of October 2, 1992 on the Use of Personal Medical Data in IT Processing: http://www.legilux.public.lu/leg/a/archives/1992/0074/a074.pdf#page=12 | Preparation Guide for the New General Data Protection Regulation (2018): https://cnpd.public.lu/fr/dossiers-thematiques/Reglement-general-sur-la-protection-des-donnees/responsabilite-accrue-des-responsables-du-traitement/guide-preparation-rqpd.html |

Macedonia

Note: All websites and documents are available in Macedonian.

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| <i>Drugs and Devices</i> | <i>Drugs</i> 1. Ministry of Health of Republic of Macedonia: http://moh.gov.mk/ 2. Drug and Devices Register: https://lekovi.zdravstvo.gov.mk/ 3. Drug Agency http://malmed.gov.mk/ | 1. Law on Medicinal Products and Medical Devices (Official Gazette No.106/2007) and Laws Amending and Supplementing the Law (2010-2018, according to year of amendment): Click on file folder 1., then open sub-folders: https://lekovi.zdravstvo.gov.mk/documents/2 2. Health Care Law (Official Gazette No. 43/2012) and Laws Amending and Supplementing the Law, Article 275: http://www.fzo.org.mk/default.asp?ItemID=37115BDC6DEF524D877A8C36F95A85F6 | 1. Rulebook on the Manner and the Procedure for Clinical Trials on Medicinal Products and the Documentation Contents (2009): https://lekovi.zdravstvo.gov.mk/documents.documentcomponent:downloadfile/817325622?t:ac=1/1 1.1. Rulebook on Amending and Supplementing the Rulebook on the Manner and the Procedure for Clinical Trials on Medicinal Products and the Documentation Contents (2010): https://lekovi.zdravstvo.gov.mk/documents.documentcomponent:downloadfile/879452170?t:ac=1/1 | 1. Guideline for the Clinical Trial Applicant (Annex 3) (Sub-folder 23.2) (2012): https://lekovi.zdravstvo.gov.mk/documents/1/1 2. Guideline for Good Clinical Practice, Official Gazette No.62/2009, Document No. 19: https://lekovi.zdravstvo.gov.mk/documents/1/1 |
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| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs and Devices</i> | | | <p>1.2. Rulebook on Amending and Supplementing the Rulebook on the Manner and the Procedure for Clinical Trials on Medicinal Products and the Documentation Contents (2012): https://lekovi.zdravstvo.gov.mk/documents.documentcomponent:downloadfile/880033320?t:ac=1/1</p> <p>1.3. Rulebook on Amending and Supplementing the Rulebook on the Manner and the Procedure for Clinical Trials on Medicinal Products and the Documentation Contents and Annex No.3 (Guideline for the Clinical Trial Applicant) (Document No. 23.3) (2012): https://lekovi.zdravstvo.gov.mk/documents/1/1</p> <p>1.4. Rulebook on Supplementing the Rulebook on the Manner and the Procedure for Clinical Trials on Medicinal Products and the Documentation Contents (2016) (Document No. 23.4): https://lekovi.zdravstvo.gov.mk/documents/1/1</p> <p>2. Regulation on the Manner of Reporting, Contents of the Reporting Form for Adverse Reactions to Medicinal Products and the Manner of Organisation of Pharmacovigilance System (2012): https://lekovi.zdravstvo.gov.mk/documents.documentcomponent:downloadfile/880287913?t:ac=1/1</p> | |
| | <p><i>Devices</i></p> <p>1. Ministry of Health of Republic of Macedonia: http://moh.gov.mk/</p> <p>2. Drug and Devices Register: https://lekovi.zdravstvo.gov.mk/</p> <p>3. Drug Agency</p> | <p>Law on Medicinal Products and Medical Devices (Official Gazette No.106/2007) and Laws Amending and Supplementing the Law (2010-2018): Click on file folder 1., then open</p> | <p>1. Rulebook for the Required Documentation and the Method of Application for Clinical Trials on Medical Devices and the Amendments, and Reporting of Drug Adverse Reactions and</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs and Devices</i> | http://malmed.gov.mk/ | sub-folders: https://lekovi.zdravstvo.gov.mk/documents/2 | Events (Official Gazette No. 62/2010): https://lekovi.zdravstvo.gov.mk/documents.documentcomponent:downloadfile/844338380?t:ac=1/2 2. Rulebook on the Manner of Reporting Adverse Effects During the Use of Medical Devices, Types of Reactions they Cause, the Actions of Health Workers and Suppliers, As Well as the Manner of Organizing the System of Monitoring Adverse Effects and Reactions to Medical Devices (Official Gazette No.100/2016) (Document No.8): https://lekovi.zdravstvo.gov.mk/documents/1/2 | |
| <i>Research Injury</i> | 1. Ministry of Health of Republic of Macedonia: http://moh.gov.mk/ 2. Drug Agency: http://malmed.gov.mk/ | | Rulebook on the Manner and the Procedure for Clinical Trials on Medicinal Products and Documentation Contents (2009): https://lekovi.zdravstvo.gov.mk/documents.documentcomponent:downloadfile/817325622?t:ac=1/1 | |
| <i>Privacy/Data Protection</i> | Directorate for Personal Data Protection: www.dzlp.mk | 1. Law on Ratification on Convention for the Protection of Individuals with Regard to Automatic Processing of Personal Data (2005): http://www.dzlp.mk/sites/default/files/pdf/Zakon_za_ratifikacija_na_Konvencijata_108.pdf 2. Law on Ratification on Additional Protocol to the Convention for the Protection of Individuals with Regard to Automatic Processing of Personal Data (2008): http://www.dzlp.mk/sites/default/files/pdf/Dopolnitelen_protokol_Konvencija_108.pdf 3. Law on Personal Data Protection, Consolidated (2016): http://www.dzlp.mk/sites/default/files/u4/ZZLP_konsolidiran_tekst_2016.pdf | Regulations on Protection of Personal Data: http://www.dzlp.mk/mk/podzakonski_akti | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | 1. Ministry of Health of the Republic of Macedonia: http://moh.gov.mk/ 2. Health Insurance Fund of Republic of Macedonia: http://www.fzo.org.mk | 1. Law on Ratification of the Convention for the Protection of Human Rights and Dignity of the Human Being with Regard to the Application of Biology and Medicine, With Additional Protocol on the Prohibition of Cloning Human Beings and Additional Protocol Concerning Transplantation of Organs and Tissues of Human Origin (2009): http://www.pravo.org.mk/document/Detail.php?id=5543 2. Law on Health Protection: (Official Gazette No. 43/2012) and Laws Amending and Supplementing the Law (2012-2016): http://zdravstvo.gov.mk/zakon-za-zdravstvenata-zashtita/ 3. Law on Taking and Transplanting of Human Body Organs (Official Gazette No. 47/2011) and Laws Amending and Supplementing the Law (2011-2016): http://zdravstvo.gov.mk/zakon-za-zemanje-i-presaduvanje-na-delovi-na-chovechkoto-telo-zaradi-lekuvanje/ 4. Sub-Law Acts : http://www.fzo.org.mk/default.asp?ItemID=6541AC10FFC3C5498F0887C57131D996 | Regulations for Transplantation of Tissues and Organs: http://www.fzo.org.mk/default.asp?ItemID=6541AC10FFC3C5498F0887C57131D996 | Regulation on Criteria Relating to Space, Personnel and Equipment for Collection, Transplantation and Exchange of Organs and Tissues, the Necessary Space, Equipment and Staff Required to be Provided by the Health Institution for the Collection, Transfer, Exchange and Storage of Organs and Tissues from Human Body for Treatment Purposes (2012): http://zdravstvo.gov.mk/wp-content/uploads/2012/12/za_pobliskite_kriteriumi_vo_odnos_na_prostorot_kadarot_i_opremata_za_zemawe_presaduvawe_i_razmenuvawe_na_organite_i_tkivata_za_potrebniot_pr.pdf |
| <i>Genetic Research</i> | Ministry of Health of the Republic of Macedonia: http://moh.gov.mk/ | Law on Patient Rights Protections, Article 21: Action on Human Genome (2012): http://zdravstvo.gov.mk/wp-content/uploads/2012/12/zakon-za-zastita-na-pravata-na-pacientite-precisten.pdf | | |
| <i>Embryos, Stem Cells, and Cloning</i> | Ministry of Health of the Republic of Macedonia: http://moh.gov.mk/ | Law on Ratification of the Convention for the Protection of Human Rights and Dignity of the Human Being with Regard | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | | to the Application of Biology and Medicine, With Additional Protocol on the Prohibition of Cloning Human Beings and Additional Protocol Concerning Transplantation of Organs and Tissues of Human Origin (2009): http://www.pravo.org.mk/document/Detail.php?id=5543 | | |
| Malta | | | | |
| <i>General</i> | Bioethics Committee: http://health.gov.mt/en/regcounc/Bioethics-Committee/Pages/CommitteeMembers.aspx | | | Various: http://health.gov.mt/en/regcounc/Bioethics-Committee/Pages/Opinions.aspx |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> Medicines Authority: http://medicinesauthority.gov.mt/ | 1. Medicines Act, 2003: http://justiceservices.gov.mt/DownloadDocument.aspx?app=lom&itemid=8924&l=1 2. Subsidiary Legislation, 458.43, Clinical Trials Regulations, 2004: http://justiceservices.gov.mt/DownloadDocument.aspx?app=lom&itemid=11281&l=1 3. Subsidiary Legislation, 458.47, Good Clinical Practice and Requirements for Manufacturing or Import Authorisation of Investigational Medicinal Products (Human Use) Regulations, 2004: http://justiceservices.gov.mt/DownloadDocument.aspx?app=lom&itemid=11285&l=1 | | Guidance Notes on Good Clinical Practice (2010): http://medicinesauthority.gov.mt/clinicaltrials.htm |
| | <i>Devices</i> 1. Medicines Authority: http://medicinesauthority.gov.mt/ 2. Malta Competition and Consumer Affairs Authority, Technical Regulations Division, Regulatory Affairs Directorate: http://www.mccaa.org.mt/en/regulatory-affairs-directorate | 1. Product Safety Act, 2001: http://justiceservices.gov.mt/DownloadDocument.aspx?app=lom&itemid=8893&l=1 2. Subsidiary Legislation, 427.16, <i>In Vitro</i> Diagnostic Medical Devices Regulations, 2003 | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | http://justiceservices.gov.mt/DownloadDocument.aspx?app=lom&itemid=10756&l=1 3. Subsidiary Legislation, 427.44, Medical Devices Regulations, 2010: http://justiceservices.gov.mt/DownloadDocument.aspx?app=lom&itemid=10781&l=1 4. Subsidiary Legislation, 427.10, Active Implantable Medical Devices Regulations, 2010: http://justiceservices.gov.mt/DownloadDocument.aspx?app=lom&itemid=10753&l=1 | | |
| <i>Privacy/Data Protection</i> | Office of the Information and Data Protection Commissioner: http://idpc.gov.mt/index.aspx | 1. Data Protection Act, 2002: http://justiceservices.gov.mt/DownloadDocument.aspx?app=lom&itemid=8906&l=1 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | | |

Moldova

For an overview of human subject protections in Moldova, see “Ethical Review of Biomedical Research in the CIS Countries,” Chapter 3, Section 7:

http://www.unesco.org/new/fileadmin/MULTIMEDIA/FIELD/Moscow/pdf/ethical_review_cis_book_kubar_english.pdf

Note: All websites and documents are in Moldovian.

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| <i>General</i> | Ministry of Health, National Committee for Ethical Expertise of Clinical Trials: http://ms.gov.md/?q=comitetul-national-etica | Oviedo Convention on Human Rights and Biomedicine (2002) | | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health , National Committee for Ethical Expertise of Clinical Trials: http://ms.gov.md/?q=comitetul-national-etica 2. Medicines and Medical Devices Agency: http://www.amed.md/ | 1. Law No. 1409 Dated 17.12.1997 on Medicines, Articles 11 and 12: http://lex.justice.md/index.php?action=view&view=doc&lang=1&id=311586 2. Law No. 263 Dated 27.10.2005 on Patients’ Rights and Responsibilities. Articles 9, 10, 11, 12, 13, and 14: http://lex.justice.md/index.php?action=view&view=doc&lang=1&id=313060 | MOH: 1. Government Decision No. 5/18.01.2016 Regarding the National Committee for Ethical Expertise of Clinical Trial: http://lex.justice.md/md/362783/ 2. Order No.648/12.08.2016 Concerning the Regulation of Authorizing the Conduct of Clinical Trials in the Republic of Moldova: http://amed.md/sites/default/files/Legislatie/Ordine%20ale%20MS/Ordinul%20MS%20nr.%20648.pdf | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|-----------------------------------|--|---|---|-------------------|
| <i>Research Injury</i> | Ministry of Health (MOH): http://www.ms.gov.md/ | Law No. 411-XIII Dated 28.03.1995 on Health: http://lex.justice.md/viewdoc.php?action=view&view=doc&id=312823&lang=1 | 1. Government Decision No. 5/18.01.2016 Regarding the National Committee for Ethical Expertise of Clinical Trials: http://lex.justice.md/md/362783/ 2. Order No. 648/12.08.2016 Concerning the Regulation of Authorizing the Conduct of Clinical Trials in the Republic of Moldova: http://amed.md/sites/default/files/Legislatie/Ordine%20ale%20MS/Ordinul%20MS%20nr.%20648.pdf | |
| <i>Privacy/Data Protection</i> | National Center for Personal Data Protection of the Republic of Moldova: http://www.datepersonale.md/en/start/ | 1. Convention No. 108 for the Protection of Individuals with Regard to Automatic Processing of Personal Data (1981): http://datepersonale.md/en/international003/ 2. Decision of Parliament No. 483-XIV Dated 02.07.1999 on Ratification of Convention No. 108: http://lex.justice.md/index.php?action=view&view=doc&lang=1&id=309121 3. Law No. 982 Dated 11.05.2000 on Access to Information: http://lex.justice.md/index.php?action=view&view=doc&lang=1&id=311759 4. Law No.133 Dated 08.07.2011 on the Protection of Personal Data: http://lex.justice.md/index.php?action=view&view=doc&lang=1&id=340495 5. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 6. LP143 Din 19.07.18, MO309-320/17.08.18 Article 482 | Decision of Government No. 1123 Dated 14.12.2010 on the Approval of the Requirements for the Assurance of Personal Data Security at their Processing within the Information Systems of Personal Data: http://www.datepersonale.md/file/hotariri/cerinte_securitate%20eng_101228.pdf | |
| <i>Human Biological Materials</i> | 1. Ministry of Health (MOH): http://www.ms.gov.md/ 2. Transplant Agency | 1. Law No. 42 Dated 06.03.2008 on Transplantation of Organs, Tissues and Human Cells: | MOH: Order No.648/12.08.2016 Concerning the Regulation of | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | http://lex.justice.md/md/334622 | http://lex.justice.md/index.php?action=view&view=doc&lang=1&id=327709 2. LP79 Din 24.05.18, MO195-209/15.06.18 Article 338 | Authorizing the Conduct of Clinical Trials in Republic of Moldova: http://amed.md/sites/default/files/Legislatie/Ordine%20ale%20MS/Ordinul%20MS%20nr.%20648.pdf | |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Ministry of Health (MOH): http://www.ms.gov.md/ 2. National Commission on Biological Security: http://lex.justice.md/index.php?action=view&view=doc&lang=1&id=303353 | 1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being, on the Prohibition of Cloning Human Beings (2002) 2. Law No. 42 Dated 06.03.2008 on Transplantation of Organs, Tissues and Human Cells: http://lex.justice.md/index.php?action=view&view=doc&lang=1&id=327709 3. LP79 Din 24.05.18, MO195-209/15.06.18 Article 338 | | |
| Montenegro | | | | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health of Montenegro: http://www.mzd.gov.me/en/ministry?alphabet=lat 2. Agency for Medicines and Medical Devices: https://www.calims.me/Portal/faces/glavna?_adf.ctrl-state=rsbe35pln_83 | 1. Law on Medicines (“Official Gazette of Montenegro”, No. 56/2011 and 06/13): https://www.calims.me/Portal/faces/servlet?putanja=CG_Zakon_o_ljekovima.pdf&_afWindowMode=0&_afLoop=3654755254077715&_adf.ctrl-state=13nzchbscd_171 2. Law on Medical Devices (“Official Gazette of Montenegro” No. 79/2004, 53/09, and 40/11): https://www.calims.me/Portal/faces/servlet?putanja=CG-Zakon%2520o%2520medicinski%2520sredstvima.pdf&_afWindowMode=0&_afLoop=3654994298177994&_adf.ctrl-state=13nzchbscd_181 | Rulebook on More Detailed Conditions and Documentation Required for Approval and Conduct of Clinical Trials of Medicines for Human Use (2013): https://www.calims.me/Portal/faces/servlet?_afLoop=26656243505641585&_afWindowMode=0&putanja=Rulebook%2520on%2520Clinical%2520trials.pdf&_adf.ctrl-state=wdqo8wvwo_214 | |
| <i>Research Injury</i> | 1. Ministry of Health of Montenegro: http://www.mzd.gov.me/en/ministry?alphabet=lat 2. Agency for Medicines and Medical Devices: https://www.calims.me/Portal/faces/glavna | 1. Law on Medicines (“Official Gazette of Montenegro”, No. 56/2011 and 06/13): https://www.calims.me/Portal/faces/servlet?putanja=CG_Zakon_o_ljekovima.pdf&_afWindowMode=0&_afLoop=3654755254077715&_adf.ctrl-state=13nzchbscd_171 | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | a?_adf.ctrl-state=rsbe35pln_83 | afrLoop=3654755254077715&_adf.ctrl-state=13nzchbscd_171 2. Law on Medical Devices (“Official Gazette of Montenegro” No. 79/2004, 53/09, and 40/11): https://www.calims.me/Portal/faces/servlet1?putanja=CG-Zakon%2520o%2520medicinskim%2520sredstvima.pdf&_afrWindowMode=0&_afrLoop=3654994298177994&_adf.ctrl-state=13nzchbscd_181 | | |
| <i>Privacy/Data Protection</i> | National Security Agency: http://www.anb.gov.me/en/Home?alphabet=lat | Law on the Protection of Personal Data (Official Gazette of Montenegro No. 79/08, 70/09, 44/12): http://www.azlp.me/docs/zajednicka/zakoni/zakon-o-zastiti-podataka-o-licnosti.pdf | | |
| <i>Human Biological Materials</i> | Ministry of Health of Montenegro: http://www.mzd.gov.me/en/ministry?alphabet=lat | Law on the Collection and Use of Biological Samples (Official Gazette of Montenegro No. 14/2010): http://www.mzdravlja.gov.me/ResourceManager/FileDownload.aspx?rid=57491&rType=2&file=ZAKON%20O%20UZIMANJU%20I%20KORI%20C5%A0%C4%86ENJU%20BIOLO%20C5%A0KIH%20UZORAKA.pdf | | |
| <i>Genetics</i> | Ministry of Health of Montenegro: http://www.mzd.gov.me/en/ministry?alphabet=lat | Law on the Protection of Genetic Data (Official Gazette of Montenegro No. 25/2010): http://www.mzdravlja.gov.me/ResourceManager/FileDownload.aspx?rid=57566&rType=2&file=ZAKON%20O%20ZA%20C5%A0TITI%20GENETI%20C4%8CKIH%20PODATAKA%20.pdf | | |
| <i>Embryos, Stem Cells, and Cloning</i> | Ministry of Health of Montenegro: http://www.mzd.gov.me/en/ministry?alphabet=lat | | Rulebook on the Collection, Storage, and Use of Stem Cells (2012): http://www.mzdravlja.gov.me/ResourceManager/FileDownload.aspx?rid=222783&rType=2&file=Pravilnik%20o%20postupku%20prikupljanja.%20 | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | | | %C4%8Duvanja%20i%20upotrebe%20mati%C4%8Dnih%20%C4%87elija%2056-2012.pdf | |
| Netherlands | | | | |
| <i>General</i> | Central Committee for Research Involving Human Subjects (CCMO): http://www.ccmo.nl/en/ | 1. Population Screening Act (1996): http://wetten.overheid.nl/BWBR0005699/geldigheidsdatum_24-09-2015 2. Medical Research Involving Human Subjects Act (2012) 2006 English version: http://www.ccmo.nl/attachments/files/wmo-engelse-vertaling-29-7-2013-afkomstig-van-vws.pdf | 1. Concerning the Use of a Special Form (2002) 2. Concerning Requirements of Expertise of Accredited Review Board Members (2002) 3. Concerning the Organization and Working Method of Accredited Review Board Members (2003) 4. External Review Guideline (2004) 5. Research Contract Review Guideline (2009) | Various: http://www.ccmo.nl/en/publications-of-the-ccmo |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health, Welfare, and Sport (VWS): http://www.government.nl/ministries/vws/#ref-minvws 2. Central Committee for Research Involving Human Subjects (CCMO): http://www.ccmo.nl/en/ 3. Medicines Evaluation Board (MEB): http://english.cb-g-meb.nl/ | Medicines Act (2007): http://wetten.overheid.nl/BWBR0021505 | VWS: 1. Medicines Act Decree (2007): http://www.ccmo.nl/attachments/files/eng-decree-on-scientific-research-with-medicinal-products.pdf 2. Medicines Act Regulation (2007): http://wetten.overheid.nl/BWBR0022160 | CCMO: Clinical Research with Medicinal Products in the Netherlands: Instructional Manual (2005): http://www.vumc.nl/afdelingen-themas/1646433/7876770/7876776/7955410/Clinical_research_with_medi1.pdf |
| <i>Clinical Trials Registry</i> | 1. Netherlands Trial Register: http://www.trialregister.nl/trialreg/index.asp 2. Central Committee Register (Dutch): https://www.toetsingonline.nl/to/ccmo_research.nsf/Searchform?OpenForm | | | |
| <i>Research Injury</i> | Ministry of Health, Welfare and Sport: http://www.government.nl/ministries/vws/#ref-minvws | Medical Research Involving Human Subjects Act, Article 7 (2006): http://www.ccmo.nl/attachments/files/wmo-engelse-vertaling-29-7-2013-afkomstig-van-vws.pdf | Regulation on Mandatory Insurance Regarding Medical Research Involving Human Subjects (2003): https://zoek.officielebekendmakingen.nl/stb-2014-477.html | |
| <i>Social-Behavioral Research</i> | National Ethics Council for Social and Behavioural Sciences: http://www.nethics.nl/ | | | Ethical Code (2018): http://www.nethics.nl/Gedragscode-Ethical-Code/ |
| <i>Privacy/Data Protection</i> | 1. Dutch Data Protection Authority: https://cbpweb.nl/en 2. Central Committee for Research | 1. Personal Data Protection Act (2004): http://wetten.overheid.nl/BWBR001 | | CCMO: 1. General Data Protection Regulation (2018): http://www.ccmo.nl/en/algemene- |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Privacy/Data Protection</i> | Involving Human Subjects (CCMO): http://www.ccmo.nl/en/ 3. Federation of Biomedical Scientific Societies (FMWV): http://www.federa.org/ | 1468 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | | verordening-gegevensbescherming?5ad0a79c-a970-44d7-8c78-6de7c35ff8ba 2. Adaptations to the Trial Information Form Due to New European Privacy Legislation (2018) FMWV: 1. Code for Adequate Secondary Use of Data (2004): http://www.federa.org/sites/default/files/bijlage_n/coreon/code_of_conduct_for_medical_research_1.pdf 2. Explanatory Report Accompanying the Code (2004): http://www.federa.org/sites/default/files/bijlage_n/coreon/explanatory_report1.pdf |
| <i>Human Biological Materials</i> | Federation of Biomedical Scientific Societies (FMWV): http://www.federa.org/ | Civil Code, Article 467 (1994): http://www.ccmo.nl/attachments/files/wgbo-pdf.pdf | | Code for Proper Secondary Use of Human Tissue in the Netherlands (2002): http://www.federa.org/sites/default/files/bijlage_n/coreon/codepropersecondaryuseofhumantissue1_0.pdf |
| <i>Genetic Research</i> | 1. Ministry of Infrastructure and the Environment (IenM): http://www.government.nl/ministries/ienm 2. Dutch Health Care Inspectorate (IGZ): http://www.igz.nl/english/ 3. Central Committee for Research Involving Human Subjects (CCMO): http://www.ccmo.nl/en/ | Medical Research Involving Human Subjects Act (2006): http://www.ccmo.nl/attachments/files/wmo-engelse-vertaling-29-7-2013-afkomstig-van-vws.pdf | | IenM, VWS, and CCMO: Guidelines for Researchers and Sponsors with Regard to the Assessment by Official Bodies of Clinical Research Involving Gene Therapeutics in the Netherlands (2012): http://www.ggo-vergunningverlening.nl/dsresource?type=pdf&objectid=rivmp:193539&versionid=&subobjectname= |
| <i>Embryos, Stem Cells, and Cloning</i> | Central Committee for Research Involving Human Subjects (CCMO): http://www.ccmo.nl/en/ | 1. Foetal Tissue Act (2001) (Dutch): http://wetten.overheid.nl/BWBR0012983/ 2. Embryos Act (2002): http://www.ccmo.nl/attachments/files/embryos-act.pdf | | |
| Norway | | | | |
| <i>General</i> | National Committee for Medical and Health Research Ethics (NEM): http://www.etikkom.no/en/In-English/Committee-for-Medical-and-Health-Research/ | 1. Oviedo Convention on Human Rights and Biomedicine (2006) 2. Law regarding Ethics and Integrity in Research (2006): http://www.ub.uio.no/ujur/ulovdata/lov-20060630-056-eng.pdf 3. Act on Health Care Research | | 1. Guidelines for Research on Persons with Impaired Informed Consent Capacity (2005) 2. Payment for Research Participants in Medical and Health Research (2009) 3. Guidelines for Research Ethical and Scientific Evaluation of Qualitative |

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| <i>General</i> | | (2008): http://www.lovdata.no/cgi-wift/wiftldles?doc=/usr/www/lovdata/all/nl-20080620-044.html&emne=helseforskningslov*& | | Research Projects in Medical and Health Research (2009): https://www.etikk.no/forskningsetiske-retningslinjer/Medisin-og-helse/Kvalitativ-forskning/ |
| | National Committee for Research Ethics in the Social Sciences and the Humanities: http://www.etikk.no/en/In-English/ | | | Guidelines for Research Ethics in the Social Sciences, Law, and the Humanities (2001) |
| | National Committee for Research Ethics in Science and Technology: https://www.etikk.no/en/In-English/Committee-for-Research-Ethics-in-Science-and-Technology/ | | | Research Ethics Guidelines for Science and Technology (2007): https://www.etikk.no/Forskningsetikk/Etiske-retningslinjer/Naturvitenskap-og-teknologi/ |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | | | |
| | Norwegian Medicines Agency: http://www.legemiddelverket.no/English/Sider/default.aspx | | Regulation Relating to Clinical Trials on Medicinal Products for Human Use (2009): http://lovdata.no/dokument/SF/forskrift/2009-10-30-1321?q=forskrift+om+kliniske+utpr%C3%B8vning | Guidelines for the Regulations Concerning Clinical Trials of Human Drugs (1999): http://www.legemiddelverket.no/Godkjenning-og-regelverk/Klinisk-utproving/Regelverk%20og%20veiledninger/Documents/Veiledning%20-%20revidert%20versjon%202.2%2006.11.2012.pdf |
| | <i>Devices</i> | | | |
| 1. Norwegian Directorate of Health: http://www.helsedirektoratet.no/kvalitet-planlegging/medisinsk-utstyr/klinisk-utprovning/Sider/default.aspx 2. Regional Committees for Medical and Health Research Ethics: https://helseforskning.etikk.no/ikbVier/page/forside | Act of 12 January 1995 No. 6 Relating to Medical Devices (1995): http://lovdata.no/dokument/NL/lov/1995-01-12-6?q=lov+om+medisinsk+utstyr | Regulation of December 15th 2005 No. 1690 Relating to Medical Devices (2005): http://lovdata.no/dokument/SF/forskrift/2005-12-15-1690?q=forskrift+medisinsk+utstyr | Guidelines on Notification for Clinical Investigation of Medical Devices in Norway (2010): https://helsedirektoratet.no/Documents/Medisinsk%20utstyr/Guidance%20for%20completing%20the%20Notification%20form.pdf | |
| <i>Research Injury</i> | | Convention on Human Rights and Biomedicine (Convention of Oviedo), Article 24, ETS No. 164 (2007) | | |
| <i>Social-Behavioral Research</i> | National Committee for Research Ethics in the Social Sciences and the Humanities | | | Guidelines for Research Ethics in the Social Sciences, Law, and the Humanities (2016): https://www.etikk.no/globalassets/documents/english-publications/60127_fek_guidelines_nesh_digital_corr.pdf |
| | Norwegian National Research Ethics Committees | | | Ethical Guidelines for Internet Research (2014): https://www.etikk.no/en/ethical- |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Social-Behavioral Research</i> | | | | guidelines-for-research/ethical-guidelines-for-internet-research/ |
| <i>Privacy/Data Protection</i> | Data Inspectorate: http://www.datatilsynet.no/English | 1. Personal Data Act No. 31 (2000): http://lovdata.no/dokument/NL/lov/2000-04-14-31 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | Regulations on the Processing of Personal Data (2003) | |
| <i>Human Biological Materials</i> | 1. Ministry of Health and Care Services (MHCS): https://www.regjeringen.no/en/dep/hod/id421/ 2. Ministry of Education and Research (MER): http://www.regjeringen.no/en/dep/kd.html?id=586 | 1. Act on Biobanks (February 21, 2003, No. 12): http://lovdata.no/dokument/NL/lov/2003-02-21-12?q=biobank 2. Act Relating to the Application of Biotechnology in Human Medicine, etc. (December 5, 2003, No. 100) 3. Act on Health Care Research (2008): http://www.lovdata.no/cgi-wift/wiftldles?doc=/usr/www/lovdat a/all/nl-20080620-044.html&emne=helseforskningslov*&& | MHCS: Guidelines for the Norwegian Act on Biobanks (2003) | |
| <i>Genetic Research</i> | 1. Ministry of Health and Care Services (MHCS): https://www.regjeringen.no/en/dep/hod/id421/ 2. Norwegian Biotechnology Advisory Board: http://www.bion.no/english/ 3. Regional Committees for Medical Research Ethics (REK): https://www.etikkom.no/en/In-English/Committee-for-Medical-and-Health-Research/ | Act Relating to the Application of Biotechnology in Human Medicine, Etc. (December 5, 2003, No. 100): https://www.regjeringen.no/globalassets/upload/kilde/hod/red/2005/0081/ddd/pdfv/242718-biotechnology_act_master.pdf | | |
| <i>Embryos, Stem Cells, and Cloning</i> | Directorate for Health and Social Affairs: http://www.helsedirektoratet.no/kvalitet-planlegging/biogenteknologi/Sider/default.aspx | 1. Revised Act Relating to the Application of Biotechnology in Human Medicine (June 15, 2007) Regarding Changes in the Act Related to Stem Cell Research and Pre-implantation Diagnostics (2007) 2. Norwegian Law on the Human-Medical Use of Biotechnology, Chapter 3 | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Poland | | | | |
| <i>General</i> | <p>1. Ministry of Health, Bioethics Appeals Commission (MOH) Bioethics Appeals Commission (MOH): https://www.gov.pl/zdrowie/odwolawcza-komisja-bioetyczna</p> <p>2. Center of Bioethics, Polish Chamber of Physicians and Dentists (NIL): http://www.nil.org.pl/dzialalnosc/orodek-bioetyki</p> | <p>1. Constitution of the Republic of Poland, Article 39 (1997): http://www.sejm.gov.pl/prawo/konsult/polski/kon1.htm</p> <p>2. Medical Profession Act, Articles 21-29 (1997): http://isap.sejm.gov.pl/Download?id=WDU19970280152&type=3</p> | <p>MOH: Order of the Minister of Health and Social Welfare on How to Establish, Finance, and the Mode of Action of Bioethics Committees (1999): http://isap.sejm.gov.pl/DetailsServlet?id=WDU19990470480</p> | <p>NIL: Code of Medical Ethics, Chapter II (2003): http://www.nil.org.pl/dokumenty/kodeks-etyki-lekarskiej</p> |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | <p>1. Pharmaceutical Law (2017): http://prawo.sejm.gov.pl/isap.nsf/DocDetails.xsp?id=WDU20170002211</p> <p>2. Law of 20/04/2004 on Amendment of the Pharmaceutical Law, L, and Regulations</p> | <p>Decree of the Minister of Health on Clinical Trials on Minors (2004)</p> | |
| | <i>Devices</i> | <p>1. Act on Medical Devices: http://isap.sejm.gov.pl/DetailsServlet?id=WDU20170000211&min=1</p> | <p>1. Regulation of the Minister of Health on Detailed Conditions to be Met for Clinical Evaluation of Medical Devices or Active Implantable Medical Devices (2011): http://prawo.sejm.gov.pl/isap.nsf/DocDetails.xsp?id=WDU20110630331</p> <p>Various: http://www.urpl.gov.pl/pl/wyroby-medyczne/akty-prawne/przepisy-rp</p> | |
| <i>Research Injury</i> | | <p>Pharmaceutical Law, Chapter 36b: http://isap.sejm.gov.pl/DetailsServlet?id=WDU20160002142</p> | <p>1. Order of the Minister of Finance Concerning the Mandatory Civil Liability Insurance of Researchers and Sponsors (2004): http://isap.sejm.gov.pl/DetailsServlet?id=WDU20041011034</p> <p>2. Order of the Minister of Finance Amending the Regulation Concerning the Mandatory Civil Liability Insurance of Researchers and Sponsors (2005):</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|-----------------------------------|---|--|---|------------|
| <i>Research Injury</i> | | | http://isap.sejm.gov.pl/DetailsServlet?id=WDU20051010845 3. Order of the Minister of Finance Concerning the Mandatory Civil Liability Insurance of Researchers and Sponsors in Clinical Trials of Medicinal Products (2010): http://isap.sejm.gov.pl/DetailsServlet?id=WDU20101941290 | |
| <i>Privacy/Data Protection</i> | Personal Data Protection Office: https://uodo.gov.pl/en | 1. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj 2. Act on the Protection of Personal Data (2018): https://uodo.gov.pl/pl/131/262 | | |
| <i>Human Biological Materials</i> | | 1. Act of 22 August 1997 on the Public Blood Service: http://prawo.sejm.gov.pl/isap.nsf/DocDetails.xsp?id=WDU20170001371 2. July 1, 2005 Act Regarding Sampling, Storage, and Transplanting of Cells, Tissues, and Organs: http://prawo.sejm.gov.pl/isap.nsf/DocDetails.xsp?id=WDU20170001000 | | |

