# ClinVar Report

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$\mathbf{S}_{0}$	urci	ng ClinVar input from: clinvar_2015-02-03.vcf	
$\mathbf{Se}$	ndin	ng output to: Report_2015-02-03.pdf	

# 1 Collect and Merge ClinVar Data

# 1.1 Import ClinVar VCF

## Processed ClinVar data frame 99460 x 14 (selected rows/columns):

# 1.2 Merge ClinVar with 1000 Genomes and ExAC

#### ## Breakdown of ClinVar Variants

Subset_ClinVar	Number_of_Variants
Total ClinVar	99460
LP/P-ClinVar	22071
LP/P-ClinVar & ACMG	4073
LP/P-ClinVar & ACMG & ExAC	619
LP/P-ClinVar & ACMG & 1000	120
Genomes	

#### ## Breakdown of ACMG-1000 Genomes Variants

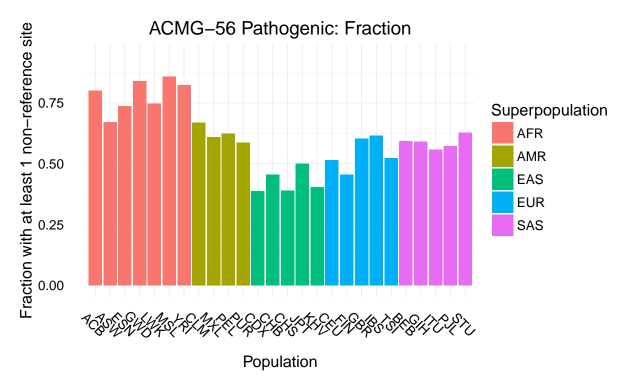
Subset_1000_Genomes	Number_of_Variants
Total 1000_Genomes & ACMG	139335
1000_Genomes & ACMG & ClinVar	2531
1000_Genomes & ACMG &	120
LP/P-ClinVar	

#### ## Breakdown of ACMG-ExAC Variants

Subset_ExAC	Number_of_Variants
Total ExAC & ACMG	58873
ExAC & ACMG & ClinVar	5139
ExAC & ACMG & LP/P-ClinVar	619

# 2 Summary Statistics

# 2.1 Fraction of Individuals with Pathogenic Non-Reference Sites



ACMG-56 Pathogenic: Mean in ExAC

Superpopulation

AFR

AMR

EAS

Population

AFR

SAS

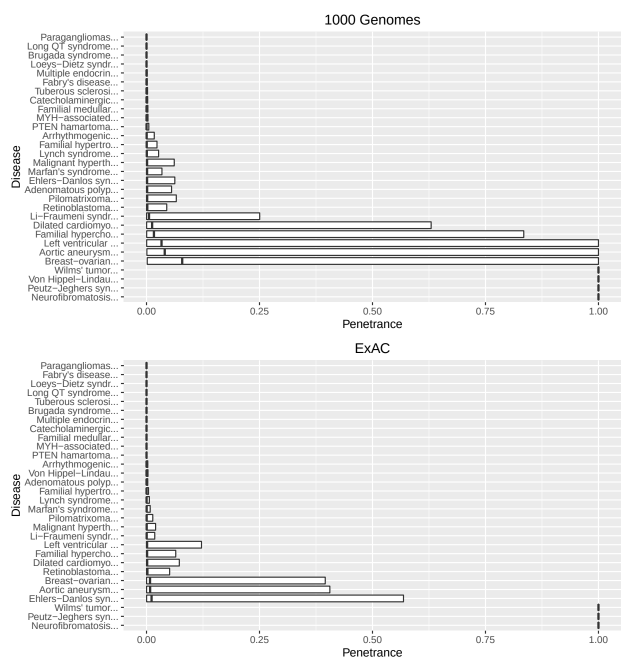
AMR

SAS

#### 3 Penetrance Estimates

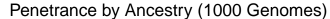
### 3.1 Max/Min Penetrance as a Function of P(D) and P(V|D)

