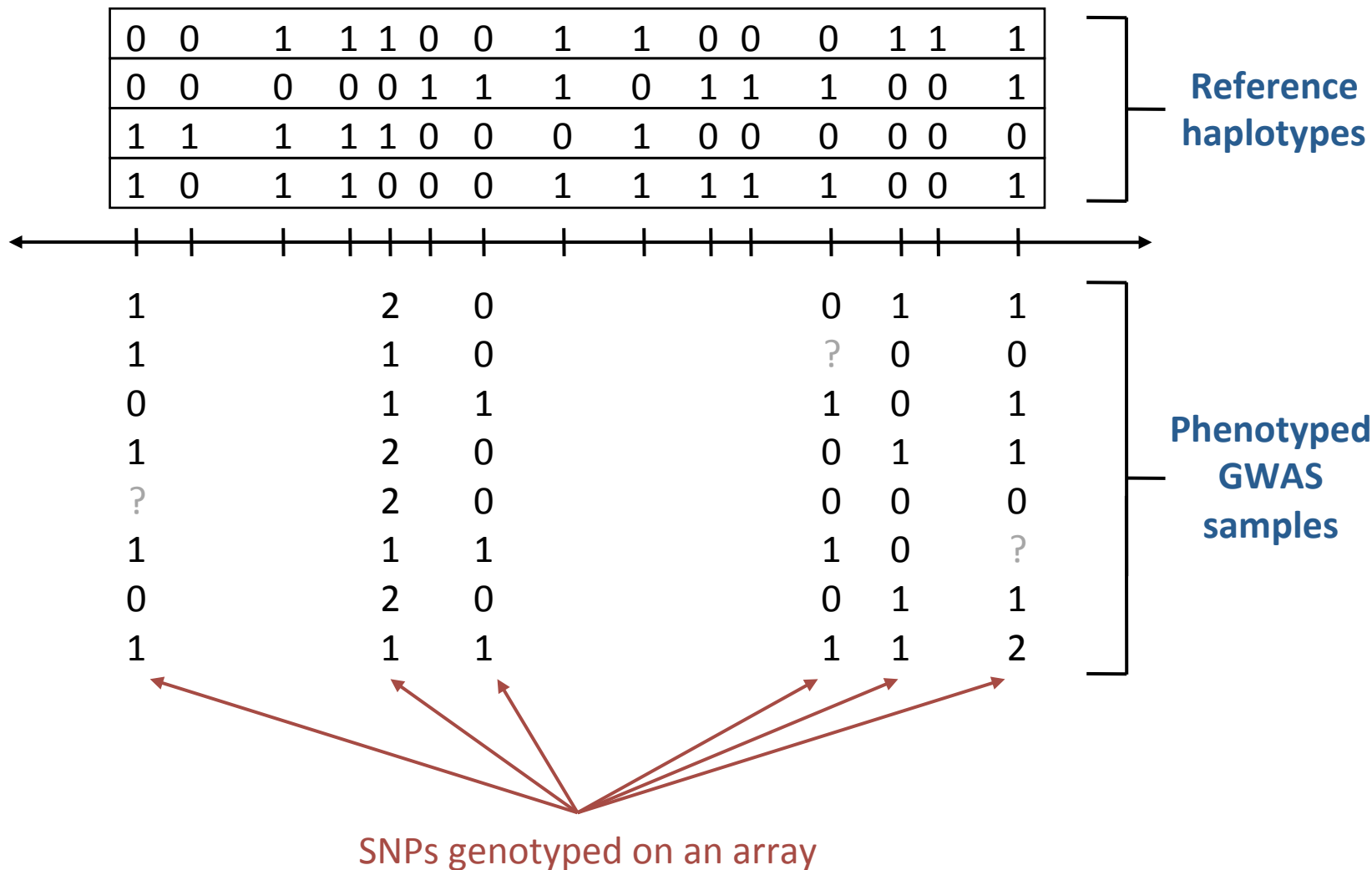


# Using 1,000 Genomes data for imputation in genome-wide association studies

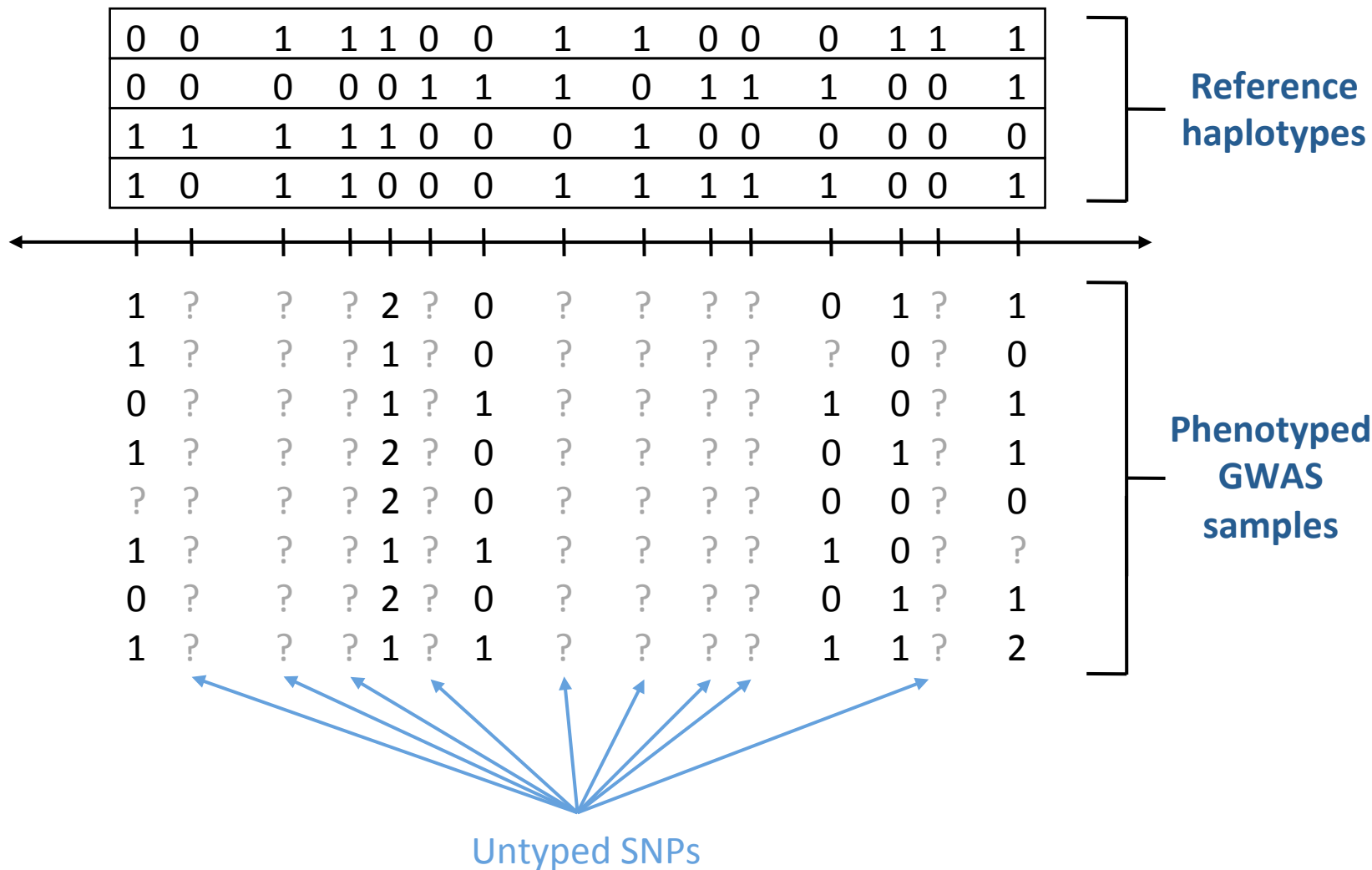
1,000 Genomes Data Tutorial  
ICHG 2011, Montreal