| Portugal | | | | |
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| <i>General</i> | National Council of Ethics for the Life Sciences: http://www.cnecev.gov.pt/cnecev/en/ | Oviedo Convention on Human Rights and Biomedicine (2001) | | Various: http://www.cnecev.gov.pt/cnecev/en/opinions/ |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. National Institute of Pharmacy and Medicines: http://www.infarmed.pt/portal/page/portal/INFARMED/ENGLISH 2. Ethics Commission for Clinical Research (CEIC): http://www.infarmed.pt/portal/page/portal/INFARMED/MEDICAMENTOS_USO_HUMANO/CEIC | 1. Approval of the Applicable Legal Standards for the Conduct of Clinical Trials of Medicines for Human Use, Law No. 46/2004 2. Approval of the Composition, Operations, and Financing of the Ethics Commission for Clinical Research, Decree No. 57/2005: http://www.infarmed.pt/portal/page/portal/INFARMED/LEGISLACAO/LEGISLACAO_FARMACEUTICA_COMPILADA/TITULO_III/TITU | Decree-Law No. 102/2007 of April 2 | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | LO III CAPITULO I/portaria 57-2005.pdf | | |
| | <i>Devices</i> National Institute of Pharmacy and Medicines: http://www.infarmed.pt/portal/page/portal/INFARMED/DISPOSITIVOS_MEDICOS | Various: http://www.infarmed.pt/portal/page/portal/INFARMED/LEGISLACAO/LEGISLACAO_FARMACEUTICA_COMPILADA/TITULO_V/TITULO_V_CAPITULO_II | | Various: http://www.infarmed.pt/portal/page/portal/INFARMED/DISPOSITIVOS_MEDICOS/NOTAS_INFORMATIVAS |
| <i>Research Injury</i> | | Convention on Human Rights and Biomedicine (Convention of Oviedo), Article 24, ETS No. 164 (2001) | | |
| <i>Privacy/Data Protection</i> | National Data Protection Commission: http://www.cnpd.pt/english/index_en.htm | 1. Constitution, Article 35 (1997) 2. Act on the Protection of Personal Data, No. 67/98 (1998): http://www.cnpd.pt/english/bin/legislation/Law6798EN.HTM 3. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | | FAQs: Consent (2018): https://www.cnpd.pt/bin/faqs/faqs.htm#consentimento |
| <i>Genetic Research</i> | Ministry of Health: http://www.portugal.gov.pt/en/the-ministries/ministry-of-health.aspx | Law 12/2005 | | |
| <i>Embryos, Stem Cells, and Cloning</i> | National Council of Ethics for the Life Sciences: http://www.cnecev.gov.pt/cnecev/en/ | 1. Oviedo Convention on Human Rights and Biomedicine, Additional Protocol on Prohibition of Human Cloning (2001) 2. Portuguese Law on Assisted Reproductive Technologies, Articles 7 and 9 (2006) | | 1. Opinion 15/CNECV/95 on Embryo Research (1995) 2. Opinion 47/CNECV/2005 on Stem Cell Research (2005): http://www.cnecev.gov.pt/NR/rdonlyres/F13B34FD-F9F7-4C9D-96DC-419999D9B693/0/47CNECV2005.pdf 3. Opinion 48/CNECV/2006 on Human Cloning (2006): http://www.cnecev.gov.pt/NR/rdonlyres/770EA390-9326-4FF9-B28D-D70A7E9AD961/0/p048_en.pdf |
| Romania | | | | |
| <i>General</i> | Ministry of Health (MOH): http://www.ms.ro/ | Oviedo Convention on Human Rights and Biomedicine (2001) | Ordinance No. 57/16.08.2002 (2002): http://www.research.ro/ro/articol/1021/despre-ancs-legislatie | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health (MOH): http://www.ms.ro/ 2. National Agency for Medicines and | | MOH: Order 904/25July 2006 on Approval of Rules Relating to | MOH: Guideline for Clinical Trials in Pediatric Populations (CPMP/ICH/2711/99) (1999) |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | Medical Devices: https://www.anm.ro/en/ 3. National Bioethics Committee for Medicines and Medical Devices: http://www.bioetica-medicala.ro/ | | the Implementation of Good Clinical Practice in the Conduct of Clinical Trials on Medicinal Products for Human Use -- Transposition of 2001/20/EC Directive Access: https://www.anm.ro/en/medicament-de-uz-uman/legislatie/legi-ordonante-si-hotarari-de-guvern/ | |
| <i>Research Injury</i> | 1. National Agency for Medicines and Medical Devices: http://www.anm.ro/anmdm/en/index.html 2. National Bioethics Committee for Medicines and Medical Devices: http://www.bioetica-medicala.ro/ | Convention on Human Rights and Biomedicine (Convention of Oviedo), Article 24, ETS No. 164 (2001) | | |
| <i>Privacy/Data Protection</i> | National Supervisory Authority for Personal Data Processing: http://www.dataprotection.ro/index.jsp?page=documents&lang=en | 1. Law No. 667/2001 On the Protection of Individuals with Regard to the Processing of Personal Data and on the Free Movement of Such Data: http://www.dataprotection.ro/servlet/ViewDocument?id=174 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | | The New General Data Protection Regulation (2018): http://www.dataprotection.ro/?page=Regulamentul_nr_679_2016 |
| <i>Human Biological Materials</i> | Ministry of Health (MOH): http://www.ms.ro/ | Law No. 95/2006 Regarding the Reform in Health Field. Title VI. Performing of Sampling and Transplant of Organs, Tissues and Human Origin Cells with Therapeutic Purpose: http://www.transplant.ro/Lege/Lege-2006-95.pdf | 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: http://europa.eu/legislation_summaries/public_health/threats_to_health/sp0008_ro.htm | |
| <i>Embryos, Stem Cells, and Cloning</i> | | 1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with Regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (2001) 2. Law No. 301 from 2004 Penal | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | | Code – Chapter IV – Crimes and Felonies Regarding Genetic Manipulation: http://www.codpenal.ro/legislatie/document/lege-301-din-2004-codul-penal-capitol-4-crime-si-delicte-privind-manipularea-genetica-1260-63259.html | | |
| Russia For an overview of human subject protections in Russia, see http://www.unesco.org/new/fileadmin/MULTIMEDIA/FIELD/Moscow/pdf/ethical_review_cis_book_kubar_english.pdf | | | | |
| <i>General</i> | 1. Ministry of Healthcare of the Russian Federation (MOH): http://www.rosminzdrav.ru 2. Federal Service on Surveillance in Healthcare (Roszdravnadzor): http://www.roszdravnadzor.ru/ 3. Russian Committee for Bioethics: http://www.bioethics.ru/eng/ | 1. Constitution of the Russian Federation, Article 21 (1993): http://www.constitution.ru/en/10003000-03.htm 2. Federal Law #FZ 323 “On Foundations of Protection of Citizen’s Health in the Russian Federation” (2011): http://acto-russia.org/en/index.php?option=com_content&task=view&id=105 3. Federal Law #FZ55 “On Introduction of Changes in FZ “On Foundations of Protection of Citizens’ Health in the Russian Federation” with Regard to Questions of Organization of Medical Aid Administered in the Course of Clinical Approbation of the Methods of Prevention, Diagnostics, Treatment and Rehabilitation” (2015): http://www.consultant.ru/document/cons_doc_LAW_176159 | | MOH: 1. Ministry of Health Order 433n (July 10, 2015) “On Adoption of the Regulations on Organization of Clinical Approbation of the Methods of Prevention, Diagnostics, Treatment and Rehabilitation (Including Order of Patients’ Assignment for Administering Such Medical Help), Standard Form of Protocol for Clinical Approbation of the Methods of Prevention, Diagnostics, Treatment, and Rehabilitation”: http://base.consultant.ru/cons/cgi/online.cgi?req=doc;base=LAW;n=183847 2. Ministry of Health Order 435h “On Ethics Committee of the Ministry of Health of the Russian Federation” (July 10, 2015): http://base.consultant.ru/cons/cgi/online.cgi?req=doc;base=LAW;n=183677 |
| <i>Drugs, Biologics, and Devices</i> | 1. Council of Ethics of the Ministry of Healthcare of the Russian Federation (MOH): http://www.grls.rosminzdrav.ru/ 2. Association of Clinical Trials Organizations: http://acto-russia.org/en/ 3. Federal Agency for Technical Regulation and Metrology (GOST): Main">http://www.gost.ru/wps/portal/pages.en>Main | Federal Law #61FZ “On Circulation of Medicines” (2011): http://acto-russia.org/files/zakon_ob_obr_ls_en.docx | MOH: 1. Ministry of Health Order No. 753n (August 26, 2010) “On Assertion of Order of Organization and Carrying out of Ethical Review...” (Russian): http://base.garant.ru/12178437/ 2. Ministry of Health Order No. 774n (August 31, 2010) “On | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | Council of Ethics” (Russian): http://www.rg.ru/2013/02/22/etika-dok.html 3. Ministry of Health Order of April 1, 2016 No. 200H "On Approval of the Rules of Good Clinical Practice.” http://acto-russia.org/files/prikaz_200n.docx GOST: Good Clinical Practice. GOST-R 52379-2005 (September 27, 2005): http://acto-russia.org/index.php?option=com_content&task=view&id=17 | |
| <i>Research Injury</i> | | Federal Law #61FZ “On Circulation of Medicines” (2011), Art. 38-44: http://acto-russia.org/files/zakon_ob_obr_ls_en.docx | | |
| <i>Privacy/Data Protection</i> | | 1. Federal Law of the Russian Federation on Information, Information Technologies, and Protection of Information (2006): http://www.consultant.ru/document/cons_doc_LAW_165971/ 2. Federal Law of the Russian Federation No. 152-FZ on Personal Data (2006): http://base.garant.ru/12148567/ | | |
| <i>Genetic</i> | Interdepartmental Commission on Genetic-Engineering Activity | Federal Law of July 5, 1996, N OF 8'-FZ “About the State Control in the Area of Genetic-Engineering Activity.” http://base.garant.ru/10135402/ | Order of the Ministry of Education and Science of the Russian Federation #154: “Statute of the Inter-Departmental Commission on Genetic-Engineering Activity” (2005): http://www.zakonprost.ru/content/ba/se/part/438157 | |
| <i>Embryos, Stem Cells, and Cloning</i> | | Federal Law #30-FZ “On Introduction of Change in Art. 1 of the Federal Law “On | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | | Temporary Ban on Human Cloning” (2010): http://base.garant.ru/184467/ | | |
| San Marino | | | | |
| <i>General</i> | San Marino Bioethics Committee (Italian): http://www.sanita.sm/online/home/comitato-bioetica/comitato-sammarinese-di-bioetica.html | Oviedo Convention on Human Rights and Biomedicine (1998) | | |
| <i>Research Injury</i> | | Oviedo Convention on Human Rights and Biomedicine, Article 24, ETS No. 164 (1998) | | |
| Serbia | | | | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health (MOH): http://www.zdravlje.gov.rs/ 2. Serbian Drug Agency http://www.alims.gov.rs/eng/ | Law on Medicines and Medical Devices, Official Gazette of RS No. 30/2010 and 107/2012: https://www.alims.gov.rs/eng/files/2013/04/Law-on-Medicines-and-Medical-Devices-2010.pdf | MOH: 1. Regulation on Content of Requests and Documents for Approval of Clinical Trials and Procedures for Conducting Clinical Trials, Official Gazette of RS, 64/2011 and 91/2013: https://www.alims.gov.rs/ciril/files/2017/11/pravilnik-ki-91-2013.pdf 2. Regulation on Amendment to Regulation on Content of Requests and Documents for Approval of Clinical Trials and Procedures for Conducting Clinical Trials, Official Gazette of RS, 60/2016: https://www.alims.gov.rs/ciril/files/2018/04/L-Klinicka-izm60-16.docx | |
| <i>Research Injury</i> | 1. Ministry of Health (MOH): http://www.zdravlje.gov.rs/index.php? 2. Serbian Drug Agency http://www.alims.gov.rs | Law on Medicines and Medical Devices, Article 72 (2013): https://www.alims.gov.rs/eng/files/2013/04/Law-on-Medicines-and-Medical-Devices-2010.pdf | MOH: 1. Regulation on Content of Requests and Documents for Approval of Clinical Trials and Procedures for Conducting Clinical Trials, Official Gazette of RS, 64/2011 and 91/2013: https://www.alims.gov.rs/ciril/files/2017/11/pravilnik-ki-91-2013.pdf 2. Law on Patients' Rights, Article 25 Official Gazette of RS, 45/13: http://www.zdravlje.gov.rs/downloads/2013/Jun/Jul2013ZakonOPravimaPacijenata.pdf | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|--|--|---|-------------------|
| <i>Privacy/Data Protection</i> | Commissioner for Information of Public Importance and Personal Data Protection: https://www.poverenik.rs/en/ | Law on the Protection of Personal Data, Official Gazette 97/08, 104/09, 68/20 and 107/12: http://www.minrzs.gov.rs/files/doc/porodica/ostali/Zakon%20o%20zastiti%20podataka%20o%20licnosti.pdf | | |
| <i>Genetics</i> | Ministry of Health (MOH): http://www.zdravlje.gov.rs/index.php? | Law on the Prevention and Diagnosis of Genetically Conditioned Diseases, Genetically Caused Anomalies and Rare Diseases (2015): http://www.parlament.gov.rs/upload/archive/files/lat/pdf/zakoni/2015/2245-14%20lat.pdf | | |
| <i>Embryos, Stem Cells, and Cloning</i> | National Health Insurance Fund: http://www.rfzo.rs/ | 1. Law on Organ Transplantation, Official Gazette No. 57/2018: https://www.paragraf.rs/propisi_download/zakon-o-presadjivanju-ljudskih-organa.pdf 2. Law on Human Cells and Tissues, Official Gazette No. 57/2018: https://www.paragraf.rs/propisi_download/zakon-o-ljudskim-celijama-i-tkivima.pdf | | |
| Slovakia | | | | |
| <i>General</i> | 1. Ministry of Health (Slovak): http://www.health.gov.sk/ 2. Institute of Medical Ethics and Bioethics: http://www.bioethics.sk/ | 1. Oviedo Convention on Human Rights and Biomedicine (1998) 2. Additional Protocol on Biomedical Research (2005) 3. Act No. 576/2004 Coll on Health Care, As Amended by Acts No. 350/2005, 282/2006, 662/2007, 345/2009 Coll. | | |
| <i>Drugs, Biologics, and Devices</i> | State Institute for Drug Control: http://www.sukl.sk/en | Act No. 140/1998 Coll. on Drugs and Medical Devices, as amended by Acts No. 9/2004 and 542/2006, 489/2008, and 402/2009 Coll. | Ministerial Regulation No. 239/2004 Coll. on Requirements for Clinical Trials and Good Clinical Practice, as Amended by Ministerial Regulation No. 148/2009 Coll. | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | | Law 277/1994 on Health Care, Section 44 | | |
| <i>Privacy/Data Protection</i> | Office for Personal Data Protection: https://dataprotection.gov.sk/uouu/en | 1. Act No. 428/2002 Coll. on Protection of Personal Data, as amended by Act No. 90/2005 Coll. 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | | GDPR Regulation (2018): https://dataprotection.gov.sk/uouu/sk/main-content/nariadenie-gdpr |
| <i>Human Biological Materials</i> | | 1. Act No. 576/2004 Coll. on Health Care, Sections 35-39. 2. Act No. 489/2008 Coll. on Drugs and Medical Devices, Section 18 (29b). | Governmental Regulation No. 20/2007 Coll. on Tissue and Cell Collection | |
| <i>Embryos, Stem Cells, and Cloning</i> | | 1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (1998) 2. Act No. 576/2004 Coll. on Health Care, Section 26.10.a. | | |
| Slovenia | | | | |
| Note: All websites and documents are in Slovenian. | | | | |
| <i>General</i> | Republic of Slovenia National Medical Ethics Committee (NMEC): http://www.kme-nmec.si/ | 1. Act Ratifying the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine: Convention on Human Rights and Biomedicine, and of the Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (1998): http://pisrs.si/Pis.web/pregledPredpisa?id=ZAKO1168 2. Decree Ratifying the Additional Protocol to the | | Slovenian Code of Medical Deontology, Articles 47-50 (1997) |

