ISoOR-ISOB Accreditation Checklist for Organoid Biobanks

Step 1: Eligibility Check

☐ Biobank legally registered in China (valid business license)
Complies with national biosafety regulations (e.g. Biosafety Law, WS233-2002, GB19489-2008)
☐ Institutional Review Board (IRB) approval in place for human-derived organoids
☐ Data governance protocols ensure donor privacy and sample information security
\square Processes align with ISO 20387 and international biobanking best practices (WHO, ISBER OECD)
☐ Self-assessment completed using the ISoOR-ISOB checklist
☐ Any gaps in documentation, facilities, or staff training have been addressed
Step 2: Application Submission
☐ Filled out the ISoOR-ISOB Pre-Assessment Questionnaire, covering:
- Institutional info (name, registration, structure, contacts)
- Organoid workflows (acquisition, processing, expansion, QC, storage, distribution, data management)
- Facility layout, environmental monitoring, and equipment lists
- Personnel qualifications, roles, and training records
- SOPs for all organoid-related procedures and data handling
- Ethical approvals, consent frameworks (IRB, consent types)

- Data traceability: LIMS, audit trails, backup/security systems

☐ Quality Manual detailing QC, risk management, and internal audit processes
☐ Submitted all documents via ISoOR portal (plus hard copies, if requested)
☐ Application fee paid and scope of activities clearly specified
\square All submissions made within six months of eligibility confirmation
Step 3: Documentation Review
☐ SOPs meet ISO 20387/international standards for biobanking
☐ Staff training records are complete and current
\square IRB, consent forms, and ethics documentation compliant with Chinese and international regulations
☐ Biosafety documentation demonstrates compliance with national standards
\square Data management systems validated for security, traceability, and privacy
\square If a Corrective Action Request (CAR) is issued, responses and revisions submitted within 3 months
Step 4: On-Site Audit
☐ Organoid workflows inspected and validated on site
☐ Sample traceability (barcoding, LIMS) verified
☐ Biosafety protocols, facilities, and PPE observed
\square Staff interviews and qualification verification conducted
\square Equipment calibration and maintenance confirmed (e.g. freezers, incubators)
\square Data systems audited for security and privacy compliance
$\hfill\square$ Non-conformities (major/minor) documented and corrective action plans in place with 30–90 day remediation

Step 5: Accreditation Decision

\square All major issues closed within 30 days for full accreditation (3-year validity)
☐ Minor issues resolved in 3–6 months for conditional accreditation
☐ Corrective action evidence submitted via portal
☐ Await official ISoOR-ISOB certificate
Step 6: Post-Accreditation Monitoring
☐ Annual self-assessment report submitted (SOP updates, internal audits, staff training, key QC metrics)
\square Surveillance audits conducted yearly (onsite or remote), possibly unannounced
\square Quality management team assigned to oversee ongoing compliance
☐ Full re-accreditation every 3 years (repeat Steps 2–5)
Step 7: Global Recognition
☐ Certificate received with unique accreditation number
\square Listed in the ISoOR International Registry of organoid biobanks
\square Accreditation promoted via publications, grant proposals, and communications
$\hfill\square$ Network and collaboration opportunities leveraged (ISoOR events, partnerships, funding programs)
Additional Compliance Aspects
☐ Meets China's Human Genetic Resources Management Regulations (2019)

Adheres to pathogens-related biosafety regulations (e.g. Biosafety Law, pathogen lab regulations)
\square Follows ethical governance and waste disposal norms (e.g. 2003 biomedical waste rules)
$\hfill \Box$ Aligns with global interoperability via ISO 20387, IAF/ILAC MRAs, and APAC recognition
Recommendations
\square Kick off with a self-assessment using this checklist to identify critical gaps.
$\hfill\square$ Map out SOPs, Quality Manual, and IRB/consent documentation well in advance.
\square Train staff systematically and calibrate equipment pre-audit.
$\hfill\square$ Engage with ISoOR community events and stay updated on international best practices.
☐ Use ISoOR's templates and guidance, and reach out to the Accreditation Committee for support.