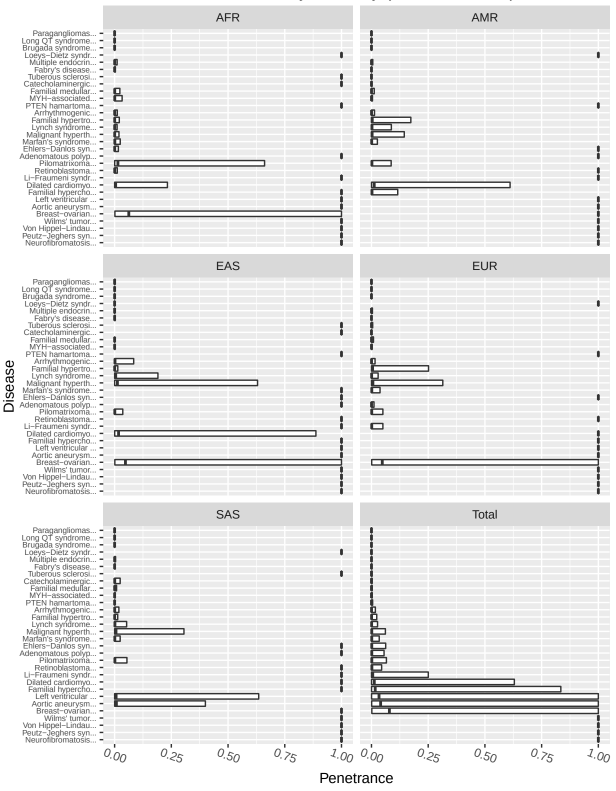
The left end of the boxplot indicates P(D) AND P(V|D) = lower value, the bold line in the middle indicates P(D) AND P(V|D) = geometric\_mean(values), the right end of the boxplot indicates P(D) AND P(V|D) = upper value.



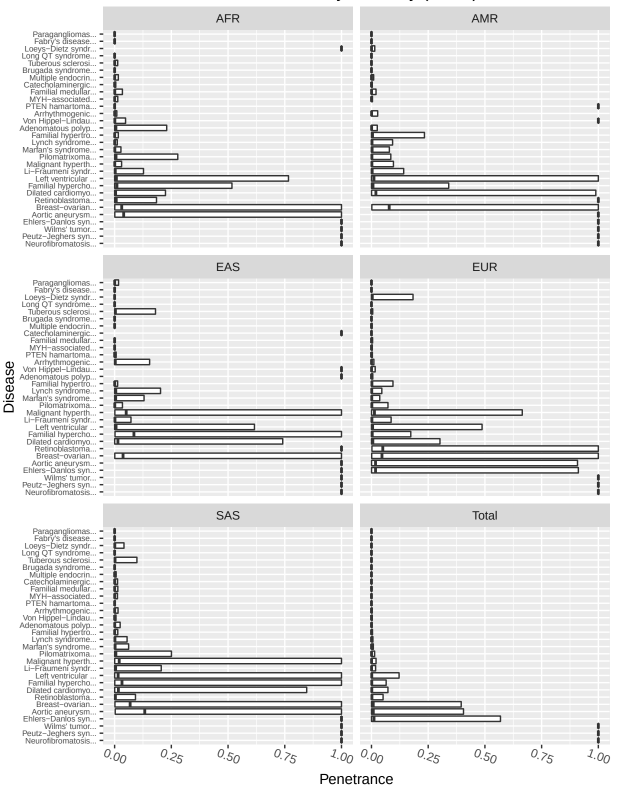
Note: Prevalence ranges of 5x were assumed for all point estimates of prevalence. For example: a point estimate of 0.022 would be given the range 0.01-0.05.

#### 3.2 Penetrance Estimates by Ancestry

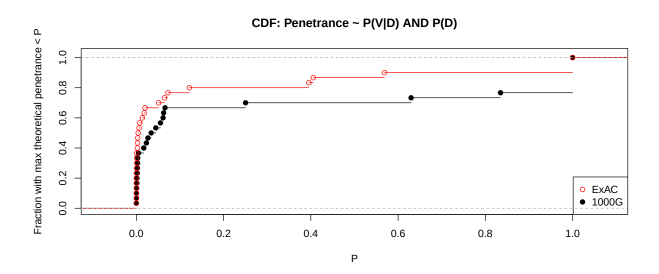




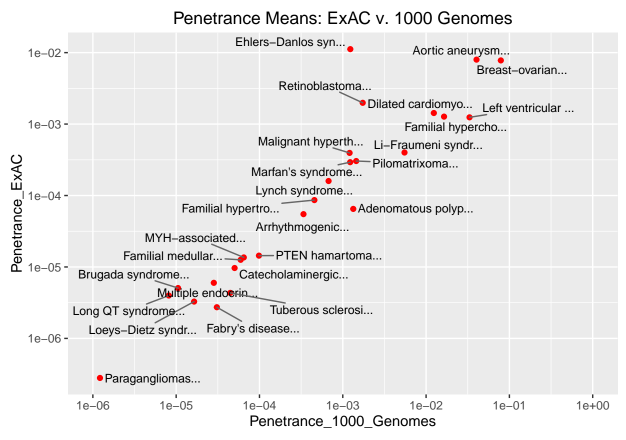
### Penetrance by Ancestry (ExAC)



#### 3.3 Empirical CDFs for All Penetrance Plots



#### 3.4 Comparing Mean Penetrance between ExAC and 1000 Genomes



The Pearson correlation is 0.6. Max penetrance values computed using 1000 Genomes are 3.6-fold larger than those computed using ExAC.