| Country | Key Organizations | Legislation | Regulations | Guidelines | |
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| <i>General</i> | | Convention on Human Rights and Biomedicine, Concerning Biomedical Research (2005): http://pisrs.si/Pis.web/pregledPredpisa?id=URED3728 3. Patient Rights Act, Official Gazette No. 15/2008 55/2017: http://pisrs.si/Pis.web/pregledPredpisa?id=ZAKO4281 and https://www.uradni-list.si/glasilo-uradni-list-rs/vsebina/2017-01-2526?sop=2017-01-2526 4. Mental Health Act, Official Gazette Nos. 77/2008 and 46/2015: http://pisrs.si/Pis.web/pregledPredpisa?id=ZAKO2157 | | | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | 1. Ministry of Health of the Republic of Slovenia http://www.mz.gov.si/ 2. Republic of Slovenia National Medical Ethics Committee (NMEC): http://www.kme-nmec.si/ 2. Agency for Medicinal Products and Medical Devices (JAZMP): http://www.jazmp.si/ | 1. Medicinal Products Act, Official Gazette No. 17/2014: http://www.uradni-list.si/1/objava.jsp?sop=2014-01-0539 2. EU Clinical Trials Regulation No. 536/2014: http://eur-lex.europa.eu/legal-content/EN/TXT/HTML/?uri=CELEX:32014R0536&from=EN | 1. Rules on Clinical Testing of Medicinal Products, Official Gazette, No. 54/2006 and 17/2014: http://www.pisrs.si/Pis.web/pregledPredpisa?id=PRAV6611 2. Rules on the Composition, Duties, Responsibilities, and Working Methods of the Commission for Medical Ethics, Official Gazette No. 21/2018: http://pisrs.si/Pis.web/pregledPredpisa?id=PRAV13345 | |
| | <i>Devices</i> | 1. Ministry of Health of the Republic of Slovenia: http://www.mz.gov.si/ 2. Agency for Medicinal Products and Medical Devices (JAZMP): http://www.jazmp.si/ 3. Republic of Slovenia National Medical Ethics Committee (NMEC): http://www.kme-nmec.si/ | 1. Medical Devices Act, Official Gazette No. 98/2009: http://www.pisrs.si/Pis.web/pregledPredpisa?id=ZAKO5503 | Rules on Medical Devices, Official Gazette Nos. 37/2010 and 66/2012: http://www.pisrs.si/Pis.web/pregledPredpisa?id=PRAV9508 | |
| <i>Research Injury</i> | 1. Ministry of Health of the Republic of Slovenia: http://www.mz.gov.si/ 2. Agency for Medicinal Products and Medical Devices (JAZMP): | 1. Medicinal Products Act, Official Gazette No. 17/2014: http://www.uradni-list.si/1/objava.jsp?sop=2014-01-0539 | 1. Rules on Clinical Testing of Medicinal Products, Official Gazette, No. 54/2006 and 17/2014: http://www.pisrs.si/Pis.web/pregledPredpisa?id=PRAV6611 | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | http://www.jazmp.si/ 3. Republic of Slovenia National Medical Ethics Committee (NMEC): http://www.kme-nmec.si/ | 2. Medical devices Act Official Gazette No. 98/2009 http://www.pisrs.si/Pis.web/pregledPredpisa?id=ZAKO5503 3. Convention on Human Rights and Biomedicine (Convention of Oviedo), Article 24, ETS No. 164 (1999) 4. Additional Protocol to the Convention on Human Rights and Biomedicine, Concerning Biomedical Research (2005) Decree ratifying the Additional Protocol to the Convention on Human Rights and Biomedicine, Concerning Biomedical Research: http://pisrs.si/Pis.web/pregledPredpisa?id=URED3728 | redpisa?id=PRAV6611 2. Rules on Medical Devices, Official Gazette Nos. 37/2010 and 66/2012: http://www.pisrs.si/Pis.web/pregledPredpisa?id=PRAV9508 | |
| <i>Privacy/Data Protection</i> | Information Commissioner of the Republic of Slovenia: http://www.ip-rs.si/ | 1. Personal Data Protection Act No. 94/2007: http://pisrs.si/Pis.web/pregledPredpisa?id=ZAKO3906 2. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | | |
| <i>Human Biological Materials</i> | 1. Ministry of Health of the Republic of Slovenia: http://www.mz.gov.si/ 2. Agency for Medicinal Products and Medical Devices (JAZMP): http://www.jazmp.si/ 3. Republic of Slovenia National Medical Ethics Committee (NMEC): http://www.kme-nmec.si/ 4. Institute for transplantation of Organs and Tissues of the Republic of Slovenia: http://www.slovenija-transplant.si/index.php?id=predstavitev&L=2 5. Institute Service of Slovenia for Transfusion Medicine: http://www.ztm.si/en/ | 1. Additional Protocol to the Convention on Human Rights and Biomedicine Concerning Transplantation of Organs and Tissues of Human Origin (2006) 2. Act on Quality and Safety of Human Tissues and Cells, for the Purposes for Medical Treatment, Official Gazette No. 61/2007: http://www.uradni-list.si/1/objava.jsp?sop=2007-01-3297 3. Rules on Donation and Procurement of Human Tissues and Cells, Official Gazette Nos. 70/2008, 67/2014, and 79/2014 4. Act Regulating the Collection and Transplantation of Human Body Parts for the Purposes of | | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 21-22 (1999) 2. On Interventions into the Human Corpse Which are Not Part of the Routine Autopsy and on Handling with Biologic Material of Human Origin (2004): http://bswww.mf.uni-lj.si/pls/bs/BS_full_rec?lang=SLO&c_docid=105859 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | | Medical Treatment, Official Gazette No. 56/2015: http://www.uradni-list.si/1/objava.jsp?sop=2015-01-2357 | | |
| <i>Genetic</i> | Republic of Slovenia National Medical Ethics Committee (NMEC): http://www.kme-nmec.si/ | Additional Protocol to the Convention on Human Rights and Biomedicine concerning Genetic Testing for Health Purposes (2009) | | |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Republic of Slovenia National Medical Ethics Committee (NMEC): http://www.kme-nmec.si/ 2. Agency for Medicinal Products and Medical Devices (JAZMP): http://www.jazmp.si/ | 1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (1998) 2. Infertility Treatment and Procedures of Biomedically-Assisted Procreation Act, Official Gazette No. 70/2000, Section 9 (Slovenian): http://www.uradni-list.si/1/objava.jsp?sop=2000-01-3307 3. Act on Quality and Safety of Human Tissues and Cells, for the Purposes for Medical Treatment, Official Gazette No. 61/2007 (Slovenian): http://www.uradni-list.si/1/objava.jsp?sop=2007-01-3297 4. Rules on Donation and Procurement of Human Tissues and Cells, Official Gazette Nos. 70/2008, 67/2014, and 79/2014 | | |
| Spain | | | | |
| Note: Many of the 17 Spanish autonomous regions have their own laws and regulations on human subject protections. | | | | |
| <i>General</i> | 1. Spanish Bioethics Committee: http://www.comitedebioetica.es/?lang=en_US 2. Coordinating Center for Ethical Committees on Clinical Research (Spanish): | 1. Oviedo Convention on Human Rights and Biomedicine (1999): http://www.coe.int/t/dg3/healthbioethic/texts_and_documents/ETS164Spanish.pdf | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | http://www.msc.es/profesionales/farmacia/ceic/home.htm 3. Institute of Health Carlos III, Ministry of Science and Innovation http://www.isciii.es/htdocs/en/index.jsp | 2. Law 14/2007 on Biomedical Research: http://www.catedraderechoygenoma humano.es/images/novedades/SpanshLawonBiomedicalResearchEnglis h.pdf | | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> Spanish Agency of Medicines and Medical Devices: http://www.aemps.gob.es/en/investigacionClinica/medicamentos/home.htm | | 1. Order SCO/362/2008 that Modifies Order SCO/256/2007: http://www.aemps.gob.es/legislacion/espana/investigacionClinica/docs/rc1_2008_410.pdf 2. Order SAS/3470/2009 on Drugs Post Authorization Research: http://www.aemps.gob.es/legislacion/espana/medicamentosUsoHumano/docs/farmacovigilancia/rc1_2009_2577.pdf 3. Royal Decree 1015/2009: Drug Availability for Special Purposes: http://www.boe.es/boe/dias/2009/07/20/pdfs/BOE-A-2009-12002.pdf 4. Royal Decree 577/2013 Regulating Pharmacovigilance in Human Use Medicines: http://www.boe.es/diario_boe/txt.php?id=BOE-A-2013-8191 5. Royal Decree 1090/2015 Regulating Clinical Trials with Medicinal Products, Ethics Committees for Investigation with Medicinal Products and the Spanish Clinical Studies Registry: http://noticias.juridicas.com/base_datos/Admin/565124-rd-1090-2015-de-4-dic-regula-los-ensayos-clinicos-con-medicamentos-los-comites.html | |
| | <i>Devices</i> Spanish Agency of Medicines and Medical Devices: http://www.aemps.gob.es/en/investigacionClinica/productosSanitarios/home.htm | Royal Decree 1591/2009, Regulating Sanitary Devices: http://www.ont.es/infesp/Legislacion/RD_1591_2009.pdf | Various: http://www.aemps.es/actividad/pschb/implantables1.htm#circulares | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | Spanish Agency of Medicines and Medical Devices: http://www.aemps.gob.es/en/home.htm | 1. Law 14/2007 on Biomedical Research, Article 18: http://www.catedraderechoygenoma humano.es/images/novedades/SpanishLawonBiomedicalResearchEnglish.pdf 2. Regulation No. 536/2014 of the European Parliament and of the Council on Clinical Trials on Medicinal Products for Human Use, Repealing Directive 2001/20/EC: http://eur-lex.europa.eu/legal-content/EN/TXT/HTML/?uri=CELEX:32014R0536&from=EN 3. Royal Decree 1090/2015 Regulating Clinical Trials with Medicinal Products, Ethics Committees for Investigation with Medicinal Products and the Spanish Clinical Studies Registry: https://www.aemps.gob.es/legislacion/espana/investigacionClinica/docs/Royal-Decree-1090-2015-4-December.pdf | | |
| <i>Privacy/Data Protection</i> | 1. Spanish Data Protection Authority: https://www.agpd.es/portalweb/index-ides-idphp.php 2. Spanish Agency of Medicines and Medical Devices (AEMPS): http://www.aemps.gob.es/en/investigacionClinica/medicamentos/home.htm | 1. Organic Law 15/1999 of December 13 on the Protection of Personal Data: http://www.legislationline.org/documents/id/9044 2. Law 14/2007 on Biomedical Research, Title I, Article 5: http://www.catedraderechoygenoma humano.es/images/novedades/SpanishLawonBiomedicalResearchEnglish.pdf 3. EU General Data Protection Regulation (2018): https://eur-lex.europa.eu/legal-content/EN/TXT/HTML/?uri=CELEX:32016R0679&from=EN | 1. Royal Decree 1720/2007: http://www.davara.com/documentos/relacionados/proteccion/RD_1720-2007_english.pdf 2. Royal Decree of 19 January 2008 | AEMPS: Revised Instructions for Updating the Section “Protection of Personal Data in the Subject Information Sheet (HIP /CI) Regarding the Regulation (EU) No. 2016/679 General Data Protection (2018): https://www.aemps.gob.es/investigacionClinica/medicamentos/docs/anexo8c-Ins-AEMPS-EC.pdf |
| <i>Human Biological Materials</i> | Ministry of Health, Consumer Affairs, and Social Welfare: http://www.msssi.gob.es/en/home.htm | Law 14/2007 of July 3 on Biomedical Research, Title I, Article 11; Title III, Article 37; Title V: | 1. Royal Decree 65/2006 of Requirements for the Import and Export of Biological Samples: http://www.boe.es/boe/dias/2006/02/ | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | | http://www.catedradercheygenoma humano.es/images/novedades/SpanishLawonBiomedicalResearchEnglish.pdf | <p>07/pdfs/A04626-04636.pdf</p> <p>2. Royal Decree 1723/2012 Regarding Activities of Collection, Clinical Use and Territorial Coordination of Human Organs for Transplants and Establishing Their Quality and Safety Requirements: http://noticias.juridicas.com/base_datos/Admin/rd1716-2011.html</p> <p>3. Royal Decree 1716/2011 on Biobanks: http://www.comitede bioetica.es/normativa/docs/RD%201716_2011_de%20autorizacion%20y%20funcionamiento%20de%20los%20biobancos.pdf</p> <p>4. Royal Decree 9/2014 on Quality and Security Rules Regarding Donating, Gathering, Evaluation, Processing, Storage, Preservation, and Distribution of Human Cells and Tissues and Rules Regarding Coordination and Functioning of their Use in Human Beings: http://www.boe.es/buscar/doc.php?id=BOE-A-2014-7065</p> | |
| <i>Genetic</i> | Spanish Bioethics Committee: http://www.comitede bioetica.es/?lang=en_US | Law 14/2007 of July 3 on Biomedical Research, Title I, Articles 6-9; Title V: http://www.catedradercheygenoma humano.es/images/novedades/SpanishLawonBiomedicalResearchEnglish.pdf | | |
| <i>Embryos, Stem Cells, and Cloning</i> | <p>1. Spanish Bioethics Committee: http://www.comitede bioetica.es/?lang=en_US</p> <p>2. National Commission for the Donation and Use of Embryos, Cells, and Human Tissues for Biomedical Research: http://www.isciii.es/ISCIII/es/contenidos/fd-el-instituto/organizacion.shtml</p> <p>3. National Biobank Register: http://www.isciii.es/ISCIII/es/contenidos/f</p> | <p>1. Additional Protocol to the Convention for the Protection of Human Rights and Dignity of the Human Being with regard to the Application of Biology and Medicine, on the Prohibition of Cloning Human Beings (2000)</p> <p>2. Law 14/2006 on Methods of Assisted Human Reproduction, Chapters IV and V</p> <p>3. Law 14/2007 of July 3 on</p> | Royal Decree 1527/2010 By Which the Guarantees Commission for the Donation and Use of Human Cells and Tissues and Registration Research Projects is Regulated: http://www.boe.es/diario_boe/txt.php?id=BOE-A-2010-18654 | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | d-el-instituto/organizacion.shtml 4. National Stem Cell Bank: http://www.isciii.es/ISCIII/es/contenidos/fd-el-instituto/fd-organizacion/fd-estructura-directiva/fd-subdireccion-general-investigacion-terapia-celular-medicina-regenerativa/fd-centros-unidades/banco-nacional-lineas-celulares.shtml | Biomedical Research, Title III: http://www.catedraderechoygenoma.humano.es/images/novedades/SpanishLawonBiomedicalResearchEnglish.pdf | | |
| Sweden | | | | |
| For an overview of human subject protections in Sweden, see CODEX: Rules and Guidelines for Research: http://www.codex.uu.se/en/index.shtml | | | | |
| <i>General</i> | Central Ethical Review Board: http://www.epn.se/en/start/ | Act No. 460 on the Ethical Review of Research Involving Humans (2003): http://www.epn.se/media/2348/the_ethical_review_act.pdf | 1. Ordinance No. 615 Concerning the Ethical Vetting of Research Involving Humans (2003): http://www.epn.se/media/1204/2003_615.pdf 2. Statute No. 2007:1068 Containing Instructions for the Central Ethical Review Board (2007): http://www.epn.se/media/1202/1068.pdf 3. Statute No. 2007:1069 Containing Instructions for Regional Ethical Review Boards (2007): http://www.epn.se/media/1203/1069.pdf | Information for Research Participants: https://www.epn.se/media/1210/information_for_research_participants.pdf |
| | Swedish Research Council: http://www.vr.se/english | | Regulations and General Counsel VRFS 2012:1 on Ethical Vetting of Human Subjects Research: https://www.vr.se/download/18.514d156f1639984ae0744e60/1529480567023/VRFS+2012.1.pdf | 1. Guidelines for the Ethical Evaluation of Medical Research on Humans (2003) 2. Policy Statement Regarding the Assessment of Scientific Studies in which Patients or Healthy Subjects are to Undergo Invasive Operations (2003) 3. Good Research Practice (2017): https://www.vr.se/english/analysis-and-assignments/we-analyse-and-evaluate/all-publications/publications/2017-08-31-good-research-practice.html |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> Medical Products Agency: https://lakemedelsverket.se/english/ | 1. Pharmaceuticals Act No. No 2015:315: https://open.karnovgroup.se/halso-och-sjukvard/lakemedelslagen | MPA Regulations on Clinical Trials in Humans -- LVFS 2011:19: http://www.lakemedelsverket.se/upload/lvfs/LVFS_2011_19.pdf | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | <i>Devices</i> Medical Products Agency: http://www.lakemedelsverket.se/english/product/Medical-devices/Clinical-Investigations/ | 1. Swedish Medical Devices Act (SFS 1993:584): http://www.notisum.se/rnp/sls/lag/19930584.htm 2. Medical Devices Ordinance (SFS1993:876): http://www.riksdagen.se/sv/dokument-lagar/dokument/svensk-forfattningssamling/forordning-1993876-om-medicintekniska_sfs-1993-876 | Swedish Implementation of Directive 93/42/EEC -- LVFS 2003:11 with Amendment LVFS 2004:11: https://lakemedelsverket.se/upload/lvfs/konsoliderade/LVFS_2003_11_konsoliderad_tom_2011_13.pdf | |
| <i>Social-Behavioral Research</i> | Swedish Research Council | | | Good Research Practice: Observational Studies Conducted Through Participating, Observing, and Recording (2017): https://www.vr.se/english/analysis-and-assignments/we-analyse-and-evaluate/all-publications/publications/2017-08-31-good-research-practice.html |
| <i>Privacy/Data Protection</i> | Swedish Data Protection Agency (SDPA): http://www.datainspektionen.se/in-english/ | 1. Patient Data Act: SFS 2008:355: http://www.notisum.se/rnp/sls/lag/20080355.htm 2. SFS 2009:400 - Public Access to Information and Secrecy Act: http://www.notisum.se/rnp/sls/lag/20090400.htm 3. Act on Certain Health Research Registers, SFS 2013:794: http://www.notisum.se/Pub/Doc.aspx?url=/rnp/sls/lag/20130794.htm 4. EU General Data Protection Regulation (2016): https://eur-lex.europa.eu/eli/reg/2016/679/oj | SFS 2009:641 - Public Access to Information and Secrecy Ordinance: http://www.notisum.se/rnp/sls/lag/20090641.htm | 1. General Data Protection Regulation (2018): https://www.datainspektionen.se/lagar-regler/dataskyddsförordningen/ 2. Transmission to Third Countries (2018): https://www.datainspektionen.se/lagar-regler/dataskyddsförordningen/tredjelandsöverföring/ |
| <i>Human Biological Materials</i> | 1. National Board of Health and Welfare (SOS): http://www.socialstyrelsen.se/english 2. Biobank Sweden: http://biobanksverige.se/ | 1. Biobanks in Medical Care Act No. 297 (2002): https://www.riksdagen.se/sv/dokument-lagar/dokument/svensk-forfattningssamling/lag-2002297-om-biobanker-i-halso--och_sfs-2002-297 2. Regulation No. 746 (2002): http://www.notisum.se/rnp/sls/lag/20020746.htm | SOS: Consolidated Regulations: http://www.socialstyrelsen.se/sosfs/2002-11 | SRC: Research Ethics Guidelines for Using Biobanks (2003) http://www.vr.se/download/18.6b2f98a910b3e260ae28000350/Riktlinjer_Biobanker_11.pdf |
| <i>Genetic Research</i> | 1. Medical Products Agency: https://lakemedelsverket.se/english/ | Act on Genetic Integrity | Drug Administration Regulations and Guidelines (LVFS 2004:10) on | SGTAB: |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | 2. The Swedish Gene Technology Advisory Board (SGTAB): https://www.genteknik.se/ | (2006:351): http://www.notisum.se/rnp/sls/lag/20060351.htm | the Intentional Release of Clinical Trials of Medicinal Products Containing or Consisting of Genetically Modified Organisms: http://www.lakemedelsverket.se/upplad/lvfs/LVFS_2004-10.pdf | Advice for Ethical Assessments: https://www.genteknik.se/wp-content/uploads/2017/09/072_2010-Etisk-v%C3%A4gledning.pdf |
| <i>Embryos, Stem Cells, and Cloning</i> | National Board of Health and Welfare (SOS): http://www.socialstyrelsen.se/english | Act on Genetic Integrity (2006:351): http://www.notisum.se/rnp/sls/lag/20060351.htm | 1. Legal Regulation of Stem Cell Research 2002:119: http://www.regeringen.se/sb/d/108/a/2717 2. Regulations and Guidelines for the Use of Tissues and Cells in Healthcare and Clinical Research - SOSFS 2009:32: http://www.socialstyrelsen.se/sosfs/2009-32 | |
| Switzerland | | | | |
| For an overview of human subject protections in Switzerland, see: http://kofam.ch/en/home/ | | | | |
| <i>General</i> | 1. Federal Office of Public Health (FOPH): http://www.bag.admin.ch/index.html?lang=en 2. Federal Office of Public Health, Portal for Human Research (FOPH): http://kofam.ch/en/home/ 3. National Advisory Commission on Biomedical Ethics (NEK-CNE): http://www.nek-cne.ch/en/homepage/ 3. Swiss Ethics Committees on Research Involving Humans: http://www.swissethics.ch/index_e.html | 1. Council of Europe Convention on Human Rights and Biomedicine of 4 April 1997, ETS No. 164, Articles 15-18: http://conventions.coe.int/Treaty/Commun/QueVoulezVous.asp?NT=164&CM=7&DF=9/15/2008&CL=ENG 2. Federal Constitution of the Swiss Confederation of 18 April, 1999, RS 101, Article 118b: http://www.admin.ch/opc/en/classified-compilation/19995395/index.html 3. Federal Act of 30 September 2011 on Research Involving Human Beings (Human Research Act, HRA), RS 810.30: http://www.admin.ch/opc/en/classified-compilation/20061313/index.html | 1. Ordinance of 20 September 2013 on Clinical Trials in Human Research with the Exception of Clinical Trials (Human Research Ordinance, HRO), RS 810.301: http://www.admin.ch/opc/en/classified-compilation/20121177/index.html 2. Ordinance of 20 September 2013 on Clinical Trials in Human Research (Clinical Trials Ordinance, ClinO), RS 810.305: https://www.admin.ch/opc/en/classified-compilation/20121176/index.html 3. Ordinance of 20 September 2013 on Organizational Aspects of the Human Research Act (HRA Organisational Ordinance, OrgO-HRA), RS 810.308: https://www.admin.ch/opc/en/classified-compilation/20121179/index.html | Swiss Clinical Trial Organisation, Guidelines for Good Operational Practice (GGOP) (2017): https://www.scto.ch/en/publications/guidelines.html <i>Access:</i> http://www.scto.ch/en/News.html |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. Swiss Agency for Therapeutic Products (Swissmedic): http://www.swissmedic.ch/index.html?lan | 1. Federal Act of 15 December 2000 on Medicinal Products and Medical Devices | 1. Ordinance of 20 September 2013 on Human Research with the Exception of Clinical Trials | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | g=en 2. Federal Office of Public Health (FOPH): http://www.bag.admin.ch/index.html?lang=en | (Therapeutic Products Act, TPA), RS 812.21, Articles 53-54: http://www.admin.ch/opc/en/classified-compilation/20002716/index.html 2. Federal Act of 30 September 2011 on Research Involving Human Beings (Human Research Act, HRA), RS 810.30: http://www.admin.ch/opc/en/classified-compilation/20061313/index.html | (Human Research Ordinance HRO), RS 810.301, Article 7 (2014): http://www.admin.ch/opc/en/classified-compilation/20121177/index.html 2. Ordinance of 20 September 2013 on Clinical Trials in Human Research (Clinical Trials Ordinance, ClinO), RS 810.305: https://www.admin.ch/opc/en/classified-compilation/20121176/index.html 3. Ordinance of 20 September 2013 on Organisational Aspects of the Human Research Act (HRA Organisational Ordinance, OrgO-HRA), RS 810.308, Articles 6-7: https://www.admin.ch/opc/en/classified-compilation/20121179/index.html | |
| | <i>Devices</i> Swiss Agency for Therapeutic Products (Swissmedic): http://www.swissmedic.ch/index.html?lang=en | 1. Federal Act of 15 December 2000 on Medicinal Products and Medical Devices (Therapeutic Products Act, TPA), RS 812.21, Articles 1-2, 45-67: https://www.admin.ch/opc/en/classified-compilation/20002716/index.html 2. Federal Act of 30 September 2011 on Research involving Human Beings, (Human Research Act, HRA), RS. 810.30: https://www.admin.ch/opc/en/classified-compilation/20061313/index.html | 1. Ordinance of 20 September 2013 on Human Research with the Exception of Clinical Trials (Human Research Ordinance HRO), RS 810.301, Article 7: https://www.admin.ch/opc/en/classified-compilation/20121179/index.html 2. Ordinance of 20 September 2013 on Clinical Trials in Human Research (Clinical Trials Ordinance, ClinO), RS 810.305 articles 20, 32, 37, 42-45 and Annexes 1, 3 and 4: https://www.admin.ch/opc/en/classified-compilation/20121176/index.html 3. Ordinance of 20 September 2013 on Organisation Aspects of the Human Research Act (HRA Organisation Ordinance, OrgO-HRA), RS 810.308, Articles 6-7: https://www.admin.ch/opc/en/classified-compilation/20121179/index.html | Swissmedic Guide to the Regulation of Medical Devices: https://www.swissmedic.ch/medizinprodukte/00287/index.html?lang=en |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Clinical Trials Registry</i> | Swiss National Clinical Trials Portal: http://kofam.ch/en/swiss-clinical-trials-portal/ | Federal Act on Research Involving Human Beings, Articles 56, 64, 65, and 67 (2014): https://www.admin.ch/opc/en/classified-compilation/20061313/index.html | | |
| <i>Research Injury</i> | 1. Swiss Agency for Therapeutic Products (Swissmedic): http://www.swissmedic.ch/index.html?lang=en 2. Federal Office of Public Health (FOPH): http://www.bag.admin.ch/index.html?lang=en | Federal Act of 30 September 2011 on Research Involving Human Beings (Human Research Act, HRA), RS 810.30, Articles 19-20: http://www.admin.ch/opc/en/classified-compilation/20061313/index.html | 1. Ordinance of 20 September 2013 on Human Research with the Exception of Clinical Trials (Human Research Ordinance HRO), RS 810.301, Articles 8, 12, 13, and 15, and Annexes 1-2: https://www.admin.ch/opc/en/classified-compilation/20121179/index.html 2. Ordinance of 20 September 2013 on Clinical Trials in Human Research (Clinical Trials Ordinance ClinO), RS 810.305, Articles 7, 10-13, 25, and 71, and Annexes 2-3: https://www.admin.ch/opc/en/classified-compilation/20121176/index.html | |
| <i>Privacy/Data Protection</i> Note: Most Swiss cantons have enacted laws regarding data collection in the public sector that are similar to the Federal Act on Data Protection. | Federal Data Protection and Information Commissioner (FDPIC): http://www.edoeb.admin.ch/index.html?lang=en | 1. Federal Act of 19 June 1992 on Data Protection (FADP), RS 235.1: http://www.admin.ch/opc/en/classified-compilation/19920153/index.html 2. Federal Act of 30 September 2011 on Research Involving Human Beings (Human Research Act, HRA), RS 810.30, Articles 2, 3, 8, 16-18, 31-35, 41-45, 47, 49, 58-60, and 63: http://www.admin.ch/opc/en/classified-compilation/20061313/index.html | 1. Ordinance of 20 September 2013 on Human Research with the Exception of Clinical Trials (Human Research Ordinance, HRO), RS 810.301, Articles 5 - 8, 10, 15, 21, 24-34, 37-39, 41, and 44-45, and Annex 2: http://www.admin.ch/opc/en/classified-compilation/20121177/index.html 2. Ordinance of 20 September 2013 on Clinical Trials in Human Research (Clinical Trials Ordinance, CLinO), RS 810.305 Articles 5, 7, 9, 12, 16-18, and 25, and Annexes 2-3: https://www.admin.ch/opc/en/classified-compilation/20121176/index.html | |
| <i>Human Biological Materials</i> | 1. Federal Office of Public Health (FOPH): http://www.bag.admin.ch/index.html?lang=en | Federal Act of 30 September 2011 on Research Involving | 1. Ordinance of 14 February 2007 on Human Genetic Testing, | SAMS: Biobanks: Obtainment, Preservation and |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | =en 2. Swiss Academy of Medical Sciences (SAMS): http://www.samw.ch/en/News/News.html | Human Beings (Human Research Act, HRA), RS 810.30, Articles 2, 3, 17, 18, 31, 32 - 35, 41-43, 45, 47, 49, and 63: http://www.admin.ch/opc/en/classified-compilation/20061313/index.html | RS 810.122.1: http://www.admin.ch/opc/fr/classified-compilation/20051790/index.html 2. Ordinance of 20 September 2013 on Human Research with the Exception of Clinical Trials (Human Research Ordinance, HRO), RS 810.301 Articles 5 - 8, 10, 15, 21, 24-30, 33-34, 37 - 39, 41, 44-45 and Annex 2): http://www.admin.ch/opc/en/classified-compilation/20121177/index.html 3. Ordinance of 20 September 2013 on Clinical Trials in Human Research (Clinical Trials Ordinance, CLinO), RS 810.305, Articles 7, 9, 12, 16 - 18 and Annex 2: http://www.admin.ch/opc/en/classified-compilation/20121176/index.html | Utilization of Human Biological Material (2006): http://www.samw.ch/en/Ethics/Guidelines/Archive.html |
| <i>Genetic Research</i> | Federal Office of Public Health (FOPH): http://www.bag.admin.ch/index.html?lang=en | 1. Federal Constitution of the Swiss Confederation of 18 April 1999, RS 101, Article 119: http://www.admin.ch/opc/en/classified-compilation/19995395/index.html 2. Federal Act of 8 October 2004 on Human Genetic Testing (HGTA), RS 810.12: http://www.admin.ch/opc/en/classified-compilation/20011087/index.html 3. Federal Act of 30 September 2011 on Research Involving Human Beings (Human Research Act, HRA), RS 810.30, Articles 3, 32 - 35, 42, and 49: http://www.admin.ch/opc/en/classified-compilation/20061313/index.html | 1. Ordinance of 14 February 2007 on Human Genetic Testing, RS 810.122.1 (French): http://www.admin.ch/opc/fr/classified-compilation/20051790/index.html 2. Ordinance of 20 September 2013 on Human Research with the Exception of Clinical Trials (Human Research Ordinance, HRO), RS 810.301, Articles 28 - 32: http://www.admin.ch/opc/en/classified-compilation/20121177/index.html 3. Ordinance of 20 September 2013 on Clinical Trials in Human Research (Clinical Trials Ordinance, CLinO), RS 810.305 Articles 22 and 35, and Annexes 3 and 4: http://www.admin.ch/opc/en/classified-compilation/20121176/index.html | |
| <i>Embryos, Stem Cells, and Cloning</i> | Swiss National Advisory Commission on Biomedical Ethics (NEK-CNE): http://www.nek-cne.ch/en/homepage/ | <i>Embryos in Vivo:</i> Federal Act of 30 September 2011 on Research Involving Human Beings (Human | <i>Embryos in Vivo:</i> 1. Ordinance of 2 February 2005 on Research involving Embryonic Stem Cells (Stem | NEK-CNE: 1. Pre-Implantation Genetic Diagnosis, 2007/9: http://www.nek-cne.ch/fileadmin/nek-cne-dateien/Themen/Stellungnahmen/en/pid_en.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|--|--|--|---|
| <i>Embryos, Stem Cells, and Cloning</i> | | <p>Research Act, HRA), RS 810.30 Articles 2, 25 - 27, 39, 40, 44, and 62: http://www.admin.ch/opc/en/classified-compilation/20061313/index.html</p> <p><i>Others:</i> Federal Act of 19 December 2003 on Research Involving Embryonic Stem Cells (Stem Cell Research Act, StRA), RS 810.31: http://www.admin.ch/opc/en/classified-compilation/20022165/index.html</p> | <p>Cell Research Ordinance, SCRO), RS 810.311: http://www.admin.ch/opc/en/classified-compilation/20042542/index.html</p> <p>2. Ordinance of 20 September 2013 on Human Research with the Exception of Clinical Trials (Human Research Ordinance, HRO), RS 810.301, Articles 44 – 46, and Annex 2: http://www.admin.ch/opc/en/classified-compilation/20121177/index.html</p> <p>3. Ordinance of 20 September 2013 on Clinical Trials in Human Research (Clinical Trials Ordinance, CLinO), RS 810.305, Articles 49, 53, 55, and 56, and Annexes 3 and 4: http://www.admin.ch/opc/en/classified-compilation/20121176/index.html</p> | <p>2. Research Involving Human Embryos and Fetuses. Opinion No. 11/2006: http://www.nek-cne.ch/fileadmin/nek-cne-dateien/Themen/Stellungnahmen/en/embryonen_en.pdf</p> <p>3. Pre-Implantation Genetic Diagnosis II, Opinion No. 14/2007: http://www.nek-cne.ch/fileadmin/nek-cne-dateien/Themen/Stellungnahmen/PID_II_d.pdf</p> <p><i>Access:</i> http://www.nek-cne.ch/en/topics/opinions/</p> |
| Ukraine | | | | |
| <i>General</i> | <p>Ukrainian Ministry of Health: http://www.moz.gov.ua/en/</p> | <p>1. Constitution of Ukraine Art. 28 (1996) 2. Health Care Law, Article 45 (1992) 3. Criminal Code of Ukraine 2001, Article 141 and 142</p> | | |
| <i>Drugs, Biologics, and Devices</i> | <p>1. Ministry of Health of Ukraine State Expert Center: http://www.dec.gov.ua 2. National Academy of Sciences Bioethics Committee: http://biomed.nas.gov.ua/index-en/bioethics-committee</p> | <p>1. Ministry of Health Act On Procedure of Clinical Trials and Basic Statute of Ethics Commission 23.09/2009 No. 690 (2014): http://zakon5.rada.gov.ua/laws/show/z1010-09 2. On Medicines, Articles 7 and 8 No. 123/96BP (2014): http://zakon4.rada.gov.ua/laws/show/123/96-%D0%B2%D1%80</p> | <p>1. Ukrainian Ministry of Health Order No. 95 About Approval of Documents Related to the Quality Assurance of Medicines (2009): http://zakon5.rada.gov.ua/rada/show/v0095282-09 2. Ministry of Health Act 14.12.2009 N 944 on Approval of the Clinical Trial and Expertise of Clinical Trials: http://zakon4.rada.gov.ua/laws/show/z0053-10</p> | <p>Bioethics Committee: 1. Information Letters on Ethics Questions of Clinical Trials and Implementation of Medicines (2006) 2. Ethics Expertise of Clinical Trials Medicines (2007) 3. Methodological Aspects of Central EC Activity of Ukrainian Ministry of Health (2007) 4. Ethical Aspects of Placebo Controlled Clinical Trials in Patients with MS (2008) 5. Optimization of Local Ethics Committee Activities (2009)</p> <p>Ministry of Health: Guidelines for Pre-Clinical and Clinical Trials: http://www.dec.gov.ua/index.php/ekspertiza-</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | | materialiv-doklinichnikh-ta-klinichnikh-viprobuvan/metodichni-rekomendatsiji-shchodo-provedennya-doklinichnikh-ta-klinichnikh-viprobuvan |
| <i>Research Injury</i> | Ukrainian Ministry of Health: http://www.moz.gov.ua/en/ | On Medicines, Article 8 No. 123/96BP (2014): http://zakon4.rada.gov.ua/laws/show/123/96-%D0%B2%D1%80 | | |
| <i>Privacy/Data Protection</i> | 1. State Service of Ukraine on Personal Data Protection 2. Ukrainian Parliament Commissioner for Human Rights: www.ombudsman.gov.ua | 1. Convention for the Protection of Individuals with Regard to Automatic Processing of Personal Data (2010) 2. On Protection of Personal Data Act, 01.06.2010 with changes from 19.10.2017: http://zakon3.rada.gov.ua/laws/show/2297-17 | | |
| <i>Human Biological Materials</i> | Ukrainian Ministry of Health: http://www.moz.gov.ua/en/ | 1. Cabinet Ministry of Ukraine Act No. 286 on 02.03.2016 License Conditions on Providing Activities of Banks of Cord Blood and Other Human Tissues and Cells: http://zakon2.rada.gov.ua/laws/show/286-2016-%D0%BF 2. Ministry of Health Act 20.04.12 No. 276 On Approving the List of Human Tissues and Cells, Allowing the Use of Banks of Cord Blood and Other Human Tissues and Cells: http://zakon3.rada.gov.ua/laws/show/z1124-12 | Ukrainian Ministry of Health Order No. 630 Regarding Approval of the Procedure for the Conduct of Clinical Trials of Tissue and Cell Transplants and Expert Evaluation of Materials of Clinical Trials (2007) with Changes from 23.09.2009 No. 690: http://zakon1.rada.gov.ua/laws/show/z1206-07 | |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. National Academy of Sciences Bioethics Committee: http://biomed.nas.gov.ua/index-en/bioethics-committee 2. Ukrainian Ministry of Health: http://www.moz.gov.ua/en/ | 1. Act on the Banning of Human Reproductive Cloning (2004): http://zakon0.rada.gov.ua/laws/show/2231-15 2. Act on the Transplantation on Human Using Anatomic Materials (2019): http://zakon.rada.gov.ua/laws/show/2427-19 | 1. Ukrainian Ministry of Health Order No. 630 Regarding Approval of the Procedure for the Conduct of Clinical Trials of Tissue and Cell Transplants and Expert Evaluation of Materials of Clinical Trials (2007): http://zakon1.rada.gov.ua/laws/show/z1206-07 2. Ukrainian Ministry of Health Order No. 787 on Approval of the Use of Reproductive | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Embryos, Stem Cells, and Cloning | | | Technologies in Ukraine 09.09.2013: http://zakon4.rada.gov.ua/laws/show/z1697-13 | |
| <p>United Kingdom Unless otherwise noted, all laws, regulations, and guidelines listed for England also apply to the entire United Kingdom. For an overview of clinical research regulations in the United Kingdom, see the ClinRegs report: http://clinregs.niaid.nih.gov/single_country.php?c_id=226</p> | | | | |
| General | <p><i>England:</i> Health Research Authority (HRA): http://www.hra.nhs.uk/</p> <p>Department of Health and Social Care (DHSC): https://www.gov.uk/government/organisations/department-of-health-and-social-care</p> <p>Medical Research Council (MRC): https://www.mrc.ac.uk/</p> | <p>1. Mental Capacity Act (2005) (England and Wales only): http://www.legislation.gov.uk/ukpga/2005/9/contents</p> <p>2. Health and Social Care Act (2012): http://www.legislation.gov.uk/ukpga/2012/7/contents/enacted</p> <p>3. Care Act (2014): http://www.legislation.gov.uk/ukpga/2014/23/contents/enacted/data.htm</p> <p>4. Ionising Radiation (Medical Exposure) Regulations (2017): http://www.legislation.gov.uk/uksi/2017/1322/contents/made</p> | <p>1. Research Governance Framework for Health and Social Care UK Policy Framework for Health and Social Care Research (2018): https://www.hra.nhs.uk/planning-and-improving-research/policies-standards-legislation/uk-policy-framework-health-social-care-research/</p> <p>2. Governance Arrangements for Research Ethics Committees (2018): https://www.hra.nhs.uk/planning-and-improving-research/policies-standards-legislation/governance-arrangement-research-ethics-committees/</p> | <p>1. HRA Guidance: https://www.hra.nhs.uk/planning-and-improving-research/</p> <p>2. Integrated Research Application System: https://www.myresearchproject.org.uk/</p> <p>1. Research Involving Human Participants in Developing Societies (2004): https://www.mrc.ac.uk/publications/browse/research-involving-human-participants-in-</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | | | | developing-societies/ 2. Medical Research Involving Children (2004): https://www.mrc.ac.uk/documents/pdf/medical-research-involving-children/ 3. Medical Research Involving Adults Who Cannot Consent (2007): http://www.mrc.ac.uk/documents/pdf/medical-research-involving-adults-who-cannot-consent/ 4. Good Research Practice: Principles and Guidelines (2012): https://www.mrc.ac.uk/publications/browse/good-research-practice-principles-and-guidelines/ |
| | <i>Scotland:</i> | | | |
| | 1. NHSScotland, Chief Scientist Office (CSO): http://www.cso.scot.nhs.uk/ 2. NHS Research Scotland: http://www.nhsresearchscotland.org.uk/ | Adults with Incapacity (Scotland) Act 2000, Section 51: http://www.scotland.gov.uk/Topics/Justice/law/awi/legislation | Adults with Incapacity (Ethics Committee) (Scotland) Regulations (2002): http://www.scotland-legislation.hmso.gov.uk/legislation/scotland/ssi2002/20020190.htm | CSO: Research Governance Framework for Health and Community Care (2006): http://www.cso.scot.nhs.uk/wp-content/uploads/2013/02/RGF-Second-Edition-February-06.pdf |
| | <i>Wales:</i> | | | |
| Health and Care Research Wales: http://www.healthandcareresearch.gov.wales/ | | | Research Governance Framework for Health and Social Care in Wales Second Edition (2009): http://www.wales.nhs.uk/sites3/Documents/952/Research%20Governance%20Framework%202009%20%28English%291.pdf | |
| | <i>Northern Ireland:</i> | | | |
| | 1. Department of Health, Social Services and Public Safety: http://www.dhsspsni.gov.uk/ 2. Office for Research Ethics Committees Northern Ireland: http://www.hscbusiness.hscni.net/orecni.htm | Ionising Radiation (Medical Exposure) (Northern Ireland) Regulations (2018): http://www.legislation.gov.uk/nisr/2018/17/contents/made | | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | | | |
| | 1. Medicines and Healthcare Products Regulatory Agency (MHRA): https://www.gov.uk/government/organisations/medicines-and-healthcare-products-regulatory-agency 2. Administration of Radioactive Substances Advisory Committee (ARSAC) (UK): https://www.gov.uk/government/organisations/administration-of-radioactive- | Medicines Act (1968): http://www.legislation.gov.uk/ukpga/1968/67/contents | 1. Medicines for Human Use (Clinical Trials) Regulations, Statutory Instrument No. 1031 (2004): http://www.legislation.gov.uk/uksi/2004/1031/contents/made 2. Amendment Regulations (SI 2006/1928) http://www.legislation.gov.uk/uksi/2006/1928/contents/made | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | <p>substances-advisory-committee</p> <p>3. Department of Environment, Food & Rural affairs (DEFRA) https://www.gov.uk/government/organisations/department-for-environment-food-rural-affairs</p> <p>4. Health and Safety Executive (HSE) http://www.hse.gov.uk/</p> | | <p>3. Amendment to the Medicines for Human Use (Clinical Trials) Regulations 2004 and Adults with Incapacity (Scotland) Act 2000 to Facilitate Clinical Research in Emergency Settings (SI 2006/2984): http://www.legislation.gov.uk/uksi/2006/2984/pdfs/uksi_20062984_en.pdf</p> <p>4. SI 2008 No.941 The Medicines for Human Use (Clinical Trials) and Blood Safety and Quality Amendment Regulations 2008: http://www.legislation.gov.uk/uksi/2008/941/contents/made</p> <p>5. Genetically Modified Organisms (Deliberate Release) Regulations 2002: http://www.legislation.gov.uk/uksi/2002/2443/contents/made</p> <p>6. Genetically Modified Organisms (Contained Use) Regulations 2014 (England, Scotland and Wales): http://www.legislation.gov.uk/uksi/2014/1663/part/1/made</p> <p>7. Genetically Modified Organisms (Contained Use) Regulations (Northern Ireland) 2015: http://www.legislation.gov.uk/nisr/2015/339/contents/made</p> | |
| | <p>Association of the British Pharmaceutical Industry (ABPI): http://www.abpi.org.uk</p> | | | <p>Guidelines for Phase I Clinical Trials (2012): http://www.abpi.org.uk/our-work/library/guidelines/Pages/phase-1-trials-2012.aspx</p> |
| | <p>National Institute for Health Research: http://www.nihr.ac.uk/</p> | | | <p>Clinical Trials Toolkit: http://www.ct-toolkit.ac.uk/</p> |
| | <p>Health Research Authority (HRA): http://www.hra.nhs.uk/</p> | | | <p>Clinical Trials of Investigational Medicinal Products (CTIMPs) – Resource page: http://www.hra.nhs.uk/resources/before-you-apply/types-of-study/clinical-trials-of-investigational-medicinal-products/</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | <i>Devices</i> Medicines and Healthcare Products Regulatory Agency (MHRA): https://www.gov.uk/government/collections/regulatory-guidance-for-medical-devices | | 1. Medical Devices Regulations (2002): http://www.opsi.gov.uk/si/si2002/20020618.htm 2. Medical Devices (Amendment) Regulations 2008 No 2936: http://www.legislation.gov.uk/ukxi/2008/2936/contents/made | 1. Clinical Trials for Medical Devices: https://www.gov.uk/government/collections/regulatory-guidance-for-medical-devices 2. Notify MHRA About a Clinical Investigation for a Medical Device: https://www.gov.uk/guidance/notify-mhra-about-a-clinical-investigation-for-a-medical-device |
| | Health Research Authority (HRA): http://www.hra.nhs.uk/ | | | Medical Devices Guidance: http://www.hra.nhs.uk/resources/before-you-apply/types-of-study/medical-devices-research-2/ |
| <i>Clinical Trials Registry</i> | 1. ISRCTN: http://www.isrctn.com/ 2. Health Research Authority (HRA): http://www.hra.nhs.uk/ | | | ISRCTN: FAQs: http://www.isrctn.com/page/faqs HRA: Transparency: Researchers' Responsibilities: https://www.hra.nhs.uk/planning-and-improving-research/research-planning/research-registration-and-research-project-identifiers/ |
| <i>Research Injury</i> | Medicines and Healthcare Products Regulatory Agency (MHRA): https://www.gov.uk/government/organisations/medicines-and-healthcare-products-regulatory-agency | | Medicines for Human Use (Clinical Trials) Regulations, Statutory Instrument No. 1031, Regulation 15(5)(i)(j)(k) and Schedule 3 Part 1, Paragraphs 1(g) and 3(c) (2004): http://www.legislation.gov.uk/ukxi/2004/1031/contents/made | |
| | Department of Health (DH): https://www.gov.uk/government/organisations/department-of-health | | | NHS Indemnity Arrangements for Clinical Negligence Claims in the NHS: www.nhs.uk/clinicalnegligence/claims/Documents/NHS%20Indemnity.pdf |
| | Association of the British Pharmaceutical Industry (ABPI): http://www.abpi.org.uk | | | 1. Insurance and Compensation in the Event of Injury in Phase I Clinical Trials (2012): http://www.abpi.org.uk/our-work/library/guidelines/Pages/clinical-trials-insurance.aspx 2. Clinical Trial Compensation Guidelines (2014): http://www.abpi.org.uk/our-work/library/guidelines/Pages/ct-compensation.aspx |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | Association of the British Healthcare Industry (ABHI): http://www.abhi.org.uk/ | | | Clinical Investigations Compensation Guidelines (2014): http://www.abhi.org.uk/multimedia/groups/clinical-investigations/ci_compensationguidelines.doc |
| <i>Social-Behavioral Research</i> | Economic and Social Research Council | | | ESRC Framework for Research Ethics (2015): http://www.esrc.ac.uk/files/funding/guidance-for-applicants/esrc-framework-for-research-ethics-2015/ |
| | UK Research Integrity Office | | | Good Practice in Research: Internet-Mediated Research (2016): http://ukrio.org/wp-content/uploads/UKRIO-Guidance-Note-Internet-Mediated-Research-v1.0.pdf |
| <i>Privacy/Data Protection</i> | <i>United Kingdom:</i> | | | |
| | Information Commissioner's Office: https://ico.org.uk/ | Data Protection Act (2018): http://www.legislation.gov.uk/ukpga/2018/12/contents/enacted | | 1. Guide to the General Data Protection Regulation (2018): https://ico.org.uk/for-organisations/guide-to-the-general-data-protection-regulation-gdpr/ 2. International Transfers (2018): https://ico.org.uk/for-organisations/guide-to-the-general-data-protection-regulation-gdpr/international-transfers/ |
| | Health Research Authority: https://www.hra.nhs.uk | | | 1. GDPR Guidance: https://www.hra.nhs.uk/planning-and-improving-research/policies-standards-legislation/data-protection-and-information-governance/gdpr-guidance/ 2. Consent in Research (2018): https://www.hra.nhs.uk/planning-and-improving-research/policies-standards-legislation/data-protection-and-information-governance/gdpr-guidance/what-law-says/consent-research/ |
| | Medical Research Council (MRC): http://www.mrc.ac.uk/ | | | 1. Using Information About People in Health Research (2017): https://mrc.ukri.org/documents/pdf/using-information-about-people-in-health-research-2017/ 2. Data and Tissues Tool Kit: http://www.dt-toolkit.ac.uk/home.cfm |
| | <i>England and Wales:</i> | | | |
| 1. Health Research Authority (HRA) (England): http://www.hra.nhs.uk/ 2. Confidentiality Advisory Group (CAG): | Health Service (Control of Patient Information) Regulations 2002 (HS (CPI) Regs): http://www.legislation.gov.uk/uksi/ | | 1. Research Data and Tissue Resources: https://www.hra.nhs.uk/planning-and-improving-research/research-planning/research-data-and-tissue-resources/ | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Privacy/Data Protection | http://www.hra.nhs.uk/about-the-hra/our-committees/section-251 | 2002/1438/made?view=plain | | 2. Section 251 and the Confidentiality Advisory Group (CAG): http://www.hra.nhs.uk/about-the-hra/our-committees/section-251/ |
| Human Biological Materials | <i>United Kingdom:</i> Human Tissue Authority (HTA): http://www.hta.gov.uk/ | 1. Human Tissue Act (2004): http://www.legislation.gov.uk/ukpga/2004/30/contents (Applies to England, Wales, and Northern Ireland. Section 45 also applies in Scotland.) 2. Statutory Instrument 2006 No. 1260: The Human Tissue Act 2004 (Ethical Approval, Exceptions from Licensing and Supply of Information about Transplants) Regulations 2006: http://www.legislation.gov.uk/uksi/2006/1260/contents/made (Applies to England, Wales, and Northern Ireland.) 3. Statutory Instrument 2006 No. 1659: The Human Tissue Act 2004 (Persons who Lack Capacity to Consent and Transplants) Regulations 2006 (Different provisions apply to England, Wales, Northern Ireland, and/or Scotland.): http://www.legislation.gov.uk/uksi/2006/1659/contents/made | | Guidance for Professionals: https://www.hta.gov.uk/guidance-professionals |
| | Medical Research Council (MRC): https://www.mrc.ac.uk/ | | | 1. Human Tissue and Biological Samples for Use in Research (2014): https://mrc.ukri.org/publications/browse/human-tissue-and-biological-samples-for-use-in-research/ 2. Data and Tissues Tool Kit: http://www.dt-toolkit.ac.uk/home.cfm |
| | <i>Scotland:</i> Healthcare Improvement Scotland: http://www.healthcareimprovementscotland.org/our_work/governance_and_assurance/human_tissue_banks.aspx | Human Tissue (Scotland) Act 2006: http://www.legislation.gov.uk/asp/2006/4/contents | | |
| | 1. Public Health Genetics Foundation: http://www.phgfoundation.org/ | | | |
| Genetics Research | | | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | 2. Gene Therapy Advisory Committee: http://www.hra.nhs.uk/resources/applying-to-recs/gene-therapy-advisory-committee-gtac/ 3. Genomics England: https://www.genomicsengland.co.uk/ | | | |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Human Fertilisation and Embryology Authority: http://www.hfea.gov.uk/ 2. Human Tissue Authority (HTA): https://www.hta.gov.uk/regulated-sectors | 1. Human Fertilisation and Embryology Act (1990): http://www.legislation.gov.uk/ukpga/1990/37/contents 2. HFE Act (2008): http://www.legislation.gov.uk/ukpga/2008/22/contents | Human Fertilisation and Embryology Regulation and Chronology: https://www.hfea.gov.uk/about-us/how-we-regulate/ | HFEA Code of Practice 9th Edition (2018): https://www.hfea.gov.uk/media/2609/june-2018-code-of-practice-9th-edition-draft.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| ASIA/PACIFIC | | | | |
| Australia | | | | |
| <i>General</i> | 1. National Health and Medical Research Council (NHMRC): http://www.nhmrc.gov.au/ 2. Australian Research Council (ARC): http://www.arc.gov.au | National Health and Medical Research Council Act 1992 (2014): http://www.comlaw.gov.au/Details/C2014C00364 | National Health and Medical Research Regulation 2016: https://www.legislation.gov.au/Details/F2016L00682 | NHMRC: 1. Ethical conduct in Research with Aboriginal and Torres Strait Islander Peoples and Communities: Guidelines for Researchers and Stakeholders (2018): https://nhmrc.gov.au/about-us/publications/ethical-conduct-research-aboriginal-and-torres-strait-islander-peoples-and-communities 2. Keeping Research on Track II (2018): https://nhmrc.gov.au/about-us/publications/keeping-research-track-ii NHMRC, ARC, and Universities Australia: 1. Australian Code for the Responsible Conduct of Research (2018): https://nhmrc.gov.au/research-policy/research-integrity/release-2018-australian-code-responsible-conduct-research 2. National Statement on Ethical Conduct in Human Research, 2007 (2018): https://nhmrc.gov.au/research-policy/ethics/national-statement-ethical-conduct-human-research |
| | Australian Institute of Aboriginal and Torres Strait Islander Studies (AIATSIS): http://aiatsis.gov.au/ | | | Guidelines for Ethical Research in Australian Indigenous Studies (2012): http://www.aiatsis.gov.au/research/ethics/GERAIS.html |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> Therapeutic Goods Administration (TGA): http://www.tga.gov.au | Therapeutic Goods Act 1989 (2016): https://www.legislation.gov.au/Details/C2016C00269 | Therapeutic Goods Regulations 1990 (2016): https://www.legislation.gov.au/Details/F2016C00801 | TGA: Australian Clinical Trial Handbook (2018): https://www.tga.gov.au/publication/australian-clinical-trial-handbook Australian States and Territories: National Mutual Acceptance of Scientific and Ethical Review of Multi-Centre Human Research (2017): https://www.australianclinicaltrials.gov.au/ethi |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | | cal-review-process-each-australian-state-and-territory |
| | <i>Devices</i> | | | |
| | Therapeutic Goods Administration: http://www.tga.gov.au/industry/devices.htm | Therapeutic Goods Act 1989 (2016): https://www.legislation.gov.au/Details/C2016C00269 | Therapeutic Goods (Medical Devices) Regulations 2002 (2016): https://www.legislation.gov.au/Details/F2016C00801 | Australian Regulatory Guidelines for Medical Devices (ARGMD) (2011): http://www.tga.gov.au/industry/devices-argmd.htm |
| <i>Clinical Trials Registry</i> | 1. National Health and Medical Research Council and the Department of Industry, Innovation, and Science: https://www.australianclinicaltrials.gov.au 2. Australian New Zealand Clinical Trials Registry: http://www.anzctr.org.au/ | | | 1. National Statement on Ethical Conduct in Human Research, 3.1.7 (2018): https://nhmrc.gov.au/research-policy/ethics/national-statement-ethical-conduct-human-research 2. FAQs: http://www.anzctr.org.au/Faq.aspx |
| <i>Research Injury</i> | 1. Therapeutic Goods Administration (TGA): http://www.tga.gov.au/ 2. Medicines Australia https://medicinesaustralia.com.au 3. National Health and Medical Research Council (NHMRC): http://www.nhmrc.gov.au | | | TGA: Guidance on Good Clinical Practice (CPMP/ICH-135/95). Paragraphs 5.8.1, 5.11.1, 8.2.5 , 8.2.7 (2018): https://www.tga.gov.au/publication/note-guidance-good-clinical-practice Medicines Australia: Industry Standard Compensation Guidelines (2012): https://medicinesaustralia.com.au/policy/clinical-trials/indemity-and-compensation-guidelines/ NHMRC, ARC, and UA: National Statement on Ethical Conduct in Human Research. Paragraphs 5.1.38 and 5.1.39 (2018): https://nhmrc.gov.au/research-policy/ethics/national-statement-ethical-conduct-human-research |
| <i>Social-Behavioral Research</i> | National Health and Medical Research Council (NHMRC): http://www.nhmrc.gov.au | | | National Statement on Ethical Conduct in Human Research, Chapter 3.1 (2015): https://www.nhmrc.gov.au/book/chapter-3-1-qualitative-methods |
| <i>Privacy/Data Protection</i> | Office of the Australian Information Commissioner: http://www.oaic.gov.au/ | Privacy Act 1988 (2016): https://www.legislation.gov.au/Details/C2016C00838 | 1. Australian Privacy Principles Guidelines (2014): http://www.oaic.gov.au/privacy/privacy-act/australian-privacy-principles 2. Guidelines under Section 95 of the Privacy Act 1988 (2014): https://nhmrc.gov.au/about- | 1. Australian Privacy Principles Guidelines (2015): http://www.oaic.gov.au/privacy/privacy-act/australian-privacy-principles 2. Guidelines under Section 95 of the Privacy Act 1988 (2014): http://www.nhmrc.gov.au/guidelines/publicatio |
| Note: All Australian states and territories have privacy/data protection laws: | | | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| http://www.oaic.gov.au/privacy/other-privacy-jurisdictions/state-and-territory-privacy-law | | | us/publications/guidelines-under-section-95-privacy-act-1988 3. Guidelines Approved under Section 95A of the Privacy Act 1988 (2014): https://www.nhmrc.gov.au/about-us/publications/guidelines-approved-under-section-95a-privacy-act-1988 4. Guidelines Approved under Section 95AA of the Privacy Act 1988 (2014): https://www.nhmrc.gov.au/about-us/publications/guidelines-approved-under-section-95aa-privacy-act-1988 5. Privacy Regulation 2013 (2016): https://www.legislation.gov.au/Details/F2016C00599 | ns/pr1 3. Guidelines Approved under Section 95A of the Privacy Act 1988 (2014): http://www.nhmrc.gov.au/guidelines/publications/pr2 4. Guidelines Approved under Section 95AA of the Privacy Act 1988 (2014): http://www.nhmrc.gov.au/guidelines/publications/pr3 |
| <i>Human Biological Materials</i> Note: All Australian states and territories also have laws on human biological materials. | 1. National Health and Medical Research Council (NHMRC): http://www.nhmrc.gov.au/ 2. Therapeutic Goods Administration (TGA): http://www.tga.gov.au/ | | | NHMRC, ARC, and UA: National Statement on Ethical Conduct in Human Research (2018): Chapter 3.2: https://www.nhmrc.gov.au/research-policy/ethics/national-statement-ethical-conduct-human-research TGA: Australian Regulatory Guidelines for Biologicals (2017): http://www.tga.gov.au/industry/biologicals-argb.htm |
| <i>Genetic Research</i> | 1. National Health and Medical Research Council (NHMRC): http://www.nhmrc.gov.au/ 2. Office of the Gene Technology Regulator: http://www.ogtr.gov.au/ | Gene Technology Act 2000 (2016): https://www.legislation.gov.au/Details/C2016C00792 | Gene Technology Regulations 2001 (2016): https://www.legislation.gov.au/Details/F2016C00615 | NHMRC, ARC, and UA: National Statement on Ethical Conduct in Human Research, Chapter 3.3 (2018): http://www.nhmrc.gov.au/guidelines/publications/e72 |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. National Health and Medical Research Council (NHMRC): http://www.nhmrc.gov.au/ 2. National Health and Medical Research Council: Embryo Research Licensing Committee: https://nhmrc.gov.au/embryo-research-licensing-committee | 1. Prohibition of Human Cloning for Reproduction Act 2002 (2008): http://www.comlaw.gov.au/Details/C2008C00694 2. Research Involving Human Embryos Act 2002 (2014): http://www.comlaw.gov.au/Details/C2014C00605 | Research Involving Human Embryos Regulations (2017): https://www.legislation.gov.au/Details/F2017L01213 | NHMRC, ARC, and UA: National Statement on Ethical Conduct in Human Research, Chapter 3.2 (2018): https://www.nhmrc.gov.au/research-policy/ethics/national-statement-ethical-conduct-human-research NHMRC: Ethical Guidelines on the Use of Assisted Reproductive Technology in Clinical Practice and Research (2017): |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| | | | | https://nhmrc.gov.au/about-us/publications/ethical-guidelines-use-assisted-reproductive-technology |
| Bangladesh | | | | |
| <i>General</i> | Bangladesh Medical Research Council, National Research Ethics Committee: http://www.bmrcbd.org | | | 1. Ethical Guidelines for Conducting Research Studies Involving Human Subjects: https://www.bmrcbd.org/application_form/EthicalGuideline 2. Standard Operating Procedures (SOPs): https://www.bmrcbd.org/application_form/SOPs |
| <i>Drugs, Biologics, and Devices</i> | Bangladesh Directorate of Drug Administration: http://www.dgda.gov.bd/ | 1. The Drugs Act (1964) 2. Drugs (Control) Ordinance 1982, Ordinance No. VIII: http://bdlaws.minlaw.gov.bd/pdf_art.php?id=623 | | Good Clinical Practice (GCP) Guidelines: http://www.dgda.gov.bd/index.php/2013-03-31-05-16-29/registered-medical-device-list-4/129-good-clinical-practice-gcp-guidelines/file |
| <i>Human Biological Materials</i> | Bangladesh Medical Research Council, National Research Ethics Committee: http://www.bmrcbd.org | | | Guidelines for Transfer of Human Biological Materials Abroad for Research Purposes (2004) |
| China, People's Republic of | | | | |
| For an overview of clinical research regulations in China, see the ClinRegs report: http://clinregs.niaid.nih.gov/single_country.php?c_id=44 | | | | |
| <i>General</i> | 1. National Health Commission of the People's Republic of China (NHC): http://en.nhfpc.gov.cn 2. China Food and Drug Administration: http://eng.sfda.gov.cn/WS03/CL0755/ | Law on Practicing Doctors (June 26, 1998), Articles 26 and 37: http://www.gov.cn/banshi/2005-08/01/content_18970.htm | | NHFPC: Guidelines on Ethical Review of Biomedical Research Involving Human Subjects (2016): http://www.moh.gov.cn/fzs/s3576/201610/84b33b81d8e747eaaf048f68b174f829.shtml NHFPC, CFDA, and State Administration of TCM: Management Guidelines for Conducting Clinical Research at Medical/Health Institutions (Mandarin) (2014): http://www.nhfpc.gov.cn/yzygj/s3593g/201410/9bd03858c3aa41ed8aed17467645fb68.shtml |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> China Food and Drug Administration: http://eng.sfda.gov.cn/WS03/CL0755/ | Drug Administration Law of the People's Republic of China (2001): http://eng.sfda.gov.cn/WS03/CL0766/61638.html | 1. Regulations for Implementation of the Drug Administration Law of the People's Republic of China (2002): http://eng.sfda.gov.cn/WS03/CL0767/61640.html | 1. Guideline for HIV Vaccine Research Technology (2003): http://samr.cfda.gov.cn/WS01/CL0237/15705.html 2. Guideline for Vaccine Research Technology (2004): http://samr.cfda.gov.cn/WS01/CL0055/10307.h |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <p><i>Drugs, Biologics, and Devices</i></p> | | | <p>2. Chinese Good Clinical Practice (2003): http://www.sfda.gov.cn/WS01/CL0053/24473.html</p> <p>3. Special Review and Approval Procedure for Drug Registration of the State Food and Drug Administration (2005): http://eng.sfda.gov.cn/WS03/CL0768/61646.html</p> <p>4. Provisions for Drug Registration (2007): http://eng.sfda.gov.cn/WS03/CL0768/61645.html</p> <p>5. Qualification and Evaluation of Clinical Trial Sites/Institutions (2008): http://samr.cfda.gov.cn/WS01/CL0121/29571.html</p> <p>6. Rules on the Administration of Report and Supervision of Adverse Drug Reactions (2011): http://samr.cfda.gov.cn/WS01/CL1031/62621.html</p> | <p>tml</p> <p>3. Guidelines on Ethical Review of Drug Clinical Trials (2010): http://samr.cfda.gov.cn/WS01/CL0058/55613.html</p> <p>4. Interim Guidelines on International Multi-Regional Drug Clinical Trials (2015): http://samr.cfda.gov.cn/WS01/CL0087/114002.html</p> |
| | <p><i>Devices</i></p> <p>China Food and Drug Administration: http://eng.sfda.gov.cn/WS03/CL0755/</p> | | <p>CFDA and NHFPC: Good Clinical Practice on Medical Device Clinical Trials (2016): http://samr.cfda.gov.cn/WS01/CL0053/148101.html</p> | <p>CFDA:</p> <p>1. Guiding Principles of the Clinical Trail Technology on In Vitro Diagnostic (IVD) Reagents (2014): http://samr.cfda.gov.cn/WS01/CL0087/106241.html</p> <p>2. Templates for Medical Device Clinical Trials – Ethical Application and Approval:</p> <ol style="list-style-type: none"> 1. Ethical Review Application And Review Form 2. Informed Consent Form 3. CRF Template 4. Protocol Template 5. Report Template 6. Required Documents List <p><i>Access:</i> http://samr.cfda.gov.cn/WS01/CL0087/148126.html</p> |
| <p><i>Clinical Trials</i></p> | <p>Chinese Clinical Trial Registry: http://www.chictr.org.cn/index.aspx</p> | | | <p>FAQs: http://www.chictr.org.cn/question.aspx</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Registry</i> | | | | |
| <i>Privacy/Data Protection</i> | <i>Mainland:</i> | | | |
| | National Information Security Standardization Technical Committee: https://www.tc260.org.cn/ | People's Republic of China Cyber Security Law (2016): http://www.npc.gov.cn/npc/xinwen/2016-11/07/content_2001605.htm | | Information Security Technology-Personal Information Security Specification (2017, GB/T 35273-2017): https://www.tc260.org.cn/front/postDetail.html?id=20180124211617 |
| <i>Privacy/Data Protection</i> | <i>Hong Kong:</i> | | | |
| | 1. Privacy Commissioner for Personal Data, Hong Kong: http://www.pcpd.org.hk 2. eHealth Electronic Health Record Sharing System: https://www.ehealth.gov.hk/en/about_ehrs/ehr_office/index.html | Personal Data (Privacy) Ordinance (2013): https://www.elegislation.gov.hk/hk/cap486:en@2018-04-20T00:00:00?xid=ID_1438403261084_001 | | 1. Electronic Health Record Sharing System and Your Personal Data Privacy (10 Privacy Protection Tips): https://www.pcpd.org.hk/english/data_privacy_law/electronic_health_record_sharing_system/files/eHRSS_10_Tips_ENG.pdf 2. Personal Data (Privacy) Ordinance and Electronic Health Record Sharing System: https://www.pcpd.org.hk/english/data_privacy_law/electronic_health_record_sharing_system/files/eHRSS_Points_to_Notes_ENG.pdf |
| <i>Research Injury</i> | 1. National Health Commission of the People's Republic of China (NHC): http://www.nhfpc.gov.cn/ 2. Chinese Food and Drug Administration (CFDA): http://eng.sfda.gov.cn/WS03/CL0755/ | Chinese Good Clinical Practice, Article 43 (2003): http://www.sda.gov.cn/WS01/CL0053/24473.html | NHFPC: 1. Regulations on Recall of Medical Devices (Interim), Article 37 (2011): http://www.gov.cn/flfg/2011-06/13/content_1882686.htm 2. Guidelines on Ethical Review of Biomedical Research Involving Human Subjects (2016), Articles 18.5, 20.8, 36.6, 37: http://www.moh.gov.cn/fzs/s3576/201610/84b33b81d8e747eaaaf048f68b174f829.shtml CFDA and NHFPC: Good Clinical Practice on Medical Device Clinical Trials (2016), Articles 10, 22, 33, and 48: http://samr.cfda.gov.cn/WS01/CL0053/148101.html | SFDA: 1. Guideline on Vaccine Clinical Trials, Part 6 (2004): http://samr.cfda.gov.cn/WS01/CL0844/10307.html 2. Guideline on Ethical Review of Drug Clinical Trials, Appendix 1, Section 6.10 (2010): http://samr.cfda.gov.cn/WS01/CL0058/55613.html |
| <i>Genetic Research</i> | 1. National Health Commission of the People's Republic of China (NHC): http://www.nhfpc.gov.cn/ 2. Ministry of Science and Technology of the People's Republic of China (MOST): | | NHFPC and MOST: 1. Interim Measures for the Administration of Human Genetic Resources (1998): http://www.most.gov.cn/bszn/new/rlyc/wjxz/200512/t20051226_55327.htm | MOST: Service Guidelines for the Collection, Selling, Export and Admission Application of Human Genetic Resources (2015): http://www.most.gov.cn/tztg/201507/t2015070 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | http://www.most.cn/eng/ | | m 2. Regulations for the Administration of Human Genetic Resources (2012, public comment version): http://www.gov.cn/gzdt/2012-10/31/content_2254379.htm | 3_120547.htm |
| <i>Embryos, Stem Cells, and Cloning</i> | <i>Mainland:</i> 1. National Health Commission of the People's Republic of China (NHC): http://www.nhfpc.gov.cn/ 2. Ministry of Science and Technology of the People's Republic of China (MOST): http://www.most.cn/eng/ | | NHFPC: 1. Ethical Principles and Conduct Norms for Human Assisted Reproductive Technologies. (2003): http://www.moh.gov.cn/qjjys/s3581/200805/f69a925d55b44be2a9b4ada7fcdec835.shtml 2. Regulation on the Clinical Application of Medical Technique (2009) http://www.moh.gov.cn/yzygi/s3589/201308/0c579ba3babf47dc8f0e811810d438a2.shtml NHFPC and CFDA Interim Measures for the Management of Stem Cell Clinical Research (2015): http://www.nhfpc.gov.cn/qjjys/s3581/201508/28635ef99c5743e294f45e8b29c72309.shtml | NHFPC and MOST: Ethical Guidelines for Research on Human Embryo Stem Cells (2003): http://www.most.gov.cn/fggw/zfwj/zfwj2003/200512/t20051214_54948.htm |
| | <i>Hong Kong:</i> Legislative Council of the Hong Kong Special Administrative Region of the People's Republic of China: http://www.legco.gov.hk/index.html | | Human Reproductive Technology (Amendment) Ordinance 2016: http://www.legco.gov.hk/yr15-16/english/ord/ord020-2016-e.pdf | |
| India For an overview of the clinical research regulations in India, see the ClinRegs report: http://clinregs.niaid.nih.gov/single_country.php?c_id=100 | | | | |
| <i>General</i> | Indian Council of Medical Research (ICMR): http://www.icmr.nic.in/ | | | 1. National Ethical Guidelines For Biomedical and Health Research Involving Human Participants (2017): http://icmr.nic.in/guidelines/ICMR_Ethical_Guidelines_2017.pdf 2. National Ethical Guidelines for Biomedical Research Involving Children |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | | | | (2017): http://icmr.nic.in/guidelines/National_Ethical_Guidelines_for_BioMedical_Research_Involving_Children.pdf |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | Schedule Y of the Drugs and Cosmetics Act (2016): http://www.cdsc.nic.in/writereaddata/2016Drugs%20and%20Cosmetics%20Act%201940%20&%20Rules%201945.pdf | DCGI: 1. Good Clinical Practices for Clinical Research in India (2001): http://rgcb.res.in/wp-content/uploads/2014/07/Good-Clinical-Practice-Guideline.pdf 2. Permission for Clinical Trials: General Statutory Rules 63(E) (2013) 3. Ethics Committee Registration: General Statutory Rules 72(E) (2013) 4. A/V Consent – General Statutory Rules 611 (E) (2015) 5. Phytopharmaceutical Drug: General Statutory Rules 918(E) (2015) 6. Exemption for Academic Research and Animal Toxicity: General Statutory Rules 313(E) (2016) | ICMR: National Ethical Guidelines for Biomedical and Health Research Involving Human Participants, Section 7 (2017): http://icmr.nic.in/guidelines/ICMR_Ethical_Guidelines_2017.pdf |
| | <i>Devices</i> | Drugs & Cosmetics Act, 1940 (2005): https://cdsco.gov.in/opencms/opencms/en/Notifications/Gazette-Notifications/ | CDSCO: 1. Medical Devices Rules, 2017. 2. General Statutory Rules 78(E) http://134t7045rwgf19lpbh29libk9d3.wpengine.netdna-cdn.com/wp-content/uploads/sites/11/2017/07/India-Medical-Device-Rules.pdf | ICMR: National Ethical Guidelines For Biomedical and Health Research Involving Human Participants, Section 7.7 (2017): http://icmr.nic.in/guidelines/ICMR_Ethical_Guidelines_2017.pdf |
| <i>Clinical Trials Registry</i> | 1. Clinical Trials Registry – India: http://ctri.nic.in/ 2. Office of Drugs Controller General | | | Clinical Trials Registry – India: FAQs: http://ctri.nic.in/Clinicaltrials/faq.php Office of Drugs Controller General: Registration of Clinical Trial in ICMR Clinical Trial Registry: http://www.cdsc.nic.in/writereaddata/CTRegistration.doc |
| <i>Research Injury</i> | 1. Central Drugs Standard Control Organization (CDSCO): https://cdsco.gov.in/opencms/opencms/ | Drugs & Cosmetics Act, 1940 (2005): http://www.cdsc.nic.in/writereaddata/ | DCGI: CDSCO: 1. Compensation: General | ICMR: National Ethical Guidelines For Biomedical and Health Research |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Social-Behavioral Research</i> | Indian Council of Medical Research (ICMR): http://www.icmr.nic.in/ | | | National Ethical Guidelines For Biomedical and Health Research Involving Human Participants, Section 9 (2017): http://icmr.nic.in/guidelines/ICMR_Ethical_Guidelines_2017.pdf |
| <i>Privacy/Data Protection</i> | Indian Council of Medical Research (ICMR): http://www.icmr.nic.in/ | | | National Ethical Guidelines for Biomedical and Health Research Involving Human Participants, Sections 2 and 11 (2017): http://icmr.nic.in/guidelines/ICMR_Ethical_Guidelines_2017.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | Ministry of Health and Family Welfare: https://mohfw.gov.in | | Govt. of India Office Memorandum (O.M. No.19015/53/1997 - IH Pt.) 19 th November, 1997 on Exchange of Human Biological Material for Biomedical Research Purposes | National Ethical Guidelines For Biomedical and Health Research Involving Human Participants, Section 11 (2017): http://icmr.nic.in/guidelines/ICMR_Ethical_Guidelines_2017.pdf |
| <i>Genetic Research</i> | 1. Department of Biotechnology (DBT): http://dbtindia.nic.in/ 2. Indian Council of Medical Research (ICMR): http://www.icmr.nic.in/ | Environmental Protection Act (1986) | | DBT: 1. Recombinant DNA Safety Guidelines (1990) 2. Ethical Policies on the Human Genome, Genetic Research, and Services (2002) ICMR: National Ethical Guidelines For Biomedical and Health Research Involving Human Participants, Section 10 (2017): http://icmr.nic.in/guidelines/ICMR_Ethical_Guidelines_2017.pdf |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Indian Council of Medical Research (ICMR): http://www.icmr.nic.in/ 2. Central Drugs Standard Control Organization, Office of Drugs Controller General of India (DCGI): https://cdsco.gov.in | | | National Guidelines for Stem Cell Research (2017): http://icmr.nic.in/guidelines/Guidelines_for_stem_cell_research_2017.pdf |
| Indonesia | | | | |
| For an overview of health research ethics, see: http://www.fercap-sidcer.org/newsletter/2013/12/PPT/04%20Suriadi%20Guwanan-PPT.pdf | | | | |
| <i>General</i> | Ministry of Health, National Institute of Health Research and Development: http://indonesia.go.id/en | Indonesian Health Act No. 23/1992 Section on Health Research, Article 69 | 1. Regulation No. 39/1995 on Health Research and Development 2. Presidential Decree No. 100/1993: Research by Foreigners | National Guidelines on Ethics in Health Research (2003) |
| <i>Drugs, Biologics, and Devices</i> | National Agency of Drug and Food Control: http://www.pom.go.id/index.php/home/en | | 1. Ministry of Health Decree No. 56/2000: Guidelines on Clinical Trials of Traditional Drugs 2. Guidelines on Good Clinical Practice (2001) | |
| <i>Human Biological Materials</i> | | | National Guidelines on Use of Stored Biological Materials (2005) | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Japan | | | | |
| <i>General</i> | 1. Ministry of Education, Culture, Sports, Science, and Technology (MEXT): http://www.mext.go.jp/english/ 2. Ministry of Health, Labor, and Welfare (MHLW): http://www.mhlw.go.jp/english/index.html | | | MEXT and MHLW: Ethics Guidelines for Medical and Health Research Involving Human Subjects (2017): http://www.lifescience.mext.go.jp/files/pdf/n1859_01.pdf 2015 version (English): http://www.lifescience.mext.go.jp/files/pdf/n1500_01.pdf |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | 1. Pharmaceuticals, Medical Devices, and Other Therapeutic Products Act (2016): http://law.e-gov.go.jp/htmldata/S35/S35HO145.html 2. Clinical Trials Act (2017): https://www.mhlw.go.jp/file/06-Seisakujouhou-10800000-Iseikyoku/0000213334.pdf | MHLW: Ministerial Ordinance on Good Clinical Practice for Drugs (2016): http://elaws.e-gov.go.jp/search/elawsSearch/elaws_search/lsg0500/detail?lawId=409M50000100028&openerCode=1 | |
| | <i>Devices</i> | 1. Pharmaceuticals, Medical Devices, and Other Therapeutic Products Act (2016): http://law.e-gov.go.jp/htmldata/S35/S35HO145.html 2. Clinical Trials Act (2017): https://www.mhlw.go.jp/file/06-Seisakujouhou-10800000-Iseikyoku/0000213334.pdf | MHLW: Ministerial Ordinance on Good Clinical Practice for Medical Devices (2016): http://law.e-gov.go.jp/htmldata/H17/H17F19001000036.html 2009 version (English): https://www.pmda.go.jp/files/000153732.pdf | |
| <i>Clinical Trials Registry</i> | 1. Ministry of Health, Labor, and Welfare (MHLW): http://www.mhlw.go.jp/english/index.html 2. National Institute of Public Health: https://www.niph.go.jp/index_en.html 3. Japan Registry of Clinical Trials: https://jrct.niph.go.jp/ | Clinical Trials Act (2017): https://www.mhlw.go.jp/file/06-Seisakujouhou-10800000-Iseikyoku/0000213334.pdf | | NIPH Clinical Trials Search: http://rctportal.niph.go.jp/en/ |
| <i>Privacy/Data Protection</i> | 1. Personal Information Protection Commission: http://www.ppc.go.jp/en/ 2. Office of Healthcare Policy of the | 1. Amended Act on the Protection of Personal Information (2017): https://www.ppc.go.jp/files/pdf/Act | 1. Amendment to the Cabinet Order to Enforce the Act on the Protection of Personal Information (2016): | Guidelines for Act on the Protection of Personal Information (2016): https://www.ppc.go.jp/files/pdf/guidelines01.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Privacy/Data Protection</i> | Cabinet Secretariat: http://www.kantei.go.jp/jp/singi/kenkouiryou/en/ | on the Protection of Personal Information.pdf 2. Act Regarding Anonymized Medical Data to Contribute to R&D in the Medical Field (2017): http://www.kantei.go.jp/jp/singi/kenkouiryou/jisedai_kiban/pdf/170310shiryu3.pdf | https://www.ppc.go.jp/files/pdf/Cabinet_Order.pdf 2. Enforcement Rules for the Act on the Protection of Personal Information (2016): https://www.ppc.go.jp/files/pdf/PPC_rules.pdf 3. Regulation for Enforcement of the Clinical Trials Act, Article 20 (2018): https://www.mhlw.go.jp/file/06-Seisakujouhou-10800000-Iseikyoku/0000195391.pdf | https://www.ppc.go.jp/files/pdf/guidelines02.pdf https://www.ppc.go.jp/files/pdf/guidelines03.pdf https://www.ppc.go.jp/files/pdf/guidelines04.pdf |
| <i>Research Injury</i> | Ministry of Health, Labor, and Welfare (MHLW): http://www.mhlw.go.jp/english/index.html | 1. Pharmaceuticals, Medical Devices, and Other Therapeutic Products Act (2016): http://law.e-gov.go.jp/htmldata/S35/S35HO145.html 2. Clinical Trials Act (2017): https://www.mhlw.go.jp/file/06-Seisakujouhou-10800000-Iseikyoku/0000213334.pdf | 1. Ministerial Ordinance on Good Clinical Practice for Drugs, Article 14 (2016): http://law.e-gov.go.jp/htmldata/H09/H09F03601000028.html 2. Ministerial Ordinance on Good Clinical Practice for Medical Devices, Article 14 and 23 (2016): http://law.e-gov.go.jp/htmldata/H17/H17F19001000036.html | Ethics Guidelines for Medical and Health Research Involving Human Subjects, Chapter 2, No. 5, 1-(3) and No. 6, 2-(2) (2017): http://www.lifescience.mext.go.jp/files/pdf/n1859_01.pdf |
| <i>Human Biological Materials</i> | Ministry of Health, Labor, and Welfare (MHLW): http://www.mhlw.go.jp/english/index.html | | | On Research and Development Utilizing Human Tissues Removed by Surgery and Other Procedures (1998) (Japanese): https://www.mhlw.go.jp/www1/shingi/s9812/s1216-2_10.html |
| <i>Genetic Research</i> | 1. Council for Science, Technology, and Innovation (CSTI): http://www8.cao.go.jp/cstp/english/index.html 2. Ministry of Education, Culture, Sports, Science, and Technology (MEXT): http://www.mext.go.jp/english/ 3. Ministry of Health, Labor, and Welfare (MHLW): http://www.mhlw.go.jp/english/index.html 4. Ministry of Economy, Trade, and Industry (METI): http://www.meti.go.jp/english/ | | | CSTI: Fundamental Principles of Research on the Human Genome (2000) MEXT, MHLW, and METI: Ethics Guidelines for Human Genome/Gene Analysis Research (2017): http://www.lifescience.mext.go.jp/files/pdf/n1859_03r2.pdf 2008 version (English): http://www.lifescience.mext.go.jp/files/pdf/n796_00.pdf MHLW: Guidelines for Clinical Research in Gene Therapy and Others (2017): |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | | | | https://www.mhlw.go.jp/file/06-Seisakujouhou-10600000-Daijinkanboukouseikagakuka/0000161224.pdf |
| <i>Embryos, Stem Cells, and Cloning</i> | <p>1. Council for Science, Technology, and Innovation (CSTI): http://www8.cao.go.jp/cstp/english/index.html</p> <p>2. Ministry of Education, Culture, Sports, Science, and Technology (MEXT): http://www.mext.go.jp/english/</p> | <p>1. Act on Regulation of Human Cloning Techniques (2014): http://law.e-gov.go.jp/htmldata/H12/H12HO146.html</p> <p>2000 version (English): http://www.cas.go.jp/jp/seisaku/hourei/data/htc.pdf</p> <p>2. Act on Safety of Regenerative Medicine (2013): http://www.mhlw.go.jp/file/06-Seisakujouhou-10800000-Iseikyoku/0000030847.pdf</p> | <p>1. Ordinance for Enforcement of Act on Regulation of Human Cloning Techniques (2009): http://www.lifescience.mext.go.jp/files/pdf/n1564_01.pdf</p> <p>2. Ordinance for Enforcement of Act on Safety of Regenerative Medicine (2014): https://www.mhlw.go.jp/file/06-Seisakujouhou-10800000-Iseikyoku/0000059294.pdf</p> <p>3. Rules for Enforcement of Act on Safety of Regenerative Medicine (2014): http://www.mhlw.go.jp/file/06-Seisakujouhou-10800000-Iseikyoku/0000065532.pdf</p> | <p>CSTP: Fundamental Philosophy on Handling of Human Embryo (2004): http://www.lifescience.mext.go.jp/files/pdf/6_28.pdf</p> <p>MEXT: 1. Guidelines on the Handling of a Specified Embryo (2009): http://www.lifescience.mext.go.jp/files/pdf/n1564_02.pdf</p> <p>2. Guidelines on the Derivation of Human Embryonic Stem Cells (2014): http://www.lifescience.mext.go.jp/files/pdf/n1553_01.pdf</p> <p>3. Guidelines on the Distribution and Utilization of Human Embryonic Stem Cells (2014): http://www.lifescience.mext.go.jp/files/pdf/n1553_02r2.pdf</p> <p>4. Guidelines on Research on Producing Germ Cells from Human Induced Pluripotent Stem Cells or Human Tissue Stem Cells (2015): http://www.lifescience.mext.go.jp/files/pdf/n1492_01r2.pdf</p> <p>2010 version (English): http://www.lifescience.mext.go.jp/files/pdf/n1567_02r2.pdf</p> <p>MEXT and MHLW: Ethical Guidelines for Research on Assisted Reproductive Technology to Develop Human Fertilized Embryos (2017): http://www.lifescience.mext.go.jp/files/pdf/n1859_05.pdf</p> <p>2010 version (English): http://www.lifescience.mext.go.jp/files/pdf/n1567_01r2.pdf</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Kazakhstan | | | | |
| Note: For an overview of human subject protections in Kazakhstan, see “Ethical Review of Biomedical Research in the CIS Countries,” Chapter 3, Section 5: http://www.unesco.org/new/fileadmin/MULTIMEDIA/FIELD/Moscow/pdf/ethical_review_cis_book_kubar_english.pdf | | | | |
| <i>General</i> | Ministry of Healthcare and Social Development, Central Commission on Research Ethics: http://www.mzsr.gov.kz/en | | | 1. Guidelines on Ethics in Health Research. (2007) 2. Local Ethics Committees: Policy, Rules and Procedures (2014) 3. Guidelines on Ethics in Biomedical Research (2015) |
| <i>Drugs, Biologics, and Devices</i> | Ministry of Healthcare and Social Development, Control Committee of Medical and Pharmacy Activity: https://www.mzsr.gov.kz/en/taxonomy/term/674 | Code of the Republic of Kazakhstan "On People's Health and the Health Care System" (18.09.2009 No.193-IV), Articles 74 and 180 (2015): http://online.zakon.kz/Document/?doc_id=30479065#pos=1;-8 | 1. Order of the MHS D of the RK Dated 12.11.2009 No. 697 on the Approval of Regulations on the Medical-Biological Experiments, Preclinical (Non-Clinical) and Clinical Trials 2. Order of the MHS D of the RK dated 19.11.2009 No. 744 on the Approval of Regulations on the Conduct of Clinical Trials and/or Trials on Pharmaceutical and Drug Products, Medical Devices, and Medical Equipment 3. Order of the MHS D Dated 20.05.2014 No.272 on the Approval of Regulations on the Implementation of the New Methods of Diagnostic, Treatment, and Rehabilitation | Guidelines on Clinical Trials in Kazakhstan (2003) |
| <i>Privacy/Data protection</i> | Ministry of Healthcare and Social Development: http://www.mzsr.gov.kz/en | Code of the Republic of Kazakhstan “On People's Health and the Health Care System” (18.09.2009 No.193-IV), Article 28 (2015): http://online.zakon.kz/Document/?doc_id=30479065#pos=1;-8 | | |
| Korea | | | | |
| Note: All documents are in Korean. | | | | |
| <i>General</i> | Ministry of Health and Welfare: http://www.mohw.go.kr/eng/index.jsp | Bioethics and Safety Act No. No. 15188 (2017.12): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=46341&lang=ENG | 1. Enforcement Decree of Pharmaceutical Affairs Act No. 28821 (2017.7): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=2&query=BIOETHICS%20AND%20SAFETY%20ACT#liBgcolor8 | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | | | 2. Enforcement Rule of Bioethics and Safety Act No. 143 (2009.12): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=2&query=BIOETHICS%20AND%20SAFETY%20ACT#liBgcolor11 | |
| <i>Drugs, Biologics, and Devices</i> | <p data-bbox="346 344 420 370"><i>Drugs</i></p> <p data-bbox="346 383 714 467">Ministry of Food and Drug Safety (MFDS) (2013): http://www.mfds.go.kr/eng/index.do</p> <p data-bbox="346 1110 436 1133"><i>Devices</i></p> <p data-bbox="346 1156 714 1211">Ministry of Food and Drug Safety: http://www.mfds.go.kr/eng/index.do</p> | <p data-bbox="783 383 1129 496">Medical Device Act No. 15486 (2018.3): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=48691&lang=ENG</p> <p data-bbox="783 1156 1129 1247">Medical Device Act No. 15486 (2018): http://www.law.go.kr/ 법령/의료기기법</p> | <p data-bbox="1157 383 1528 522">1. Enforcement Decree of Pharmaceutical Affairs Act No. 27673 (2016.12): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=40268&lang=ENG</p> <p data-bbox="1157 529 1528 782">2. Regulation on Safety of Medicinal Products, etc. No. 1089(2014.8): http://www.mfds.go.kr/eng/brd/m_18/view.do?seq=69740&srchFr=&srchTo=&srchWord=&srchTp=&itm_seq_1=0&itm_seq_2=0&multi_itm_seq=0&company_cd=&company_nm=&page=3</p> <p data-bbox="1157 789 1528 1107">3. Regulations for Clinical Trial Personnel Education and Certification for the Educational Institution No. 2017-79 (2017): https://www.mfds.go.kr/brd/m_207/view.do?seq=13405&srchFr=&srchTo=&srchWord=%EC%A2%85%EC%82%AC%EC%9E%90&srchTp=0&itm_seq_1=0&itm_seq_2=0&multi_itm_seq=0&company_cd=&company_nm=&page=1</p> <p data-bbox="1157 1156 1528 1474">1. Enforcement Decree of the Medical Device Act No. 28224 (2017.8): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=44278&lang=ENG</p> <p data-bbox="1157 1302 1528 1474">2. Enforcement Regulations of the Medical Device Act No. 18 (2010.9): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=1&query=Medical+Device+&x=0&y=0#liBgcolor5</p> | <p data-bbox="1556 383 2020 821">MFDS: 1. Guidelines on Human Research Protection Program 0053-01 (2014.3) 2017-.5.30 告示: http://www.mfds.go.kr/brd/m_210/view.do?seq=12203 2. IND regulations No 2018-42 (2018): http://www.law.go.kr/admRulSc.do?menuId=1&query=%EC%9D%98%EC%95%BD%ED%92%88%20%EC%9E%84%EC%83%81%EC%8B%9C%ED%97%98%20%EA%B3%84%ED%9A%8D%20%EC%8A%B9%EC%9D%B8%EC%97%90%20%EA%B4%80%ED%95%9C%20%EA%B7%9C%EC%A0%95#liBgcolor0</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Clinical Trials Registry</i> | Korea Centers for Disease Control and Prevention (KCDC), Clinical Research Information Service: https://cris.nih.go.kr/cris/en/index.jsp?mobile= | | | |
| <i>Research Injury</i> | Ministry of Food and Drug Safety (MFDS): http://www.mfds.go.kr/eng/index.do | | Regulation on Safety of Medicinal Products, etc. No. 1089(2014.8): http://www.mfds.go.kr/eng/brd/m_18/view.do?seq=69740&srchFr=&srchTo=&srchWord=&srchTp=&itm_seq_1=0&itm_seq_2=0&multi_itm_seq=0&company_cd=&company_nm=&page=3 | Guidelines for Clinical Trial Indemnity and Its Process 0053-01 (2013.10) 2017.6.1 고시: http://www.mfds.go.kr/brd/m_210/view.do?seq=13069&srchFr=&srchTo=&srchWord=%EB%B3%B4%EC%83%81&srchTp=0&itm_seq_1=0&itm_seq_2=0&multi_itm_seq=0&company_cd=&company_nm=&page=1 2. Guidance for Sponsors; Safety Reporting Requirements 0785-01 (2017.8) 2017.8.31 고시: http://www.mfds.go.kr/brd/m_210/view.do?seq=13317&srchFr=&srchTo=&srchWord=%EC%95%88%EC%A0%84%EC%84%B1&srchTp=0&itm_seq_1=0&itm_seq_2=0&multi_itm_seq=0&company_cd=&company_nm=&page=1 |
| <i>Privacy/Data Protection</i> | 1. Ministry of the Interior and Safety (MOIS): http://www.mois.go.kr/eng/a01/engMain.do 2. Ministry of Health and Welfare (MOHW): http://www.mohw.go.kr/eng/index.jsp | MOIS: Personal Information Protection Act No. 14839 (2017): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=46731&lang=ENG MOHW: Bioethics and Safety Act No. 15188 (2017.12): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=46341&lang=ENG | MOIS: 1. Enforcement Rules to Personal Information Protection Act No. 1 (2013.3): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=1&query=personal&x=0&y=0#liBgcolor21 2. Enforcement Decrees to Personal Information Protection Act No. 28355 (2017.10): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=45683&lang=ENG MOHW: Enforcement Rule of Bioethics and Safety Act No. 143 (2009.12): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=1&query=BIOETHI | MOIS: Act on the Protection of Personal Information maintained by Public Institutions No. 8871 (2008.2): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=1&query=personal+information+protection&x=13&y=29#liBgcolor9 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Privacy/Data Protection</i> | | | CS+AND+SAFETY+ACT&x=0&y=0#liBgcolor11 | |
| <i>Human Biological Materials</i> | 1. Ministry of Health and Welfare (MOHW): http://www.mohw.go.kr/eng/index.jsp 2. Ministry of Food and Drug Safety (MFDS): http://www.mfds.go.kr/eng/index.do | MOHW: Bioethics and Safety Act No. 15188 (2017.12): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=46341&lang=ENG | 1. Enforcement Decree of Bioethics and Safety Act No. 28211 (2017): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=45482&lang=ENG 2. Enforcement Rule of Bioethics and Safety Act No. 143 (2009.12): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=1&query=BIOETHICS+AND+SAFETY+ACT&x=0&y=0#liBgcolor11 | |
| <i>Genetic Research</i> | 1. Ministry of Health and Welfare (MOHW): http://www.mohw.go.kr/eng/index.jsp 2. Ministry of Food and Drug Safety (MFDS): http://www.mfds.go.kr/eng/index.do | MOHW: Bioethics and Safety Act No. 15188 (2017.12): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=46341&lang=ENG | MOHW: 1. Enforcement Decree of Bioethics and Safety Act No. 28211 (2017): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=45482&lang=ENG 2. Enforcement Rule of Bioethics and Safety Act No. 143 (2009.12): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=1&query=BIOETHICS+AND+SAFETY+ACT&x=0&y=0#liBgcolor11 | MFDS: Guidelines on the Evaluation of Quality, Safety, and Efficacy of Recombinant Protein Products 0324-01 (2014.12) 2017.6.1 고시: http://www.mfds.go.kr/brd/m_210/view.do?seq=12542&srchFr=&srchTo=&srchWord=%EC%9E%AC%EC%A1%B0%ED%95%A9&srchTp=0&itm_seq_1=0&itm_seq_2=0&multi_itm_seq=0&company_cd=&company_nm=&page=1 |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Ministry of Health and Welfare (MOHW): http://www.mohw.go.kr/eng/index.jsp 2. Ministry of Food and Drug Safety (MFDS): http://www.mfds.go.kr/eng/index.do | Bioethics and Safety Act No. 15188 (2017.12): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=46341&lang=ENG | MOHW: 1. Enforcement Decree of Bioethics and Safety Act No. 28211 (2017): http://elaw.klri.re.kr/kor_service/lawView.do?hseq=45482&lang=ENG 2. Enforcement Rule of Bioethics and Safety Act No. 143 (2009.12): http://www.law.go.kr/LSW/eng/engLsSc.do?menuId=1&query=BIOETHICS+AND+SAFETY+ACT&x=0&y=0#liBgcolor11 | MFDS: Guideline on Sponsor-Investigator Trials of Cell Therapy Products for Academic Purpose 0307-01 (2014.12) 2014.12.30 고시: http://www.mfds.go.kr/brd/m_210/view.do?seq=12490&srchFr=&srchTo=&srchWord=%EC%84%B8%ED%8F%AC%EC%B9%98%EB%A3%8C&srchTp=0&itm_seq_1=0&itm_seq_2=0&multi_itm_seq=0&company_cd=&company_nm=&page=1 |
| Kyrgyzstan | | | | |
| Note: All websites and documents are in Russian. | | | | |
| <i>General</i> | 1. Government of the Kyrgyz Republic: http://www.gov.kg 2. Ministry of Health: http://www.med.kg | 1. Constitution of Kyrgyz Republic, Chapter II, Article 22 (2010): http://www.gov.kg/?page_id=263&1 | 1. Code of Professional Ethics of Medical Worker of the Kyrgyz Republic (2004): http://old.med.kg/index.php/ru/doku | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | 3. Ministry of Justice of the Kyrgyz Republic: http://cbd.minjust.gov.kg | ang=ru 2. Law on Health Protection of the Kyrgyz Republic (Sept. 1, 2005, No. 6): Articles 34 and 72: http://www.pharm.kg/ru/legislation | menty-2/kodex-prof-etiki-2.html 2. Code of Administrative Responsibility of the Kyrgyz Republic №114 from 04.08.1998r. (Updated June 11, 2008 N 115 and June 23, 2008 N 136) Chapters 7 and 10: http://www.pharm.kg/ru/legislation/ | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health, Department of Drug and Medical Devices Provision (DDMDP): http://www.pharm.kg 2. Ministry of Health, National Bioethics Committee 3. Pharmaceutical Union of Kyrgyzstan, Ethics Committee: http://farmunion.kg/o-nas/eticheskij-komitet/ | Law on the Circulation of Medicinal Products of the Kyrgyz Republic, as amended by the Law of the Kyrgyz Republic of May 3, 2018 N 44, Chapter VII, Articles 24-25: http://cbd.minjust.gov.kg/act/view/ru-ru/111672 | DDMDP: 1. National Standard KMC 1195:2010: Medical Devices: Rules for Clinical Trials (2010): http://www.pharm.kg/ru/legislation/ 2. Technical Regulations on the Safety of Medical Products for Medical Application, Approved by the Governmental Order # 74 from February 1, 2012: http://www.pharm.kg/ru/legislation/ | |
| <i>Research Injury</i> | 1. Ministry of Health, Department of Drug and Medical Devices Provision (DDMDP): http://www.pharm.kg 2. Ministry of Health, National Bioethics Committee | Law on the Circulation of Medicinal Products of the Kyrgyz Republic, as amended by the Law of the Kyrgyz Republic of May 3, 2018 N 44, Chapter VII, Articles 24-25: http://cbd.minjust.gov.kg/act/view/ru-ru/111672 | DDMDP: National Standard KMC 1195:2010: Medical Devices, Rules for Clinical Trials, Paragraphs 3, 4, and 6 (2010): http://www.pharm.kg/ru/legislation/ | |
| <i>Human Biological Materials</i> | 1. Ministry of Health, Department of Drug and Medical Devices Provision: http://www.pharm.kg 2. Ministry of Health, National Bioethics Committee | Law on Health Protection of the Kyrgyz Republic (09.01.2005 No. 6): Article 39: http://www.pharm.kg/ru/legislation | Technical Regulations on the Safety of Medical Products for Medical Application, Approved by the Governmental Order #74 from February 1, 2012: http://www.pharm.kg/ru/legislation/ | |
| <i>Social-Behavioral Research</i> | Ministry of Justice of the Kyrgyz Republic: http://minjust.gov.kg/ru/ | Law On the Protection of Traditional Knowledge, as amended by the Law of the Kyrgyz Republic of July 18, 2014 No. 144): http://cbd.minjust.gov.kg/act/view/ru-ru/202149/20?cl=ru-ru | | |
| <i>Privacy/Data Protection</i> | 1. Ministry of Health, Department of Drug and Medical Devices Provision (DDMDP): http://www.pharm.kg 2. Ministry of Health, National | Law on Health Protection of the Kyrgyz Republic (09.01.2005 No. 6): Article 91: http://www.pharm.kg/ru/legislation | DDMDP: 1. National Standard KMC 1195:2010: Medical Devices, Rules for Clinical Trials, | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Privacy/Data Protection</i> | Bioethics Committee | | Paragraphs 3, 4, and 6 (2010): http://www.pharm.kg/ru/legislation/ 2. Technical Regulations on the Safety of Medical Products for Medical Application, Approved by the Governmental Order #74 from February 1, 2012: http://www.pharm.kg/ru/legislation/ | |
| Malaysia | | | | |
| <i>Drugs, Biologics, and Devices</i> | National Committee for Clinical Research: http://www.nccr.gov.my/ | | | 1. Guidelines for Ethical Review of Clinical Research or Research Involving Human Subjects (2006): http://www.nccr.gov.my/view_file.cfm?fileid=16 2. Malaysian Guidelines of Good Clinical Practice (2011): https://mrc.ukri.org/documents/pdf/malaysian-guidelines-for-good-clinical-practice/ |
| <i>Privacy/Data Protection</i> | | Act 709: Personal Data Protection Act 2010: http://www.pdp.gov.my/images/LAWS_OF_MALAYSIA_PDPA.pdf | | |
| <i>Human Biological Materials</i> | National Committee for Clinical Research: http://www.nccr.gov.my/ | 1. Act 130: Human Tissues Act (1974): http://www.agc.gov.my/Akta/Vol.%203/Act%20130.pdf 2. Act 699: DNA Identification Act 2009. Malaysian Government Gazette of 3 September 2009 | DNA Identification Regulations 2012. Malaysian Government Gazette of 30 Aug 2012. | Guideline on the Use of Human Biological Tissues for Research (2006): http://www.nccr.gov.my/index.cfm?menuid=25&parentid=17 |
| <i>Genetic Research</i> | Malaysian Medical Council: http://www.mmc.gov.my/v1/ | | | Medical Genetics and Genetic Services. MMC Guidelines 010/2006: http://www.mmc.gov.my/v1/docs/Medical%20Genetics%20&%20Genetic%20Services.pdf |
| <i>Embryos, Stem Cells and Cloning</i> | Ministry of Health, Medical Research and Ethics Committee | | | Checklist for Research on Stem Cell and Cell-Based Therapies (2014): http://www.nih.gov.my/mrec/documents/Research_On_Stem_cell_and_Cell_based_Therapies.pdf |
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| <i>General</i> | Nepal Health Research Council, Ethical Review Board: http://www.nhrc.org.np/ | Nepal Health Research Council Act, 1991, Section 3(1): http://www.lawcommission.gov.np/en/documents/2015/08/nepal-health-research-council-act-2047-1991.pdf | | 1. National Ethical Guidelines for Health Research in Nepal and Standard Operating Procedure (2011): http://nhrc.org.np/guidelines 2. Guidelines for Institutional Review Committees (IRCs) for Health Research in Nepal (2016): http://nhrc.org.np/guidelines |
| <i>Drugs, Biologics, and Devices</i> | Nepal Health Research Council: http://www.nhrc.org.np/ | | | National Guidelines on Clinical Trials with the Use of Pharmaceutical Products (2005): http://nhrc.org.np/guidelines |
| New Zealand | | | | |
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| | <i>Devices</i> | New Zealand Medicines and Medical Devices Safety Authority (Medsafe): http://www.medsafe.govt.nz | | Medicines (Database of Medical Devices) Regulations (2003): http://www.legislation.govt.nz/regulation/public/2003/0325/latest/DLM224223.html | 1. Standard Operating Procedures for Health and Disability Ethics Committees (2012): http://www.ethics.health.govt.nz/operating-procedures 2. Various: http://medsafe.govt.nz/regulatory/DevicesNew/13ConductingClinicalTrials.asp |
| | | Australian New Zealand Clinical Trials Registry: http://www.anzctr.org.au/ | | | FAQs: http://www.anzctr.org.au/Faq.aspx |
| <i>Privacy/Data Protection</i> | Privacy Commissioner: http://www.privacy.org.nz/ | 1. Official Information Act 1982 (2012) 2. Public Records Act (2005) 3. Privacy Act 1993 (2012) | Health Information Privacy Code 1994: http://www.privacy.org.nz/assets/Files/Codes-of-Practice-materials/Health-Information-Privacy-Code-1994-including-Amendment.pdf | | |

