Biotechnology — Biobanking — General requirements for biobanking

Biotechnologie — «Biobanking» — Exigences générales relatives au «biobanking»
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Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

The procedures used to develop this document and those intended for its further maintenance are described in the ISO/IEC Directives, Part 1. In particular, the different approval criteria needed for the different types of ISO documents should be noted. This document was drafted in accordance with the editorial rules of the ISO/IEC Directives, Part 2 (see www.iso.org/directives).

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights. Details of any patent rights identified during the development of the document will be in the Introduction and/or on the ISO list of patent declarations received (see www.iso.org/patents).

Any trade name used in this document is information given for the convenience of users and does not constitute an endorsement.

For an explanation of the voluntary nature of standards, the meaning of ISO specific terms and expressions related to conformity assessment, as well as information about ISO’s adherence to the World Trade Organization (WTO) principles in the Technical Barriers to Trade (TBT) see www.iso.org/iso/foreword.html.

This document was prepared by Technical Committee ISO/TC 276, Biotechnology.

Any feedback or questions on this document should be directed to the user’s national standards body. A complete listing of these bodies can be found at www.iso.org/members.html.
Introduction

This document has been developed with the objective of promoting confidence in biobanking. It contains requirements to enable biobanks to demonstrate competent biobank operation and the ability to provide biological material and associated data of appropriate quality for research and development.

This is intended to be achieved by the planning and implementation of policies, processes and procedures covering the life cycle of biological materials and their associated data. The use of this document facilitates cooperation, fosters exchange, and assists in the harmonization of practices among biobanks, researchers and other parties.

In this document, the following verbal forms are used:

— "shall" indicates a requirement;
— "should" indicates a recommendation;
— "may" indicates a permission;
— "can" indicates a possibility or a capability.

Further details can be found in the ISO/IEC Directives, Part 2.
Biotechnology — Biobanking — General requirements for biobanking

1 Scope

This document specifies general requirements for the competence, impartiality and consistent operation of biobanks including quality control requirements to ensure biological material and data collections of appropriate quality.

This document is applicable to all organizations performing biobanking, including biobanking of biological material from multicellular organisms (e.g. human, animal, fungus and plant) and microorganisms for research and development.

Biobank users, regulatory authorities, organizations and schemes using peer-assessment, accreditation bodies, and others can also use this document in confirming or recognizing the competence of biobanks.

This document does not apply to biological material intended for food/feed production, laboratories undertaking analysis for food/feed production, and/or therapeutic use.

NOTE 1 International, national or regional regulations or requirements can also apply to specific topics covered in this document.

NOTE 2 For entities handling human materials procured and used for diagnostic and treatment purposes ISO 15189 and other clinical standards are intended to apply first and foremost.

2 Normative references

The following documents are referred to in the text in such a way that some or all of their content constitutes requirements of this document. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO 8601, Data elements and interchange formats — Information interchange — Representation of dates and times

3 Terms and definitions

For the purposes of this document, the following terms and definitions apply.

ISO and IEC maintain terminological databases for use in standardization at the following addresses:

— ISO Online browsing platform: available at https://www.iso.org/obp

3.1 accessioning
logging
documenting the addition of a new biological material and/or associated data to a biobank

3.2 acquisition
act of obtaining possession and/or custody of biological material and/or associated data
3.3 
**associated data**
any information affiliated with biological material including but not limited to research, phenotypic, clinical, epidemiologic, and procedural data

3.4 
**authentication**
process by which biological material is characterized to a defined level of specification using appropriate technology/documentation to establish a conclusive basis for accepting the material as genuine

3.5 
**biobank**
legal entity or part of a legal entity that performs biobanking (3.6)

3.6 
**biobanking**
process of acquisition (3.2) and storing, together with some or all of the activities related to collection, preparation, preservation, testing, analysing and distributing defined biological material as well as related information and data

3.7 
**biological material**
any substance derived or part obtained from an organic entity such as a human, animal, plant, microorganism(s) or multicellular organism(s) that is(are) neither animal nor plant (e.g. brown seaweed, fungi)

3.8 
**biosafety**
containment principles, technologies and practices that are implemented to prevent unintentional exposure to pathogens and toxins, or their accidental release


3.9 
**biosecurity**
institutional and personal security measures and procedures designed to prevent the loss, theft, misuse, diversion or intentional/unintentional release of pathogens, genetically modified organisms, toxin-producing organisms, or parts thereof, as well as such toxins that are held, transferred and/or supplied by the biobank

3.10 
**catalogue**
systematically arranged list or record often including descriptive information

3.11 
**cataloguing**
act of creating and maintaining a systematically arranged list or record often including descriptive information

3.12 
**chain of custody**
responsibility for or control of materials and associated data as they move through each step of a process

3.13 
**competence**
ability to apply knowledge, experience, and skills to achieve intended results

[SOURCE: ISO 17100:2015, 2.4.9]
3.14 complaint
expression of dissatisfaction other than appeal by any person or organization to a biobank (3.5), relating to the activities, products or results of that biobank where a response is expected

Note 1 to entry: The wording “activities, products or results” includes biological material and/or associated data.

Note 2 to entry: “Appeal” is defined in ISO/IEC 17000:2004, 6.4.

[Source: ISO/IEC 17000:2004, 6.5 modified — “conformity assessment body or accreditation body” has been replaced by “biobank”, the words “products or results” have been added, Note 1 to entry and Note 2 to entry have been added]

3.15 conformity
fulfillment of a requirement

Note 1 to entry: In English the word “conformance” is synonymous but deprecated. In French the word “compliance” is synonymous but deprecated.

Note 2 to entry: This constitutes one of the common terms and core definitions for ISO management system standards given in Annex SL of the Consolidated ISO Supplement to the ISO/IEC Directives, Part 1. The original definition has been modified by adding Note 1 to entry.

[Source: ISO 9000:2015, 3.6.11]

3.16 critical
having a potential impact on the fitness for the intended purpose of biological material and/or associated data

3.17 dedicated area
space containing the biological material kept by the biobank (3.5) or where the biobank activities take place

3.18 destruction
process of eliminating biological material and/or deleting associated data, beyond any possible reconstruction

3.19 disposal
act of removing a biological material and/or associated data usually for scrapping, destruction or returning to provider/donor

3.20 distribution
process of providing selected biological material and/or associated data to recipient(s)/user(s)

3.21 documented information
information required to be controlled and maintained by an organization and the medium on which it is contained

Note 1 to entry: Documented information can be in any format and media and from any source.

Note 2 to entry: Documented information can refer to:
— the management system, including related processes;
— information created in order for the organization to operate (documentation);
— evidence of results achieved.

Note 3 to entry: This constitutes one of the common terms and core definitions for ISO management system standards given in Annex SL of the Consolidated ISO Supplement to the ISO/IEC Directives, Part 1.


3.22 donor
organic entity such as a human, animal, plant etc. from which the biological material and/or associated data is collected for biobanking (3.6)

Note 1 to entry: A human donor can also be a provider (3.41).

3.23 examination
set of operations having the objective of determining the value or characteristics of a property

[SOURCE: ISO 15189:2012, 3.7, modified — Notes to entry have been deleted]

3.24 fit for purpose
fitness for the intended purpose

in line with prearranged requirements for an intended use

Note 1 to entry: The definition of such requirements can take place within the biobank itself and/or in collaboration with users and should consider analytical and other relevant criteria.

3.25 governance
leadership that sets policy and management of operations and can advise/decide on scientific, administrative, technical, financial and other issues

3.26 impartiality
presence of objectivity

Note 1 to entry: Objectivity means that conflicts of interest do not exist, or are resolved so as not to adversely influence the activities of the biobank.

Note 2 to entry: Other terms that are useful in conveying the element of impartiality include “independence”, “freedom from conflicts of interests”, “freedom from bias”, “lack of prejudice”, “neutrality”, “fairness”, “open-mindedness”, “even-handedness”, “detachment”, “balance”.