Bryan Howie  
University of Chicago

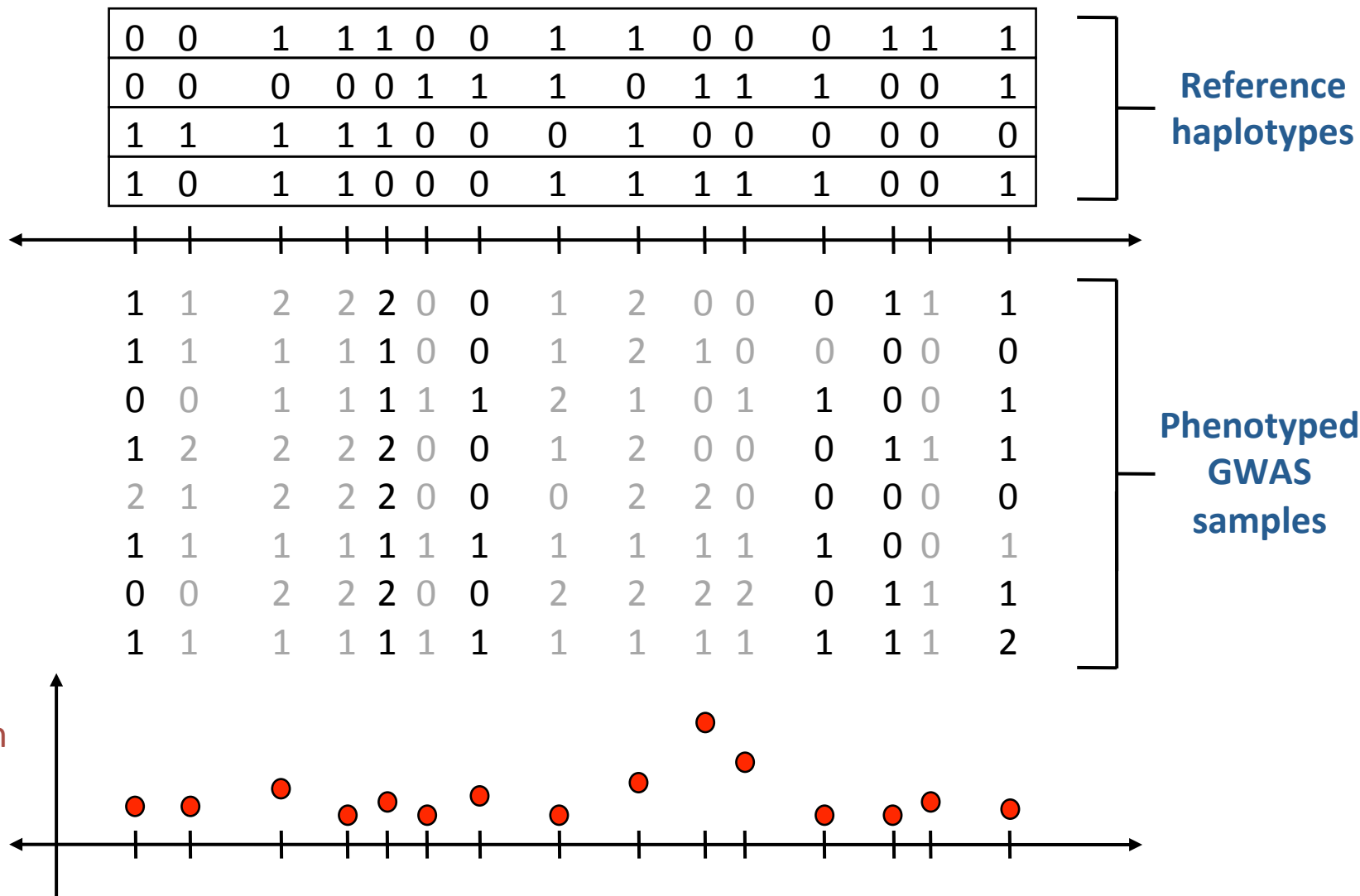
# Genotype imputation background



# Genotype imputation background



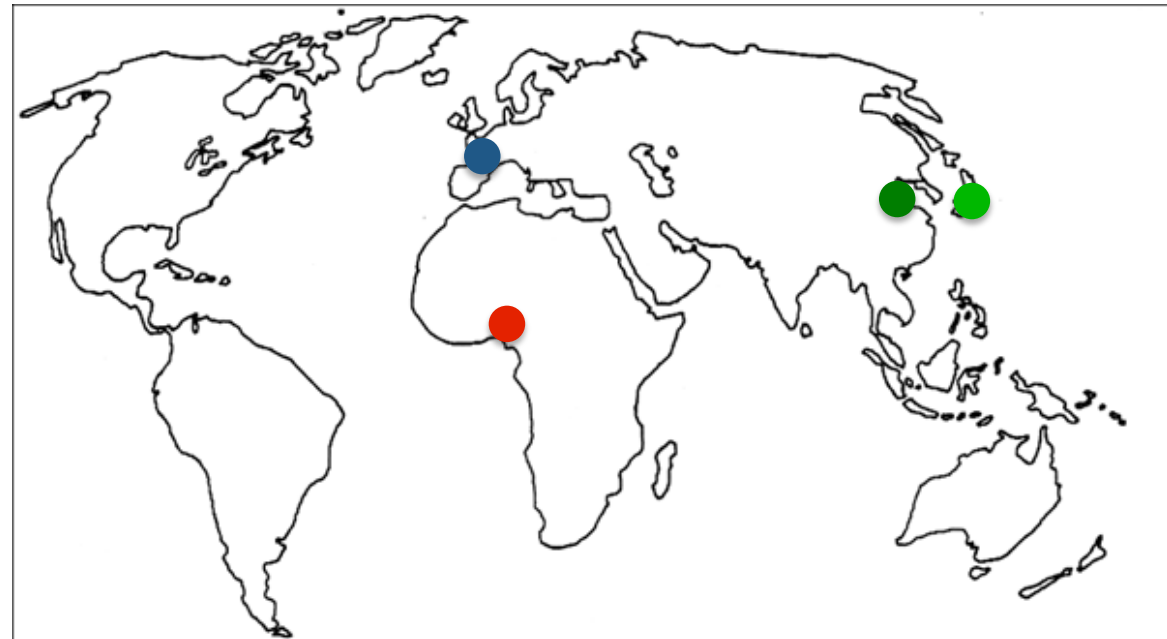
# Genotype imputation background



# A brief history of imputation reference panels:

HapMap 2, HapMap 3, and  
the 1,000 Genomes Project

# HapMap 2 (2007)



CEU  
CHB  
JPT  
YRI

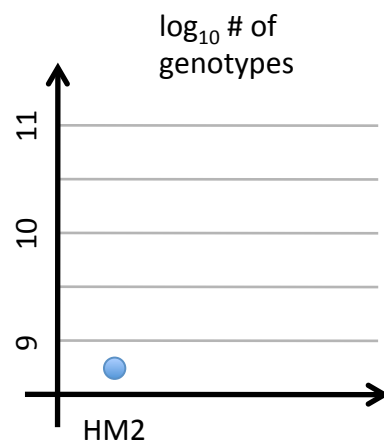
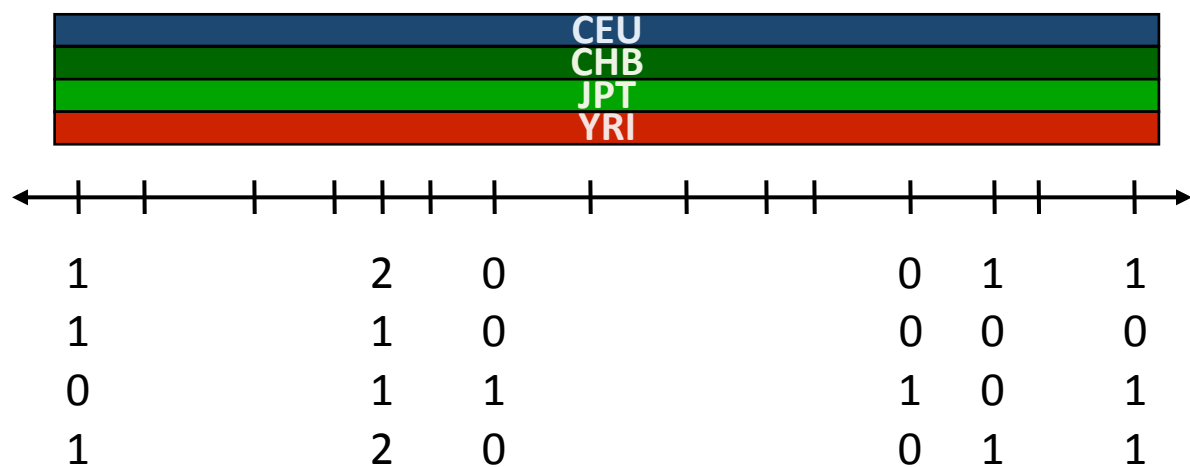
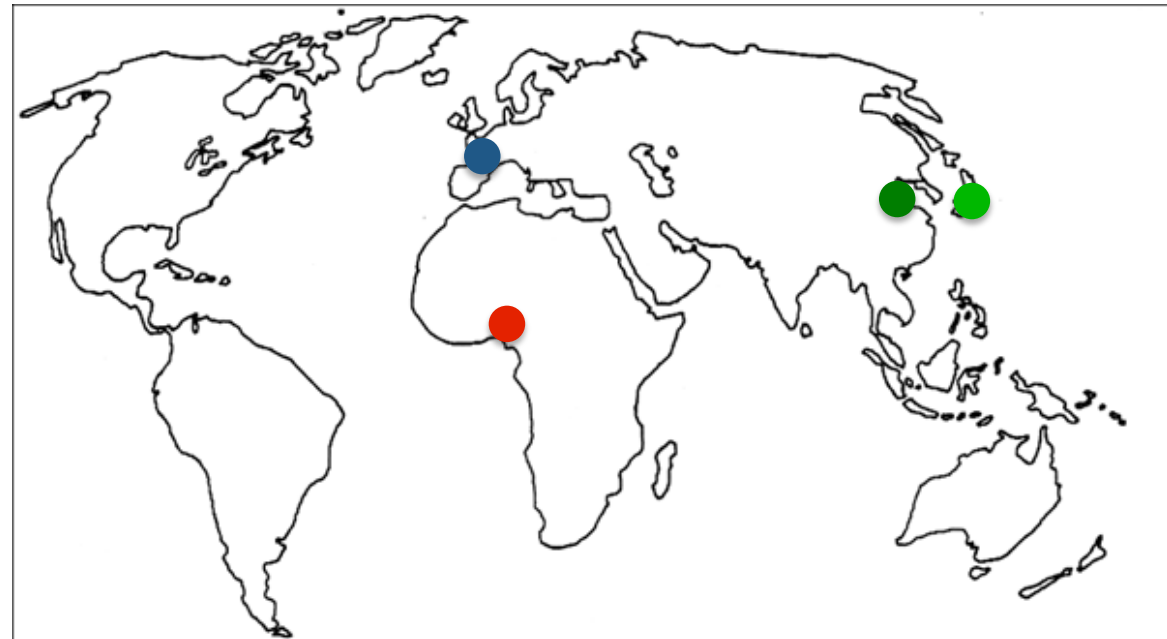
Reference  
panels