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| <i>Genetic Research</i> | 1. Environmental Protection Authority: http://www.epa.govt.nz/ 2. Health Research Council (HRC), Gene Technology Advisory Committee: http://www.hrc.govt.nz/about-us/committees/gene-technology-advisory-committee-gtac | Hazardous Substances and New Organisms Act 1996 (2012) | | |
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| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | | | | Access: https://acart.health.govt.nz/publications-and-resources/guidelines-and-advice-issued-ecart |
| Pakistan | | | | |
| <i>General</i> | National Bioethics Committee: http://nbcPakistan.org.pk/ | | | Various: http://nbcPakistan.org.pk/guidelines.html |
| <i>Drugs, Biologics, and Devices</i> | National Bioethics Committee: http://nbcPakistan.org.pk/ | | | Guidelines For Healthcare Professionals Interaction with Pharmaceutical Trade and Industry (PPI Guidelines): http://nbcPakistan.org.pk/?page_id=61 |
| <i>Human Biological Materials</i> | National Bioethics Committee: http://nbcPakistan.org.pk/ | | | Ethical Guidelines for Collection, Usage, Storage, and Export of Human Biological Materials (HBM): http://nbcPakistan.org.pk/assets/hbm-nbc-guidelines-final-18june-2016.pdf |
| <i>Embryos, Stem Cells, and Cloning</i> | National Bioethics Committee: http://nbcPakistan.org.pk/ | | | Protocol/Guidelines for Stem Cell Research/Regulation in Pakistan: http://nbcPakistan.org.pk/?page_id=61 |
| Philippines | | | | |
| <i>General</i> | <ol style="list-style-type: none"> Philippine Health Research Ethics Board (PHREB): www.ethics.healthresearch.ph Department of Science and Technology (DOST): http://www.dost.gov.ph/ Department of Health (DOH): http://www.doh.gov.ph/ Commission of Higher Education (CHED): www.ched.gov.ph/ National Commission for Indigenous Peoples (NCIP): www.ncip.gov.ph | <p>Republic Act No. 10532: An Act Institutionalizing the Philippine National Health Research System (2013): http://www.gov.ph/2013/05/07/repulic-act-no-10532/</p> | <p>PHREB: <ol style="list-style-type: none"> PNHRS Act Implementing Rules and Regulations: http://www.ethics.healthresearch.ph/index.php/component/content/article/2-uncategorised/214-implementing-rules-of-pnhrs Memorandum: Registration and Accreditation of all Ethics Review Committees in the Philippines (2015): http://www.ethics.healthresearch.ph/index.php/orders-and-memorandums/10-orders-and-memos/226-phreb-memo </p> <p>DOST: <ol style="list-style-type: none"> Administrative Order 001 Series 2007: Requirement for Review of All Research Involving Human Subjects/Participants (2007): http://ethics.healthresearch.ph/index.php/component/content/article/2-uncategorised/112-ao-001-2007 Administrative Order 001 </p> | <p>PHREB: National Ethical Guidelines for Health and Health-Related Research (2017): http://www.ethics.healthresearch.ph/index.php/phoca-downloads/category/4-neg?download=96:2017-national-ethical-guidelines-for-health-and-health-related-research</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> Food and Drug Administration (FDA): http://www.fda.gov.ph/ | | FDA: 1. Rules and Regulations on the Registration, Including Approval and Conduct of Clinical Trials, and Lot or Batch Release Certification of Vaccines and Biologic Products(Administrative Order No. 47-a) (2001) 2. FDA Circular 2015-026: Adoption of the ICH Harmonised Tripartite Guideline, Quality of Biotechnological Products: Stability Testing of Biotechnological/Biological Products Q5C: http://www.fda.gov.ph/attachments/article/118205/FC2013-026.pdf DOST, DOH, CHED, and UPM: Joint Memorandum Order 001 Series of 2012: http://www.ethics.healthresearch.ph/index.php/component/content/article/10-orders-and-memos/215-joint-memo-01 DOST, DOH, CHED, and UPM: Joint Administrative Order No. 001: The Implementing Rules and Regulations of Republic Act 10532 Otherwise Known as “The Philippine National Health Research System Act of 2013:” http://www.ethics.healthresearch.ph/index.php/component/content/article/2-uncategorised/214-implementing-rules-of-pnhrs | National Ethical Guidelines for Health and Health-Related Research, Page 70 (2017): http://www.ethics.healthresearch.ph/index.php/phoca-downloads/category/4-neg?download=96:2017-national-ethical-guidelines-for-health-and-health-related-research |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Clinical Trials Registry</i> | Philippine Health Research Registry: http://registry.healthresearch.ph/ | | | FAQs: http://registry.healthresearch.ph/index.php?option=com_content&view=article&id=7&Itemid=185 |
| <i>Research Injury</i> | 1. Department of Science and Technology (DOST): http://www.dost.gov.ph/ 2. Philippine Health Research Ethics Board (PHREB): www.ethics.healthresearch.ph | | | PHREB: National Ethical Guidelines for Health and Health-Related Research (2017): http://www.ethics.healthresearch.ph/index.php/phoca-downloads/category/4-neg?download=96:2017-national-ethical-guidelines-for-health-and-health-related-research |
| <i>Social-Behavioral Research</i> | 1. Philippine Health Research Ethics Board (PHREB): www.ethics.healthresearch.ph 2. Philippine Social Science Council (PSSC): http://pssc.org.ph/ | | | National Ethical Guidelines for Health and Health-Related Research, Pages 108-118. (2017): http://www.ethics.healthresearch.ph/index.php/phoca-downloads/category/4-neg |
| <i>Privacy/Data Protection</i> | | Republic Act No. 10173: Data Privacy Act of 2012: http://www.officialgazette.gov.ph/2012/08/15/republic-act-no-10173/ | Data Privacy Act Implementing Rules and Regulations (2016): https://privacy.gov.ph/implementing-rules-and-regulations-of-republic-act-no-10173-known-as-the-data-privacy-act-of-2012/ | |
| <i>Embryos, Stem Cells, and Cloning</i> | Philippine Health Research Ethics Board (PHREB): www.ethics.healthresearch.ph | | | National Ethical Guidelines for Health and Health-Related Research, Pages 91, 157 and 163 (2017): http://www.ethics.healthresearch.ph/index.php/phoca-downloads/category/4-neg?download=96:2017-national-ethical-guidelines-for-health-and-health-related-research |
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| <i>Drugs, Biologics, and Devices</i> | <p><i>Drugs</i></p> <p>1. Health Sciences Authority of Singapore (HSA): http://www.hsa.gov.sg 2. Ministry of Health, National Medical Ethics Committee (NMEC): https://www.moh.gov.sg/content/moh_web/home/Publications/guidelines/national_medical_ethics_committee_guidelines.html</p> | <p>Medicines Act (1975): http://statutes.agc.gov.sg/</p> | <p>Medicines (Clinical Trials) Regulations (2000): http://statutes.agc.gov.sg/aol/search/display/view.w3p;orderBy=date-rev.loadTime;page=0;query=Id%3A7e3c748b-8089-4699-a4b2-9f66af6f7820;rec=0</p> | <p>HSA: Singapore Guideline for Good Clinical Practice (1990): http://www.pacra.org/dev-pacra/images/pdf-files/singapore/sg-gcp.pdf</p> <p>NMEC: Recommendations on Clinical Trials: Update Focusing On Phase I Trials (2007): https://www.moh.gov.sg/content/moh_web/home/Publications/guidelines/national_medical_ethics_committee_guidelines/2007.html</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | 1. Health Sciences Authority: http://www.hsa.gov.sg 2. Ministry of Health, National Medical Ethics Committee (NMEC): https://www.moh.gov.sg/content/moh_web/home/Publications/guidelines/national_medical_ethics_committee_guidelines.html | 1. Medicines Act (1975): http://statutes.agc.gov.sg/ 2. Radiation Protection Act (2007): http://statutes.agc.gov.sg/ | Medicines (Clinical Trials) Regulations (2000): http://statutes.agc.gov.sg/aol/search/display/view.w3p;orderBy=date-rev.loadTime;page=0;query=Id%3A7e3c748b-8089-4699-a4b2-9f66af6f7820;rec=0 | HSA: Singapore Guideline for Good Clinical Practice (1999): http://www.pacra.org/dev-pacra/images/pdf-files/singapore/sg-gcp.pdf NMEC: Recommendations on Clinical Trials: Update Focusing on Phase I Trials (2007) |
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| <i>Human Biological Materials</i> | 1. Ministry of Health (MOH): http://www.moh.gov.sg/ 2. Health Sciences Authority: http://www.hsa.gov.sg 3. Bioethics Advisory Committee (BAC): http://www.bioethics-singapore.org | 1. Medical (Therapy, Education, and Research) Act (1973): http://statutes.agc.gov.sg/ 2. Medicines Act (1975): http://statutes.agc.gov.sg/ 3. Human Biomedical Research Bill No. 25/2015, Part 6: http://statutes.agc.gov.sg/aol/search/display/view.w3p;orderBy=date-rev.loadTime;page=0;query=Id%3A1f615627-01d3-4250-a720-de776cd4f794;rec=0 | Medicines (Clinical Trials) Regulations (2000): http://statutes.agc.gov.sg/aol/search/display/view.w3p;orderBy=date-rev.loadTime;page=0;query=Id%3A7e3c748b-8089-4699-a4b2-9f66af6f7820;rec=0 | BAC: Human Tissue Research (2002): http://www.bioethics-singapore.org/index/publications/reports/173-human-tissue-research.html |
| <i>Genetic Research</i> | 1. Ministry of Health, National Medical Ethics Committee (NMEC) 2. Bioethics Advisory Committee (BAC): http://www.bioethics-singapore.org | | | NMEC: Ethical Guidelines for Gene Technology (2001): https://www.moh.gov.sg/content/moh_web/home/Publications/guidelines/national_medical_e |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | 1. Ministry of Health (MOH): http://www.moh.gov.sg/ 2. Bioethics Advisory Committee (BAC): http://www.bioethics-singapore.org/ | Human Cloning and Other Prohibited Practices Act (2004): http://statutes.agc.gov.sg/ | Licensing Terms and Conditions on Assisted Reproduction Services (2011): http://www.moh.gov.sg/content/dam/moh_web/Publications/Guidelines/Private%20healthcare%20institutions/2011/AR_LTCs_260411.pdf | BAC: 1. Ethical, Legal and Social Issues in Human Stem Cell Research, Reproductive and Therapeutic Cloning (2002): http://www.bioethics-singapore.org/index/publications/reports/86-reports/174-stem-cell-research.html 2. Donation of Human Eggs for Research (2008): http://www.bioethics-singapore.org/index/publications/reports/86-reports/168-donation-of-human-eggs-for-research.html 3. Human-Animal Combinations in Stem-Cell Research (2010): http://www.bioethics-singapore.org/index/publications/reports/86-reports/167-human-animal-combinations-in-stem-cell-research.html |
| Sri Lanka | | | | |
| <i>Drugs and Devices</i> | Cosmetics, Devices, and Drugs Regulatory Authority, Subcommittee on Clinical Trials: http://www.cdda.gov.lk/index.php?option=com_content&view=article&id=78&Itemid=115&lang=en | National Medicines Regulatory Authority Act of 2015: http://www.cdda.gov.lk/images/stories/new/pdf/legislations/5e_nmdra.pdf | | Guidelines for the Conduct of Clinical Trials in Sri Lanka (2014): http://www.cdda.gov.lk/images/pdf/clinical%20trials%20guidelines_oct2014.pdf |
| <i>Clinical Trials Registry</i> | Sri Lanka Clinical Trials Registry: http://www.slctr.lk/ | | | FAQs: http://slctr.lk/faq |
| Taiwan | | | | |
| <i>General</i> | Ministry of Health and Welfare: http://www.mohw.gov.tw/EN/Ministry/Index.aspx | 1. Human Subjects Research Act (2011): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020176 2. Medical Care Act (2014): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=L0020021 | 1. Regulations on Human Trials (2009): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020162 2. Enforcement Rules of the Medical Care Act (2010): http://law.moj.gov.tw/LawClass/LawContent.aspx?PCODE=L0020023 3. Regulations Governing the Organization and Operational Management of the Institutional | Regulations Governing the Organization and Operational Management of the Institutional Review Board for Human Subject Research (2012): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=L0020179 |

