ClinVar Report

James Diao

November 4, 2016

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| \mathbf{So} | ourci | ng ClinVar input from: clinvar_2013-01-18.vcf | |
| \mathbf{Se} | ndin | ng output to: Report_2013-01-18.pdf | |

1 Collect and Merge ClinVar Data

1.1 Import ClinVar VCF

Processed ClinVar data frame 67080×14 (selected rows/columns):

1.2 Merge ClinVar with 1000 Genomes and ExAC

Breakdown of ClinVar Variants

| Subset_ClinVar | Number_of_Variants |
|----------------------------|--------------------|
| Total ClinVar | 67080 |
| LP/P-ClinVar | 12656 |
| LP/P-ClinVar & ACMG | 916 |
| LP/P-ClinVar & ACMG & ExAC | 206 |
| LP/P-ClinVar & ACMG & 1000 | 65 |
| Genomes | |

Breakdown of ACMG-1000 Genomes Variants

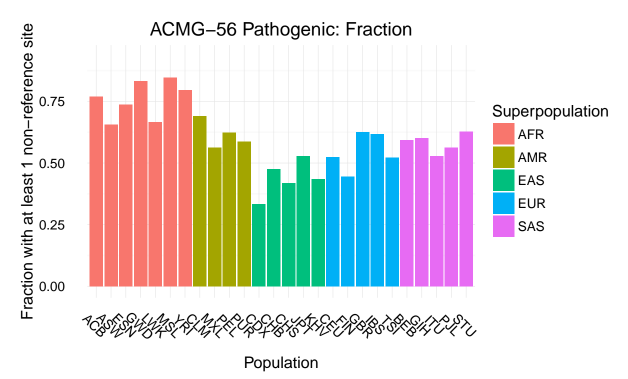
| Subset_1000_Genomes | Number_of_Variants |
|-------------------------------|--------------------|
| Total 1000_Genomes & ACMG | 139335 |
| 1000_Genomes & ACMG & ClinVar | 1214 |
| 1000_Genomes & ACMG & | 65 |
| LP/P-ClinVar | |

Breakdown of ACMG-ExAC Variants

| Subset_ExAC | Number_of_Variants |
|----------------------------|--------------------|
| Total ExAC & ACMG | 58873 |
| ExAC & ACMG & ClinVar | 2255 |
| ExAC & ACMG & LP/P-ClinVar | 206 |

