

Investigator Site File

Study name

Physical exercise during neoadjuvant chemotherapy for breast cancer as a means to increase pathological complete response rates: the randomized Neo-ACT trial

Investigator Site File, ISF, Table of content GCP & EMA: According to chronological order

1.	Study Contact List
2.	Protocol
۷.	- Current protocol (with Signature page)
	- Superseded protocols (signed and dated)
	Patient Information
3.	- Patient screening & enrolment log
	- Subject identification log
	- Informed consent (blank copies all versions and patient information version log)
	- Signed informed consents (indicate if in a separate folder)
	- Other Patient Facing Material: Questionnaires
4.	Safety
4.	- SAE Form
	- SAE Guidelines
5.	Ethics Committee
J.	- CTA form/ Application
	- Approval/favorable opinion (including members list)
	- Amendments and approvals
	- Correspondence
6.	Financial and legal
0.	- Clinical trial agreement (site agreement)
	- Data processing agreement (if applicable)
	- Data Protection registration (if applicable)
7.	Investigational site
	- Site signature and delegation list
	- CV: s (staff on delegation list)
	- Training (e.g., certificates, attendance log)
8.	Clinical supplies
	- Inventory log
	- Shipment Information
9.	Study procedure manuals
	- Study Lab Manual
	- Handling of samples
	- Vitala App Manual
10.	Laboratory and Biobank
	- Lab normal ranges and lab accreditation
	 Log of blood and tissue samples retained at site
	Dialand. Annance of
	- Biobank Agreement



Investigator Site File

investigator one i ne
Monitoring
- Monitoring visit log
- Monitor's secrecy agreement
- Initiation visits report (original)
Source Data & eCRF
- List of Source Data (Källdatalista)
- CRF Completion Guidelines
Note to File
- Sponsor generated
- Site generated
Correspondence
- To-From-Sponsor
- Newsletters
Study reporting
- Publications
- Clinical study report