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| <i>General</i> | | | Review Board for Human Subject Research (2012): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020179 4. Exempt Review Categories for Human Research (2012): http://gazette.nat.gov.tw/egFront/eng/EngIndex.jsp 5. Informed Consent Exemptions for Human Research (2012): http://gazette.nat.gov.tw/egFront/eng/EngIndex.jsp 6. Expedited Review Categories for Human Research (2012): http://gazette.nat.gov.tw/egFront/eng/EngIndex.jsp 7. Partial Amended Articles of Enforcement Rules of Medical Care Act (2016) http://gazette.nat.gov.tw/egFront/eng/EngIndex.jsp | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health and Welfare (MOHW): http://www.mohw.gov.tw/EN/Ministry/Index.aspx 2. Taiwan Food and Drug Administration (FDA): http://www.fda.gov.tw/EN/index.aspx | MOHW: Medical Care Act (2014): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020021 FDA: Pharmaceutical Affairs Act (2015): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0030001 | MOHW: 1. Regulations on Human Trials (2009): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020162 2. Pharmaceutical Affairs Act Enforcement Rules (2012): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0030002 3. Regulations for Drug Safety Monitoring (2013) http://mohwlaw.mohw.gov.tw/Chi/EngContent.asp?msgid=516&KeyWord ≡ 4. Guideline for Good Clinical Practice (2014): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=L0030056 5. Regulations for Governing the Management of Medical Devices (2014): http://mohwlaw.mohw.gov.tw/Chi/EngContent.asp?msgid=528&KeyWord ≡ | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | 6. Regulations for Bioavailability and Bioequivalence Studies (2015): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=L0030065 | |
| <i>Research Injury</i> | 1. Ministry of Health and Welfare (MOHW): http://www.mohw.gov.tw/EN/Ministry/Index.aspx 2. Food and Drug Administration (FDA), MOHW: http://www.fda.gov.tw/EN/index.aspx | Medical Care Act (2014): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=L0020021 | FDA: Guideline for Good Clinical Practice (2014): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=L0030056 | |
| <i>Social-Behavioral Research</i> | Ministry of Health and Welfare | | Exempt Review Categories for Human Research (2012) http://www.mohw.gov.tw/EN/Ministry/ | |
| <i>Privacy/Data Protection</i> | Ministry of Justice: http://www.moj.gov.tw/mp095.html | Personal Information Protection Act (2015): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=I0050021 | | |
| <i>Human Biological Materials</i> | Ministry of Health and Welfare: http://www.mohw.gov.tw/EN/Ministry/Index.aspx | 1. Human Subjects Research Act (2011): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020176 2. Human Biobank Management Act (2012): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020164 3. Medical Care Act (2015): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020021 | 1. Regulations on Human Trials (2009): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020162 2. Administrative Regulations on the Establishment of Human Biobanks (2011) http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?PCODE=L0020173 | 1. Good Tissue Practice (2002) (Chinese): http://www.fda.gov.tw/TC/includes/GetFile.aspx?id=1153&chk=342a5c73-c206-4756-ade9-9c63265c859d&mid=46&name=fdContent 2. Guidelines for Collection and Use of Human Specimens for Research (2006): http://www.fda.gov.tw/TC/includes/GetFile.aspx?id=1598&chk=6056f7dd-eb0a-48bf-ae7e-8a2a5875e6e0&mid=46&name=fdContent |
| <i>Genetic Research</i> | 1. Ministry of Health and Welfare (MOHW): http://www.mohw.gov.tw/EN/Ministry/Index.aspx 2. Food and Drug Administration (FDA): http://www.fda.gov.tw/EN/index.aspx 3. Ministry of Science and Technology: https://www.most.gov.tw/en/public | MOHW: Human Biobank Management Act (2012): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020164 | MOHW: 1. Regulations on Commercial Benefit Feedback of Human Biobanks (2010): http://law.moj.gov.tw/LawClass/LawContentIf.aspx?PCODE=L0020170 2. Administrative Regulations on the Establishment of Human Biobanks (2011): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0020173 | MOHW: Guidance for Information Safety of Human Biobank (2010): http://mohwlaw.mohw.gov.tw/Chi/FLAW/FLAWDAT0202.asp |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Embryos, Stem Cells, and Cloning</i> | Health Promotion Administration, MOHW: http://www.hpa.gov.tw/BHPNet/English/Index.aspx | Artificial Reproduction Act (2007): http://law.moj.gov.tw/Eng/LawClass/LawContent.aspx?pcode=L0070024 | | |
| Tajikistan | | | | |
| Note: For an overview of human subject protections in Tajikistan, see “Ethical Review of Biomedical Research in the CIS Countries,” Chapter 3, Section 9: http://www.unesco.org/new/fileadmin/MULTIMEDIA/FIELD/Moscow/pdf/ethical_review_cis_book_kubar_english.pdf | | | | |
| Note: All websites and documents are in Russian. | | | | |
| <i>General</i> | Ministry of Public Health: http://www.health.tj/ | | Order of the Ministry of Public Health of the Republic Tajikistan of 10 March, 2005 No. 118: About the Assertion of the Normative Documents of Republic Committee on Medical Ethics | |
| Thailand | | | | |
| For an overview of the clinical research regulations in Thailand, see: https://clinregs.niaid.nih.gov/single_country.php?c_id=213 | | | | |
| <i>General</i> | 1. National Research Council of Thailand (NCRT): http://en.nrct.go.th/en/home.aspx 2. Medical Council of Thailand (MCT): http://www.tmc.or.th/en_home.php 3. Forum for Ethical Review Committees in Thailand (FERCIT): http://www.fercit.org/ | Medical Professions Act (2009), Articles 47-50: http://www.fercit.org/SIDCER-FERCAP/Handout_10/4.%20Accreditation-update_surveyor_aj.Sopit.pdf | NCRT: Regulation on the Permission of Foreign Researchers (1982): http://www.dnp.go.th/otec/eng_laws_regs/NRCT_Reg2525E.pdf MCT: Rule of the Medical Council on the Observance of Medical Ethics (1983): http://thailaws.com/law/t_laws/tlaw0510.pdf | MCT: National Guideline for Ethical Research on Human Subjects (2002) FERCIT: Ethical Guidelines for Research on Human Subject in Thailand (2007): http://www.fercit.org/file/Guideline_English_version.pdf |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | 1. Food and Drug Administration, Drug Control Division: http://www.fda.moph.go.th/eng/index.stm 2. Medical Council of Thailand (MCT): http://www.tmc.or.th/en_home.php | Consumer Protection Act (2007) | MCT: Thailand Good Clinical Practice Guidelines (2002) |
| | <i>Devices</i> | Food and Drug Administration, Medical Device Control Division: http://www.fda.moph.go.th/eng/medical/p_re.stm | 1988 Medical Device Act: http://www2.fda.moph.go.th/Exporters/law/Document/Mdc/36-MEDICAL%20DEVICE%20ACT.htm | |
| <i>Clinical Trials Registry</i> | Thai Clinical Trials Registry: http://www.clinicaltrials.in.th/ | | | FAQs: http://www.clinicaltrials.in.th/index.php?meun |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Clinical Trials Registry</i> | | | | =home&smenu=4&task=home&task1=openpage&task2=view&topid=4 |
| <i>Privacy/Data Protection</i> | Office of the Information Commission: http://www.oic.go.th/content_eng/default_eng.asp | Official Information Act, B.E. 2540 (1997): http://www.oic.go.th/content_eng/act.htm | | |
| Uzbekistan | | | | |
| Note: All websites and documents are in Uzbek and Russian. | | | | |
| <i>General</i> | 1. Government of the Republic of Uzbekistan: http://www.gov.uz 2. Ministry of Health: http://www.minzdrav.uz | 1. Constitution of Republic of Uzbekistan, Articles 24, 26, 40, 44 (1992): http://www.gov.uz 2. Law on Protection of Citizens' Health (1997): http://www.minzdrav.uz | | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health, Pharmacological Committee of the Central Department for Quality Control of Pharmaceuticals and Medical Equipment: http://www.minzdrav.uz 2. Ministry of Health, National Ethics Committee 3. Scientific Boards of Medical Institutes | 1. Law on Protection of Citizens' Health (1997): http://www.minzdrav.uz 2. Law on Drugs and Pharmaceutical Activity (1997) 3. Law on Narcotic and Psychoactive Drugs (2000) | 1. Guidelines on Conducting Clinical Trials and Determining Clinical Sites (2001) 2. National Standard of Uzbekistan: Good Clinical Practice (2013) | |
| <i>Human Biological Materials</i> | 1. Ministry of Health, Pharmacological Committee of the Central Department for Quality Control of Pharmaceuticals and Medical Equipment: http://www.minzdrav.uz 2. Ministry of Health, National Ethics Committee 3. Scientific Boards of Medical Institutes | | 1. Guidelines on Conducting Clinical Trials and Determining Clinical Sites (2001) 2. National Standard of Uzbekistan: Good Clinical Practice (2013) | |
| Vietnam | | | | |
| For an overview of the clinical research regulations in Vietnam, see the ClinRegs report: https://clinregs.niaid.nih.gov/single_country.php?c_id=233 | | | | |
| <i>General</i> | 1. Ministry of Health (MOH): http://www.moh.gov.vn/homebyt/en/portall/index.jsp 2. Ministry of Health, Independent Ethics Committee (MOH): http://iecmoh.vn | MOH: Decision No. 111/QD-BYT – On Promulgation of Regulation on Organization and Operation of Council of Ethics in Biomedical Research at Grass-Roots Level, Chapter I (Articles 3 and 4), Chapter II, and | MOH: Decision No. 460/QD-BYT – On the Promulgation of Regulations on Organization and Operation of Ethical Evaluation Committee in Biomedical Research of the Ministry of Health, Period 2012-2017, Chapters I-III (2012): | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | | Chapter III (2013): http://clinregs.niaid.nih.gov/documents/vietnam/DecisionNo111-QD-BYT.pdf | http://clinregs.niaid.nih.gov/documents/vietnam/DecisionNo460-QD-BYT.pdf | |
| <i>Drugs, Biologics, and Devices</i> | Ministry of Health: http://www.moh.gov.vn/homeby/en/porta/index.jsp | 1. Law on Pharmacy (No. 34/2005/QH11), Chapter II (Section III, Article 20), Chapter VIII (Articles 54 and 59) (2005): http://www.vertic.org/media/National%20Legislation/Vietnam/VN_Law_on_Pharmacy.pdf 2. Decision No. 799/QD-BYT on the Issuance of Guideline on Good Clinical Practice, Chapter III, Articles 1 and 2 (2008): http://clinregs.niaid.nih.gov/documents/vietnam/DecisionNo799-QD-BYT.pdf | 1. Decision No. 799/QD-BYT of the Minister of Health on the Promulgation of the Guidelines on Good Clinical Practice of Clinical Trials (2008): http://clinregs.niaid.nih.gov/documents/vietnam/DecisionNo799-QD-BYT.pdf 2. Circular – Guidelines for Clinical Trials on Drugs (C-ClinDrugTrial), Articles 2, 4, 5, 9, 17, 18, 31, and 39 (2012): http://clinregs.niaid.nih.gov/documents/vietnam/C-ClinDrugTrial.pdf | Guidelines for Clinical Trials of Drugs, Chapter III, Articles 10, 16, and 17 (2012): https://clinregs.niaid.nih.gov/documents/vietnam/C-ClinDrugTrial.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| MIDDLE EAST/NORTH AFRICA | | | | |
| Egypt | | | | |
| <i>General</i> | Medical Professionals Union | Constitution of the Arab Republic of Egypt, Article 43: http://www.sis.gov.eg/Newvr/Dustor-en001.pdf | Professional Ethics Regulations: Conducting Medical Research on Human Beings, Articles 52-61 (2003) | |
| <i>Drugs, Biologics, and Devices</i> | Egyptian Drug Authority: http://www.eda.mohp.gov.eg/ | | | |
| Iran | | | | |
| <i>General</i> | Ministry of Health and Medical Education, Office for the Study of Humanistic and Islamic Science in Medicine and Medical Ethics: http://www.mohme.gov.ir/ | | Protection Code for Human Subjects in Medical Research (1999) | |
| <i>Clinical Trials Registry</i> | Iranian Registry of Clinical Trials: http://www.irct.ir/ | | | FAQs: http://www.irct.ir/faq.php |
| Israel | | | | |
| <i>General</i> | Ministry of Health: http://www.health.gov.il/english/ | | Public Health Regulations (Medical Experiments Involving Human Subjects) (1999) | |
| <i>Drugs, Biologics, and Devices</i> | Ministry of Health, Pharmaceutical Administration: http://www.health.gov.il/English/MinistryUnits/HealthDivision/MedicalTechnologies/Drugs/Pages/default.aspx | Public Health Order (1940) | 1. Public Health Regulations (Clinical Studies in Human Subjects) – 1980 2. 1990 Amendment 3. 1992 Amendment 4. 2005 Amendment | Guidelines for Clinical Trials in Human Subjects (2006): https://firstclinical.com/regdocs/doc/?db=INT-Israel_Clinical_Trials |
| <i>Privacy/Data Protection</i> | Israeli Law, Information, and Technology Authority: http://www.justice.gov.il/MOJEng/ILITA/ | 1. Privacy Protection Act No. 5741 (1981): http://www.justice.gov.il/NR/rdonlyres/6A5EC09A-BDBC-419F-8007-5FD6A6B8E0A5/18334/ProtectionofPrivacyLaw57411981unofficialtranslation.pdf 2. Protection of Privacy Law No. 5741, as Amended by Law No. 5745 (1985) | | |
| <i>Genetic Research</i> | Ministry of Health: http://www.health.gov.il/english/ | Genetic Information Law (2000): http://www.moital.gov.il/NR/exeres/66F4DD4E-FA4A-4B76-94BC-DC29543471DE.htm | | 1. Instruction of the Supreme Committee for Clinical Studies on Humans Regarding Establishment and Usage of Genetic Samples Reservoir (2005) 2. Amendment (2007) |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|---|---|---|---|------------|
| <i>Embryos, Stem Cells, and Cloning</i> | | Genetic Intervention Prohibition Law (Human Cloning and Genetic Changes in Reproduction Cells) (1999) | | |
| Jordan Note: All documents are in Arabic. | | | | |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health: http://www.moh.gov.jo/en/Pages/default.aspx 2. Jordan Food and Drug Administration: http://www.jfda.jo/Default.aspx | 1. Law of Clinical Studies, Law No. 2 (2011) http://www.jfda.jo/EchoBusV3.0/SystemAssets/PDF/AR/LawsAndRegulation/Drug/PharmaceuticalStudies/50_211.pdf 2. Drug and Pharmacy Law No. 12 (2013) http://www.jfda.jo/EchoBusV3.0/SystemAssets/PDF/AR/LawsAndRegulation/Drug/DrugDirectorate/%D9%82%D8%A7%D9%86%D9%88%D9%86%20%D8%A7%D9%84%D8%AF%D9%88%D8%A7%D8%A1%20%D9%88%D8%A7%D9%84%D8%B5%D9%8A%D8%AF%D9%84%D8%A9.pdf 3. Narcotic and Psychotropic Law No. 23 (2016) http://www.jfda.jo/EchoBusV3.0/SystemAssets/PDF/AR/LawsAndRegulation/Drug/DrugsAndPsychotropicSubstances/%D9%82%D8%A7%D9%86%D9%88%D9%86%20%D8%A7%D9%84%D9%85%D8%AE%D8%AF%D8%B1%D8%A7%D8%AA%20%D9%88%D8%A7%D9%84%D9%85%D8%A4%D8%AB%D8%B1%D8%A7%D8%AA%20%D8%A7%D9%84%D8%B9%D9%82%D9%84%D9%8A%D8%A9.pdf | | |
| <i>Research Injury</i> | | | Regulations for Insurance on Research-Related Injury (2013): http://www.jfda.jo/EchoBusV3.0/SystemAssets/PDF/AR/LawsAndRegulati | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | | | on/Drug/PharmaceuticalStudies/22_252.pdf | |
| <i>Embryos, Stem Cells, and Cloning</i> | | Stem Cell By-law No. 10 (2014) | | |
| Kuwait | | | | |
| <i>General</i> | Ministry of Health, Kuwait Institute for Medical Specialization: http://www.kims.org.kw/ | | | Ethical Guidelines for Biomedical Research: http://www.kims.org.kw/Ethical%202.doc |
| Qatar | | | | |
| <i>General</i> | Ministry of Public Health, Research Division: https://www.moph.gov.qa/about-us/Pages/research.aspx | Policies, Regulations and Guidelines for Research Involving Humans: https://www.moph.gov.qa/about-us/Documents/research/Policies%2c%20Regulations%20and%20Guidelines%20for%20Research%20Involving%20Human.pdf | | 1. IRB - IEC Registration Application: https://www.moph.gov.qa/about-us/Documents/research/IRB%20-%20IEC%20Registration%20Application.pdf 2. Assurance Application for the Protection of Human Subjects Involved in Research: https://www.moph.gov.qa/about-us/Documents/research/Local%20-%20Foreign%20IRB%20-%20IEC%20Assurance%20Application%20for%20the%20Protection%20of%20Human%20Subjects%20Involved%20in%20Research.pdf 3. Guidelines on Reviewing and Reporting Adverse Events: https://www.moph.gov.qa/about-us/Documents/research/Guidelines%20on%20Reviewing%20and%20Reporting%20Unanticipated%20Problems%20Involving%20Risks%20to%20Subject%20or%20Others%20and%20Adverse%20Events.pdf |
| <i>Human Biological Materials</i> | Ministry of Public Health, Research Division: https://www.moph.gov.qa/about-us/Pages/research.aspx | | | Guidance for the Use of Stored Data and Biological Specimens in Human Research: https://www.moph.gov.qa/about-us/Documents/research/Guidance%20for%20the%20Use%20of%20Stored%20Data%20and%20Biological%20Specimens%20in%20Human%20Research.pdf |
| <i>Genetic Research</i> | Ministry of Public Health, Research Division: https://www.moph.gov.qa/about-us/Pages/research.aspx | | | 1. Guidance for the Design, Ethical Review, and Conduct of Genomic Research in Qatar: https://www.moph.gov.qa/about-us/Documents/research/Guidance%20for%20the%20Design%2c%20Ethical%20Review%2c%20and%20Conduct%20of%20Genomic%20Research%20in%20Qatar.pdf 2. Guidelines for Gene Transfer Research |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | | | | in Humans: https://www.moph.gov.qa/about-us/Documents/research/Guidelines%20for%20Gene%20Transfer%20Research%20in%20Humans.pdf |
| <i>Embryos, Stem Cells, and Cloning</i> | Ministry of Public Health, Research Division: https://www.moph.gov.qa/about-us/Pages/research.aspx | | | Guidance for Research Involving Human Embryonic Stem Cells Germ Cells: https://www.moph.gov.qa/about-us/Documents/research/Guidance%20for%20Research%20Involving%20Human%20Embryonic%20Stem%20Cells%20Germ%20Cells%20and%20Cells%20Obtained%20From%20Cord%20Blood.pdf |
| Saudi Arabia | | | | |
| <i>General</i> | National Committee of BioEthics: http://bioethics.kacst.edu.sa/?lang=en-US | Law of Ethics of Research on Living Creatures: http://bioethics.kacst.edu.sa/getattachment/4bd0d4e2-1b93-4c32-b483-57902227fae2/Bioethic-Rgl-fin-bks.aspx | Implementing Regulations of the Law of Ethics of Research on Living Creatures (2016): http://prod.kau.edu.sa/Med/ali/files/Publications/Guide/National_Committee_of_BioEthics-Regulations_of_the_Law_of_Ethics_of_Research_on_Living_Creatures.pdf | |
| <i>Social-Behavioral Research</i> | National Committee of BioEthics | | Implementing Regulations of the Law of Ethics of Research on Living Creatures, Expedited Research (Article 10.18g) and Categories of Social-Behavioral Research That do not Require Continuing Review (Article 10.32) (2016): http://prod.kau.edu.sa/Med/ali/files/Publications/Guide/National_Committee_of_BioEthics-Regulations_of_the_Law_of_Ethics_of_Research_on_Living_Creatures.pdf | |
| Sudan | | | | |
| <i>General</i> | Federal Ministry of Health: http://www.fmoh.gov.sd/ | | | 1. National Guidelines for Ethical Conduct of Research Involving Human Subjects (2008): http://sites.google.com/site/healthresearchlibrary/national-guidelines 2. National Accreditation Guidelines for Ethics Committees (2016) 3. Operation Guidelines, Functions, and Procedures (2016) |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
|--------------------------------------|--|---|---|---|
| <i>Drugs, Biologics, and Devices</i> | National Medicines and Poisons Board: http://www.nmpb.gov.sd/en/ | Act on Pharmaceuticals and Poisons (2009) (Arabic): http://www.nmpb.gov.sd/index.php/2015-08-05-11-05-04/regulations/113-laws2009 | | |
| <i>Human Biological Materials</i> | 1. Federal Ministry of Health: http://www.fmoh.gov.sd/ 2. National Council on Biosafety | 1. Human Organs and Tissues Transplant Legislation, Chapter 2, Articles 3 and 4 (1978) Act on Biosafety (2010) | | |
| <i>Genetic Research</i> | University of Khartoum, Institute of Endemic Diseases: http://iend.uofk.edu/index.php?lang=en | | | Guidelines for Genetics Research on Sudanese Subjects (2005) |
| Tunisia | | | | |
| <i>Drugs, Biologics, and Devices</i> | Ministry of Public Health, Institut Pasteur: www.pasteur.tn | | Conditions of Contract and Specifications Related to Medical or Scientific Experimentation of Medicines Intended for Humans | Disposals and Director's Principles Related to Good Practices in Clinical Trials |
| Turkey | | | | |
| <i>General</i> | Ministry of Health (Turkish): http://www.saglik.gov.tr/ | 1. Turkish Constitution, Article 17 2. Health Services Basic Law No. 3359 (1987) 3. Oviedo Convention on Human Rights and Biomedicine (2004) 4. Update on the Law of the Support of Research and Development Activities (2016). Official Gazette (Turkish): http://www.resmigazete.gov.tr/eskiler/2016/02/20160226.htm | 1. Regulation on Medical Deontology, Article 11 (1960) 2. Bylaw on Patient Rights No. 23420 (1998) | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> 1. Turkey Pharmaceuticals and Medical Devices Agency (TITCK): http://www.titck.gov.tr 2. Clinical Research Association (CRA): http://www.klinikarastirmalar.org.tr/en/ | Turkish Penal Law, Article 90 (2005) | 1. Fundamental Law #3359 on Health Services, Supplemental Article 10 (2011): http://www.titck.gov.tr/Default.aspx?sayfa=klirik_mevzuat&lang=tr-TR&thelawtype=1&thelawId=347 2. Regulation on Clinical Trials with Drugs and Biological Products (2015): An Update of 2014 Clinical Trials Regulation: http://www.klinikarastirmalar.org.tr/doc/file_345.docx 3. Regulation on Efficacy, Safety, | CRA: 1. GCP Guideline (2015): http://www.klinikarastirmalar.org.tr/dokuman.php?id=374 2. Various: http://www.klinikarastirmalar.org.tr/en/documents.php?dok_cat=0 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | and Clinical Trials of Cosmetic Products (2015): http://www.klinikarastirmalar.org.tr/doc/file_346.pdf 4. Update on the Regulation of the Management and Inspection of the Support of Research and Development Activities (2016). Official Gazette: http://www.resmigazete.gov.tr/eskiler/2016/08/20160810-7.htm | |
| | <i>Devices</i> | | | |
| | Turkey Pharmaceuticals and Medical Devices Agency (TITCK): http://www.titck.gov.tr | | Regulation on Research on Medical Devices (2014): http://www.klinikarastirmalar.org.tr/doc/file_318.pdf | |
| <i>Research Injury</i> | Turkey Pharmaceuticals and Medical Devices Agency (TITCK): http://www.titck.gov.tr | Convention on Human Rights and Biomedicine (Convention of Oviedo), Article 24, ETS No. 164 (2004) | | Guidance on Insuring Volunteers in a Clinical Trial (2011) |
| <i>Human Biological Materials</i> | | 1. Law on Procurement, Preservation, Grafting, and Transplantation of Organs and Tissues, No. 2238 (1979) 2. Law on Blood and Blood Products, No. 2857 (1983) | Regulation on Blood and Blood Products, No. 7314 (1983) | 1. Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 21-22 (1999) 2. Good Clinical Practice Guidelines for Advanced Therapy Medicinal Products (2011) |
| <i>Genetic Research</i> | | | Regulation on Centers for Diagnosis and Genetic Diseases, No. 23368 (1998) | Convention on Human Rights and Biomedicine (Convention of Oviedo), Articles 12-14 (1999) |
| <i>Embryos, Stem Cells, and Cloning</i> | | | 1. Regulation on Centers for Medically Assisted Procreation, No. 19551 (1987) 2. Regulation on Organ and Tissue Transplantation Services (2005) 3. Regulation on Cordon Blood Banks (2005) | 1. Circular on Research of Embryonic Stem Cells (2005) 2. Guideline on Clinical Research of Non-Embryonic Stem Cells (2006) |
| United Arab Emirates | | | | |
| <i>General</i> | Health Authority - Abu Dhabi: http://www.haad.ae/haad/ | | | Standard Operating Procedures for Research Ethics Committees (2012): http://www.haad.ae/HAAD/LinkClick.aspx?fileticket=UL7o8f5mukc%3D&tabid=820 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| LATIN AMERICA and the CARIBBEAN | | | | |
| Regionwide | | | | |
| <i>General</i> | Caribbean Public Health Agency: http://carpha.org/What-We-Do/Research-Training-and-Policy-Development | | | |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | Pan American Health Organization: http://www.paho.org/ | | Good Clinical Practices: Document for the Americas (2005): http://www.paho.org/english/ad/ths/ev/GCP-Eng-doct.pdf |
| | <i>Devices</i> | Pan American Health Organization: http://www.paho.org/ | | A Model Regulatory Program for Medical Devices: An International Guide (2001): http://www.paho.org/English/HSP/HSE/medical_devices.pdf |
| Argentina | | | | |
| Note: Several provinces have their own regulations pertaining to human subjects research. | | | | |
| <i>General</i> | Ministry of Health: http://www.msal.gov.ar | Civil and Commercial Code, Articles 26, 58, and 59 (2015): http://servicios.infoleg.gob.ar/infolegInternet/anexos/235000-239999/235975/norma.htm | Ministerial Resolution 1480/2011 Approving the Guidelines for Human Health Research and Creating the National Registry of Health Research: http://www.anmat.gov.ar/webanmat/legislacion/medicamentos/Resolucion_1480-2011.pdf | Resolution 1480/2011: Approving the Guidelines for Human Health Research and Creating the National Registry of Health Research: http://www.anmat.gov.ar/webanmat/legislacion/medicamentos/Resolucion_1480-2011.pdf |
| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> | National Administration of Drugs, Foods, and Medical Devices (ANMAT): http://www.anmat.gov.ar/index.asp | | |
| | | | 1. Provision ANMAT 6677/10: Regulatory Guideline for Good Clinical Practices in Clinical Pharmacological Studies (2010): http://www.anmat.gov.ar/Comunicados/Dispo_6677-10_en.pdf 2. Provision ANMAT 12.792/2016: Request for Import of Medication /Treatment and Materials – Procedure (2016): http://www.anmat.gov.ar/boletin_anmat/Noviembre_2016/Dispo_12792-16.pdf 3. Provision ANMAT 828/2017: Authorization of Expanded Access Programs: http://www.anmat.gov.ar/boletin_anmat/enero_2017/Dispo_0828-17.pdf | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | 4. Provision ANMAT 4008/2017: Substitution of Article 2° of Provision ANMAT N° 6677/10: http://www.anmat.gov.ar/boletin_anmat/Abril_2017/Dispo_4008-17.pdf 5. Provision ANMAT 4009/2017: Health Care Institutions: Requirements and Conditions of Authorization for Conducting Phase I and/or Bioequivalence Clinical Pharmacology Studies: http://www.anmat.gov.ar/boletin_anmat/Abril_2017/Dispo_4009-17.pdf 6. Provision 10017-E/2017 ANMAT: Promoting Cooperation Between ANMAT and Jurisdictional Health Authorities for the Evaluation and Oversight of Clinical Research Sites and Investigators: http://servicios.infoleg.gob.ar/infolegInternet/verNorma.do?id=279820 | |
| | <i>Devices</i> | | | |
| | National Administration of Drugs, Foods, and Medical Devices (ANMAT): http://www.anmat.gov.ar/index.asp | | Provision ANMAT No. 969/97 on the Regulation of Good Clinical Practice with Medical Devices (1997) | |
| <i>Clinical Trial Registries</i> | National Registry of Health Research: https://www.argentina.gob.ar/salud/registroinvestigaciones | | Resolution 1480/2011 Approving a Guide for Human Subjects Research: http://servicios.infoleg.gob.ar/infolegInternet/anexos/185000-189999/187206/norma.htm | 1. FAQs: https://sisa.msar.gov.ar/sisa/sisa/service/FileDownloadServlet?location=/docs/otros/681/2214-ReNIS_Preguntas_Frecuentes.pdf 2. Tutorial: https://sisa.msar.gov.ar/sisa/#Renis |
| <i>Privacy/Data Protection</i> | National Directorate for the Protection of Personal Data: http://www.jus.gob.ar/datos-personales.aspx | Personal Data Protection Act No. 25.326 (2000): http://www.protecciondedatos.com.ar/law25326.htm | | |
| Barbados | | | | |
| | University of the West Indies – Cave Hill / Ministry of Health: http://www.cavehill.uwi.edu/researchethics/home.aspx | | | Research Ethics Policy and Guidelines |
| Bermuda | | | | |
| <i>General</i> | Department of Health: https://www.gov.bm/department/health | | | Research Governance Framework (2008): http://citeseerx.ist.psu.edu/viewdoc/download?doi=10.1.1.592.8671&rep=rep1&type=pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Bolivia | | | | |
| <i>General</i> | 1. Ministry of Health and Sport (MHS): http://www.sns.gob.bo 2. National Bioethics Committee (NBC) | 1. Legal Decree No. 15.629 of July 18, 1978, Articles 147 and 148. 2. New Political Constitution of the State, Article 44 (2009): https://www.constituteproject.org/constitution/Bolivia_2009.pdf | 1. Regulations on Public Health Research, Chapter V (1978) 2. Rules and Regulations of the National Bioethics Committee | MHS: Guidelines for the Development of Health Research and Ethical Norms (2002) NBC: 1. Requirements for the Evaluation of Research Projects 2. Code of Ethics and Medical Deontology |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health and Sport, National Pharmacological Commission (MHS): http://www.sns.gob.bo 2. National Bioethics Committee (NBC) | | | MHS: Rule on Clinical Studies with Medicines or Products in the Clinical Investigation Stage (2005) NBC: Projects that Involve Drugs or Therapeutic Products |
| Brazil | | | | |
| For an overview of clinical research regulations in Brazil, see the ClinRegs report: http://clinregs.niaid.nih.gov/single_country.php?c_id=30 | | | | |
| <i>General</i> | 1. National Health Council (CNS): http://www.conselho.saude.gov.br/ 2. National Commission on Research Ethics (CONEP): http://conselho.saude.gov.br/web_comissoes/conep/index.html | | CNS/CONEP: 1. Resolution CNS No. 240/97 - Defining "Participating User" According to IRB: http://conselho.saude.gov.br/resolucoes/1997/reso240.doc 2. Regulation of Resolution CNS No. 292/99 on Research with Foreign Cooperation: http://conselho.saude.gov.br/web_comissoes/conep/aquivos/resolucoes/regulation_res_292_english.doc 3. Resolution CNS No. 304/2000: Rules on Research Involving Human Beings – Area of Indigenous Peoples: http://conselho.saude.gov.br/resolucoes/2000/Res304_en.pdf 4. Internal CONEP Regulation (2001): http://conselho.saude.gov.br/web_comissoes/conep/aquivos/conep/regimento.doc 5. Resolution CNS No. 346/2005 on Multicenter Research: http://conselho.saude.gov.br/resolucoes/2005/reso346.doc | CNS/CONEP: Various: http://plataformabrasil.saude.gov.br/login.jsf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | | | <p>es/2005/Res346_en.pdf</p> <p>6. Resolution CNS No. 370/07 on Registration and Accreditation or Renewal of Registration and Accreditation of CEP: http://conselho.saude.gov.br/resolucoes/2007/Reso370.doc</p> <p>7. Resolution CNS No. 446/2011 on Composition of the National Commission on Research Ethics: http://conselho.saude.gov.br/resolucoes/2012/466_english.pdf</p> <p>8. Resolution CNS No. 466/2012 on Guidelines and Rules for Research Involving humans Subjects: http://conselho.saude.gov.br/resolucoes/2012/466_english.pdf</p> <p>9. Resolution CNS N° 506/2016 Accreditation of CEP: http://conselho.saude.gov.br/resolucoes/2016/Reso_506.pdf</p> <p>10. Resolution CNS No 580/2018 on Research of Strategic Interest for the Unified Health System (SUS): http://conselho.saude.gov.br/resolucoes/2018/Reso580.pdf</p> | |
| <i>Drugs, Biologics, and Devices</i> | <p><i>Drugs and Biologics</i></p> <p>1. National Health Council (CNS): http://www.conselho.saude.gov.br/</p> <p>2. Brazilian Health Surveillance Agency (ANVISA): http://portal.anvisa.gov.br/wps/portal/anvisa-ingles</p> <p>3. Federal Council of Medicine (CFM): http://portal.cfm.org.br/</p> <p>4. National Commission on Research Ethics (CONEP): http://conselho.saude.gov.br/web_comissoes/conep/index.html</p> | <p>Law N° 9782/99 Defining the National Health Surveillance System: http://www.planalto.gov.br/ccivil_03/leis/L9782.htm</p> | <p>CNS:</p> <p>1. Resolution CNS No. 251/1997: On Complimentary Rules for Research with New Pharmaceutical Products, Medicines, Vaccines, and Diagnostic Tests: http://conselho.saude.gov.br/resolucoes/1997/Res251_en.pdf</p> <p>2. Resolution CNS No. 301, 16th March 2002: Regarding Placebos: http://conselho.saude.gov.br/resolucoes/2000/Res301_en.pdf</p> <p>CFM:</p> <p>1. Resolution CFM N° 1.885,</p> | <p>ANVISA:</p> <p>1. Manual for Submission of “Drug Clinical Development Dossier” (DDCM) (2017): http://portal.anvisa.gov.br/documents/33836/2492465/Manual+para+Submiss%C3%A3o+de+Dossi%C3%AA+de+Desenvolvimento+CI%C3%ADnico+de+Medicamento+%28DDCM%29+e+Dossi%C3%AA+Espec%C3%ADfico+de+Ensaio+CI%C3%ADnico+-+3%C2%AA+edi%C3%A7%C3%A3o/29e9c5b1-2942-4bb9-a4dd-4fccc6fccda3</p> <p>2. Manual for Submission of Modifications, Amendments, Suspensions and Cancellations on DDCM (2018): http://portal.anvisa.gov.br/documents/33836/24</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | <p>2008 – about placebo: http://www.portalmedico.org.br/resolucoes/cfm/2008/1885_2008.htm</p> <p>ANVISA: 1. Resolution ANVISA 09/15 - Regulations for Clinical Trials with Drugs: https://clinregs.niaid.nih.gov/documents/brazil/ResolutionNo9-English.pdf 2. Resolution RDC No. 9, 20 February 2015 Regarding Regulation for Realization of Clinical Trials of Medication in Brazil: https://clinregs.niaid.nih.gov/documents/brazil/ResolutionNo9-English.pdf</p> | <p>92465/Manual+Para+Submiss%C3%A3o+de+Modifica%C3%A7%C3%B5es%2C+Emendas%2C+Suspens%C3%B5es+e+Cancelamentos/+4%C2%AA+edi%C3%A7%C3%A3o/85672ffa-db76-4869-b286-ff59bc3fcf60</p> |
| | <i>Devices</i> | | <p>Resolution ANVISA 10/15 - Regulations for Clinical Trials with Medical Devices: http://pesquisa.in.gov.br/imprensa/jsp/visualiza/index.jsp?data=03/03/2015&jornal=1&pagina=73&totalArquivo=140</p> | <p>ANVISA: Manual for Submission of Modifications, Amendments, Suspensions, and Cancellations on DICD (2015): http://portal.anvisa.gov.br/documents/33912/2785629/Manual+Para+Submiss%C3%A3o+de+Modifica%C3%A7%C3%B5es%2C+Emendas%2C+Suspens%C3%B5es+e+Cancelamentos/431fa7ef-24e6-4b14-80b9-ce68bccc24d8</p> |
| <i>Clinical Trials Registry</i> | <p>Brazilian Clinical Trials Registry: http://www.ensaiosclinicos.gov.br/</p> | | | <p>FAQs: http://www.ensaiosclinicos.gov.br/assistance/faq/</p> |
| <i>Research Injury</i> | <p>1. Brazilian Health Surveillance Agency: http://portal.anvisa.gov.br/english 2. National Health Council (CNS): http://www.conselho.saude.gov.br/ 3. National Commission on Research Ethics (CONEP): http://conselho.saude.gov.br/web_comissoes/conep/index.html</p> | <p>ANVISA: Law N° 6360/76: http://www.planalto.gov.br/ccivil_03/leis/l6360.htm</p> | <p>CNS/CONEP: 1. Standards Survey of New Drugs, Medicines, Vaccines, and Diagnostic Tests Involving Human Beings - Resolution CNS No. 251/97: http://conselho.saude.gov.br/resolucoes/1997/Res251_en.pdf 2. Resolution CNS No. 346/2005 on Multicenter Research: http://conselho.saude.gov.br/resolucoes/2005/Res346_en.pdf 3. Resolution MS/CNS No. 466/2012 - Guidelines and Rules for Research Involving Human Subjects:</p> | <p>CNS/CONEP: Orientation of Adverse Event Reporting in Clinical Trials (2011): http://conselho.saude.gov.br/web_comissoes/conep/carta_circular/Informacoes_sobre_o_formulario_para_submissao_de_Eventos_Adversos_Serios_a_CONEP.pdf</p> <p>ANVISA: 1. Manual of Adverse Event Notification and Safety Monitoring in Clinical Trials Involving Drugs (2016): http://portal.anvisa.gov.br/documents/33836/2492465/Manual+para+Notifica%C3%A7%C3%A3o+de+Eventos+Adversos+e+Monitoramento+de+Seguran%C3%A7a+em+Ensaio+Cl%C</p> |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | | | http://conselho.saude.gov.br/resolucoes/2012/466_english.pdf | 3%ADnicos+-+1%C2%AA+Edi%C3%A7%C3%A3o/04a68574-8aac-43c9-b0b2-7b7cd80831c4 2. Manual of Adverse Event Notification and Safety Monitoring in Clinical Trials Involving Medical Devices (2016): http://portal.anvisa.gov.br/documents/33912/2785629/MANUAL+PARA+NOTIFICA%C3%87%C3%83O+DE+EVENTOS+ADVERSOS+E+MONITORAMENTO+DE+SEGURAN%C3%87A+EM+ENSAIOS+CL%C3%8DNICOS+ENVOLVENDO+DISPOSITIVOS+M%C3%89DICOS+EM+INVESTIGA%C3%87%C3%83O/df22b9ac-688d-4e6a-8207-faf862a05994 |
| <i>Social-Behavioral Research</i> | National Commission on Research Ethics (CONEP): http://conselho.saude.gov.br/web_comissoes/conep/index.html | | Resolution No. 510 of April 7, 2016: http://conselho.saude.gov.br/resolucoes/2016/Reso510.pdf | |
| <i>Privacy/Data Protection</i> | 1. National Health Council (CNS): http://www.conselho.saude.gov.br/ 2. National Commission on Research Ethics (CONEP): http://conselho.saude.gov.br/web_comissoes/conep/index.html 3. Federal Council of Medicine: http://portal.cfm.org.br | | Resolution CFM N° 1.821, 23 November 2007: http://www.portalmedico.org.br/resolucoes/cfm/2007/1821_2007.htm | |
| <i>Human Biological Materials</i> | 1. National Health Council (CNS): http://www.conselho.saude.gov.br/ 2. National Commission on Research Ethics (CONEP): http://conselho.saude.gov.br/web_comissoes/conep/index.html 3. National Secretary on Science, Technology and Innovation: http://www.mctic.gov.br/mctic/opencms/institucional/paginaInstitucional.html | Ordinance No. 2.201/11: Establishing the National Guidelines for Biobanks of Human Biological Material for Research Purposes (2011): http://www2.inca.gov.br/wps/wcm/connect/8b19d5804eb688ee9cb39ef11fae00ee/portaria_2201_de_14_de_set_2011.pdf?MOD=AJPERES&CACHEID=8b19d5804eb688ee9cb39ef11fae00ee | CONEP: 1. Resolution CNS No. 441 of 12 May 2011: http://conselho.saude.gov.br/web_comissoes/conep/aquivos/resolucoes/Resolucao441_English_contribuicao_pesquisadora.doc 2. Resolution – RDC No. 20 of 10 April 2014: http://www.saude.pr.gov.br/arquivos/File/RDC_20_de_10_de_abril_2014_Transporte_de_material_Biologico.pdf | |
| <i>Genetic Research</i> | 1. National Commission on Research Ethics (CONEP): http://conselho.saude.gov.br/web_comissoes/conep/index.html 2. National Biosafety Technical Commission (CTNBio): http://ctnbio.mcti.gov.br/inicio 3. National Health Council (CNS): | 1. Biosafety Law 11.105/05 (2005): http://www.planalto.gov.br/ccivil_03/ato2004-2006/2005/lei/111105.htm 2. Decree No. 5.591, of November 22, 2005: http://www.planalto.gov.br/ccivil_03/ | CTNBio: 1. Instruction CTNBio No. 8 of 9 July 1997: http://ctnbio.mcti.gov.br/instrucoes-normativas/-/asset_publisher/3dOuwS2h7LU6/content/instrucao-normativa-ctnbio-n%C2%BA-8-de-09-07-97 | 1. Guidance to Researchers and Ethics Committees about the Item V.1.a of CNS Resolution 340 2004: http://conselho.saude.gov.br/web_comissoes/conep/aquivos/documentos/Carta_Circular_041_Orientacoes_pesquisadores_comites.pdf 2. Statement on Pharmacogenetic Studies in Brazil N° 011/2012/CONEP, 12 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | http://www.conselho.saude.gov.br/ | <p>3/ ato2004-2006/2005/Decreto/D5591.htm</p> <p>3. Law 13.123/2015 (2015): Brazilian Legislation on Biodiversity Access (Genetic Heritage): http://www.planalto.gov.br/ccivil_03/_Ato2015-2018/2015/Lei/L13123.htm</p> <p>4. Law Decree No 8.772/2016 (2016), Regulating Law No. 13.123/2015: http://www.planalto.gov.br/ccivil_03/_ato2015-2018/2016/decreto/D8772.htm</p> | <p>2. Instruction CTNBio No. 9 of 10 October 1997: http://www.agrobiobrasil.org.br/wp-content/uploads/2014/06/CTNBio-Normative-Instruction-n%C2%BA-9-of-October-10-1997.pdf</p> <p>3. Resolution CNS No. 340/2004: On Research on Human Genetics (2004): http://conselho.saude.gov.br/resolucoes/2004/Res340_en.pdf</p> | <p>January 2012: http://www.fcm.unicamp.br/fcm/sites/default/files/11_-_Comunicado_sobre_estudos_farmacogeneticos_no_Brasil.pdf</p> |
| <i>Embryos, Stem Cells, and Cloning</i> | <p>1. National Biosafety Technical Commission: http://ctnbio.mcti.gov.br/inicio</p> <p>2. National Commission on Research Ethics (CONEP): http://conselho.saude.gov.br/web_comissoes/conep/index.html</p> <p>3. National Health Council (CNS): http://www.conselho.saude.gov.br/</p> | <p>1. Biosafety Law 11.105/05 (2005): http://www.planalto.gov.br/ccivil_03/_ato2004-2006/2005/lei/111105.htm</p> <p>2. Decree No. 5.591, of November 22, 2005: http://www.planalto.gov.br/ccivil_03/_ato2004-2006/2005/Decreto/D5591.htm</p> | <p>1. Resolution RDC No. 9, 14 March 2011: http://bvsms.saude.gov.br/bvs/saudelegis/anvisa/2011/prt0009_14_03_2011.html</p> <p>2. Resolution RDC No. 29, 12 May 2008: http://bvsms.saude.gov.br/bvs/saudelegis/anvisa/2008/rdc0029_12_05_2008.html</p> | |
| Chile | | | | |
| Note: All websites and documents are in Spanish. | | | | |
| <i>General</i> | <p>1. Ministry of Health: http://www.minsal.cl</p> <p>2. Institute of Public Health: http://www.ispch.cl</p> | <p>1. Law No. 20.120 Regarding Scientific Research in Human Beings, their Genome, and the Prohibition of Human Cloning (2006): http://www.leychile.cl/Navegar?idNorma=253478</p> <p>2. Law No. 20584. Regulating the Rights and Duties Incumbent upon Persons in Connection with Actions Linked to their Health Care (2012): http://www.leychile.cl/Navegar?idNorma=1039348</p> <p>3. Law No. 20.724 Modifying the Health Code in the Area of the Regulation of Pharmacies and Medications (2014): http://www.leychile.cl/Navegar?idNorma=1039348</p> | <p>1. Supreme Decree No. 114 of 2010: Regulation on Law No. 20.120 Regarding Scientific Research in Human Beings, the Genome, and the Prohibition of Human Cloning. Official Diary of November 19, 2011: http://www.leychile.cl/Navegar?idNorma=1032919</p> <p>2. Supreme Decree N° 30/2013 Regulation on Law N°20.120 Modifying Supreme decree N°114/2010, Regulation on Law No. 20.120 Regarding Scientific Research in Human Beings, the Genome, and the Prohibition of Human Cloning Official Diary January 14, 2013: http://www.leychile.cl/Navegar?idNorma=1032919</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <p><i>General</i></p> <p><i>Drugs, Biologics, and Devices</i></p> | <p>1. Ministry of Health : http://www.minsal.cl</p> <p>2. Institute of Public Health: http://www.ispch.cl</p> | <p>orma=1058373</p> <p>Law No. 20.724 Modifying the Health Code in the Area of the Regulation of Pharmacies and Medications (2014): http://www.leychile.cl/Navegar?idNorma=1058373</p> | <p>rma=1048008&</p> <p>1. Supreme Decree No. 114 of 2010: Regulation on Law No. 20.120 Regarding Scientific Research in Human Beings, the Genome, and the Prohibition of Human Cloning. Official Diary of November 19, 2011: http://www.leychile.cl/Navegar?idNorma=1032919</p> <p>2. Supreme Decree No. 3 of 2010. Regulation of the National Control System of Pharmaceutical Products for Human Use. Official Diary of June 25, 2011: http://www.ispch.cl/ley20285/t_activa/marco_normativo/7c/ds_minsal_3_2010.pdf</p> <p>3. Exempt Resolution 2263, July 30th 2015 Modifying Resolution N° 403 Ex. February 5, 2015 that Approves the Guidelines for Use Control of Pharmaceuticals Products in Scientific Research: http://www.leychile.cl/Navegar?idNorma=1080011</p> | |
| <p><i>Research Injury</i></p> | <p>1. Ministry of Health: http://www.minsal.cl</p> <p>2. Institute of Public Health: http://www.ispch.cl</p> | <p>Law No. 20.120 Regarding Scientific Research in Human Beings, their Genome, and the Prohibition of Human Cloning (2006): http://www.leychile.cl/Navegar?idNorma=253478</p> | <p>1. Supreme Decree No. 3 of 2010. Regulation of the National Control System of Pharmaceutical Products for Human Use. Official Diary of Jun 25, 2011: http://www.ispch.cl/ley20285/t_activa/marco_normativo/7c/ds_minsal_3_2010.pdf</p> <p>2. General Technical Rule No. 140 Regarding the National System of Pharmacovigilance of Pharmaceutical Products for Human Use. June 20, 2012: http://web.minsal.cl/portal/url/item/c4a31ad6db50e085e040010165017a39.pdf</p> <p>3. Resolution No. 441, Notification of Adverse events in Clinical Research in Chile,</p> | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | | | February 13, 2012: http://www.ispch.cl/sites/default/files/res_441.pdf | |
| <i>Privacy/Data Protection</i> | 1. Ministry of Health: http://www.minsal.cl 2. Ministry of the Secretary General of the Government: http://www.msgg.gob.cl | 1. Law for the Protection of Private Life No. 19.628 (1999): http://www.bcn.cl/leves/141599 2. Law No. 20584. Regulating the Rights and Duties Incumbent upon Persons in Connection with Actions Linked to their Health Care (2012): http://www.leychile.cl/Navegar?idNorma=1039348 | Supreme Decree No. 41 of 2012: Regulation Regarding Clinical Records of December 15, 2012: http://www.leychile.cl/Navegar?idNorma=1046753 | |
| <i>Genetic Research</i> | Ministry of Health: http://www.minsal.cl | Law No. 20.120: Scientific Research Involving Human Beings, Their Genome, and Prohibition of Human Cloning (2006): http://www.leychile.cl/Navegar?idNorma=253478 | Supreme Decree No. 114 of 2010: Regulation on Law No. 20.120 Regarding Scientific Research in Human Beings, the Genome, and the Prohibition of Human Cloning. Official Diary of November 19, 2011: http://www.leychile.cl/Navegar?idNorma=1032919 | |
| <i>Embryos, Stem Cells, and Cloning</i> | Ministry of Health: http://www.minsal.cl | Law No. 20.120: Scientific Research Involving Human Beings, Their Genome, and Prohibition of Human Cloning (2006): http://www.leychile.cl/Navegar?idNorma=253478 | Supreme Decree No. 114 of 2010: Regulation on Law No. 20.120 Regarding Scientific Research in Human Beings, the Genome, and the Prohibition of Human Cloning. Official Diary of November 19, 2011: http://www.leychile.cl/Navegar?idNorma=1032919 | |
| Colombia | | | | |
| Note: All websites and documents are in Spanish. | | | | |
| <i>General</i> | 1. Ministry of Health and Social Protection: http://www.minsalud.gov.co 2. National Institute of Drug and Food Surveillance (INVIMA): https://www.invima.gov.co/ 3. Administrative Department of Science, Technology, and Innovation (COLCIENCIAS): http://www.colciencias.gov.co/ | | Scientific, Technical, and Administrative Regulations for Health Research, Resolution No. 8430 (1993): https://www.minsalud.gov.co/sites/rid/Lists/BibliotecaDigital/RIDE/DE/DI/J/RESOLUCION-8430-DE-1993.PDF | INVIMA 1. Guide for Research Ethics Committees. Code: ASS-RSA-GU040 Version: 00 (2015): https://www.invima.gov.co/images/stories/for/matotramite/ASS-RSA-GU040.pdf 2. Guide for Assessing and Monitoring of Research Protocols. Code: ASS-RSA-GU039 Version: 03 (2018): https://www.invima.gov.co/images/stories/for/matotramite/ASS-RSA-GU039.pdf |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | <i>Drugs</i> National Institute of Drug and Food Surveillance (INVIMA): http://www.invima.gov.co/ | | 1. Resolution No. 2378 of 2008, Adapting Good Clinical Practices for Institutions that Conduct Research with Medicines in Human Beings: http://www.alcaldiabogota.gov.co/sisjur/normas/Norma1.jsp?i=31169 2. Resolution No. 2011020764 of June 10th, 2011: Regulation Related to the Content and Frequency of Adverse Event Reports in Clinical Investigation in Humans: https://www.invima.gov.co/index.php?option=com_content&view=article&id=725:resolucion-no-2011020764-del-10-de-junio-de-2011&catid=58:2011&Itemid=105 | 1. ABC Good Clinical Practice (2009) https://www.invima.gov.co/images/pdf/tecnovigilancia/buenas_practicas/ABCBCPultima_version.pdf 2. Circular No 600-5776-14: Processes of Good Clinical Practice (2014): https://www.invima.gov.co/images/pdf/tecnovigilancia/buenas_practicas/normatividad/CIRCULAR_600-5776-14-2.pdf 3. Guide of Medications and Supplies for Clinical Research, Version 1 (2018): https://www.invima.gov.co/images/stories/for_matotramite/ASS-RSA-GU045.pdf 4. Guide for the Evaluation and Follow-up of Research Protocols, Version 3 (2018): https://www.invima.gov.co/images/stories/for_matotramite/ASS-RSA-GU039.pdf 5. External Circular No. 600-2006-16: National Reporting Serious Adverse Events (2016): https://www.invima.gov.co/images/pdf/tecnovigilancia/buenas_practicas/normatividad/Circular-600-1081-16-Reporte-de-Eventos-adversos-serios-Nacionales-Febrero2016.pdf 6. External Circular No. 600-1414-16: Notification of Deviations (2016): https://www.invima.gov.co/images/pdf/tecnovigilancia/buenas_practicas/normatividad/Circular_600-2006-16_Alcance-Circular-600-1081-16_Abril2016.pdf |
| | <i>Devices</i> National Institute of Drug and Food Surveillance: http://www.invima.gov.co/ | | Scientific, Technical, and Administrative Regulations for Health Research, Resolution No. 008430, Title III, Chapters I and III (1993): https://www.minsalud.gov.co/sites/rid/Lists/BibliotecaDigital/RIDE/DE/DIJ/RESOLUCION-8430-DE-1993.PDF | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Research Injury</i> | Ministry of Health and Social Protection: http://www.minsalud.gov.co | | Scientific, Technical, and Administrative Regulations for Health Research, Resolution No. 008430, Title II, Chapter I, Art. 13 (1993): https://www.minsalud.gov.co/sites/rid/Lists/BibliotecaDigital/RIDE/DE/DIJ/RESOLUCION-8430-DE-1993.PDF | |
| <i>Privacy/Data Protection</i> | Ministry of Health and Social Protection: http://www.minsalud.gov.co | 1. Constitution of Colombia, Article 15 (2003): http://www.corteconstitucional.gov.co/inicio/Constitucion%20politica%20de%20Colombia%20-%202015.pdf 2. Law 1581 of 2012: General Regimen of Protection of Personal Data: https://www.mintic.gov.co/portal/604/articles-4274_documento.pdf | Scientific, Technical, and Administrative Regulations for Health Research, Resolution No. 008430, Title II, Chapter I, Article 8 (1993) | |
| <i>Human Biological Materials</i> | Ministry of Health and Social Protection: http://www.minsalud.gov.co | | 1. Scientific, Technical, and Administrative Regulations for Health Research, Resolution No. 008430, Title II, Chapter VI (1993): https://www.minsalud.gov.co/sites/rid/Lists/BibliotecaDigital/RIDE/DE/DIJ/RESOLUCION-8430-DE-1993.PDF 2. Requirements for the Use of Unclaimed Cadavers for Research Purposes, Resolution No. 002640, Article 21 (2005): https://www.minsalud.gov.co/sites/rid/Lists/BibliotecaDigital/RIDE/DE/DIJ/Resolución_2640_de_2005.pdf | |
| <i>Genetic Research</i> | Ministry of Health and Social Protection: http://www.minsalud.gov.co | | Scientific, Technical, and Administrative Regulations for Health Research, Resolution No. 008430, Title III, Chapter II (1993): https://www.minsalud.gov.co/sites/rid/Lists/BibliotecaDigital/RIDE/DE/DIJ/RESOLUCION-8430-DE-1993.PDF | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Costa Rica | | | | |
| Note: All websites and documents are in Spanish. | | | | |
| <i>General</i> | Ministry of Health: http://www.misalud.go.cr | | Reform Regulation to the Biomedical Research Regulatory Law: http://www.pgrweb.go.cr/scij/Busqueda/Normativa/Normas/nrm_texto_completo.aspx?param1=NRTC&nValor1=1&nValor2=81213&nValor3=103450&strTipM=TC | |
| <i>Drugs, Biologics, and Devices</i> | National Health Research Council: http://www.ministeriodesalud.go.cr/index.php/consejos/conis | Regulatory Law of Biomedical Research No. 9234 (2014): http://www.pgrweb.go.cr/scij/Busqueda/Normativa/Normas/nrm_texto_completo.aspx?param1=NRTC&nValor1=1&nValor2=77070&nValor3=96424&strTipM=TC | 1. Regulatory Decree N° 39061-S (2016) on the Regulatory Law of Biomedical Research N° 39533-S: http://www.pgrweb.go.cr/scij/Busqueda/Normativa/Normas/nrm_texto_completo.aspx?param1=NRTC&nValor1=1&nValor2=81213&nValor3=103450&strTipM=TC 2. Reforms to the Regulatory Decree No. 39533-S (2016) Regulatory Law of Biomedical Research No. 9234: http://www.pgrweb.go.cr/scij/Busqueda/Normativa/Normas/nrm_texto_completo.aspx?param1=NRTC&nValor1=1&nValor2=81213&nValor3=103450&strTipM=TC | Various: http://www.ministeriodesalud.go.cr/index.php/consejos/conis |
| <i>Clinical Trials Registry</i> | National Health Research Council (Spanish): http://www.ministeriodesalud.go.cr/index.php/consejos/conis (scroll to bottom of page to Investigaciones Registradas) | | | |
| Cuba | | | | |
| Note: All websites and documents are in Spanish. | | | | |
| <i>Drugs, Biologics, and Devices</i> | Center for State Control of Medications: http://www.cecmecd.cu/ | | | Various: http://www.cecmecd.cu/ensayos-clinicos/autorizos |
| <i>Clinical Trials Registry</i> | Public Cuban Registry of Clinical Trials: http://registroclinico.sld.cu/en/home | | | |
| Dominica | | | | |
| <i>General</i> | Ministry of Health: http://www.dominica.gov.dm/cms/index.php?q=node/21 | | | Guidelines for the Conduct of Research on Human Subjects (2005) |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Dominican Republic | | | | |
| <i>General</i> | National Council on Health Bioethics: http://conabios.gob.do/ | National Health Law 42-01, Chapter VI: https://www.dol.gov/ilab/submissions/pdf/20100408-10.pdf | Regulation for Evaluation Request for a Clinical Investigation Project: http://conabios.gob.do/index.php/reglamentos | |
| <i>Biological Materials</i> | | National Health Law 42-01, Book Five: https://www.dol.gov/ilab/submissions/pdf/20100408-10.pdf | | |
| Ecuador | | | | |
| Note: All websites and documents are in Spanish. | | | | |
| <i>General</i> | Ministry of Public Health : http://www.salud.gob.ec/ | 1. Constitution of the Republic: http://www.asambleanacional.gob.ec/sites/default/files/documents/old/constitucion_de_bolsillo.pdf 2. Organic Health Law of 22 December 2006, Articles 207- 208: http://www.vertic.org/media/National%20Legislation/Ecuador/EC_Ley_Organica_de_Salud.pdf 3. Code on Childhood and Adolescence. Law 100 Official Register 737 of January 3, 2003 (2014): http://www.funcionjudicial.gob.ec/otaip/index.php/component/phocadownload/category/1-nacional?download=283:codigo-de-la-ninez-y-adolescencia | 1. Regulation for the Approval of Ethics Committees (2014): http://instituciones.msp.gob.ec/images/Documentos/CNBS/1%20normativa/Registro%20Oficial%20Comites%20de%20Etica%20julio%202014.pdf 2. Regulation on Research, Ministerial Agreement No. 0066, Public Registry No. 292 (March 11, 2008): http://www.controlsanitario.gob.ec/wpcontent/uploads/downloads/2017/01/A.M.-66-REGLAMENTODE-PROYECTOS-ENINVESTIGACION-DESALUD.pdf | National Policy on Scientific Research. Ministerial Agreement 209, Public Registry No. 87 of August 23, 2005 |
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| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | <p>20Comites%20de%20Etica%20julio%202014.pdf</p> <p>3. Regulation on Research, Ministerial Agreement No. 0066, Public Registry No. 292 (March 11, 2008): http://www.controlsanitario.gob.ec/wp-content/uploads/downloads/2017/01/A.M.-66-REGLAMENTO-DE-PROYECTOS-EN-INVESTIGACION-DE-SALUD.pdf</p> | |
| <i>Privacy/Data Protection</i> | Ministry of Public Health: http://www.salud.gob.ec/ | Constitution of the Republic of Ecuador 2008 (Article: 92): http://www.asambleanacional.gob.ec/sites/default/files/documents/old/constitucion_de_bolsillo.pdf | Ministerial Agreement No. 005216, Public Registry No. 427, Confidential Information in National Health System (January, 29, 2015): http://instituciones.msp.gob.ec/cz6/images/lotaip/Enero2015/Acuerdo%20Ministerial%205216.pdf | |
| <i>Biological Materials</i> | National Institute on Donation and Transplantation of Organs, Tissues, and Cells: http://www.donaciontrasplante.gob.ec/indot/ | <p>1. Organic Health Law of December 22, 2006, Articles 81-86: http://www.vertic.org/media/National%20Legislation/Ecuador/EC_Ley_Organica_de_Salud.pdf</p> <p>2. Organic Law on the Donation and Transplantation of Organs, Tissues, and Cells (2011): http://www.donaciontrasplante.gob.ec/indot/wp-content/uploads/downloads/2013/11/ley_y_reglamento_a_la_ley_organica_de_donacion_y_trasplantes.pdf</p> | <p>1. Executive Order 1205, July 13, 2012: Regulation for the Organic Law on the Donation and Transplantation of Organs, Tissues, and Cells: http://www.donaciontrasplante.gob.ec/indot/wp-content/uploads/downloads/2013/11/ley_y_reglamento_a_la_ley_organica_de_donacion_y_trasplantes.pdf</p> <p>2. Import and Export of Human Biological Samples for research. Ministerial Agreement No. 0088, Public Registry No. 34, (July 12, 2017): http://www.controlsanitario.gob.ec/wp-content/uploads/downloads/2017/08/Acuerdo-Ministerial-0088-2017_Autorizaci%C3%B3n-de-importaci%C3%B3n-y-exportaci%C3%B3n-de-muestras-biol%C3%B3gicas.pdf</p> | |
| <i>Genetic Research</i> | Ministry of Public Health: http://www.salud.gob.ec/ | Organic Health Law, December 22, 2006, Articles 209-210 (2011): http://www.vertic.org/media/Nation | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Genetic Research</i> | | al%20Legislation/Ecuador/EC Ley Organica de Salud.pdf | | |
| <i>Embryos, Stem Cells, and Cloning</i> | 1. Ministry of Public Health: http://www.salud.gob.ec/ 2. National Institute of Donation and Transplantation of Organs, Tissues, and Cells: http://www.donaciontrasplante.gob.ec/indot/ | Organic Health Law of 22 December 2006, Article 214 (2011): http://www.vertic.org/media/National%20Legislation/Ecuador/EC Ley Organica de Salud.pdf | Regulation for the Organic Law on the Donation and Transplantation of Organs, Tissues, and Cells. Executive Order 1205, July 13, 2012: http://www.donaciontrasplante.gob.ec/indot/wp-content/uploads/downloads/2013/11/ley_y_reglamento_a_la_ley_organica_de_donacion_y_trasplantes.pdf | |
| El Salvador | | | | |
| <i>General</i> | National Health Research Ethics Committee: http://www.cneis.org.sv/ | | | 1. Standard Operating Procedures for the Ethical Evaluation of Health Research (2015): http://cssp.gob.sv/wp-content/uploads/2016/06/MANUAL-CNEIS-2017-03-15.pdf 2. Manual on the Functioning of the National Health Research Ethics Committee (2017): http://cssp.gob.sv/wp-content/uploads/2016/06/manual_funcionamiento_comite_nacional_etica_investigacion_en_salud.pdf |
| <i>Drugs and Devices</i> | National Directorate of Medications: http://www.medicamentos.gob.sv/index.php/es/ | | | User's Guide for the Application of Clinical Investigation Protocols: http://www.medicamentos.gob.sv/index.php/es/servicios-m/descargables/ensayos-clinicos |
| Grenada | | | | |
| <i>General</i> | St. George's University/Windward Islands Research and Education Foundation: http://www.sgu.edu/school-of-medicine/institutional-review-board.html | | | 45 CFR 46: http://www.hhs.gov/ohrp/humansubjects/guidance/45cfr46.html |
| Guyana | | | | |
| <i>General</i> | | | Medical Research Involving Human Subjects Regulations (2007) | |
| Guatemala | | | | |
| Note: All websites and documents are in Spanish. | | | | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>General</i> | Ministry of Public Health and Social Assistance: http://www.mspas.gob.gt/ | | Internal Regulations of the National Committee on Health Ethics (2018): http://www.mspas.gob.gt/images/files/acuerdosministeriales/2018/AcuerdoMinisterial1392018NormativaCNES.pdf | |
| <i>Drugs, Biologics, and Devices</i> | Ministry of Public Health and Social Assistance: http://www.mspas.gob.gt/ | | 1. Governmental Agreement 712-99, Articles 91-94 (1999): http://asisehace.gt/media/ag_712_99.pdf 2. Rules for the Regulation of Human Clinical Trials. Ministerial Accord SP-M-466-2007: https://healthresearchweb.org/?action=download&file=GUA_EnsayosClinicos_Acuerdo_SPM4662007I11.pdf | Drug Surveillance -- Clinical Trials: http://www.mspas.gob.gt/index.php/servicios/farmacovigilancia |
| Haiti | | | | |
| <i>General</i> | Ministry of Public Health and Population: http://mspp.gouv.ht/newsite/ | | | Internal Regulations (2010) |
| Honduras | | | | |
| Note: All websites and documents are in Spanish. | | | | |
| <i>General</i> | Secretariat of Health: http://www.salud.gob.hn/ | | Health Code, Decree No. 65-91, Articles 175 and 176 | |
| <i>Human Biological Materials</i> | | Law of Donation and Transplantation of Anatomical Organs in Human Beings (2014): http://www.tsc.gob.hn/leyes/Ley_donacion_transp_organos_2014.pdf n. 329-2013 | | |
| Jamaica | | | | |
| <i>General</i> | Ministry of Health, Ethics and Medico-Legal Affairs Panel: http://moh.gov.jm/ | | | Ministry of Health Guidelines for the Conduct of Research on Human Subjects (2010): http://moh.gov.jm/guidelines/guidelines-for-the-conduct-of-research-on-human-subjects/ |
| <i>Drugs, Biologics, and Devices</i> | Ministry of Health, Standards and Regulation Division: http://moh.gov.jm/divisions-agencies/divisions/standards-and-regulation-division/ | Food and Drugs Act (1975): http://www.moj.gov.jm/sites/default/files/laws/Food%20and%20Drugs%20Act%20LN%2065%20of%2075.pdf | Food and Drugs Regulations (1975): http://www.moj.gov.jm/sites/default/files/laws/Food%20and%20Drugs%20Act%20LN%2065%20of%2075.pdf | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| México Note: All websites and documents are in Spanish. | | | | |
| <i>General</i> | 1. Secretariat of Health: https://www.gob.mx/salud 2. General Health Council: http://www.csg.gob.mx/ 3. National Bioethics Commission (CNB): https://www.gob.mx/salud/conbioetica 4. Federal Commission for Protection Against Health Risks: https://www.gob.mx/cofepris | General Health Law, Title V, Chapter 1, Articles 96-103: Health Research (2014) | 1. Rule NOM-012-SSA3-2012 Establishing Criteria for the Conduct of Health Research Projects (2013): http://dof.gob.mx/nota_detalle.php?codigo=5284148&fecha=04/01/2013 2. Regulation on the General Health Law in the Matter of Health Research (2014): http://www.diputados.gob.mx/LeyesBiblio/regley/Reg_LGS_MIS.pdf | CNB: National Guidelines for the Integration and Operation of Research Ethics Committees (2016): http://www.conbioetica-mexico.salud.gob.mx/descargas/pdf/registrocomites/Guia_CEI_paginada_con_forros.pdf |
| <i>Drugs, Biologics, and Devices</i> | Federal Commission for Protection Against Health Risks (COFEPRIS): https://www.gob.mx/cofepris | General Health Law, Title V, Chapter I, Articles 96-103: Health Research (2014) | Regulation on the General Health Law in the Matter of Health Research (2014): http://www.diputados.gob.mx/LeyesBiblio/regley/Reg_LGS_MIS.pdf | 1. Guidelines to Fulfill Good Clinical Practice in Health Research: http://www.ccinshae.salud.gob.mx/descargas/5.5.-Guia_Protocolo_Inicial.pdf 2. Technical Rule No. 314 for Registration and Follow-up in the Area of Health Research (1998): https://ssj.jalisco.gob.mx/sites/ssj.jalisco.gob.mx/files/norma_tecnica_313-314-315_0.pdf |
| <i>Privacy/Data Protection</i> | Federal Institute on Access to Public Information: www.inai.org.mx/ | 1. Federal Law for the Protection of Personal Data in the Possession of Private Individuals (2017): http://www.dof.gob.mx/nota_detalle.php?codigo=5469949&fecha=26/01/2017 2. Federal Law on Transparency and Access to Public Information (2017): http://www.diputados.gob.mx/LeyesBiblio/pdf/LFTAIP_270117.pdf | | |
| <i>Human Biological Materials</i> | Secretariat of Health: https://www.gob.mx/salud | General Health Law, Title XIV, Articles 313-342 (2018): http://www.salud.gob.mx/unidades/cdi/legis/lgs/index-t14.htm | Regulation of the General Law of Health in Matter of Transplants (2014): http://www.diputados.gob.mx/LeyesBiblio/regley/Reg_LGS_MT.pdf | |
| <i>Genetic Research</i> | National Institute of Genomic Medicine: http://www.inmegen.gob.mx/ | 1. Biosafety Law on Genetically Modified Organisms (2017): http://www.dof.gob.mx/nota_detalle.php?codigo=5468449&fecha=03/01/2017 2. Regulation of the Biosafety Law on Genetically Modified | Regulation on the General Health Law in the Matter of Health Research, Title Four, Chapter Two (2014): http://www.diputados.gob.mx/LeyesBiblio/regley/Reg_LGS_MIS.pdf | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Panamá | | | | |
| <i>General</i> | 1. Ministry of Health (MINSA): http://www.minsa.gob.pa/ 2. ICGES Bioethics Research Committee (CBI): http://www.gorgas.gob.pa/index.php?option=com_content&view=article&id=54&Itemid=103&lang=es | | MINSA: 1. Resolution No. 390 Adopting the Operational Guide for Research Bioethics, Official Gazette 24,938 (2003) 2. Executive Decree N°1843 on the National Research Ethics Committee of Panama (2014): https://www.gacetaoficial.gob.pa/.../GacetaNo_27716_20150206.pdf 3. Executive Decree N° 6 on the National Research Ethics Committee of Panama (2015): https://www.gacetaoficial.gob.pa/pdfTemp/27716/GacetaNo_27716_20150206.pdf | CBI : Various: http://www.gorgas.gob.pa/index.php?option=com_content&view=article&id=54&Itemid=103&lang=es |
| <i>Drugs, Biologics, and Devices</i> | | Law 1 of 2001, Official Gazette 24,218: http://www.perezcarrera.com/leyes/ley-registro-sanitario-panama.pdf | | |
| <i>Privacy/Data Protection</i> | | Law 68 of 2003, Official Gazette 24,935 | | |
| <i>Human Biological Materials</i> | | Law 3 of 2010, Official Gazette 26,468-B on Transplant of Organs and Tissues: https://www.gacetaoficial.gob.pa/pdfTemp/26468_B/GacetaNo_26468b_20100210.pdf | | |
| <i>Embryos, Stem Cells, and Cloning</i> | | | Executive Decree No. 2 on Stem Cells (2013): http://www.gacetaoficial.gob.pa/pdfTemp/27207/40367.pdf | |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Perú | | | | |
| For an overview of clinical research regulations in Peru, see the ClinRegs report: http://clinregs.niaid.nih.gov/single_country.php?c_id=170 | | | | |
| <i>General</i> | National Institute of Health: http://www.ins.gob.pe/ | General Health Law No. 26842, Article 28 (1997): http://www.wipo.int/wipolex/en/text.jsp?file_id=203140 | | |
| <i>Drugs, Biologics, and Devices</i> | 1. National Institute of Health: http://www.ins.gob.pe/ 2. National Directorate of Drugs and Medical Devices: www.digemid.minsa.gob.pe | | Supreme Decree No. 021-2017-SA. Regulation of Clinical Trials (2017): http://busquedas.elperuano.com.pe/download/url/aprueban-reglamento-de-ensayos-clinicos-decreto-supremo-n-021-2017-sa-1538902-2 | Various: http://www.ensayosclinicos-repec.ins.gob.pe/otros-repec/216-comunicados |
| <i>Clinical Trials Registry</i> | Peruvian Registry of Clinical Trials: http://www.ensayosclinicos-repec.ins.gob.pe/en/about-repec/clinical-trial-search | | Supreme Decree No. 021-2017-SA. Regulation of Clinical Trials, Title III, Chapter I, Articles 16 - 32 (2017): http://busquedas.elperuano.com.pe/download/url/aprueban-reglamento-de-ensayos-clinicos-decreto-supremo-n-021-2017-sa-1538902-2 | |
| <i>Research Injury</i> | National Institute of Health: http://www.ins.gob.pe/ | | Regulation on Clinical Trials in Peru: Articles 26, 27 and 28: http://www.ins.gob.pe/portal/jerarquia/2/990/reglamento-de-ensayos-clinicos/jer.990 | |
| <i>Privacy/Data Protection</i> | National Directorate of Drugs and Medical Devices: www.digemid.minsa.gob.pe | 1. Law 29733 for the Protection of Personal Information: http://www.minjus.gob.pe/legislacion/ 2. Law for Electronic Medical Charts (2013): http://elperuanolegal.blogspot.com/2013/05/ley-30024-ley-que-crea-el-registro.html | | |
| Saint Lucia | | | | |
| <i>Drugs, Biologics, and Devices</i> | | Clinical Trials Act (2016): http://slugovprintery.com/template/files/document_for_sale/laws/3742/Act%2010%20of%202016.pdf | | |
| Trinidad and Tobago | | | | |
| | 1. Ministry of Health http://www.health.gov.tt/ 2. University of the West Indies (UWI), St. Augustine: https://sta.uwi.edu/research/ethics.asp | | | UWI: 1. UWI Policy on Research Ethics 2. Application Guidelines 3. Ethics Committee Protocols |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Uruguay | | | | |
| Note: All websites and documents are in Spanish. | | | | |
| <i>General</i> | Ministry of Public Health: http://www.msp.gub.uy/ | 1. Decree 189/998 http://www.mercosur.int/msweb/Normas/normas_web/Resoluciones/PT/GMC_RES_1996-129_PT_RT%20Verifica%20BPPesquisaClinica.PDF 2. Decree 379/008: http://www.elderechodigital.com.uy/smu/legisla/D0800379.html | Decree No. 370/2008: Regulation Concerning Research with Humans | |
| <i>Drugs, Biologics, and Devices</i> | Ministry of Public Health: http://www.msp.gub.uy/ | Decree 189/998: http://www.mercosur.int/msweb/Normas/normas_web/Resoluciones/PT/GMC_RES_1996-129_PT_RT%20Verifica%20BPPesquisaClinica.PDF | | |
| <i>Research Injury</i> | Ministry of Public Health: http://www.msp.gub.uy/ | 1. Decree 189/998 http://www.mercosur.int/msweb/Normas/normas_web/Resoluciones/PT/GMC_RES_1996-129_PT_RT%20Verifica%20BPPesquisaClinica.PDF 2. Decree 379/008: http://www.elderechodigital.com.uy/smu/legisla/D0800379.html | | |
| <i>Privacy/Data Protection</i> | Ministry of Public Health: http://www.msp.gub.uy/ | 1. Law 18.331: http://www0.parlamento.gub.uy/leyes/ AccesoTextoLey.asp?Ley=18331 2. Decree 379/008: http://www.elderechodigital.com.uy/smu/legisla/D0800379.html | | |
| <i>Human Biological Materials</i> | 1. Ministry of Public Health: http://www.msp.gub.uy/ 2. National Institute on Donation and Transplantation: www.indt.edu.uy | Decree 160/006: http://www.indt.edu.uy/documentos/documentacion_legal/decreto_160-006.pdf | | |
| Venezuela | | | | |
| Note: All websites and documents are in Spanish. | | | | |
| <i>General</i> | 1. National Fund on Science and Technology, Commission on Bioethics and Biosecurity | Constitution, Article 46 (3): http://www.venezuelaemb.or.kr/english/ConstitutionoftheBolivarianingles.pdf | Resolution No. 48 (1998): http://www.ivic.gob.ve/bioetica/?mod=bioeticahome.php | FONACIT: Code on Bioethics and Biosecurity (2002) |

