ClinVar Report

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	ourcing ClinVar input from: clinvar_2014-09-02.vcf ending output to: Report_2014-09-02.pdf	

1 Collect and Merge ClinVar Data

1.1 Import ClinVar VCF

1.2 Merge ClinVar with 1000 Genomes and ExAC

Breakdown of ClinVar Variants

Subset_ClinVar	Number_of_Variants
Total ClinVar	81457
LP/P	19417
ACMG LP/P	3777
ACMG LP/P in gnomAD	839
ACMG LP/P in ExAC	580
ACMG LP/P in 1000 Genomes	130

Breakdown of ACMG-gnomAD Variants

Subset_gnomAD	Number_of_Variants
ACMG in gnomAD	96742
ClinVar-ACMG in gnomAD	4447
LP/P-ACMG in gnomAD	839

Breakdown of ACMG-ExAC Variants

Subset_gnomAD	Number_of_Variants
ACMG in ExAC	59883
ClinVar-ACMG in ExAC	3687
LP/P-ACMG in ExAC	580

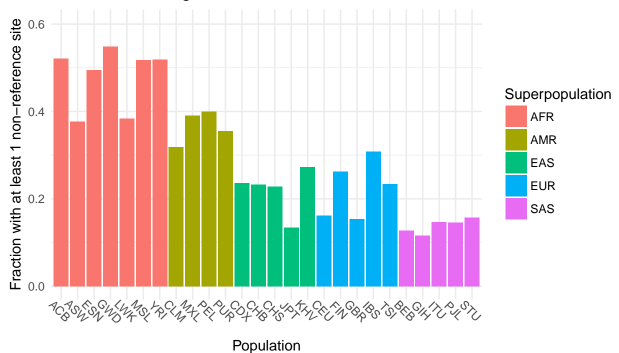
Breakdown of ACMG-1000G Variants

Subset_gnomAD	Number_of_Variants
ACMG in 1000G	141466
ClinVar-ACMG in 1000G	1960
LP/P-ACMG in 1000G	130

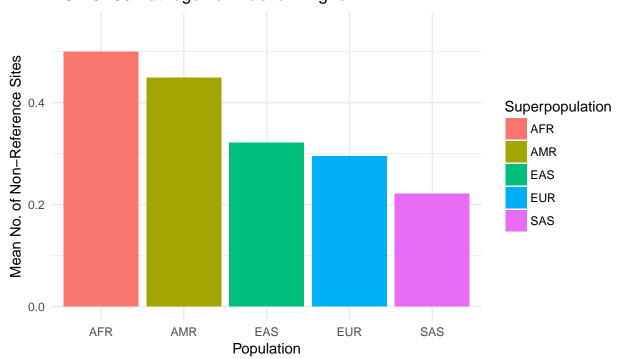
2 Summary Statistics

2.1 Fraction of Individuals with Pathogenic Non-Reference Sites

ACMG-59 Pathogenic: Fraction in 1000 Genomes



ACMG-59 Pathogenic: Fraction in gnomAD

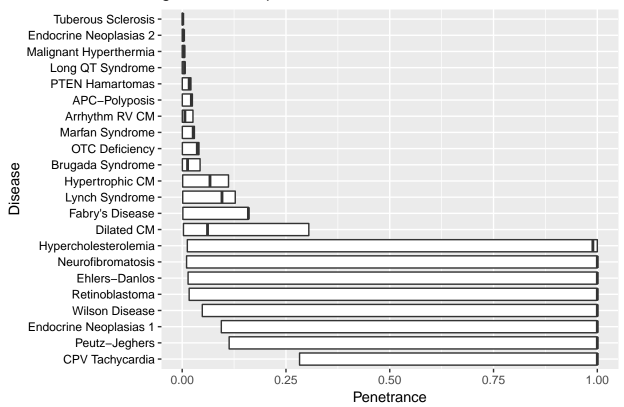


3 Penetrance Estimates

3.1 Max/Min Penetrance as a Function of P(D) and P(V|D)

The left end of the boxplot indicates P(V|D) = 0.01, the bold line in the middle indicates P(V|D) = point value, the right end of the boxplot indicates P(V|D) = 1.

