## **GENOME-WIDE ARRAY DATA VARIANT AND INDIVIDUAL FILTERING**

After "c	enotype calling	" from CEL	files to PLINK	format files (e.	a. Axiom Analy	vsis Suite 2.0	if HumanOrigins	platform)

- 0. INITIAL VARIANT FILTERS (PLINK/1.9b) Perform independently on each dataset
- 0.1. Exclude sites with genotype missingness > 5%.
- 0.2. Exclude individuals with genotype missingness > 10%
- 0.3. Filter SNPs in HW disequilibrium pvalue 10-6
- 1. RELATEDNESS ESTIMATION (PLINK/1.9b) Perform independently on each dataset
- 1.1. Remove sites with minor-allele frequency (MAF) < 1%
- 1.2. Estimate relatedness
- 1.3. Remove related individuals (the one with less missing sites)
- 2. MERGING WITH REFERENCE PANELS (PLINK/1.9b) Perform independently on each dataset
- 2.1. Merging datasets from step 0.3 (without related individuals): keeping only overlapping SNPs
- 2.2. Remove sites with minor-allele frequency (MAF) < 1%

**EXTRA filtering: Linkage disequilibrium pruning** (for allele-frequency-based analyses)

window = 200; step = 25;  $r^2 = 0.5$