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| <i>General</i> | (FONACIT): www.fonacit.gov.ve/ 2. Venezuelan Institute of Scientific Research, Bioethics Commission (IVIC): http://www.ivic.gob.ve/bioetica/?mod=home.php | | | IVIC: 1. Annex 1: General Ethical Issues in Research Involving Living Persons: http://www.ivic.gob.ve/bioetica/?mod=Anexo.php 2. Annex 2: Necessity of Establishing a Clear and Precise Study Protocol Before Starting Research: http://www.ivic.gob.ve/bioetica/?mod=Anexo.php 3. Informed Consent: http://www.ivic.gob.ve/bioetica/?mod=manual.php |
| <i>Drugs, Biologics, and Devices</i> | National Institute of Hygiene “Rafael Rangel”: http://www.inhrr.gob.ve/ | Medicines Act, Title III, Chapter II: http://www.ginecowed.com/PDF/Le-y-del-Ejercicio-de-la-Medicina.pdf | | |
| <i>Genetic Research</i> | Venezuelan Institute of Scientific Research, Bioethics Commission: http://www.ivic.gob.ve/bioetica/?mod=home.php | | | 1. Contract for Accessing Genetic Resources (2003): http://www.ivic.gob.ve/bioetica/contrato.pdf 2. Revised Outline of the International Declaration of Human Genetic Data (2003): http://www.ivic.gob.ve/bioetica/chapter3.pdf |

