

# 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
- ☐ 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
- ☐ 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
- ☐ IRB, consent forms, and ethics documentation compliant with Chinese and international regulations
- ☐ Biosafety documentation demonstrates compliance with national standards
- ☐ Data management systems validated for security, traceability, and privacy
- ☐ 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
- ☐ Staff interviews and qualification verification conducted
- ☐ Equipment calibration and maintenance confirmed (e.g. freezers, incubators)
- ☐ Data systems audited for security and privacy compliance
- ☐ Non-conformities (major/minor) documented and corrective action plans in place with 30–90 day remediation

## Step 5: Accreditation Decision

- ☐ 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)
- ☐ Surveillance audits conducted yearly (onsite or remote), possibly unannounced
- ☐ 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
- ☐ Listed in the ISoOR International Registry of organoid biobanks
- ☐ Accreditation promoted via publications, grant proposals, and communications
- ☐ 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)
- ☐ Follows ethical governance and waste disposal norms (e.g. 2003 biomedical waste rules)
- ☐ Aligns with global interoperability via ISO 20387, IAF/ILAC MRAs, and APAC recognition

## Recommendations

- ☐ Kick off with a self-assessment using this checklist to identify critical gaps.
- ☐ Map out SOPs, Quality Manual, and IRB/consent documentation well in advance.
- ☐ Train staff systematically and calibrate equipment pre-audit.
- ☐ 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.