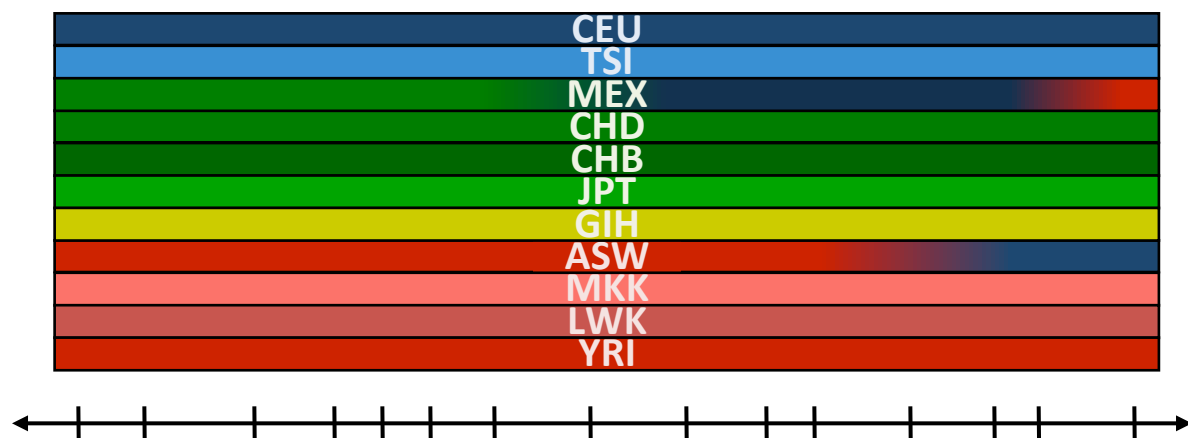
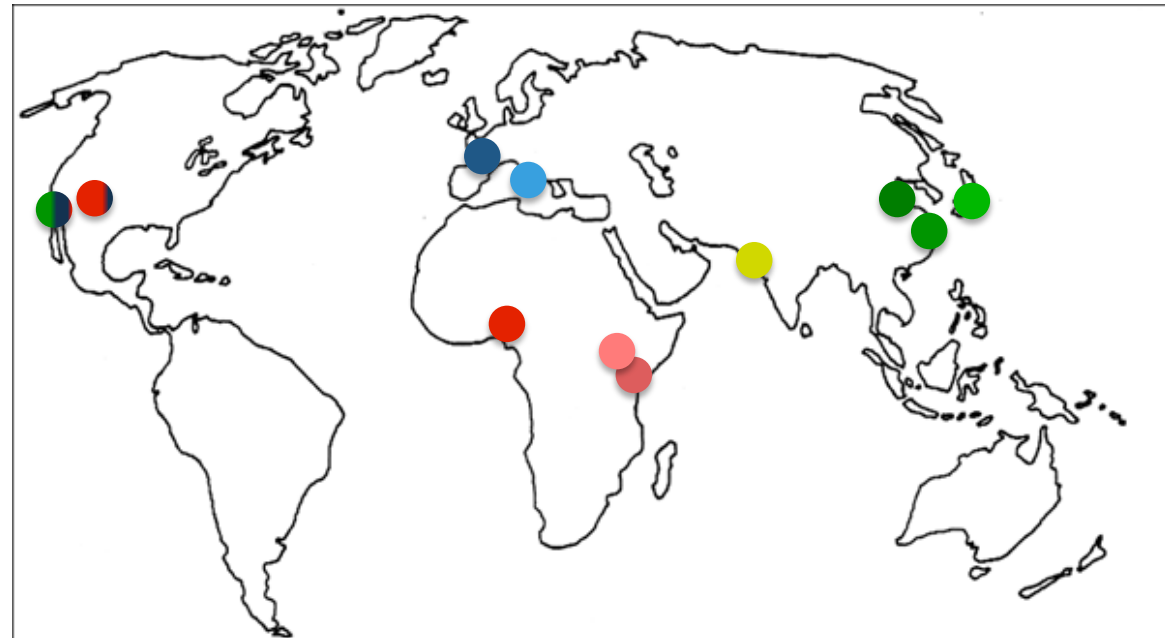
1	2	0	0	1	1
1	1	0	0	0	0
0	1	1	1	0	1
1	2	0	0	1	1

GWAS  
genotypes

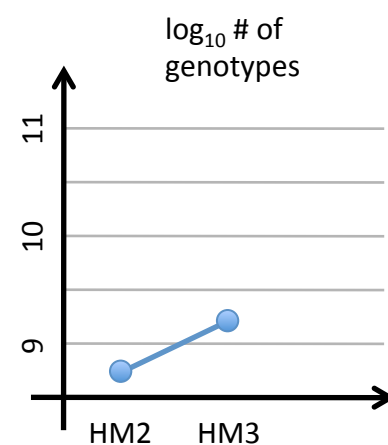
# HapMap 2 (2007)