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| AFRICA | | | | |
| Regionwide | | | | |
| <i>Clinical Trials Registry</i> | Pan African Clinical Trials Registry: http://www.pactr.org/ | | 1. Order No. 387 of July 31, 2006 Regarding Clinical Trials: http://www.ands.dz/pharmacie-med/arr%C3%AAt%C3%A9_n%C2%B0387-388_31_juil_2006.pdf 2. Order No. 00200 of July 25, 2009 Modifying Order No. 112 of October 22, 1995 Establishing Rules on Good Clinical Practice: http://www.ands.dz/pharmacie-med/arr%C3%AAt%C3%A9_n%C2%B0200%20_25_Juil_2009.pdf | FAQs: http://www.pactr.org/ATMWeb/appmanager/atm/atmregistry?_nfpb=true&_pageLabel=atmportal_page_FAQ |
| Algeria | | | | |
| <i>Drugs, Biologics, and Devices</i> | Directorate of Pharmacy and Medicine: http://www.ands.dz/ | | 1. Order No. 387 of 31 July 2006 Relating to Clinical Trials: http://www.ands.dz/pharmacie-med/arr%C3%AAt%C3%A9_n%C2%B0387-388_31_juil_2006.pdf 2. Order No. 00200 of 25 July 2009 Amending Order No. 112 of 22 October 1995 Setting the Rules of Good Clinical Practice: http://www.ands.dz/pharmacie-med/arr%C3%AAt%C3%A9_n%C2%B0200%20_25_Juil_2009.pdf | |
| Benin | | | | |
| <i>General</i> | | Law No. 2010-40 of 8 December, 2010 Regarding the Ethical Code and Duties in Health Research in the Republic of Benin: http://ethique-sante.org/pdf/loi-portant-code-ethique.pdf | | |
| Botswana | | | | |
| <i>General</i> | Ministry of Health, Research and Development Committee: http://www.moh.gov.bw/ | Anthropological Research Act 45 (1967): http://www.elaws.gov.bw/docs/statutes/Botswana%20Statute%20Law%201967.pdf | | 1. Guidelines for Application for Research Permit (2004): http://www.gov.bw/Global/OP%20Ministry/RESEARCH%20PERMIT%20GUIDELINES.pdf 2. Guide for a Consent Form (2005) 3. Guidelines for the Review of Research Proposals (2005) |