[SOURCE: ISO/IEC 17021-1:2015, 3.2, modified — In Note 1 to entry "certification body" has been replaced by "biobank"]

3.27 interlaboratory comparison
organization, performance and evaluation of measurements or tests on the same or similar items by two or more laboratories in accordance with predetermined conditions


3.28 interoperability
capability to communicate, execute programs, or transfer data among various functional units in a manner that requires the user to have little or no knowledge of the unique characteristics of those units

3.29 life cycle
consecutive and interlinked processes applied to biological material and associated data from collection, if applicable, acquisition or reception to distribution, disposal or destruction.

Note 1 to entry: This term refers to the biobanking life cycle only.

3.30 metrological traceability
property of a measurement result whereby the result can be related to a reference through a documented unbroken chain of calibrations, each contributing to the measurement uncertainty.


3.31 microorganism
entity of microscopic size

Note 1 to entry: Microorganisms include viruses, all prokaryotes (archaea and bacteria), several eukaryotic organisms (fungi including yeasts, algae, protists).

3.32 nonconforming
deviating from a particular requirement

3.33 personnel
person(s) employed by or working for the biobank (3.5)

3.34 preservation
act to prevent or retard biological or physical deterioration of biological material.

3.35 procedure
specified way to carry out an activity or a process.

3.36 processing
performing any activity on biological material and associated data during all stages of the life cycle (3.29).

3.37 preparation
activities, taking place in a laboratory after acquisitioning, to make biological material ready for further use in the life cycle (3.29), storage (3.47) or distribution (3.20).

Note 1 to entry: These activities can include, e.g. centrifuging, homogenizing, purifying, fixing, stabilizing, replicating, filtering, sorting, culturing, vacuum drying, freeze drying, freezing and thawing, tissue sectioning, fractionating, dispensing/ aliquoting, cryopreserving.

3.38 processing method
procedure, applied to biological material and/or associated data during processing (3.36), with potential to impact the intrinsic properties of the biological material and/or associated data produced as output.

3.39 process
set of interrelated or interacting activities that use inputs to deliver an intended result.
3.40 proficiency testing
evaluation of participant performance against pre-established criteria by means of interlaboratory comparisons

Note 1 to entry: For the purposes of this document, the term “proficiency testing” is taken in its widest sense and includes, but is not limited to:

a) quantitative scheme — where the objective is to quantify one or more measurands of the proficiency test item;

b) qualitative scheme — where the objective is to identify or describe one or more characteristics of the proficiency test item;

c) sequential scheme — where one or more proficiency test items are distributed sequentially for testing or measurement and returned to the proficiency testing provider at intervals;

d) simultaneous scheme — where proficiency test items are distributed for concurrent testing or measurement within a defined time period;

e) single occasion exercise — where proficiency test items are provided on a single occasion;

f) continuous scheme — where proficiency test items are provided at regular intervals;

h) sampling — where samples are taken for subsequent analysis; and

h) data transformation and interpretation — where sets of data or other information are furnished and the information is processed to provide an interpretation (or other outcome).

Note 2 to entry: Some providers of proficiency testing in the medical area use the term “External Quality Assessment (EQA)” for their proficiency testing schemes, or for their broader programs, or both.

[SOURCE: ISO/IEC 17043:2010, 3.7, modified — In Note 2 to entry, reference to Annex A and the last sentence have been deleted]

3.41 provider

provider depositor

person or entity from whom/which the biological material and/or associated data is received or acquired for biobanking (3.6)

Note 1 to entry: Proficiency testing provider and external provider are not included.

3.42 pseudonymization

pseudonymization

processing of individual data in such a manner that these data can no longer be attributed to a specific data subject without the use of additional information

Note 1 to entry: Additional information is kept separately and is subject to technical and organizational measures to ensure that the individual data are not attributed to an identified or identifiable subject.

3.43 rare biological material

rare biological material

biological material that is made precious by its scarcity

3.44 recipient

recipient

person or entity to whom/which the biological material and/or associated data is distributed

3.45 sample

sample

portion of a whole
3.46 stability
ability of a biological material, when stored under specified conditions, to maintain a specified property value within specified limits for a specified period of time

[SOURCE: ISO Guide 30:2015, 2.1.15, modified — The words “reference material” have been replaced by “biological material” and Note 1 to entry has been deleted]

3.47 storage
maintenance of biological material under specified conditions for future use

3.48 tagging
labelling of a biological material for the purpose of identification, location or to give other information

Note 1 to entry: An electronic device can be used for this purpose.

3.49 traceability
ability to trace the history, application or location of an object

Note 1 to entry: When considering a product or a service, traceability can relate to:
— the origin of materials and parts;
— the processing history;
— the distribution and location of the product or service after delivery.

Note 2 to entry: In the field of metrology, the definition in ISO/IEC Guide 99 is the accepted definition.


3.50 unique identifier
code that is associated with a single entity within a given system

Note 1 to entry: Such identifier establishes an unambiguous relationship between each biological material and its associated data.

3.51 user
customer, investigator, or other who/that receives or utilizes biobank service(s)

3.52 validation
confirmation, through the provision of objective evidence, that the requirements for a specific intended use or application have been fulfilled

Note 1 to entry: The objective evidence needed for a validation is the result of a test or other form of determination such as performing alternative calculations or reviewing documents.

Note 2 to entry: The word “validated” is used to designate the corresponding status.

Note 3 to entry: The use conditions for validation can be real or simulated.

3.53 **verification**
confirmation, through the provision of objective evidence, that specified requirements have been fulfilled

Note 1 to entry: The objective evidence needed for a verification can be the result of an inspection or of other forms of determination such as performing alternative calculations or reviewing documents.

Note 2 to entry: The activities carried out for verification are sometimes called a qualification process.

Note 3 to entry: The word "verified" is used to designate the corresponding status.


3.54 **workflow**
structured set of processes

4 General requirements

4.1 General

4.1.1 The biobank shall have procedures addressing biobanking of each type of biological material and associated data held. This includes processes such as collecting/procuring and/or acquiring and receiving, tagging, accessioning/logging, cataloguing/classifying, examining, preparing, preserving, storing, managing data, destroying, packaging as well as safeguarding, distributing and transporting. The biobank shall have procedures to ensure compliance with relevant biosecurity and biosafety requirements. The procedures shall also address risks and opportunities using a risk assessment.

4.1.2 When possible, the biobank should be aware of the minimal requirements for biological material and/or associated data destined for downstream application(s) to ensure that biological material and associated data are handled in a way to enable reproducible research.

4.1.3 The biobank’s mission should be defined and available.

4.1.4 Information relevant to biobank activities, processes and procedures shall be documented in a comprehensible format.

4.1.5 The documentation shall include relevant information generated from procedures pertaining to the quality management system (see Clause 8) as well as the management of facilities/dedicated areas.

4.1.6 The biobank shall comply with relevant regional, national and international ethical principles for biological material and associated data.

NOTE For more information and for guidance on social responsibility, see ISO 26000.

4.1.7 The biobank should document the identity of personnel performing activities encompassing procedures as referred to in 4.1.1.

4.1.8 The biobank should define the time period for retention of documented information and associated data relating to each biological material, after the complete distribution, disposal or destruction of that biological material.
4.2 Impartiality

4.2.1 Biobanking shall be structured and managed so as to safeguard impartiality.

4.2.2 The biobank management shall be committed to impartiality.

NOTE For more information and for guidance on social responsibility, see ISO 26000.

4.2.3 The biobank shall be responsible for the impartiality of its biobanking and shall not allow internal and/or external pressure(s) to compromise impartiality.

4.2.4 The biobank shall identify risks to its impartiality on an on-going basis.

NOTE A relationship that threatens the impartiality of the biobank can be based on ownership, governance, management, personnel, shared materials and associated data, finances, contracts, marketing (including branding), payment of a sales commission or other inducement for the referral of new users, etc.

4.2.5 If a risk to impartiality is identified, the biobank shall demonstrate how it eliminates or minimizes such risk.

4.3 Confidentiality

4.3.1 The biobank shall protect the confidential information and proprietary rights of providers/donors, recipients and users, particularly during storage and transmission of data.

4.3.2 The biobank shall be responsible, through legally enforceable commitments, for the management of confidential information obtained or created during the performance of biobanking. When sharing data or biological material and associated data, the biobank shall inform the provider/donor, where possible, of how their privacy and confidentiality are protected. The biobank shall only release information regarding biological material and associated data according to relevant agreements and approvals (e.g. contractual agreements, legally binding documents, ethical approvals).