# HapMap 3 (2009)



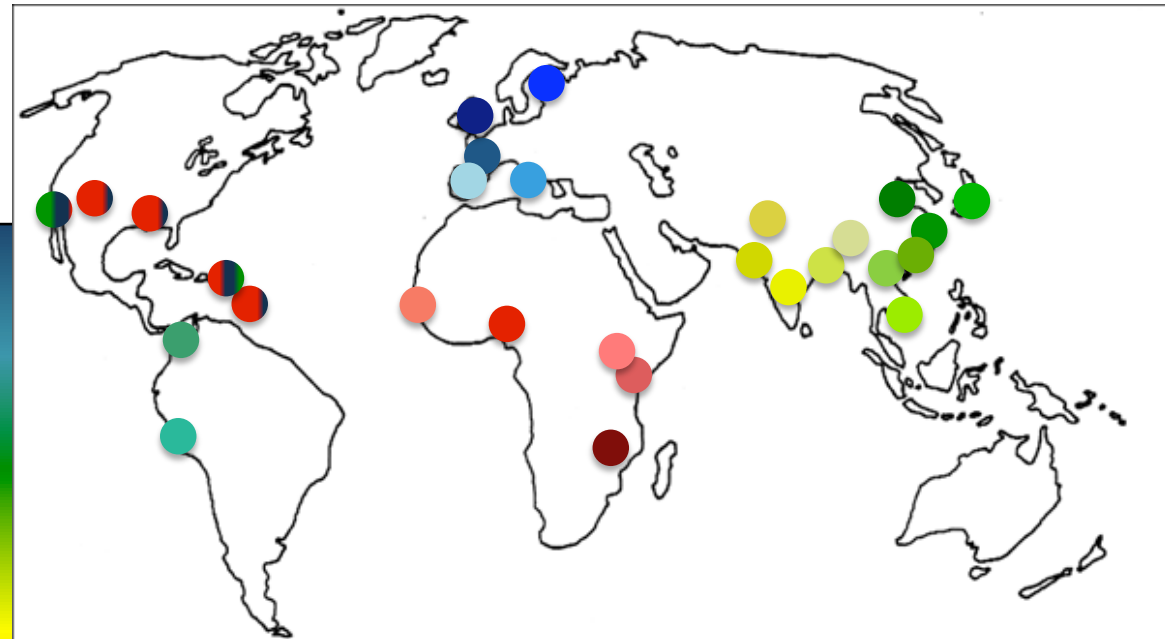
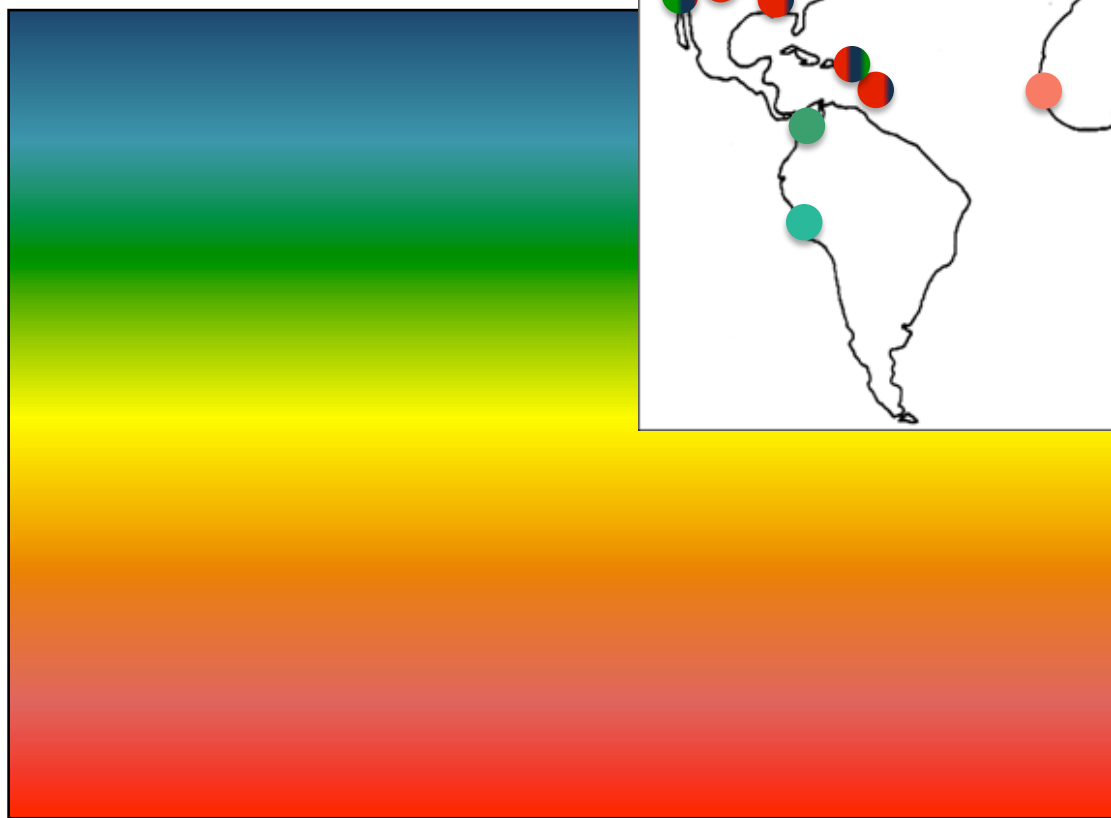
Reference  
panels



1	2	0	0	1	1
1	1	0	0	0	0
0	1	1	1	0	1
1	2	0	0	1	1

GWAS  
genotypes

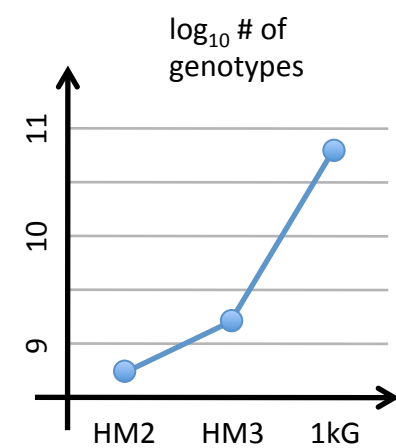
# 1,000 Genomes (2010+)



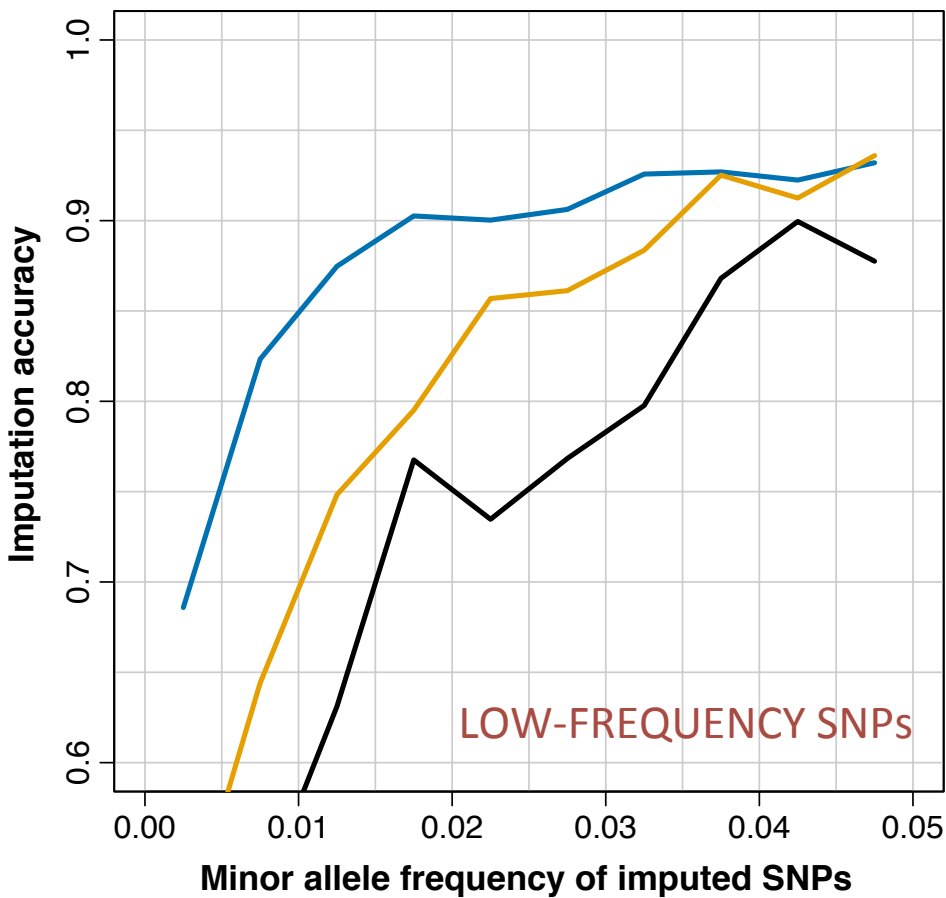
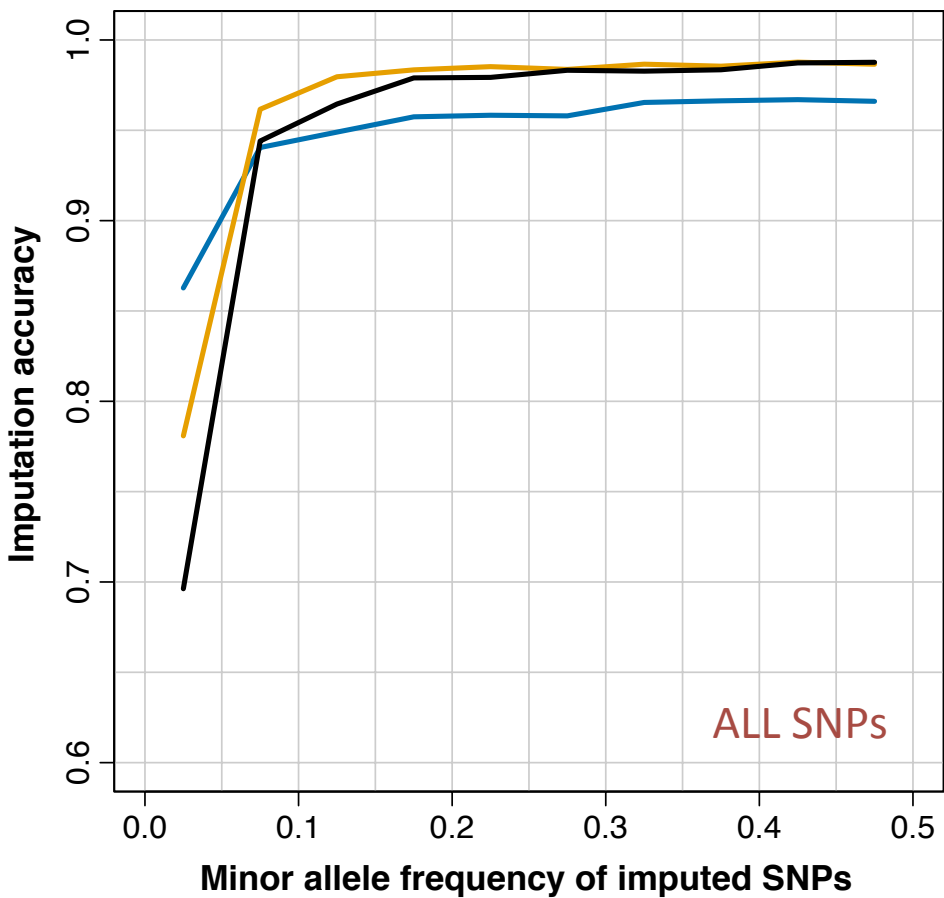
Reference  
panels