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| <i>Drugs, Biologics, and Devices</i> | Ministry of Health, Drug Regulatory Unit: http://www.moh.gov.bw/ | | Drugs and Related Substances Regulations (1993) | 1. SADC Guidelines for Regulating Clinical Trials in Human Subjects (2006) 2. Guideline for Regulating the Conduct of Clinical Trials Using Medicines in Human Participants (2012): http://www.moh.gov.bw/Publications/drug_regulation/CLINICAL%20TRIAL%20GUIDELINES%20botswana%20v4-060312.pdf |
| <i>Social-Behavioral Research</i> | Ministry of Health and Wellness, Research and Development Committee | Anthropological Research Act 45 (1967): http://webcache.googleusercontent.com/search?q=cache:A7aea2ZEMhkJ:static1.1.sqspcdn.com/static/f/723732/25889598/1422112465653/ch59-02%20BANTHROPOLOGICAL%20BRESEARCH.pdf%3Ftoken%3DTSMJNvdKWHdUJ7iPvvm7Qkzk4uU%253D+%&cd=1&hl=en&ct=clnk&gl=us | | |
| Burkina Faso | | | | |
| Note: All websites and documents are in French. | | | | |
| <i>General</i> | Ethics Committee for Health Research | | Joint Order 2004-147 / MS / MESSE of 11 May 2004 on the Organization and Functioning of the Ethics Committee for Health Research in Burkina Faso | |
| <i>Drugs, Biologics, and Devices</i> | | | Order No. 2010-292/MS /CAB of 1 October 2010 on the Conditions for Granting Authorizations for Clinical Trials: http://elearning.trree.org/pluginfile.php/34806/mod_folder/content/0/19_Arrete_autorisations_essais_cliniques.pdf?forcedownload=1 | |
| Cameroon | | | | |
| For an overview of human subject protections in Cameroon, see: http://elearning.trree.org/mod/nationalsupplement/view.php?id=227 | | | | |
| <i>General</i> | Cameroon Bioethics Initiative: www.cambin.org | | Ministerial Order No. 079/A/MSP/DS of MINSANTE (1987): http://elearning.trree.org/pluginfile.php/34735/mod_folder/content/0/cm-arrete-079-MSP-CreationComiteEthique-1987.pdf?forcedownload=1 | Operational Guidelines for Ethics Committees in Charge of the Evaluation of Biomedical Research |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| Congo, Democratic Republic of | | | | |
| <i>General</i> | | Decree-Law Framework on Public Health, Title VII: Regarding the National Medical Ethics Committee, Biomedical Research, Transplantation of Organs and Tissues, Genetic Treatment, and Cloning: https://www.mindbank.info/item/2543 | | Proposal for Ministerial Order No. 1250 Establishing the National Advisory Committee on Ethics Health (2004): https://healthresearchweb.org/?action=download&file=DRCPolicy.pdf |
| Côte-d'Ivoire | | | | |
| For an overview of human subject protections in Côte-d'Ivoire, see: http://elearning.trree.org/course/view.php?id=19 Note: All websites and documents are in French. | | | | |
| <i>Drugs, Biologics, and Devices</i> | National Committee on Ethics and Research | | Decree No 317 / SP / DSPH of 14 July 1987 on the Regulation of Drugs Before and After Marketing in Ivory Coast: http://elearning.trree.org/pluginfile.php/34816/mod_folder/content/0/20_Arrete_Regl_exp_clinique_des_substances_med.pdf?forcedownload=1 | |
| Ethiopia | | | | |
| <i>General</i> | Ethiopian Science and Technology Commission, Health Department: http://www.most.gov.et/ | Proclamation 60/1999, Section 21 | | National Health Research Ethics Review Guideline, Fourth Edition (2014): http://www.ccghr.ca/wp-content/uploads/2013/11/national-research-ethics-review-guidline.pdf |
| <i>Drugs and Devices</i> | Food, Medicine, and Health Administration and Control Authority: www.fmhaca.gov.et | | Drug Administration and Control Proclamation No. 176/1999, Article 21 | |
| <i>Human Biological Materials</i> | Ethiopian Science and Technology Commission, Health Department: http://www.most.gov.et/ | | | National Health Research Ethics Review Guideline, Fourth Edition, Chapter 9 (2005): http://www.ccghr.ca/wp-content/uploads/2013/11/national-research-ethics-review-guidline.pdf |
| Gambia | | | | |
| <i>Genetic Research</i> | MRC: Gambia Unit: http://www.mrc.gm/ | | | Guidelines of the National DNA Bank (2001) |
| Ghana | | | | |
| For an overview of the clinical trial information in Ghana, see: http://www.fdaghana.gov.gh/index.php?option=com_content&view=article&id=71&Itemid=55 | | | | |
| <i>Drugs, Biologics, and Devices</i> | Food and Drugs Authority: http://www.fdaghana.gov.gh | Public Health Act, 2012 | Act 851, Sections 150-166: http://www.fdaghana.gov.gh/images/stories/pdfs/Clinical%20Trials/REGULATION%20OF%20CLINICAL%20 | 1. Guidelines for Good Clinical Practice in Ghana (2015): http://www.fdaghana.gov.gh/images/stories/pdfs/downloads/drugs%20guidelines/Clinical%20 |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | TRIALS%20IN%20GHANA.pdf | Trials/GUIDELINES%20ON%20GOOD%20CLINICAL%20PRACTICE%20IN%20GHANA.pdf 2. Guidelines for Authorization of Clinical Trials of Medicines, Food Supplements, Vaccines, and Medical Devices (2015): http://www.fdaghana.gov.gh/images/stories/pdfs/downloads/drugs%20guidelines/Clinical%20Trials/GUIDELINES%20FOR%20AUTHORIZATION%20OF%20CLINICAL%20TRIALS%20OF%20MEDICINES.%20GHANA.pdf 3. Guidelines for Conduct of Clinical Trials in Paediatric Population (2016): http://www.fdaghana.gov.gh/images/stories/pdfs/downloads/drugs%20guidelines/Clinical%20Trials/GUIDELINES%20FOR%20CONDUCT%20OF%20CLINICAL%20TRIALS%20WITH%20PAEDIATRIC%20POPULATION%20IN%20GHANA.pdf 4. Guidelines for Conduct of Clinical Trials During Emergencies (2016): http://www.fdaghana.gov.gh/images/stories/pdfs/downloads/drugs%20guidelines/GUIDELINES%20FOR%20TRIALS%20IN%20EMERGENCY%20SITUATIONS.pdf |

Guinea

For an overview of the clinical research regulations in Guinea, see the ClinRegs report: https://clinregs.niaid.nih.gov/single_country.php?c_id=90

Note: All websites and documents are in French.

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| <i>General</i> | National Ethics Committee on Health Research (CNERs): http://cners-guinee.org/ | Public Health Code, Articles 237-316 (1997): http://www.vertic.org/media/National%20Legislation/Guinea/GN_Code_Sante_Publique.pdf | Decree No. D/218/PRG/SGG: On the Establishment, Functions and Organization of the National Ethics Committee for Research in Health (CNERs), Chapters I and II (1998): http://cners-guinee.org/wp-content/uploads/2014/02/Decret-.pdf | CNERs: Frequently Asked Questions: http://cners-guinee.org/faq/ |
| <i>Research Injury</i> | National Ethics Committee on Health Research: http://cners-guinee.org/ | Public Health Code, Articles 301-302 (1997): http://www.vertic.org/media/National%20Legislation/Guinea/GN_Code_Sante_Publique.pdf | | |

Kenya

For an overview of the clinical research regulations in Kenya, see the ClinRegs report: http://clinregs.niaid.nih.gov/single_country.php?c_id=111

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| <i>General</i> | 1. National Council for Science and Technology (NCST): http://www.nacosti.go.ke/ | 1. Science and Technology Act (2001) 2. HIV and AIDS Prevention | | MOH: National Guidelines for Ethical Conduct of Research Involving Human Subjects |
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| <i>General</i> | 2. Ministry of Health (MOH): www.health.go.ke/ | and Control Act, Chapter 14 (2006) | | (2008): https://healthresearchweb.org/?action=download&file=Final%20national%20ethical%20guidelines-last%20draft.pdf |
| <i>Drugs, Biologics, and Devices</i> | Pharmacy and Poisons Board: http://www.pharmacyboardkenya.org/ | Pharmacy and Poisons Act, Chapter 244 (2009): http://apps.who.int/medicinedocs/documents/s18245en/s18245en.pdf | MOH: Kenya National Guidelines for Research and Development of HIV/AIDS Vaccines (2005) | Guidelines for Applications to Conduct Clinical Trials in Kenya (2014): http://pharmacyboardkenya.org/downloads/?file=Clinical%20Trial%20Guidelines%202014.pdf |
| <i>Human Biological Materials</i> | Ministry of Health (MOH): www.health.go.ke/ | | Kenya National Guidelines for Research and Development of HIV/AIDS Vaccines, page 44 (2005) | |
| Liberia | | | | |
| For an overview of the clinical research regulations in Liberia, see the ClinRegs report: https://clinregs.niaid.nih.gov/single_country.php?c_id=122 | | | | |
| <i>General</i> | Ministry of Health and Social Welfare: http://www.mohsw.gov.lr/ | | 1. Institutional Review Board (IRB) Policies and Procedures Handbook (2008): http://www.ul-acre.org/wp-content/uploads/2013/03/UL-IRB-Policy-Handbook.pdf 2. Ethics Committee Guidelines: Procedures for Researchers, Section 1 (2011): http://clinregs.niaid.nih.gov/documents/liberia/G-LIBR-NHSREC.pdf | |
| <i>Drugs, Biologics, and Devices</i> | Liberia Medicines and Health Products Regulatory Authority | | | Guideline for Application to Conduct Clinical Trials in Liberia (2014): https://clinregs.niaid.nih.gov/documents/liberia/G-LibClinTrial.pdf |
| Madagascar | | | | |
| <i>Drugs and Devices</i> | | Law No. 2011-002, Article 122 Regarding Clinical Trials: https://www.ilo.org/dyn/natlex/docs/ELECTRONIC/97799/116199/F1071917999/MDG-97799.pdf | | |
| Malawi | | | | |
| For an overview of the clinical research regulations in Malawi, see the ClinRegs report: http://clinregs.niaid.nih.gov/single_country.php?c_id=129 | | | | |
| <i>General</i> | 1. National Commission for Science and Technology (NCST): http://www.ncst.mw/ 2. National Health Sciences Research Committee (NHSRC): http://www.ncst.mw/national-health-science-research-committee-nhsrc/ | 1. Presidential Decree on 30 th March 1974 2. Malawi Government Gazette, June 11, 1976, General Notice No. 398 3. Constitution of Malawi, Article 19(5) (1994) | | NCST: 1. The Framework of Guidelines for Research in the Social Sciences and Humanities in Malawi (2011) 2. Policy Requirements, Procedures and Guidelines for the Conduct and Review of Research (2012) |

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| <i>General</i> | 3. College of Medicine Research and Ethics Committee (COMREC): http://www.medcol.mw/ 4. Ministry of Health: www.malawi.gov.mw | | | 3. National Policy Measures and Requirements for the Improvement of Health Research Co-ordination in Malawi (2012) 4. National Policy Requirements and Guidance for the Provision of Insurance Cover for Research Participants in Clinical Trials in Malawi (2012) NHSRC: 1. Operational Guidelines (2001) 2. Summary Guidelines for Writing Research Proposals (2001) COMREC: General Guidelines on Health Research (2014): http://www.medcol.mw/comrec/wp-content/uploads/2014/07/comrec_guidelines.pdf |
| <i>Drugs, Biologics, and Devices</i> | Pharmacy, Medicines, and Poisons Board of Malawi | 1. Pharmacy, Medicines, and Poisons Act, Act 15 of 1988: http://www.google.com/url?sa=t&rc=t=j&q=&esrc=s&source=web&cd=1&cad=rja&uact=8&ved=0CB0QFjAAahUKEwi3qf2P2vLIAhUEqh4KHfyNBvw&url=http%3A%2F%2Fwww.malawilii.org%2Ffiles%2Fmw%2Flegislation%2Fconsolidated-act%2F35%3A01%2Fpharmacy_medicines_poisons_act_pdf_19885.pdf&usg=AFQjCNFJR-Y4F7y3eoC6DV0H7Jr77s5Msg 2. Section 42(1) of PMPB Act, 2003 Supplement | | |
| <i>Social-Behavioral Research</i> | National Committee on Research in the Social Sciences and Humanities | | | Framework of Guidelines for Research in the Social Sciences and Humanities in Malawi (2011): http://www.ncst.mw/wp-content/uploads/2014/03/NATIONAL-FRAMEWORK-OF-GUIDELINES-IN-SSH.pdf |
| <i>Human Biological Materials</i> | National Commission for Science and Technology: www.ncst.mw | | National Regulatory Requirement and Position on Accessing, Collection, Storage, and Use of Human Biological Specimens for Research (2014): https://www.ncst.mw/wp- | |

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| <i>Human Biological Materials</i> | | | content/uploads/2014/03/National-regulatory-requirement-on-human-samples.pdf | |
| <i>Genetic Research</i> | National Research Council of Malawi (NRCM): www.sdn.org.mw/nrcm/ | | Procedures and Guidelines for Access and Collection of Genetic Resources in Malawi (2002) | |
| Mali | | | | |
| For an overview of human subject protections in Mali, see: https://clinregs.niaid.nih.gov/single_country.php?c_id=132&utm_medium=GovDelivery&utm_source=ClinRegs&utm_campaign=MaliPublication#_top | | | | |
| <i>Drugs, Biologics, and Devices</i> | Directorate of Pharmacy and Medicine | Law No. 09-059 of 28 December 2009 Governing Biomedical Research on Humans: https://clinregs.niaid.nih.gov/documents/LawNo09-059.pdf | | |
| Mozambique | | | | |
| For an overview of human subject protections in Mozambique, see: http://elearning.trree.org/course/view.php?id=14&lang=en | | | | |
| <i>General</i> | | | | Science and Technology Ethics Code (2007): http://elearning.trree.org/pluginfile.php/34764/mod_folder/content/0/02-CodigoDeEtica.pdf?forcedownload=1 |
| Nigeria | | | | |
| For an overview of human subject protections in Nigeria, see: http://elearning.trree.org/mod/page/view.php?id=142 | | | | |
| <i>General</i> | National Health Research Ethics Committee: http://nhrec.net/ | National Health Act 2014 | | 1. Nigerian Code of Health Research Ethics (2007): http://nhrec.net/nhrec/wp-content/uploads/2018/10/NCHRE_Aug_07.zip 2. Policy Statement Regarding Enrollment of Children in Research in Nigeria (2016): http://nhrec.net/nhrec/Final%20NHREC%20Policy%20Statement%20on%20Enrollment%20of%20Children%20in%20Research.pdf Various: http://nhrec.net/download-guides-and-forms/ |
| <i>Drugs, Biologics, and Devices</i> | National Agency for Food, Drug Administration and Control (NAFDAC): http://www.nafdac.gov.ng/ | Decree No. 15 of 1993 | | Good Clinical Practice Guidelines (2016): http://www.nafdac.gov.ng/images/GUIDELINES/DRUG%20GUIDELINES/NAFDAC%20GOOD%20CLINICAL%20PRACTICE%20GUIDELINES%202016%20V%202013.pdf |
| <i>Clinical Trial Registries</i> | National Health Research Ethics Committee: http://nhrec.net/ | | | Frequently Asked Questions: http://nctr.nhrec.net |
| <i>Social-Behavioral Research</i> | National Health Research Ethics Committee | | | Nigerian Code of Health Research Ethics (2007): http://nhrec.net/nhrec/wp- |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Human Biological Materials</i> | National Health Research Ethics Committee: http://nhrec.net/ | | | Policy Statement on Storage of Human Samples in Biobanks and Biorepositories in Nigeria (2013): http://nhrec.net/nhrec/NHREC_Policy_Statement_on_Biobanks_FINAL.pdf |
| Rwanda | | | | |
| <i>General</i> | Ministry of Health, National Ethics Committee: http://www.moh.gov.rw/index.php?id=2 | | | Standard Operating Procedures (2009): http://www.moh.gov.rw/index.php?option=com_docman&task=doc_download&gid=126&Itemid=81 |
| Senegal | | | | |
| <i>General</i> | National Committee on Health Research Ethics | Law Supporting the Code of Ethics for Health Research (2009): http://www.sante.gouv.sn/document/1432205899.pdf | | |
| Sierra Leone | | | | |
| For an overview of the clinical research regulations in Sierra Leone, see the ClinRegs report: https://clinregs.niaid.nih.gov/single_country.php?c_id=193 | | | | |
| <i>General</i> | Sierra Leone Ethics and Scientific Review Committee | | | 1. Application Guidelines http://health.gov.sl/wp-content/uploads/2015/01/Guidelines-and-Checklist-for-Ethical-Clearance-2016.pdf 2. Application Form: https://www.healthresearchweb.org/?action=download&file=SierraLeoneEthicsandScientificReviewCommittee.docx |
| <i>Drugs, Biologics, and Devices</i> | 1. Ministry of Health: http://www.sante.gov.bf/ 2. Pharmacy Board of Sierra Leone: http://pharmacyboard.gov.sl/ | | | 1. Guidelines for Conducting Clinical Trials of Medicines, Food Supplements, Vaccines, and Medical Devices in Sierra Leone, Sections: 3.1.7 and 3.2 (2014): http://pharmacyboard.gov.sl/site/LinkClick.aspx?fileticket=9jeTGC2WIZ8%3d&tabid=316&portalid=1&mid=934 2. Guideline for Good Clinical Practice (GCP) in Sierra Leone, Sections 3.2 and 3.3 (2018): https://clinregs.niaid.nih.gov/sites/default/files/documents/sierra_leone/PBSL-GCP-Guideline-V2.pdf 3. Guideline for Conducting Clinical Trials: http://pharmacyboard.gov.sl/site/LinkClick.aspx?fileticket=YrGQkXzflP8%3d&tabid=316&portalid=1&mid=934&forcedownload=true |

| Country | Key Organizations | Legislation | Regulations | Guidelines |
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| <i>Drugs, Biologics, and Devices</i> | | | | Forms: http://pharmacyboard.gov.sl/site/Downloads/Forms.aspx |
| South Africa | | | | |
| For an overview of human subject protections in South Africa, see: http://elearning.trree.org/course/view.php?id=9&lang=en For an overview of the clinical research regulations, see the ClinRegs report: http://clinregs.niaid.nih.gov/single_country.php?c_id=199 | | | | |
| <i>General</i> | 1. Department of Health (DH): http://www.doh.gov.za 2. National Health Research Ethics Council: http://www.nhrec.org.za/ 3. Medical Research Council of South Africa (MRC): http://www.mrc.ac.za 4. Human Sciences Research Council (HSRC): http://www.hsrc.ac.za/index.phtml | 1. Constitution of South Africa, Section 12 (2) (1996) 2. National Health Act No. 61, Chapter 9 (2003): http://www.saflii.org/za/legis/consol_act/nha2003147.pdf | Regulations Relating to Research with Human Participants No. R719 (2014): http://www.google.co.za/url?url=http://www.lawsofsouthafrica.up.ac.za/index.php/browse/medical-and-health/national-health-act-61-of-2003/regulations-and-notices/61-of-2003-national-health-act-regs-gnr-719-19-sept-2014-to-date-pdf/download&rct=j&frm=1&q=&esrc=s&sa=U&ei=W6UtVOOvLa6S7Aa34YDwAg&ved=0CUBUOFjAA&usg=AFQjCNFpKA9W0jNyeWhk0n0l0Q-WxazBtg | DH: Ethics in Health Research: Principles, Structures, and Processes (2015): http://www.nhrec.org.za/docs/Documents/EthicsHealthResearchFinalAused.pdf MRC: 1. Guidelines on Ethics in Medical Research: General Principles (2002) 2. Guidelines on Ethics in the Use of Biohazards and Radiation (2003) 3. Guidelines on Ethics in HIV Vaccine Trials (2003) |
| <i>Drugs, Biologics, and Devices</i> | 1. Department of Health (DH): http://www.doh.gov.za 2. Medicines Control Council: http://www.mccza.com | Medicines and Related Substances Control Act, 101 of 1965 http://www.hpcs.co.za/Uploads/editor/UserFiles/downloads/legislations/acts/medicines_and_related_sub_act_101_of_1965.pdf | General Regulations Made in Terms of the Medicines and Related Substances Act, 1965 (2003): http://www.hpcs.co.za/Uploads/editor/UserFiles/downloads/legislations/acts/medicines_and_related_sub_act_101_of_1965.pdf | DH: Guidelines for Good Practice in the Conduct of Clinical Trials in Human Participants in South Africa (2006): http://www.nhrec.org.za/docs/trainingrequirements/gcp.pdf |
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