4.3.3 When the biobank is required by law to release confidential information, the provider/donor shall be notified of the information provided, unless prohibited by law.

4.3.4 All personnel having access to confidential data of the biobank shall be bound to confidentiality (see 6.2.1.2).

5 Structural requirements

5.1 The biobank shall be a legal entity, or a defined part of a legal entity, that is legally responsible for all its activities.

NOTE For the purpose of this document a governmental biobank is deemed to be or have equivalence of a legal entity on the basis of its governmental status.

5.2 The biobank shall identify top management that has overall responsibility for the biobank.

5.3 The biobank shall have a governance body/advisory board guiding and advising management on scientific, technical and/or administrative and other matters.

5.4 The biobank shall be responsible for activities performed in its facilities/dedicated areas.

5.5 The biobank shall have a course of action to define and address liabilities arising from its activities.
5.6 The biobank shall carry out its activities in such a way as to meet the requirements of this document, its documented agreements and/or legally binding documents, relevant authorities and organizations providing recognition.

5.7 The biobank shall define and document the range of activities for which it conforms with this document. The biobank shall only claim conformity with this document for its defined range of activities, excluding externally provided biobank activities.

5.8 The biobank shall:

a) define the governance structure, including the organization and management of the biobank, its place in any parent organization, and the relationships between management, technical operations and support services;

b) specify the responsibility, authority and interrelationship of personnel who manage, perform, validate or verify work affecting biobanking output.

5.9 The biobank shall have personnel who, irrespective of other responsibilities, have the authority and resources needed to carry out their duties, including:

a) implementation, maintenance, monitoring and improvement of the quality management system;

b) identification of deviations from the quality management system or from the procedures for performing biobanking;

c) assessment of the impact of deviations, and development and implementation of appropriate actions (see 7.11 on nonconforming outputs and 8.7 on corrective action);

d) reporting to biobank management on the performance of the quality management system and any need for improvement.

5.10 The biobank management shall ensure that:

a) changes to the quality management system are monitored and controlled;

b) communication takes place with interested parties, including personnel, regarding the performance indicators of the quality management system and any need for improvement;

c) the importance of meeting the requirements of recipient(s)/user(s), and other applicable requirements (including those described in this document) is communicated to and understood by relevant biobank personnel.

6 Resource requirements

6.1 General

6.1.1 The biobank shall have available personnel, facilities/dedicated areas, equipment, information system(s) and support services necessary to perform biobanking.

NOTE Information system(s) can be electronic or paper based.

6.1.2 The biobank shall have a documented strategy to enable continued financial viability for its activities. This strategy shall be reviewed periodically.
6.2 Personnel

6.2.1 General

6.2.1.1 All personnel of the biobank, either internal or external, who can impact biobank activities, shall act impartially (see 4.2).

6.2.1.2 All personnel having access to the biobank's confidential data shall be bound to confidentiality in respect to that data (see 4.3.4).

6.2.1.3 The biobank shall have documented procedures for personnel management and maintain documented information to indicate compliance with relevant requirements.

6.2.1.4 The biobank shall communicate to all personnel their duties, responsibilities and authorities as detailed in job descriptions.

6.2.1.5 The biobank or the legal entity of which it is a part shall ensure that health and safety requirements are established, documented, implemented and maintained. The level of safety training required shall be determined using a comprehensive risk assessment of the biological and chemical materials, processes and equipment being handled.

6.2.2 Competence and competence assessment

6.2.2.1 The biobank shall define and document the competence required for personnel involved in biobank activities.

6.2.2.2 The biobank shall ensure that all its personnel are competent on the basis of appropriate education, training, demonstrated skills and/or experience necessary to perform assigned duties and activities.

6.2.2.3 The biobank or the legal entity of which it is a part shall maintain documented information for personnel that provide evidence of all appropriate professional competence and education/training (see 6.2.3).

6.2.2.4 Personnel appointed to perform processes shall be subject to competence assessment according to the biobank's established criteria.

6.2.2.5 Personnel shall receive appropriate and relevant assessment at regular intervals to determine what is required to acquire and retain their necessary competence.

6.2.3 Training

6.2.3.1 Each personnel shall receive appropriate and relevant re-/training (internal and/or external training) with regular updates to acquire and retain the necessary competence. The training shall be documented.

6.2.3.2 Personnel undergoing training shall be supervised until the biobank confirms the personnel as competent to perform assigned tasks.

6.2.3.3 An introduction policy for the integration of new personnel shall be implemented. New personnel shall be provided with appropriate orientation to the biobank.
6.3 Facilities/dedicated areas and environmental conditions

6.3.1 The requirements for facilities/dedicated areas and the environmental conditions necessary for the performance of biobanking shall be documented.

6.3.2 The biobank or the legal entity of which it is a part shall determine, control and maintain the facilities/dedicated areas to provide the conditions required for conformity with defined quality control (QC) criteria. This includes procedures to maintain fitness for intended purpose, biosafety, and biosecurity of biological material and associated data.

6.3.3 Where necessary, there shall be effective separation between areas that host incompatible activities. Measures shall be taken to avoid cross contamination.

6.3.4 The facilities/dedicated areas and their environmental conditions shall be suitable for biobanking and should not adversely affect the fitness for the intended purpose.

NOTE Influences that can adversely affect the fitness for the intended purpose can include, but are not limited to, microbial contamination, cross contamination, dust, electromagnetic disturbances, radiation, humidity, electrical supply, temperature, sound and vibration.

6.3.5 The biobank shall measure, monitor, control and record environmental conditions in the biobanking facilities/dedicated areas, as required or where they influence the quality of the biological material and associated data, and/or the health and safety of personnel.

6.3.6 The biobank should address future expansion of its capacity to allow further addition, subdivision and/or processing of biological material.

6.3.7 The biobank shall have a contingency plan to ensure the maintenance of the required environmental conditions in the biobank facilities/dedicated areas according to the risk.

EXAMPLE It can be a contingency plan for natural and human-made disasters such as electrical failure, extreme weather conditions, earthquakes, and sabotage.

6.4 Externally provided processes, products and services

NOTE In this subclause, the term "product" encompasses items used in biobank processes except the biological material.

6.4.1.1 The biobank shall:

a) determine requirements for externally provided critical processes, products and services;
b) document and communicate these requirements to the external provider;
c) retain relevant information about such communication;
d) ensure that the externally provided processes, products and services conform to biobank requirements. Nonconformities shall be communicated to the external provider.

6.4.1.2 The biobank shall determine and apply criteria for the evaluation, selection, monitoring and re-evaluation of performance of external providers based on their ability to provide processes or products and services in accordance with requirements. The biobank shall retain documented information of these activities and any necessary actions arising from the evaluations.

6.4.1.3 The biobank shall determine which externally provided processes, or parts of these, shall be communicated to the provider/recipient/user.
6.4.1.4 The biobank shall ensure that externally provided processes, products and services do not adversely affect the biobank's ability to consistently preserve and supply authenticated biological material and associated data. The biobank shall determine and assess the risks associated with externally provided processes, products and services. Measures shall be taken to avoid negative effects on the conformity of the preservation and authentication of biological material, when necessary.

6.4.1.5 The biobank shall determine the verification, or other activities, necessary to ensure that the externally provided processes, products and services meet the biobank's requirements.

6.4.1.6 When the biobank decides to use externally provided preservation, storing and/or authentication activities, it shall ensure that:

a) the process and all the interrelated processes are validated according to the provisions of this document;

b) internal audits of these processes are planned by the external provider and performed regularly using a risk-based approach (see also ISO 19011);

c) relevant documented information related to these activities is retained.

6.5 Equipment

6.5.1 The biobank shall be furnished with or have controlled access to all equipment required for performance of biobanking.

NOTE The term “equipment” encompasses the items of equipment and associated software when applicable.

6.5.2 The biobank shall establish, document and implement procedures for controlled implementation, safe handling, transport, storage and planned maintenance of all equipment, including procedures for calibration, where necessary.

6.5.3 The biobank shall have instructions on the use and operation of all relevant equipment.

6.5.4 The biobank shall categorize equipment (including backup equipment) with potential to impact on the quality of the biological material and associated data in order to identify equipment which is critical for biobanking (e.g. using a risk-based approach).

