

Data Management and Reproducibility Standards

Abdullah Lab, IMMEI, University Hospital Bonn

2025-12-20

Protocol ID: GEN-DATA-STD-001

Version: v1

Author: Dillon Corvino

Purpose

This document defines the **mandatory standards** for data management and reproducibility in the Abdullah Lab.

These standards ensure that: - Experiments can be understood and reproduced by others - Data remain interpretable over time - Projects can be transferred without loss of context - Reagents and materials are traceable

All lab members are expected to follow these standards.

Guiding principles

- Clarity over convenience
 - Consistency across the lab
 - One experiment → one clear identifier
 - Raw data are never altered
 - Physical labels must match digital records
-

Experiment identifiers

Every experiment must have a **unique experiment ID**.

Approved formats

Experiment IDs must follow **one of the following formats**:

`<initials><YY>_<NN>`

`<initials><NN>`

Where: - `<initials>` = experimenter initials (e.g. DC) - `<YY>` = two-digit year (e.g. 25) - `<NN>` = sequential experiment number

Examples

DC25_01

DC25_02

DC01

DC02

The experiment ID must be used consistently across: - Folder names - File names - Lab notebook entries
- Analysis scripts

File and folder naming

General rules

- Lowercase letters only
- Underscores (_) only
- No spaces or special characters
- Avoid vague terms such as:
 - final
 - new
 - test

Recommended folder structure

```
experiment_id/  
  raw/  
  processed/  
  analysis/  
  figures/  
  notes/
```

File naming

Files should **almost always include the experiment ID** to ensure traceability outside the original folder context.

The experiment ID should appear at the **start of the filename**, followed by a short descriptive suffix.

Examples

```
DC25_01_gating_strategy.pdf
DC25_01_analysis.wsp
DC25_01_cell_counts.xlsx
DC25_01_analysis_notes.md
```

Files that do not include the experiment ID should be the exception and must still be unambiguous in context.

Approved data storage locations

Sciebo (approved primary storage)

Sciebo is the approved storage location for lab data.

Use Sciebo for: - Raw experimental data - Processed data - Analysis outputs - Shared documents

Local machines

- Allowed **temporarily only**
- Raw data must be transferred to Sciebo promptly
- Local machines must not be the sole copy of data

Prohibited storage

- Personal cloud services (e.g. Google Drive, Dropbox)
 - USB sticks as primary or long-term storage
-

Raw vs processed data

Raw data

- Must remain unmodified
- Must retain original file formats
- Must be stored in a dedicated **raw/** directory

Processed data

- Stored separately from raw data
 - Processing steps must be documented
 - Script-based processing is preferred
-

Reagent labelling standards

All reagents must be clearly labelled at the time of preparation or opening.

Required information on reagent labels

Each reagent container must include: - Reagent name **or** initials / identifier - Sterile or non-sterile status
- Date opened or prepared - Initials of the person who prepared/opened it

Unlabelled or ambiguously labelled reagents must not be used.

Viral stocks

Viral stocks must be labelled with sufficient information to ensure traceability.

Recommended format

virus_strain_passage_YYYY-MM-DD

Example

1cmv_we_p3_2025-02-10

Records must include: - Source - Passage number - Storage location

Cell lines

Cell lines must be labelled in a **similar manner to viral stocks**, ensuring traceability.

Required information

Cell line labels must include: - Cell line name - Passage number - Date - Initials

Example

jurkat_p18_2025-03-04_DC

A master record should document: - Cell line source - Genetic modifications (if any) - Thaw date and passage history

Tissue culture flask labelling

All tissue culture flasks must be clearly labelled.

Required information on flasks

- Cell line name
- Passage number
- Date
- Media used
- Initials

Example

jurkat | p18 | 2025-03-04 | dmem+penstrep+neaas | DC

Media and reagent modifications

Any modification to base media must be recorded.

Requirements

- Record all additives (e.g. antibiotics, supplements)
- Use consistent shorthand
- Ensure the formulation is documented digitally

Example

DMEM + Pen/Strep + NEAAs

If media composition differs from standard lab formulations, this must be clearly documented.

Record keeping

At minimum, records for each experiment must include: - Experiment ID - Date - Operator - Biological material - Reagents used - Deviations from protocol

Electronic or paper lab notebooks are acceptable, but records must be complete and legible.

Versioning

- Use sequential versions: v1, v2, v3
 - Major changes require a new version
 - Deprecated data or reagents must not be deleted without discussion
-

Responsibilities

- Each experimenter is responsible for compliance
 - Supervisors may review organisation and records
 - Issues should be corrected promptly when identified
-

Notes

These standards are living guidelines and may evolve.

Any ambiguities or gaps should be raised so this document can be updated.