

# Flowchart for use of the procedures (SOPs)

[See Introduction video to Flowchart >>](#)

The fields to the right of the form are clickable

- General
- Checklists
- Internal research permission and securing funding
- Application to authorities
- Protocol writing
- Informed consent document writing
- Contracts/ agreements
- Creating and maintaining study files
- Risk evaluation
- Training and start up
- Data management
- Handling of investigational medicinal product
- Protocol deviation handling
- Consent process
- Submissions and reports
- Reporting at the end of the trial
- Close-out and archiving