6.5.5 The biobank shall establish and maintain a register listing equipment defined under 6.5.1 and 6.5.2 including information for categorization, performance, maintenance, verification and, if applicable, validation of each item.

6.5.6 The biobank shall verify, upon installation and before use, that equipment is capable of achieving the necessary performance and complies with relevant requirements.

6.5.7 Critical equipment shall be capable of achieving the accuracy required and support compliance with specifications relevant to the processing methods or test methods concerned.

6.5.8 The biobank shall retain documented information for critical equipment, which shall include at least the following:

a) equipment and software identity;

b) the manufacturer's name, type identification, and serial number or other unique identification;

c) checks that equipment complies with specifications;

d) the current location, where appropriate;
e) the manufacturer's instructions, if available, or reference to their location;

f) results, reports, and certificates of calibrations, adjustments, acceptance criteria, and associated date(s) [documented in a standard format preferably according to ISO 8601 (see Note to 7.1.3)];

g) the due date of next calibration [documented in a standard format preferably according to ISO 8601 (see Note to 7.1.3)];

h) the maintenance plan, where appropriate, and maintenance carried out to date;

i) any damage, malfunction, modification, or repair to the equipment.

6.5.9 Critical equipment and its software shall be safeguarded from adjustments which would invalidate the process output.

6.5.10 Where applicable, the biobank shall establish and maintain metrological traceability of its measurement results by means of a documented unbroken chain of calibrations or comparisons, linking them to an appropriate reference.

6.5.11 Equipment shall be taken out of service if:

a) it is subject to overloading or mishandling;

b) it generates potentially compromised process output/results;

c) it has been shown to be defective or outside of specification limits.

It shall be isolated to prevent use or clearly labelled or marked as being out of service until repaired and shown by calibration or test to perform correctly.

6.5.12 The biobank shall examine the effect of any defect or departure from specifications using appropriate measures in accordance with 7.11.

7 Process requirements

7.1 General

7.1.1 The life cycle stages of the biological material and associated data in the biobank shall be identified, and appropriate processes shall be defined and verified. A workflow shall describe these stages followed by detailed procedures (see 4.1.1) for each relevant process (e.g. collection, accession, acquisition, identification, preservation, long-term storage, quality control, transport, disposal). All procedures shall be documented, implemented and specific to the biological material and associated data. All critical activities within each procedure shall be identified and documented (see also 7.8.2.7).

7.1.2 All procedures and processes shall be kept up to date and shall be made readily available to personnel.

7.1.3 The date of critical life cycle stages shall be documented in a standard format for all biological materials. The time of critical life cycle stages (e.g. preparation start time or duration, freezing time) should be documented in a standard format. The date and time documentation should be formatted according to ISO 8601.

NOTE The date can be expressed as YYYY-MM-DD (e.g. 2018-04-25) and time can be expressed as hh:mm:ss (e.g. 04:26:55).
7.2 Collection of biological material and associated data

7.2.1 Documented information requirements

7.2.1.1 When the biobank is responsible for collection of biological material, it shall define and document information related to the collection of the biological material. This shall include the date, place and procedure of collection, and any other information relevant to accomplish the objectives of the biobank (e.g. taxonomic information). This should also include the time of the collection of the biological material. The date and time documentation should be formatted according to ISO 8601 (see Note to 7.1.3).

7.2.1.2 When the biobank acquires biological material (i.e. the biobank is not responsible for collection), it should define required/recommended information and retain appropriate documented information related to the collection procedure.

7.2.2 Pre-acquisition information

7.2.2.1 Whenever possible, the biobank shall document and/or retain information related to stages prior to the reception of the biological material that can affect the properties of the biological material to allow the assessment of its fitness for the intended purpose. Further details and requirements are included in Annex A. Annex B gives additional information.

7.2.3 Collection procedure

7.2.3.1 The collection procedure shall be defined either by the biobank and/or the recipient/user, according to the intended use of the biological material, proven techniques, or relevant standards.

7.2.3.2 Where relevant and available, pre-analytical workflows according to existing ISO documents should be implemented (e.g. ISO 20166-1, ISO 20166-2 and ISO 20166-3, ISO 20184-1 and ISO 20184-2, ISO 20186-1, ISO 20186-2 and ISO 20186-3, ISO/TS 20658).

7.2.3.3 Qualified and authorized personnel, and/or recipient(s)/user(s) as applicable, shall collect the biological material according to defined procedures.

When biological material requiring clinical assessment and/or diagnosis is also deemed suitable for biobanking, the preparation, dissection (where appropriate), evaluation of the gross pathology and collection, shall be performed by competent personnel (e.g. specifically trained, experienced, board certified or qualified personnel). The collection of biological material and/or data for research shall never adversely affect patient care and diagnosis, or donor wellbeing.

7.2.3.4 The collection of human biological material shall be performed in accordance with relevant ethical requirements (e.g. relevant ethical approvals or waiver of consent of the patient/donor, etc.).

7.3 Reception and distribution of biological material and associated data

7.3.1 Access principles

7.3.1.1 The principles governing access to and distribution of biological material and associated data shall be defined, documented and, where relevant, published. The biobank shall ensure that documented requirements established with interested parties comply with these principles.

7.3.2 Reception

7.3.2.1 The biobank shall establish, document and implement procedures for receiving or acquiring biological material and associated data (e.g. internal transfers or external shipments/transfers).
NOTE Such procedures are sometimes referred to as accession/logging procedures.

7.3.2.2 The biobank shall define the acceptance criteria of biological material and associated data, including biosafety, biosecurity and intellectual property rights. The identification of the biological material and associated data shall be verified upon acquisition/reception according to the defined acceptance criteria.

7.3.2.3 When appropriate and applicable, (e.g. for cell lines and microorganisms), the biobank shall authenticate the biological material according to relevant and available International Standards or guidelines.

7.3.2.4 Biological material and associated data received or acquired by a biobank, whether individual, part of or an entire collection, shall be segregated (see 7.7.5) to prevent final storage until legal, ethical, documentation, and quality compliance of the biological material and associated data has been assessed and managed.

7.3.2.5 The biobank should obtain relevant documented information, in particular, information needed to assess the fitness for the intended purpose associated with the received or acquired biological material.

7.3.2.6 Where the biobank has not been responsible for collection or sampling, this shall be documented.

7.3.3 Distribution

7.3.3.1 The distribution and any exchange of biological material and associated data shall take place in accordance with the biobank’s access principles (see 7.3.1.1), reporting specifications (see 7.12) and in compliance with other relevant requirements [e.g. material transfer agreement (MTA), data transfer agreement (DTA)].

7.3.3.2 When providing biological material and associated data to a recipient/user outside the biobank, the biobank shall ensure that a documented agreement or legally binding document (e.g. contract, written and signed commitment, binding online acceptance of terms and conditions) outlining the conditions governing the provision and use of biological material and/or associated data are used. Any changes to such a document shall be documented.

7.3.3.3 The biobank shall establish, document and implement procedures for the preparation and distribution of biological material and/or associated data fulfilling the conditions of the documented agreement or legally binding document according to 7.3.3.2.

7.3.3.4 When distributing biological material and/or associated data to a recipient/user, predefined information according to 7.12 shall also be provided, unless the biobank has valid reasons for not doing so, such as data protection compliance.

7.4 Transport of biological material and associated data

7.4.1 The biobank shall establish, document and implement procedures for shipping and receiving biological material, including appropriate conditions for the continued maintenance of biological material integrity in accordance with Annex A. Examples are given in Annex B.

7.4.2 The biobank shall maintain critical chain of custody records for all biological material from point of dispatch to point of receipt. Whenever shipping can alter the quality of the biological material (or if deemed necessary), it shall be tracked and monitored for those elements pertinent to biological material integrity.
e.g. timelines/duration(s), temperature, humidity and light as appropriate to the biological material. The chain of custody records shall detail any deviations from specified parameters according to 7.11.

7.4.3 The biobank shall have procedures for safe handling, packaging, transport and reception relevant to the biological material concerned.

7.4.4 Within the biobank or legal entity of which it is a part, biological material should not be left unattended, unless in designated custody zones as indicated by relevant procedures.

