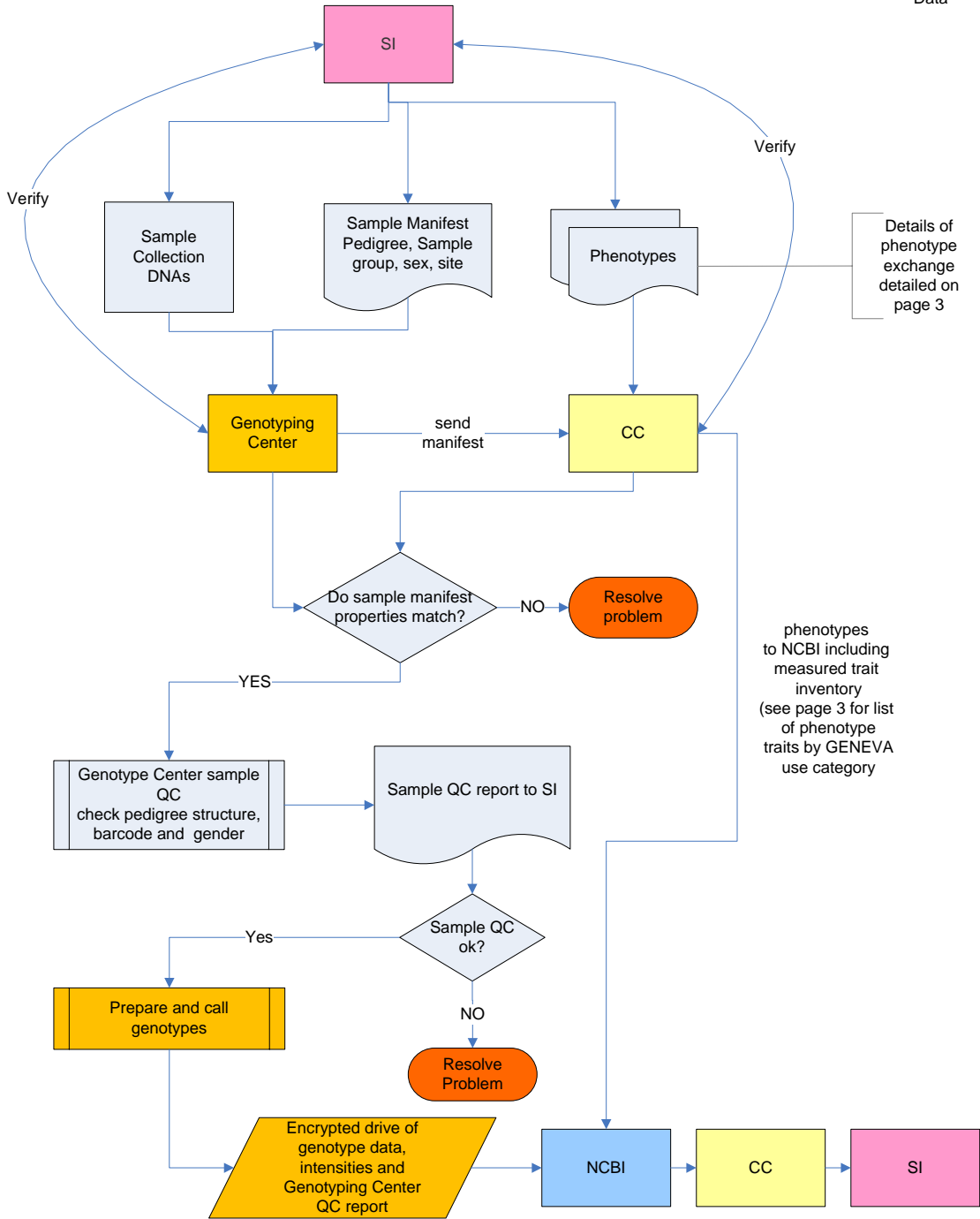


GENEVA Process Workflow

Genotype Data

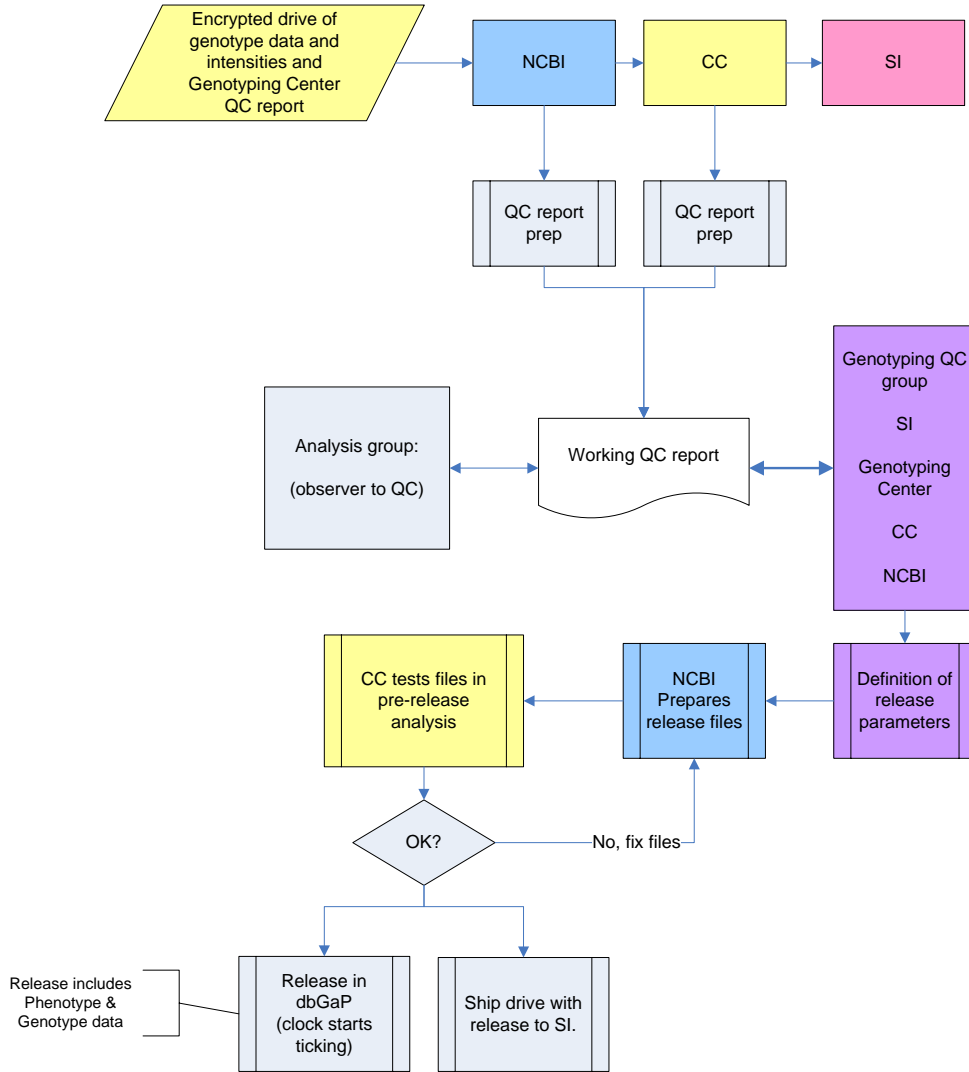


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GENEVA Process Workflow

Genotype Data
(page 2)

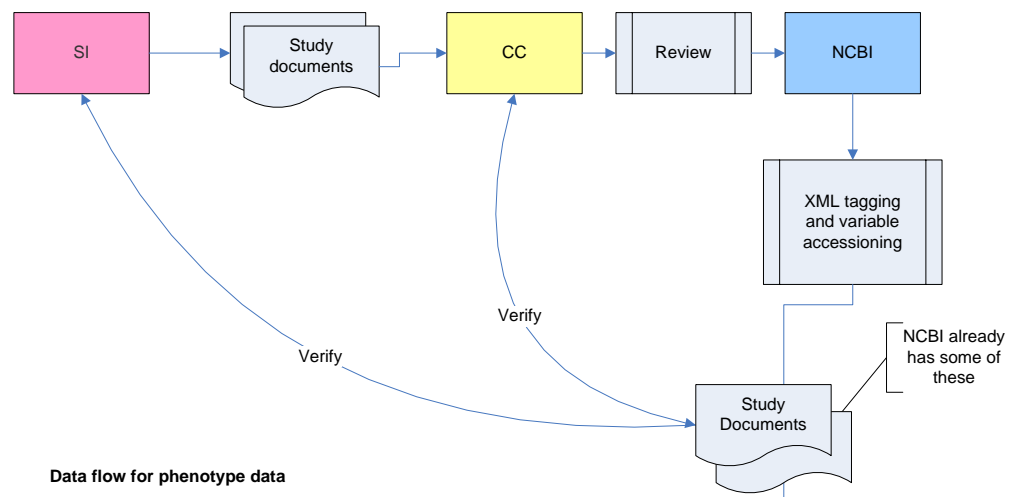


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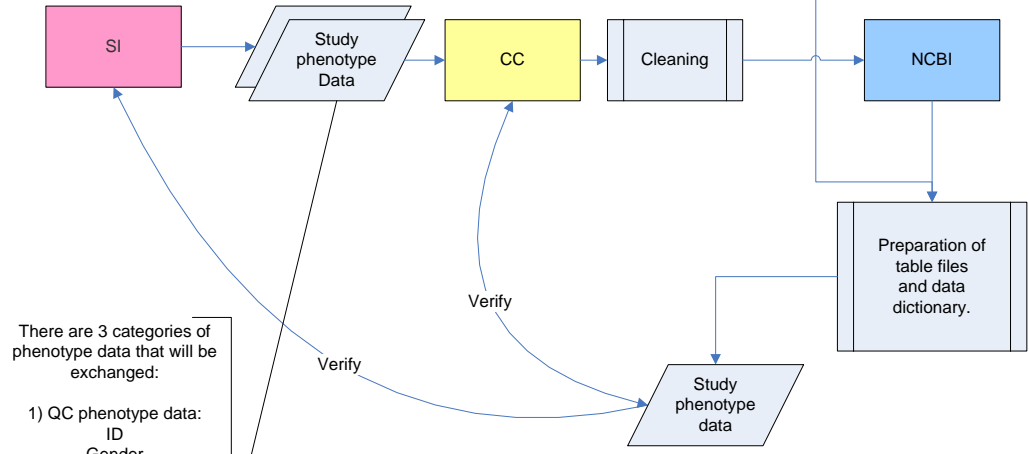
GENEVA Process Workflow

Phenotype Data

Data flow for documentation



Data flow for phenotype data



- There are 3 categories of phenotype data that will be exchanged:
- 1) QC phenotype data:
 - ID
 - Gender
 - Pedigree
 - Outcome/Affection
 - 2) Agreed Variables described in the grant submission
 - 3) Additional phenotypes Common, identified by phenotyping group and Additional data studies agree to submit beyond (2)

Symbol legend

