Towards the GWAS pipeline: (i) revising the steps

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step 1 - getting the data

- 1.get_data.sh
 - Download the data
 - Prepare the data





step 2 - filter the data

- 2.steps_filtering.sh
 - Filter genotype data:
 - MAF
 - Missing rate



step 3 - imputing missing genotypes

- 3.step_imputation.sh
 - Impute missing genotype data:
 - LHCI



the stand-alone GWAS scripts

- Run GWAS programmatically
 - gwas_rrblup.R
 - gwas_statgengwas.R
 - gwas_sommer.R



step 4 - run the GWAS

4.gwas.sh

- Run GWAS through the stand-alone script
- (try multiple scripts: gwas_rrblup.R, gwas_statgengwas.R, gwas_sommer.R)



NEXT LECTURE

Collaborative exercise