7.4.5 Only competent personnel shall prepare biological material for shipment.

7.4.6 Prior to the transfer of biological material, the requirements of 7.3.3.2 shall be fulfilled and arrangements shall be made for biological material distribution and reception with relevant parties.

7.4.7 The biobank shall establish, document and implement procedures for shipping and receiving data. The transfer of data shall be designed to ensure integrity and prevent breach of data privacy. Prior to the transfer of data, arrangements shall be made for data reception and/or distribution with relevant parties.

7.5 Traceability of biological material and associated data

7.5.1 The biobank shall ensure traceability of biological material and associated data from collection (where relevant), acquisition or reception to distribution, disposal or destruction, as follows:

a) Biological material shall be appropriately tagged so that identification is maintained throughout the life cycle under the custody of the biobank. Special attention shall be focused on persistent tagging [e.g. use of externally applied or integrated options including printed labels, barcodes, two dimension (2D) codes, radio frequency identification systems (RFID), micro electro mechanical systems (MEMS)] of biological material through the use of unique identifiers. The biobank shall have a documented tagging procedure that is also compliant with environmental requirements including relevant storage conditions.

b) Each biological material and associated data shall be linked to the documented information with detail of permissions or restrictions associated for its use.

c) An inventory or tracking system shall allow for the annotation and query of relevant information associated with any handling procedure, including collection, packaging, transportation, preparation, preservation, storing, and distribution procedures. This system should allow any deviation in biobanking procedure(s) to be flagged.

d) A link between biological material and associated data shall be established and maintained for unambiguous traceability.

e) It shall be possible to identify the location of any biological material and associated data at all times.

f) It shall be possible to identify biological material and associated data already distributed to a recipient/user or already disposed of.

7.5.2 The information should be accessible by personnel to allow querying the data as needed, e.g. upon receiving complaints or inquiries regarding distributed biological material.

7.5.3 The biobank shall establish, document and implement procedures for the disposal and transfer of biological material and/or data, both as a planned event and as a result of an emergency (see 7.1.1).
### 7.6 Preparation and preservation of biological material

**7.6.1** The method(s) of preparation and/or preservation shall be defined according to an evidence-based documented processing method (e.g. an International Standard) or as specified in agreement with the provider/recipient/user.

**7.6.2** Critical activities of the preparation and/or preservation procedure (see A.4) shall be monitored and the relevant parameters documented. Each preservation step shall be individually documented.

**7.6.3** The date of each preparation and/or preservation step shall be documented in a standard format for all biological material. The time of each related step should be documented in a standard format. The date and time documentation should be formatted according to ISO 8601 (see Note to 7.1.3).

### 7.7 Storage of biological material

**7.7.1** The biobank or the legal entity of which it is a part should establish a disaster protection plan with use of alternative methods of safeguarding to avoid loss of biological material.

**7.7.2** The biobank shall have documented procedures in place for the storage and tracking of biological material including at least:

a) the tagging information containing at least the unique identifier of the biological material;

b) the type of container and environmental conditions for the biological material storage;

c) mechanism(s) for traceability (see 7.5);

d) a short-term back-up plan for maintaining accurate storage conditions/temperatures in the case of emergency challenges in maintaining defined storage conditions.

**7.7.3** During the execution of critical activities performed during storage, relevant processing parameters shall be measured, monitored and documented. The date (see Note to 7.1.3) and, where necessary, the time(s) of critical activities during storage, and personnel (see 4.1.7) accessing the biological material shall be documented for each biological material. The documentation of the date and time should be formatted according to ISO 8601 (see Note to 7.1.3).

**7.7.4** The biobank shall document and verify the storage location of all biological material and associated data. Traceability of each biological material and each storage transaction shall be ensured at all times.

**7.7.5** The storage locations and processes shall be designed to minimize risk of contamination, and to ensure maintenance of inherent biological material integrity.

**7.7.6** The storage conditions shall comply with 6.3.

**7.7.7** The biobank should verify the biological material inventory at planned intervals by a defined procedure.

**7.7.8** When applicable, the biobank shall establish, document and implement procedures supporting the patient/donor right to withdraw consent for storage and use of biological material and associated data.
7.8 Quality control of biological material and associated data

7.8.1 General

7.8.1.1 Critical activities having an impact on the quality of the biological material and associated data shall be identified by the biobank, provider, recipient or user. The biobank shall establish, document and implement quality control (QC) procedures related to such activities.

7.8.1.2 The biobank shall provide biological material and associated data fit for purpose. The biobank shall define a minimum set of QC procedures to be performed on the biological material and associated data or a subset of it. Exceptions can be justified for rare or legacy biological material and associated data and QC procedures which lead to biological material elimination.

NOTE “Legacy biological material and associated data” refers to the biological material and associated data acquired or received by the biobank before the biobank has implemented this document.

7.8.1.3 The QC procedures shall:

a) be defined according to proven techniques and fitness for the intended purpose;

b) be regularly updated;

c) ensure that provider/recipient/user requirements are met where possible.

7.8.2 Quality control of processes

7.8.2.1 The biobank shall establish, document and implement procedures specifying QC activities throughout the biobanking processes, including QC criteria corresponding to predefined specifications, to demonstrate fitness for the intended purpose of the biological material and associated data.

7.8.2.2 The QC activities shall be performed according to planned intervals. The biobank shall retain documented information of QC activities and results.

7.8.2.3 QC data shall be analysed. If predefined criteria are not met, actions shall be taken to control reporting of invalid data and/or distribution of non-compliant biological material and associated data.

7.8.2.4 The biobank shall ensure that identified limitations are clearly documented and communicated to the recipient/user. During the biological material and associated data distribution process, it is the responsibility of the recipient/user to decide on the acceptance of receiving biological material and associated data with documented and communicated limitations.

7.8.2.5 The biobank shall ensure that information of QC results is provided to the recipient/users as specified by documented requirements.

7.8.2.6 QC results shall be periodically analysed for trends and used as input for the continuous improvement process.

7.8.2.7 The biobank shall document all process-related data in accordance with Annex A.

7.8.2.8 As part of the QC system, the biobank should have appropriate QC materials (e.g. internal control material). QC materials employed by the biobank shall be periodically examined to assess important quality characteristics of the biological material, including stability, the performance of the processing methods and the accuracy/precision of the QC procedures.
7.8.2.9 The biobank shall use approaches to provide objective evidence to demonstrate the comparability of biological material quality (the processing or testing output), where such approaches are available and appropriate. Such approaches include external quality assessment (EQA) programs, proficiency testing programs, interlaboratory comparisons or the biobank may develop its own approaches, including the use of:

a) certified reference materials, where available, produced by a reference material producer fulfilling the requirements of ISO 17034;
b) samples previously examined;
c) samples previously shared with other biobanks;
d) control materials that are tested regularly in EQA programs.

7.8.2.10 If the biobank participates in (an) interlaboratory comparison program(s), the biobank shall monitor relevant results of the interlaboratory comparison program(s), and perform and document corrective actions when predetermined performance criteria are not fulfilled.

7.8.3 Quality control of data

7.8.3.1 The biobank shall identify the critical data, and establish, document and implement QC procedures applying at least to these critical data.

7.8.3.2 The biobank shall define the type and frequency of the QC performed. QC shall focus on accuracy, completeness and consistency of data.

7.9 Validation and verification of methods

7.9.1 General

7.9.1.1 The biobank shall use validated and/or verified methods for critical activities according to 7.9.2 and 7.9.3 at all stages of the biological material life cycle.

7.9.2 Validation

7.9.2.1 When the biobank provides/applies methods for critical activities the biobank shall ensure that these methods have been validated, in order to ensure fitness for the intended purpose. When the validation is performed by the biobank, it shall document and retain for a defined period of time the results obtained, the procedure used for the validation, and a statement as to whether the method is fit for purpose.

7.9.2.2 The validation shall be as extensive as is necessary and confirm, through the provision of objective evidence (in the form of performance characteristics), that the specific requirements for the intended use have been fulfilled.

7.9.2.3 When changes are made to a validated method, the impact of such changes shall be documented and, when appropriate, a new validation shall be carried out.