GWAS  
genotypes

1	2	0	0	1	1
1	1	0	0	0	0
0	1	1	1	0	1
1	2	0	0	1	1

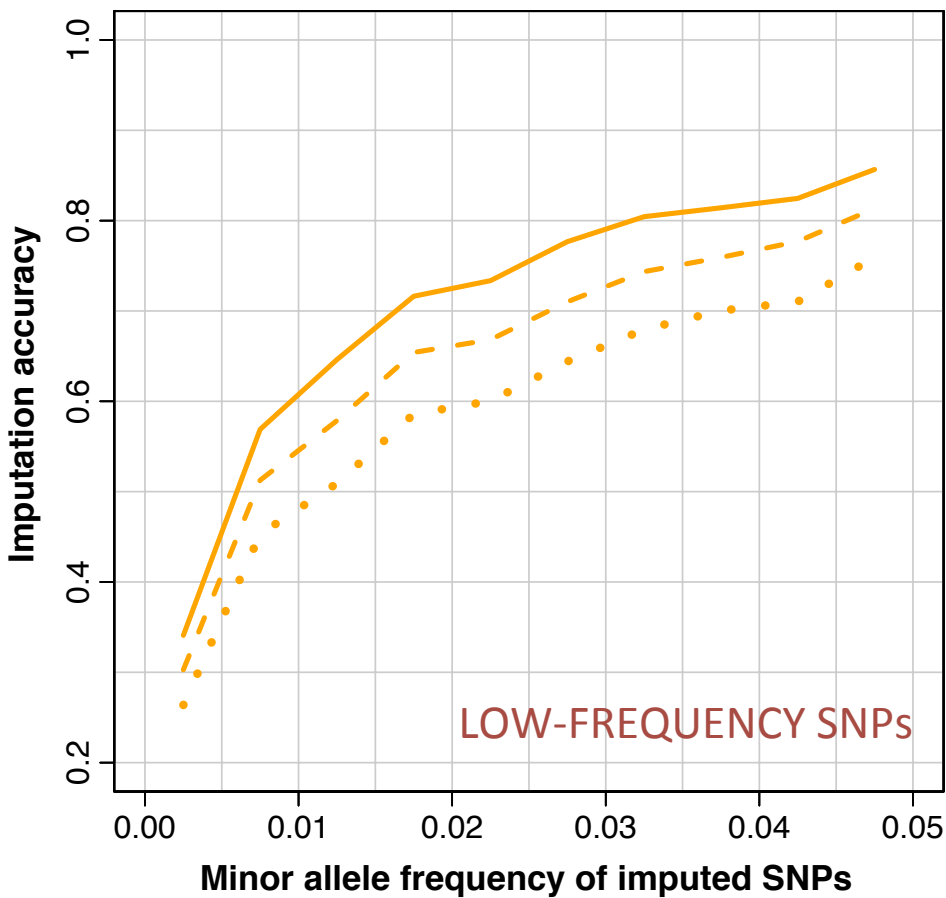
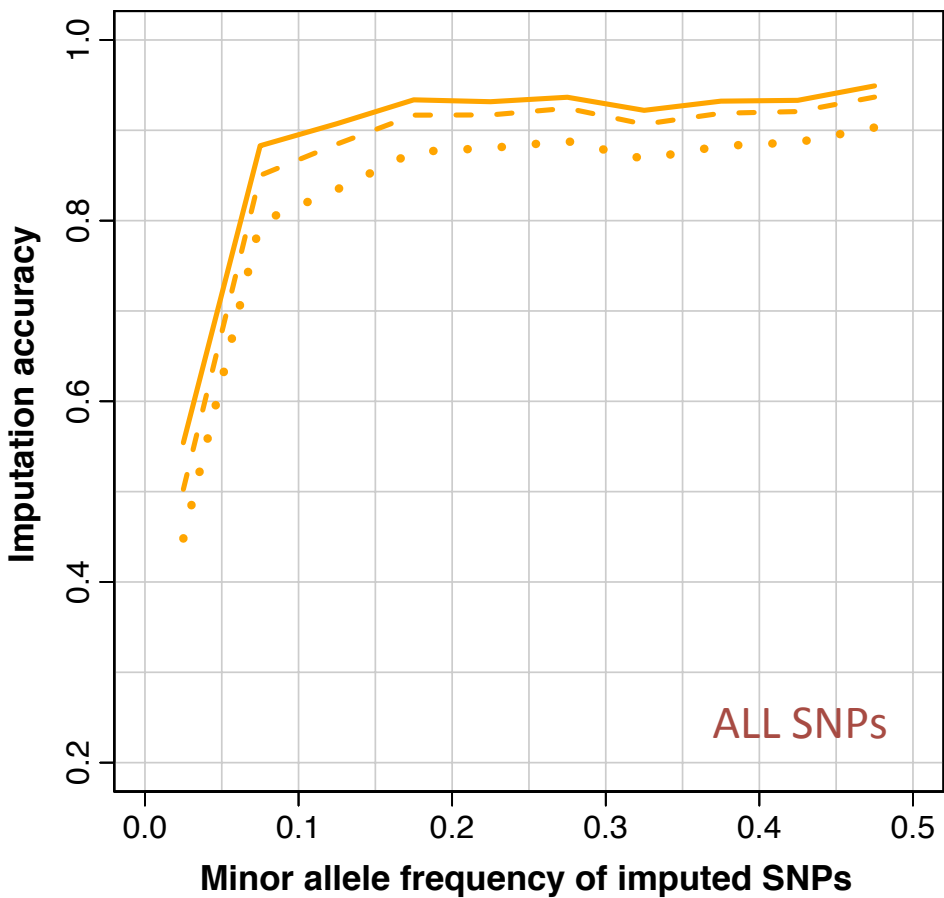


