

# SCALLOP consortium analysis plan for INF panel proteins

*Adapted from SCALLOP/CVD1 analysis plan, last updated 9/11/2018*

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Timeline for completing cohort-specific analyses and uploading the results for this project:

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## 1. Overview

The SCandinavian coLLaboration for Olink plasma Protein genetics (SCALLOP) consortium, <https://www.olink.com/scallop/>, is a collaborative framework for discovery and follow-up of genetic associations with proteins on the Olink Proteomics platform. A meta-analysis has been conducted on Olink CVD1 panel data from participating cohorts; consequent requests were sent and contributions made on the Olink INF panel. This document follows closely the SCALLOP/CVD1 analysis plan for the analysis, and in particular highlights relevant information required to facilitate the meta-analysis.

As with the CVD1 meta-analysis, the tasks will involve

- Identification of pQTLs in SCALLOP discovery cohorts
- Study of pQTLs in replication cohorts
- Investigation of the mechanistic basis of identified cis- and trans-pQTL by functional annotation
- Examination of pQTL pleiotropic effects
- Evaluation over the causal role of INF proteins disease outcomes such as CHD and stroke
- Other downstream analysis

## 2. Data and analysis

### Proteins

The Olink INFlammation panel of 92 proteins, e.g., <https://github.com/jinghuazhao/INF/blob/master/doc/olink.inf.panel.annot.tsv>.

### SNPs

- 1000 genomes imputation, build 37 (hg19) positions.
- SNPs will be filtered for imputation quality at time of meta-analysis
- Quality control on aspects such as SNP/sample call rates, gender mismatch, abnormal inbreeding coefficient, failed cryptic relatedness test, ancestry outlier, heterozygosity and Hardy-Weinberg equilibrium test.

## Association analysis

- Rank-based inverse normal transformation on the raw measurement of proteins including those below lower limit of detection, e.g., via `invnormal` function,

```
invnormal <- function(x)
  qnorm((rank(x, na.last="keep")-0.5)/sum(!is.na(x)))
```

- Multiple linear regression for all samples including sex, age, principal components and other cohort specific covariates.
- Additive genetic model
- For case-control data, cases and controls are analysed separately – results will be merged at meta-analysis stage

## Software

It is preferable to use software which account for genotype uncertainty, such as SNPTEST, QUICKTEST, and BOLT-LMM.

## 3. Descriptive statistics

Please fill out the spreadsheet as with SCALLOP/CVD1 with naming convention:

- STUDY.descriptives.DATE.xls
- Where STUDY is a short (14 characters or less) identifier for the population studied, which is the same for all files provided by your study.
- DATE is the date on which the file was prepared, in the format “DDMMYYYY”.

## 4. File formats for GWAS results

### SNP table for GWAS results

Please include the following columns. Missing values are coded as “NA”.

No	Variable name	Description of variable
1	SNPID	CHR:POS_A1_A2 (such that A1-10. The results will be replicated in independent cohorts.

## 6. Uploading of results

See the CVD1 analysis plan.

## 7. Contact information

For general questions about SCALLOP, please contact Anders Malarstig (anders.malarstig@ki.se). For technical issues about TRYGGVE, please contact Lasse Folkersen (lasfol@cbs.dtu.dk).

For questions regarding SCALLOP/INF, please contact Jing Hua Zhao (jhz22@medschl.cam.ac.uk) and James Peters (jp549@medschl.cam.ac.uk).