7.9.3 Verification

7.9.3.1 Validated methods used without modification shall be subject to verification by the biobank before being used.
7.9.3.2 The verification by the biobank shall confirm, through obtaining objective evidence (in the form of performance characteristics) that the set criteria for the method have been met.

7.9.3.3 The biobank shall document the procedure used for the verification and the results obtained.

7.10 Management of information and data

7.10.1 The biobank shall define the required information and data related to biological material and shall have a system in place for tracking. The biobank shall make reasonable efforts to support interoperability of such information and data.

7.10.2 The biobank shall address future expansion of its capacity to allow further addition and/or processing of data associated with biological material.

7.10.3 A procedure for implementation, modification and use of computer system software, hardware, and database(s) shall be in place, when used for biobanking. The procedure shall at least include data integrity, security controls and backup system to prevent loss or corruption of data.

7.10.4 The biobank shall have access to the data and information needed to provide a service specified by contractual agreements.

7.10.5 The biobank should provide interested parties with access to a catalogue of available biological material and associated data.

7.10.6 The biobank shall retain access to the appropriate data associated with the biological material, as necessary for research purposes and/or in compliance with applicable requirements and 7.3.3.2.

7.11 Nonconforming output

7.11.1 General

7.11.1.1 The biobank shall establish, document and implement procedures for management of output that does not conform to the predefined requirements of the biobank and/or the agreement with the recipient/user (see also 7.3.3.2) and/or the agreement with the provider.

7.11.1.2 The biobank shall ensure that output that does not conform to predefined requirements is identified and controlled to prevent unintended use or supply.

7.11.1.3 The biobank shall implement appropriate procedures to disclose information about nonconforming output to relevant parties and, where appropriate, to enable the recipient/user to determine the fitness for the intended purpose.

7.11.1.4 The biobank shall take appropriate corrective action (see 8.7) based on the nature of the nonconforming output and its effect on the fitness for the intended purpose or use of the output. This shall also apply to nonconforming output detected after supply of the biological material and associated data.

7.11.1.5 The procedures for nonconforming output shall address:

a) responsibilities and authorities for the management of nonconforming output;

b) evaluation of the significance of nonconforming output, including the effect on further use of the output;
c) decision on the acceptability, segregation, containment, return, suspension of provision or recall of nonconforming output;

d) persistence of nonconforming output, when
   1) remedy of the nonconformity is impossible;
   2) remedy of the nonconformity is considered impractical; or
   3) the output can have an impact on the results produced by third parties;

e) communication of nonconforming output and the authorization for acceptance by the recipient/user.

7.11.1.6 The procedures for nonconforming output shall also apply to biological material and associated data collected or acquired prior to the first adoption of this document.

7.11.2 Control of nonconforming output

7.11.2.1 The biobank shall mitigate the impacts of nonconformity, implement corrective actions in proportion to the risk(s) presented by nonconforming output, and prevent recurrence. Remedial actions appropriate to the effects shall be taken within defined limits and shall be controlled when nonconforming output is corrected (see also 8.7).

7.11.2.2 The requirements of 8.7.3 apply.

7.11.2.3 Decision(s) on recall shall be taken in a timely manner to limit the use of nonconforming output.

7.12 Report requirements

7.12.1 General

7.12.1.1 The biobank shall provide a report at least as specified in 7.12.2. This shall include the required information as agreed upon in the documented agreement or other legally binding document with the recipient/user (see 7.3.3.2).

NOTE Reports are sometimes called certificates.

7.12.1.2 A report according to 7.12.2 may be issued as hard copy or by electronic data transfer or by an electronic data entry in an accessible database.

7.12.1.3 The biobank should include a statement specifying that the report shall not be reproduced except in full.

7.12.2 Content of the report

7.12.2.1 Each report shall include at least the following, unless the biobank has documented valid reasons for not doing so:

a) a title (e.g. “Quality report” or “Material certificate”);

b) the name and address of the biobank, and the location where activities referred to in the report were carried out, if different from the address of the biobank;

c) the date of issue of the report in a standard format according to ISO 8601 (see Note to 7.1.3);
d) unique identification of the report (such as a serial number), with an identification on each page to ensure that the page is recognized as a part of the report, and a clear identification of the end of the report;
e) biological material identification or specific properties;
f) relevant quality information of the biological material and associated data;
g) method(s) used for identification or characterization of the biological material;
h) testing results with, where appropriate, the units of measurement;
i) method(s) used for testing;
j) method(s) used for collection/acquisition, preparation and/or preservation, as applicable;
k) storage conditions;
l) the name(s), function(s) of person(s) authorizing the report.

7.12.2.2 The biobank shall be responsible for all the information provided in the report, except when information is provided by the provider/recipient/user. Where the biobank has not been responsible for collection or sampling, the report shall state that it relates to the biological material as received by the biobank.

7.13 Complaints

7.13.1 The biobank shall establish, document and implement procedures to receive, evaluate and make decisions on complaints.

7.13.2 A description of the handling process for complaints shall be made available upon request. Upon receipt of a complaint, the biobank shall confirm whether the submitted complaint relates to activities for which it is responsible and, if so, shall address it. The biobank shall be responsible for all levels of complaint handling.

7.13.3 The process for handling complaints shall include at least the following elements and methods:

a) description of the process for receiving, accepting, investigating the complaint, and deciding what actions are to be taken in response to it;

b) tracking and recording complaints, including actions undertaken to resolve them;

c) ensuring that any appropriate action is taken.

7.13.4 The biobank receiving the complaint shall be responsible for gathering and verifying all necessary information to accept the complaint. The biobank shall acknowledge receipt of the complaint.

7.13.5 Whenever possible, the biobank shall provide the complainant with a progress report.

7.13.6 Impartial review shall be performed for each complaint. The outcome of this review shall be communicated to relevant parties.

7.13.7 Whenever possible, the biobank shall give formal notice of the end of the complaint handling to the complainant.
8 Quality management system requirements

8.1 Options

8.1.1 General

The biobank shall establish, document, implement and maintain a quality management system that is capable of supporting and demonstrating the consistent achievement of the requirements of this document and assuring the quality of biobanking. In addition to meeting the requirements of Clauses 4 to 7, the biobank shall implement a quality management system in accordance with option A or option B.

8.1.2 Option A

As a minimum the quality management system of the biobank shall address the following:

a) documented information for the quality management system (see 8.2);

b) control of quality management system documents (see 8.3);

c) control of records (see 8.4);

d) actions to address risks and opportunities (see 8.5);

e) improvement (see 8.6);

d) corrective action for nonconforming outputs (see 8.7);

f) internal audits (see 8.8);

g) quality management reviews (see 8.9).

8.1.3 Option B

A biobank that has established and maintains a quality management system, in accordance with the requirements of ISO 9001, and that is capable of supporting and demonstrating the consistent fulfilment of the requirements of Clauses 4 to 7 also fulfils at least the intent of the quality management system requirements specified in 8.2 to 8.9.

NOTE Explanations for option B are given in Annex C.

8.2 Documented information for the quality management system (Option A)

8.2.1 The biobank shall manage the documented information (internal and external) necessary for its planning and operation, in order to comply with applicable requirements, and to ensure its competence to perform biobanking. To do so, the biobank shall:

a) identify the information that shall be documented;

b) ensure that the documented information is appropriately created and updated;

c) ensure that the documented information is appropriately controlled.

8.2.2 The biobank management shall establish, document, and maintain policies and objectives for the fulfilment of the purpose of this document and shall ensure that the policies and objectives are acknowledged and implemented at all levels of the biobank.

8.2.3 The policies and objectives shall address the competence, impartiality and consistent operation of the biobank.
8.2.4 The biobank management shall provide evidence of commitment to the development and implementation of the quality management system and to continually improving its effectiveness.

8.2.5 All documentation, processes, systems, records, etc. related to the fulfilment of the requirements of this document shall be included in, or referenced to the quality management system.

8.2.6 All personnel involved in biobanking activities shall have access to the parts of the quality management system documentation and related information that are applicable to their responsibilities.