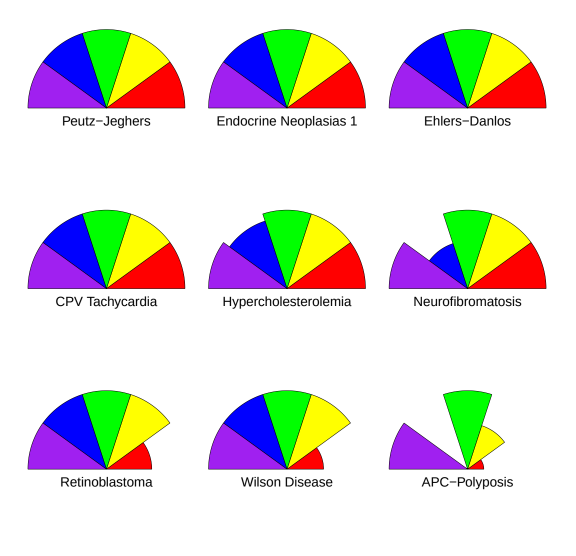
gnomAD: Barplot of Min/Point/Max Penetrance

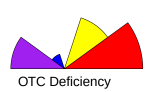


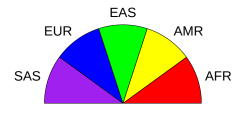
Note: Some diseases have mean theoretical penetrance = 1 because the assumed allelic heterogeneity is greater than is possible, given the observed prevalence and allele frequencies.

3.2 Penetrance Estimates by Ancestry

Radar Plot: Max Penetrance by Ancestry (gnomAD)

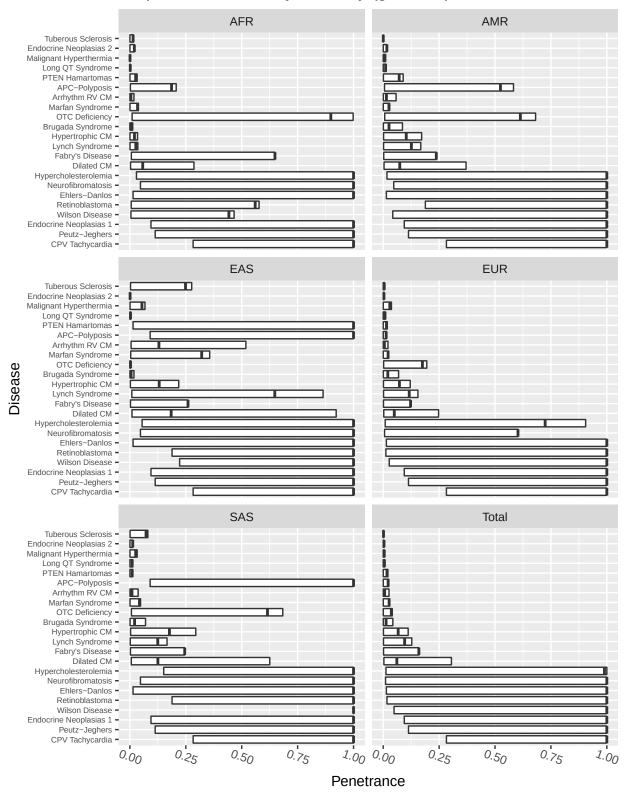




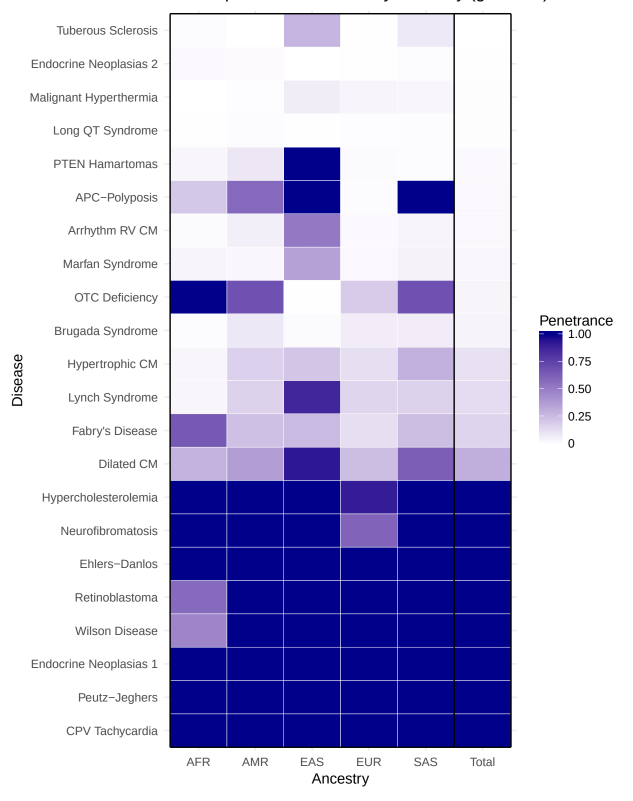


- ## [1] These are the top 10 diseases by summed allele frequencies. NULL values are not plotted.
- ## [1] Each radius is proportional to the penetrance of the disease in the given population.

Barplot: Penetrance by Ancestry (gnomAD)



Heatmap: Max Penetrance by Ancestry (gnomAD)



Dark gray boxes are NA: no associated variants discovered in that ancestral population.