# 1,000 Genomes haplotypes are highly accurate



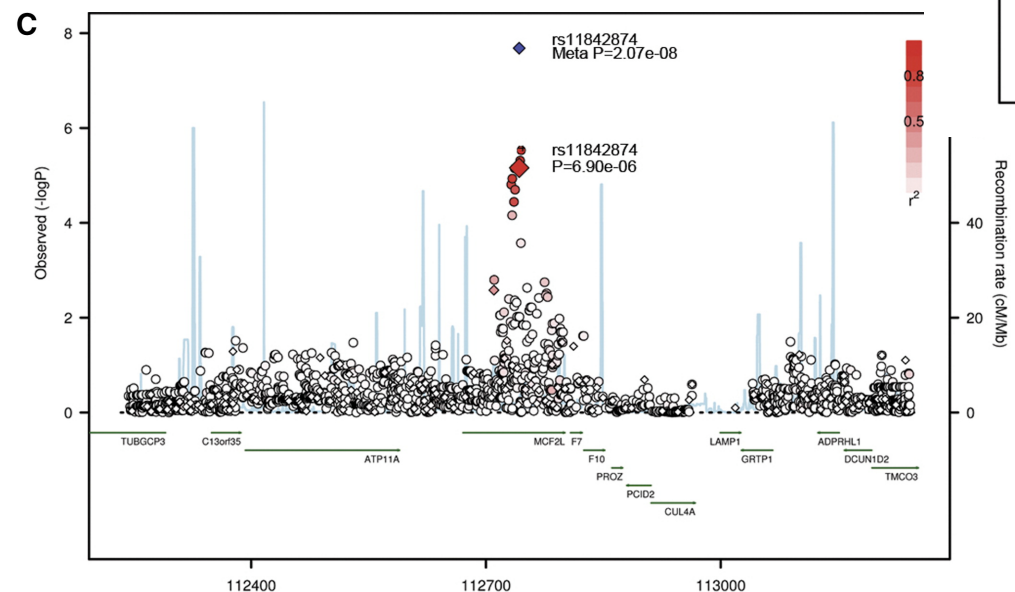
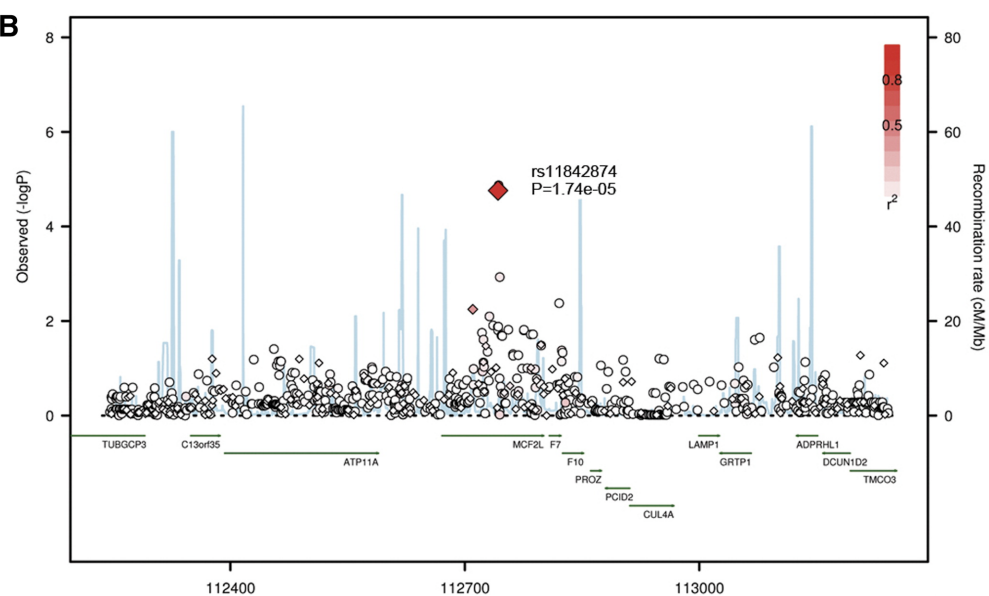
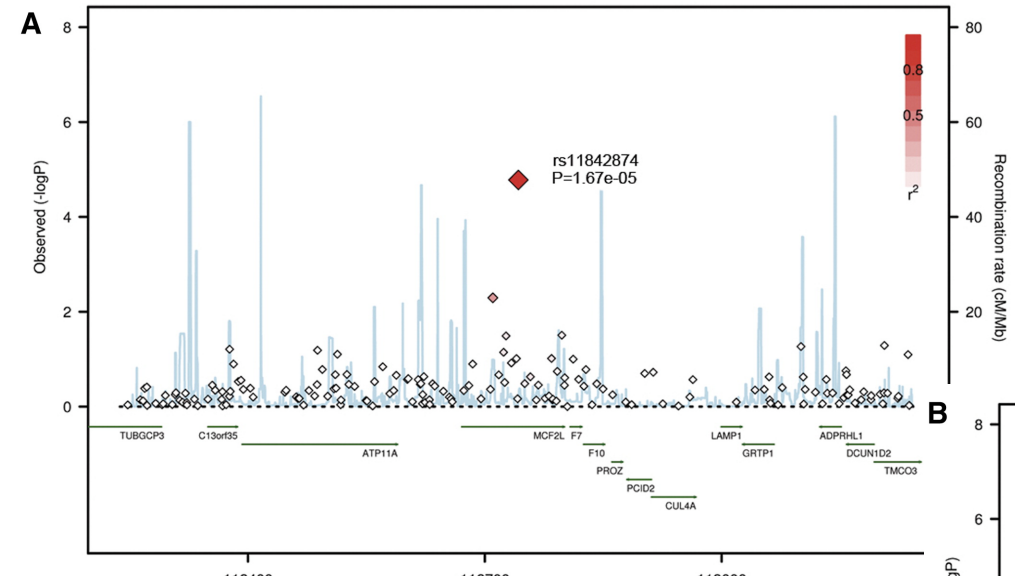
- European ancestry
- African ancestry
- Admixed (Americas)

# Imputation accuracy depends on your GWAS chip



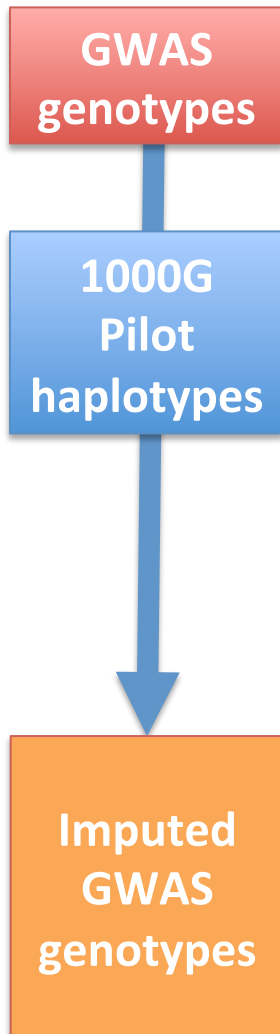
- Omni 2.5M
- - - Illumina 550k
- · · Affymetrix 500k

# Imputation from 1,000 Genomes haplotypes can strengthen association signals.



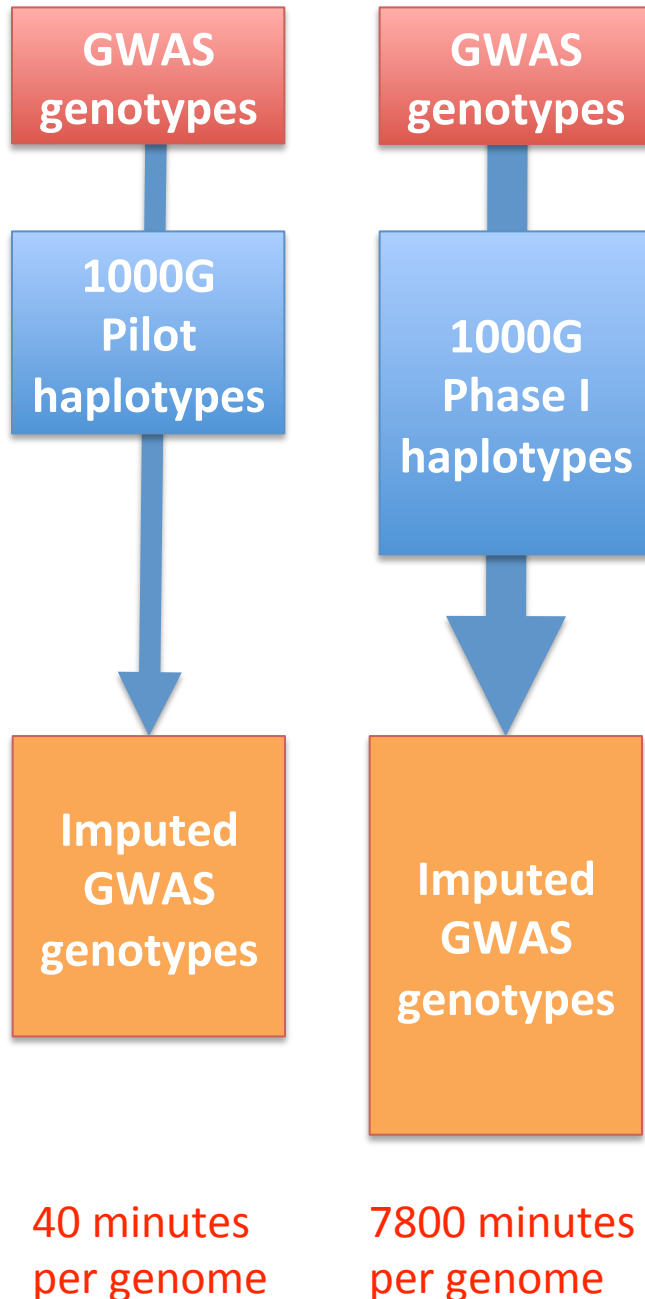
GWAS of Osteoarthritis  
Day-Williams et al. (AJHG 2011)