8.3 Control of quality management system documents (Option A)

8.3.1 The biobank shall control the documents (internal or external) that relate to the fulfilment of this document.

8.3.2 The biobank shall ensure that:

a) documents are approved for adequacy prior to issue by authorized personnel;

b) documents are periodically reviewed and updated as necessary;

c) the changes and the current revision status of documents are identified;

d) relevant versions of applicable documents are available at points of use and where necessary their distribution is controlled;

e) documents are uniquely identified;

f) the unintended use of obsolete documents is prevented, and suitable identification applied to them, if they are retained for any purpose.

8.4 Control of records (Option A)

8.4.1 The biobank shall establish and maintain legible records to demonstrate fulfilment of the requirements in this document.

8.4.2 The biobank shall implement the controls needed for the identification, storage, protection, back-up, archive, retrieval, retention time, and disposal of its records. The biobank shall retain records for a period consistent with its contractual and legal obligations.

8.4.3 Access to these records shall be consistent with the confidentiality arrangements and records shall be readily available.

8.5 Actions to address risks and opportunities (Option A)

8.5.1 The biobank shall consider the risks and opportunities associated with its biobank activities in order to:

a) give assurance that the quality management system can achieve its intended results;

b) enhance opportunities to achieve the purpose and objectives of the biobank;

c) prevent or reduce undesired impacts and potential failures in biobanking, including discontinuation of operations of the biobank;

d) achieve continuous improvement.
8.5.2 The biobank shall develop, implement and document:

a) action plan(s) to address these risks and opportunities;

b) action plan(s) to safeguard biological material and associated data in the event of a disaster;

c) action plan(s) to address discontinuation of operations in particular handling of concerned biological material and associated data;

NOTE This can be a legacy plan.

d) approach(es) to:

1) integrate and implement these actions into its quality management system;

2) evaluate the effectiveness of these actions;

3) handle the end of business in case of the biobank’s closure under any circumstances.

8.5.3 Actions taken to address risks and opportunities shall be proportionate to the potential impact on and the validity of biobanking.

NOTE 1 Options to address risks can include identifying and avoiding threats, taking risk in order to pursue an opportunity, eliminating the risk source, changing the likelihood or consequences, sharing the risk, or retaining risk by informed decision.

NOTE 2 Opportunities can lead to expanding the scope of the biobanking, addressing new recipients/users, using new technology and other possibilities to address recipient/user needs.

8.6 Improvement (Option A)

8.6.1 The biobank shall identify and select opportunities for improvement and implement any necessary actions.

NOTE Opportunities for improvement can be identified through the review of the operational procedures, the use of the policies, overall objectives, audit results, corrective actions, management review, risk assessment, analysis of data, and proficiency-testing results.

8.6.2 The biobank shall seek feedback, both positive and negative, from its provider(s)/recipient(s)/user(s). The feedback shall be analysed and used to improve the quality management system, biobanking and provider/recipient/user services.

NOTE Examples of the types of feedback include provider/recipient/user satisfaction surveys and review of reports with provider/recipients/users.

8.7 Corrective action for nonconforming output (Option A)

8.7.1 When nonconforming output according to 7.11 occurs, the biobank shall:

a) react to the nonconforming output and, as applicable:

1) take action to control and correct it;

2) deal with the consequences;

b) evaluate the need for action to eliminate the cause(s) of the nonconforming output, so that it does not recur or occur elsewhere, by:

1) reviewing and analysing the nonconforming output;
2) determining the cause(s) of the nonconforming output;
c) determine if similar nonconformities exist, or could potentially occur;
   1) developing, implementing and documenting any corrective action needed;
   2) reviewing the effectiveness of any corrective action taken;
   3) updating risks and opportunities determined during planning, if necessary;
   4) making changes to the quality management system, if necessary.

8.7.2 Corrective actions shall be appropriate to the effects of nonconforming output encountered.

8.7.3 The biobank shall retain documented information as evidence of:
a) the nature of nonconforming output, cause(s) and any subsequent actions taken;
b) the results and effectiveness of any corrective action.

8.8 Internal audits (Option A)

8.8.1 The biobank shall:
a) plan, implement and maintain an audit programme including the frequency, methods, responsibilities, planning requirements and reporting, which shall take into consideration the importance of the activities concerned, changes affecting the biobank, and the results of previous audits;
b) define the audit criteria and scope for each audit;
c) ensure that the results of the audits are reported to relevant management;
d) implement appropriate correction and corrective actions without undue delay;
e) retain records as evidence of the implementation of the audit programme and the audit results.

NOTE ISO 19011 provides guidance for internal audits.

8.8.2 The biobank shall conduct internal audits at planned intervals to provide information on whether the quality management system:
a) conforms to:
   1) the biobank's own requirements for its quality management system;
   2) the requirements of this document;
b) is effectively implemented and maintained.

8.9 Quality management reviews (Option A)

8.9.1 The biobank top management shall review its quality management system at planned intervals, in order to ensure its continuing suitability, adequacy and effectiveness, including the stated policies and objectives related to the fulfilment of this document.

8.9.2 The inputs to management review shall be documented and shall include information related to the following:
a) changes in internal and external issues that are relevant to the biobank;
b) fulfilment of objectives;
c) suitability of policies and procedures;
d) status of actions from previous management reviews;
e) outcome of recent internal audits;
f) corrective actions;
g) assessments by external bodies;
h) changes in the volume and type of work or in the range of the biobank’s activities;
i) provider/recipient/user feedback;
j) complaints;
k) effectiveness of any implemented improvements;
l) adequacy of biological material and associated data;
m) results of risk identification;
n) outcomes of the quality control;
o) other relevant factors, such as monitoring activities and training.

8.9.3 The outputs from the management review shall record decisions and actions related to:

a) the effectiveness of the quality management system and its processes;
b) improvement of the activities related to the fulfilment of the requirements of this document;
c) provision of required biological material and associated data;
d) any need for change.
Annex A
(normative)

Documentation requirements

A.1 General

The biobank shall provide documentation, which is relevant for the biological material and associated data.

This annex provides requirements for the documentation. It is not inclusive of all requirements in this document.

NOTE Annex B provides complementary information to help implement documentation requirements in this annex.

The biobank documentation, relevant for the biological material and associated data shall:

a) facilitate and verify compliance with applicable requirements, including regulations;
b) enable the biobank’s determination of fitness for the intended purpose;
c) identify the critical data having an impact on quality.

The biobank shall identify the relevant data for each biological material and associated data throughout its life cycle under the custody of the biobank.

A.2 Acquisition

In the context of the acquisition of biological material (meaning the collection or sampling of the biological material in its habitat such as e.g. in nature, in a human or animal host organism) and associated data, documentation of the following is required:

a) timestamp, i.e. date and, when appropriate, time in a standard format preferably according to ISO 8601 (see Note to 7.1.3);
b) collection site, and if relevant geographic coordinates;
c) provider/donor;
d) biological entity identification or characterization;
e) collection method;
f) biosafety and biosecurity information, as appropriate;
g) specific properties.

A.3 Transport

In the context of the internal and external transport of biological material, the transport conditions should be documented, when relevant and appropriate, and include:

a) mode of transportation/shipment specifications;
b) temperature during transport;
c) temperature or temperature range at reception;

d) transport start and end time and date for external transport in a standard format preferably according to ISO 8601 (see Note to 7.1.3);

e) specific requirements, if applicable.

A.4 Preparation/preservation

In the context of the preparation and/or preservation of biological material, documentation of the following is required:

a) preparation method and relevant data;

b) preservation method and relevant data.

A.5 Testing

In the context of the testing of biological material, documentation of the following is required:

a) testing methods;

b) requirements for testing methods;

c) validation of accuracy of testing methods.

A.6 Storage

In the context of the storage of biological material, documentation of the following is required:

a) short-term storage conditions;

b) long-term storage conditions;

c) storage traceability;

d) requirements according to 7.7.3.

A.7 Distribution and disposal

In the context of the distribution and disposal of biological material and associated data, documentation of the following is required:

a) compliance with applicable regulatory and ethical requirements;

b) verification of biological material and of associated data;

c) documented agreement(s) and legally binding document(s) (according to 7.3.3.2).
Annex B
(informative)

Implementation guidance for Annex A

B.1 General

Annex B provides complementary information to help implement documentation requirements in Annex A. These data can vary according the type of biological material and associated data.

B.2 Acquisition

<table>
<thead>
<tr>
<th>Requirement from Annex A</th>
<th>Documentation examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>timestamp</td>
<td>collection time and/or date in a standard format preferably according to ISO 8601 (see Note to 7.1.3)</td>
</tr>
<tr>
<td>collection site</td>
<td>geographical data of collection site (e.g. coordinates)</td>
</tr>
<tr>
<td></td>
<td>host/source description (e.g. farm, hospital, animal, human, forest, field)</td>
</tr>
<tr>
<td></td>
<td>environmental data of collection site</td>
</tr>
<tr>
<td>provider</td>
<td>name, address, code</td>
</tr>
<tr>
<td></td>
<td>consent information, authorization, permission</td>
</tr>
<tr>
<td></td>
<td>historical data, provenance</td>
</tr>
<tr>
<td>biological material/organic entity identification or characterization</td>
<td>consent information</td>
</tr>
<tr>
<td></td>
<td>anonymization/pseudonymization</td>
</tr>
<tr>
<td></td>
<td>taxonomy</td>
</tr>
<tr>
<td></td>
<td>phenotypic data</td>
</tr>
<tr>
<td></td>
<td>clinical data, diagnosis, treatments</td>
</tr>
<tr>
<td></td>
<td>biometric data</td>
</tr>
<tr>
<td></td>
<td>omics data</td>
</tr>
<tr>
<td></td>
<td>epidemiological data</td>
</tr>
<tr>
<td></td>
<td>life style data: smoking status, diet etc.</td>
</tr>
<tr>
<td></td>
<td>demographic data</td>
</tr>
<tr>
<td></td>
<td>unique identifier</td>
</tr>
<tr>
<td></td>
<td>sample/isolate history</td>
</tr>
<tr>
<td>collection method</td>
<td>method of sampling</td>
</tr>
<tr>
<td></td>
<td>primary container type</td>
</tr>
<tr>
<td></td>
<td>additives, stabilizers</td>
</tr>
<tr>
<td></td>
<td>final concentration of the sample</td>
</tr>
<tr>
<td></td>
<td>storage conditions prior to shipment</td>
</tr>
<tr>
<td>specific properties</td>
<td>infectiousness</td>
</tr>
<tr>
<td></td>
<td>biosafety information, radioactivity/radiation</td>
</tr>
<tr>
<td></td>
<td>transgenic/chimera/genetically modified etc.</td>
</tr>
</tbody>
</table>
### B.3 Transport

<table>
<thead>
<tr>
<th>Requirement from Annex A</th>
<th>Documentation examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>mode of transportation/shipment specifications</td>
<td>UN number (e.g. UN 3373), packing instructions (e.g. PI650), prohibitions (e.g. radiation)</td>
</tr>
<tr>
<td>temperature during transport</td>
<td>min./max. specifications</td>
</tr>
<tr>
<td>temperature at reception</td>
<td>min./max. specifications</td>
</tr>
<tr>
<td>transport start and end time and date in a standard format according to ISO 8601 (see Note to 7.1.3)</td>
<td>max. specifications</td>
</tr>
<tr>
<td>other requirements</td>
<td>humidity, light, maximum shipping time, climate/season</td>
</tr>
</tbody>
</table>

### B.4 Preparation/preservation

<table>
<thead>
<tr>
<th>Requirement from Annex A</th>
<th>Documentation examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>preparation method</td>
<td>recording of timestamps</td>
</tr>
<tr>
<td>preservation method</td>
<td>monitoring temperature for critical activities</td>
</tr>
<tr>
<td></td>
<td>cross-contamination</td>
</tr>
<tr>
<td></td>
<td>sterility</td>
</tr>
<tr>
<td></td>
<td>in-process control activities</td>
</tr>
<tr>
<td></td>
<td>storage container type</td>
</tr>
<tr>
<td></td>
<td>number of aliquots or size of batch</td>
</tr>
<tr>
<td></td>
<td>preservation technique used</td>
</tr>
<tr>
<td></td>
<td>additives/preservatives</td>
</tr>
</tbody>
</table>

### B.5 Testing

<table>
<thead>
<tr>
<th>Requirement from Annex A</th>
<th>Documentation examples</th>
</tr>
</thead>
<tbody>
<tr>
<td>testing methods with regard to</td>
<td>integrity</td>
</tr>
<tr>
<td></td>
<td>cross-contamination</td>
</tr>
<tr>
<td></td>
<td>sterility</td>
</tr>
<tr>
<td></td>
<td>concentration</td>
</tr>
<tr>
<td></td>
<td>purity</td>
</tr>
<tr>
<td></td>
<td>identity</td>
</tr>
<tr>
<td></td>
<td>viability</td>
</tr>
<tr>
<td></td>
<td>composition of biological material and associated data</td>
</tr>
<tr>
<td></td>
<td>sample homogeneity</td>
</tr>
<tr>
<td></td>
<td>quantity</td>
</tr>
<tr>
<td></td>
<td>capacity to recover the sample</td>
</tr>
<tr>
<td></td>
<td>recovery rate</td>
</tr>
<tr>
<td></td>
<td>verification of the correct genotype</td>
</tr>
<tr>
<td></td>
<td>possible danger of contamination</td>
</tr>
<tr>
<td>requirements for the testing methods</td>
<td>measurement traceability</td>
</tr>
<tr>
<td></td>
<td>monitoring of method</td>
</tr>
<tr>
<td></td>
<td>external quality assessments</td>
</tr>
</tbody>
</table>
## B.6 Storage

<table>
<thead>
<tr>
<th>Requirement from Annex A</th>
<th>Documentation examples</th>
</tr>
</thead>
</table>
| long-/short-term storage conditions | type of storage, freezer or cold room  
temperature  
timestamp  
humidity  
exposure to radiation (e.g. light)  
duration  
access procedure  
safety  
container type  
traceability |

## B.7 Distribution and disposal

<table>
<thead>
<tr>
<th>Requirement from Annex A</th>
<th>Documentation examples</th>
</tr>
</thead>
</table>
| compliance with applicable regulatory and ethical requirements | as applicable  
verification of biological material and of associated data | as applicable  
contractual information | authorized representatives of the contracting parties  
title of the project  
obligation of recognizing or citing the biobank  
requirement for feedback of information and scientific results |
Annex C
(informative)

Quality management system options

C.1 Growth in the use of quality management systems generally has increased the need to ensure that biobanks can operate a quality management system that is seen as conforming to ISO 9001, as well as to this document. As a result, this document provides two options for the requirements related to the implementation of a quality management system.

C.2 Option A (see 8.1.2) lists the minimum requirements for implementation of a quality management system in a biobank. Care has been taken to incorporate all those requirements of ISO 9001 that are relevant to the scope of biobanking that are covered by the quality management system. Biobanks that comply with Clauses 4 to 7 and implement Option A of Clause 8 will therefore also operate generally in accordance with the principles of ISO 9001.

C.3 Option B (see 8.1.3) allows biobanks to establish and maintain a quality management system in accordance with the requirements of ISO 9001, in a manner that supports and demonstrates the consistent fulfilment of Clauses 4 to 7. Biobanks that implement Option B of Clause 8 will therefore also operate in accordance with ISO 9001. Conformity of the quality management system within which the biobank operates to the requirements of ISO 9001 does not, in itself, demonstrate the competence of the biobank to produce technically valid data and outputs. This is accomplished through compliance with Clauses 4 to 7.

C.4 Both options are intended to achieve the same result in the performance of the quality management system and compliance with Clauses 4 to 7.

NOTE Documents, data and records are components of documented information as used in ISO 9001 and other quality management system standards. Control of documents is covered in 8.3. The control of records is covered in 8.4. The management of data related to biobanking is covered in 7.10.
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[16] ISO 19011, Guidelines for auditing management systems

[17] ISO 20166, (all parts), Molecular in vitro diagnostic examinations — Specifications for preexamination processes for formalin-fixed and paraffin-embedded (FFPE) tissue

[18] ISO 20184 (all parts), Molecular in vitro diagnostic examinations — Specifications for preexamination processes for frozen tissue

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[20] ISO/TS 20658, Medical laboratories — Requirements for collection, transport, receipt, and handling of samples


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Endast för Remiss-grupp 331. Får ej spridas.
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