# Standard Imputation

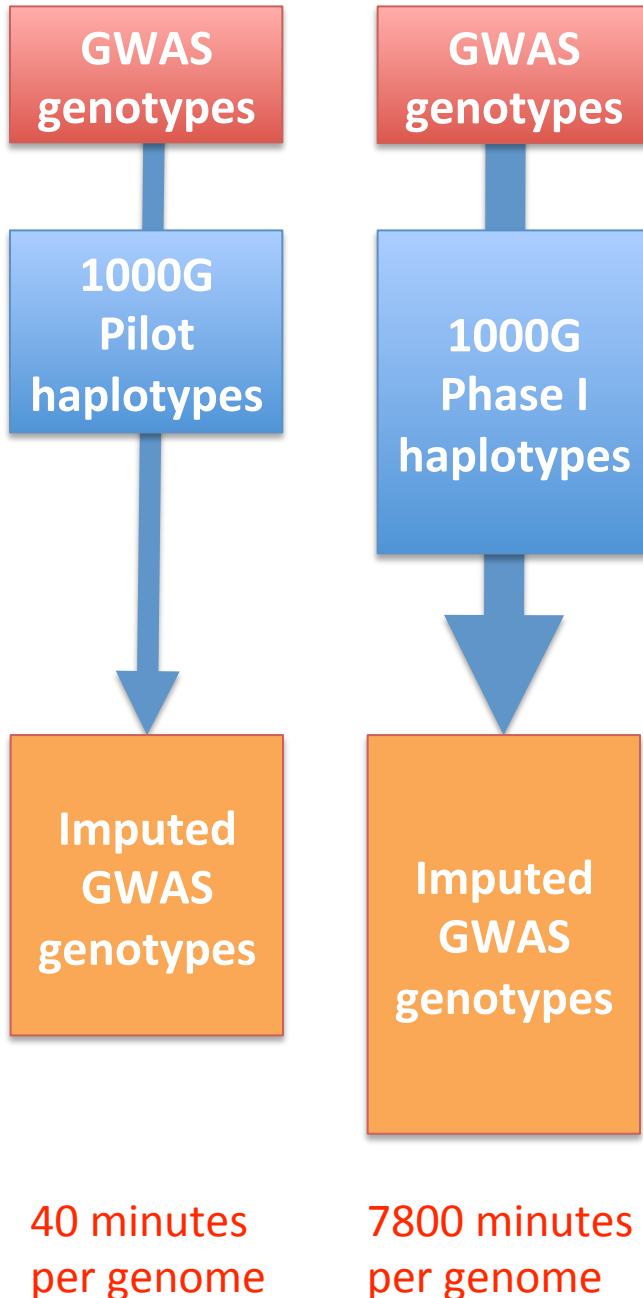


40 minutes  
per genome

# Standard Imputation



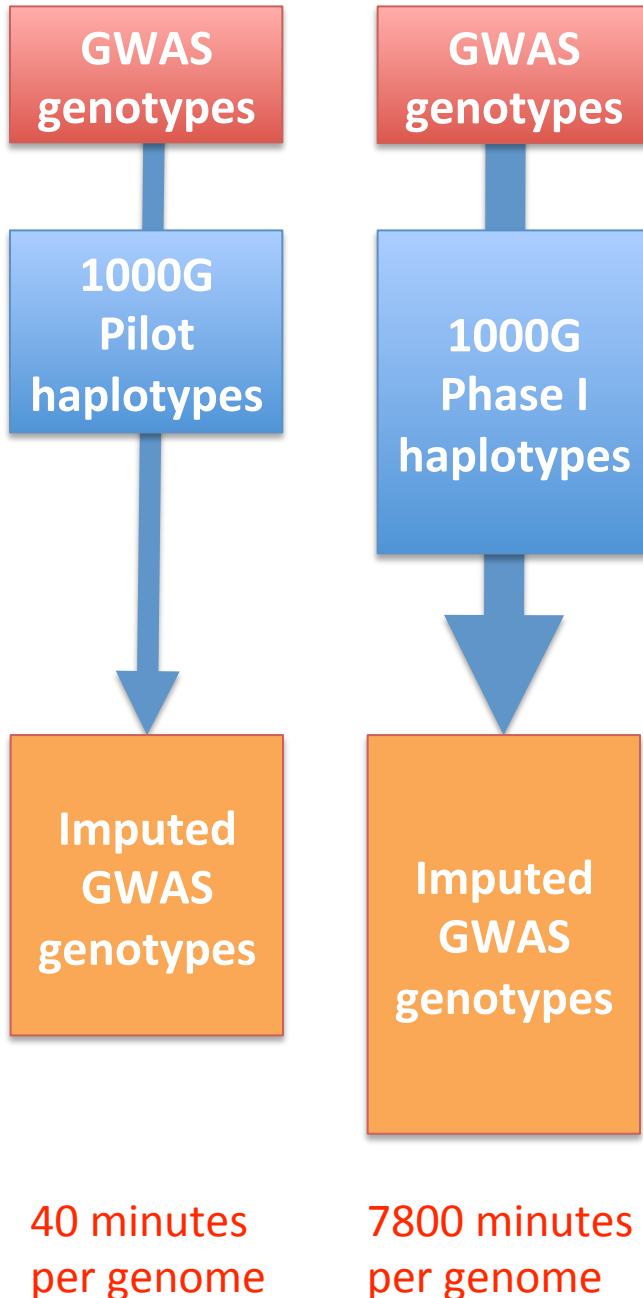
## Standard Imputation



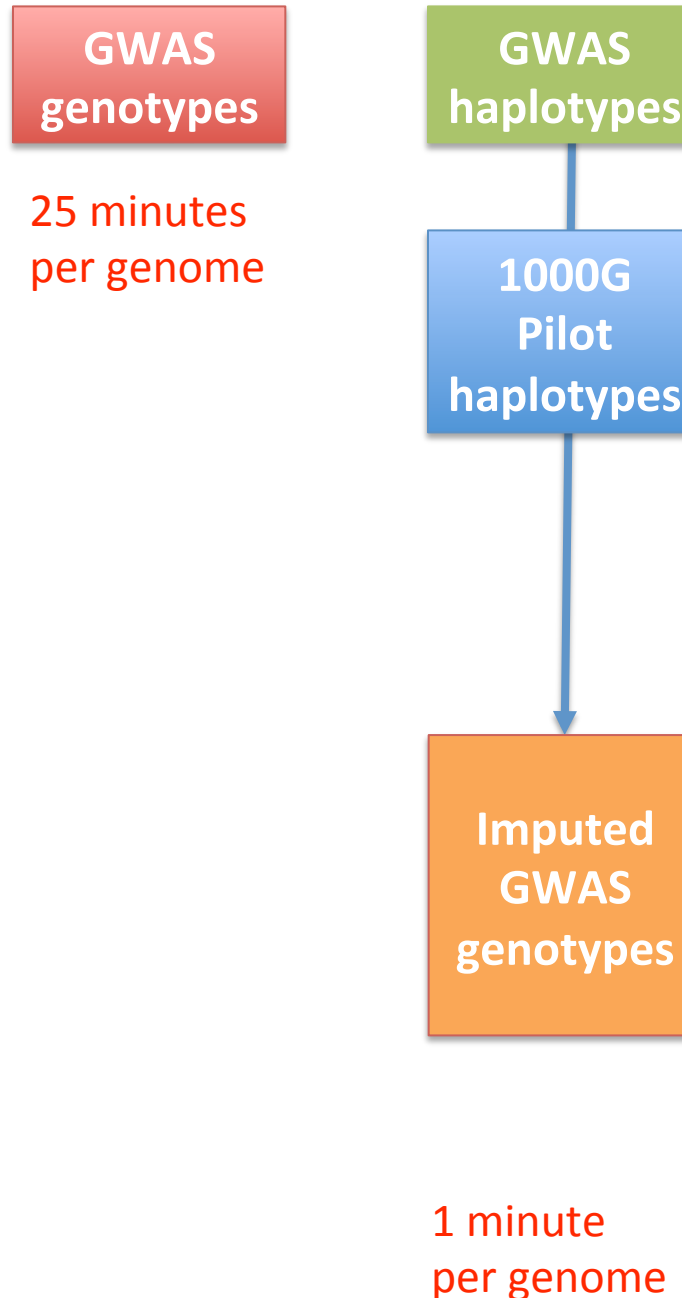
## Pre-phasing Imputation



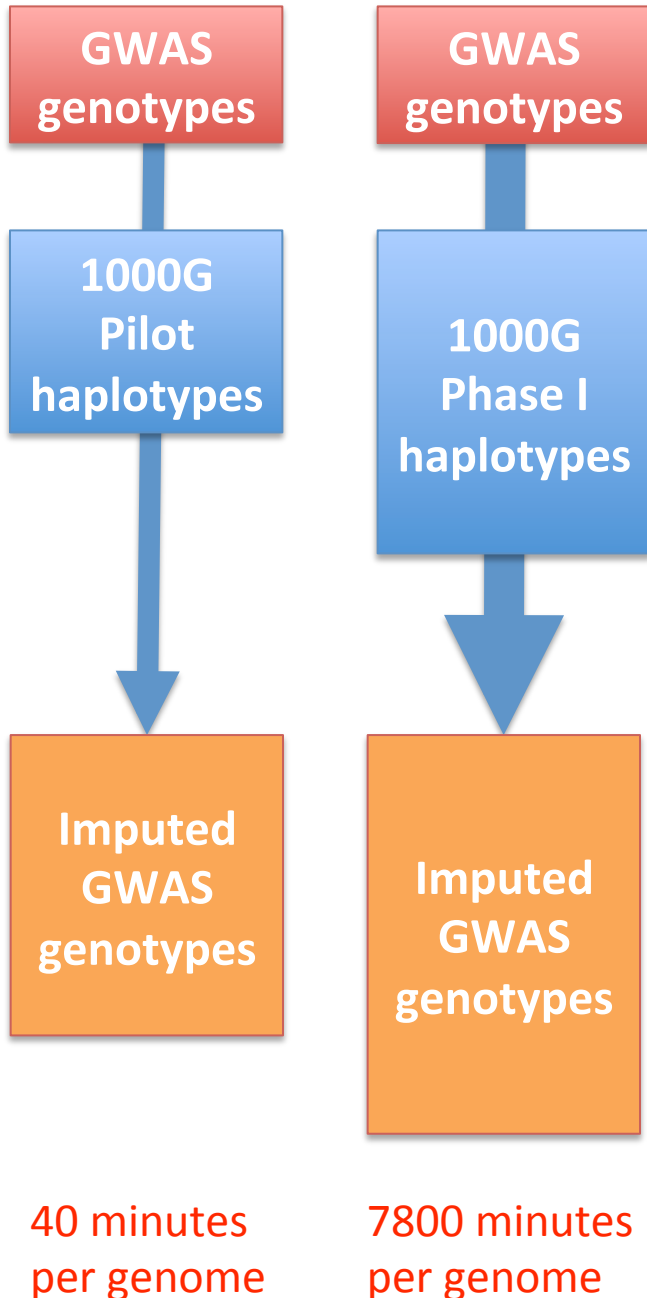
# Standard Imputation



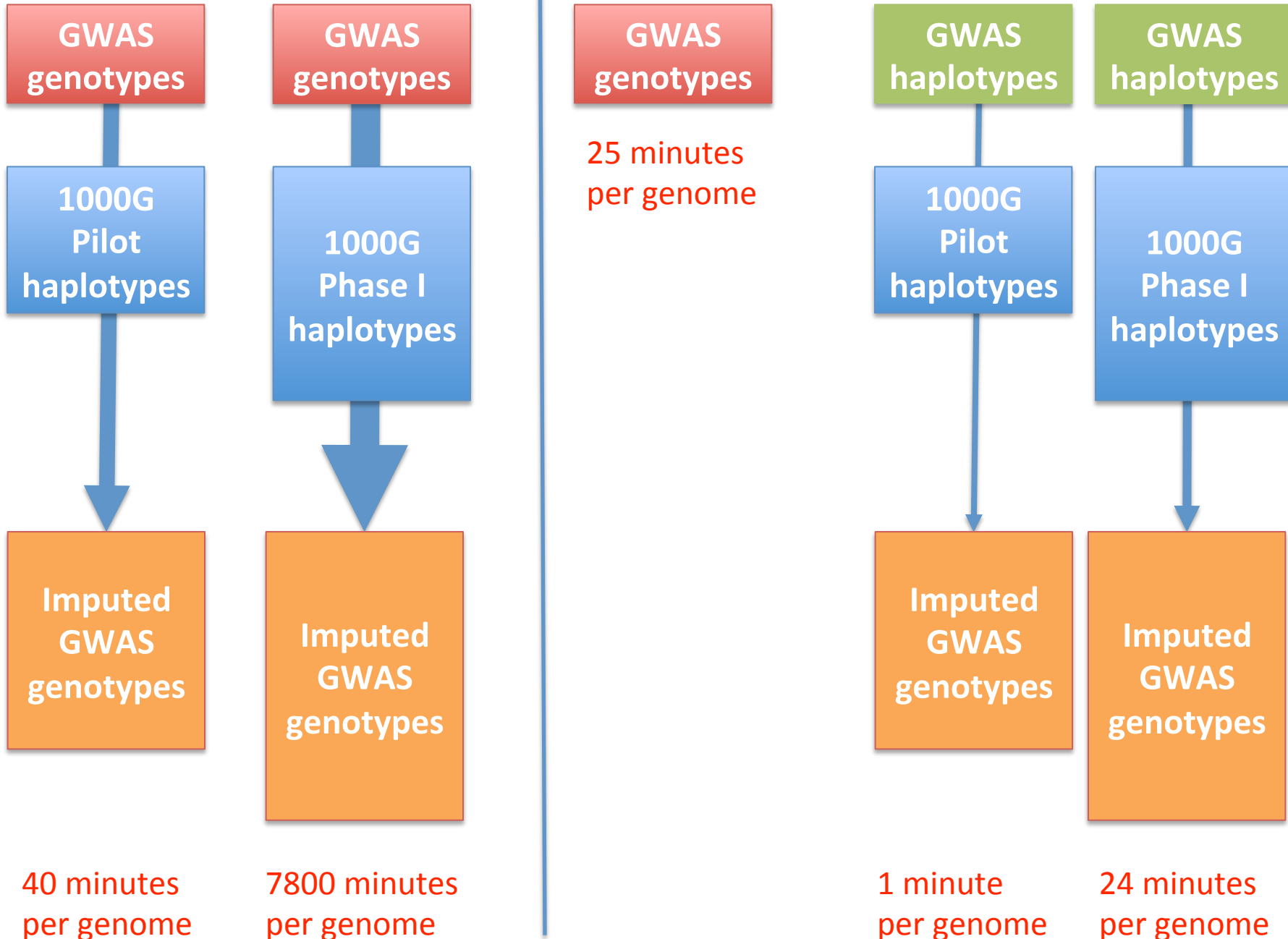
# Pre-phasing Imputation



## Standard Imputation



## Pre-phasing Imputation



## Standard Imputation

## Pre-phasing Imputation

GWAS  
genotypes

GWAS  
genotypes

GWAS  
genotypes

GWAS  
haplotypes

GWAS  
haplotypes

### Imputation Accuracy (mean $R^2$ )

1000G panel

MAF 1-3%

MAF 3-5%

MAF >5%

60 CEU

**0.66**

**0.78**

**0.88**

60 CEU

**0.65**

**0.77**

**0.87**

283 EUR

**0.73**

**0.78**

**0.92**

381 EUR

**0.83**

**0.85**

**0.94**

genotypes

GWAS  
genotypes

genotypes

GWAS  
genotypes

40 minutes  
per genome

7800 minutes  
per genome

1 minute  
per genome

24 minutes  
per genome

# Getting the latest 1,000 Genomes haplotypes

- Phase 1 haplotypes now include SNPs, INDELs, and SVs!
- 1,000 Genomes haplotypes are available in the formats required by various imputation programs. For example:
  - Beagle:  
<http://faculty.washington.edu/browning/beagle/beagle.html>
  - IMPUTE2:  
[http://mathgen.stats.ox.ac.uk/impute/impute\\_v2.html](http://mathgen.stats.ox.ac.uk/impute/impute_v2.html)
  - MaCH/minimac:  
<http://www.sph.umich.edu/csg/abecasis/MACH/download/>
- Thanks for coming!