2 Summary Statistics

2.1 Fraction of Individuals with Pathogenic Non-Reference Sites



ACMG-56 Pathogenic: Mean in ExAC

Superpopulation

AFR

AMR

EAS

Population

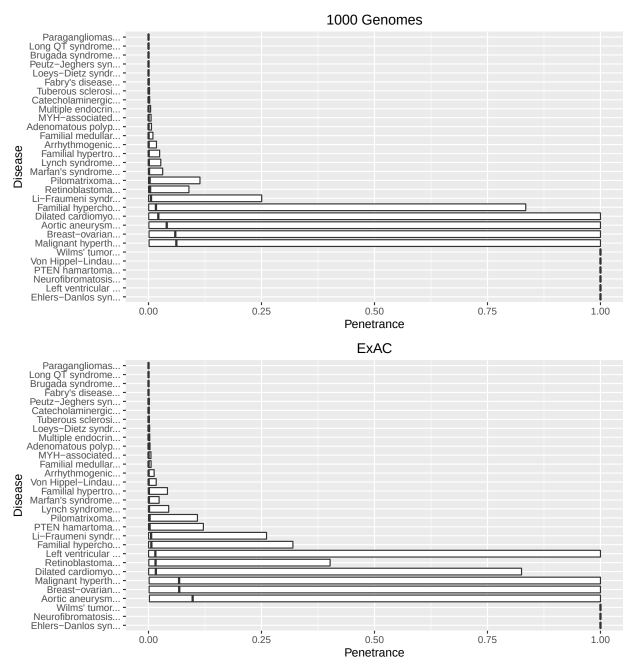
AFR

SAS

3 Penetrance Estimates

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

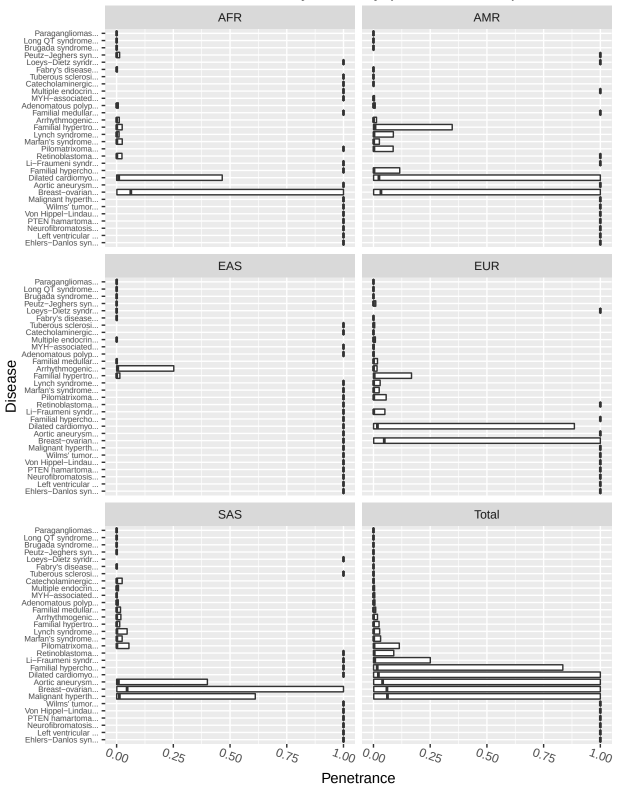
The left end of the boxplot indicates P(D) AND P(V|D) = lower value, the bold line in the middle indicates P(D) AND P(V|D) = geometric_mean(values), the right end of the boxplot indicates P(D) AND P(V|D) = upper value.



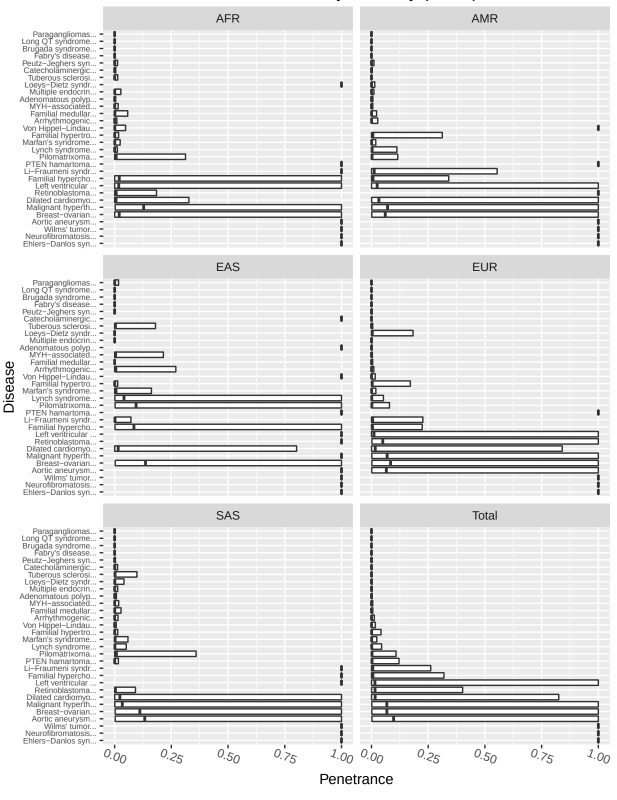
Note: Prevalence ranges of 5x were assumed for all point estimates of prevalence. For example: a point estimate of 0.022 would be given the range 0.01-0.05.

3.2 Penetrance Estimates by Ancestry

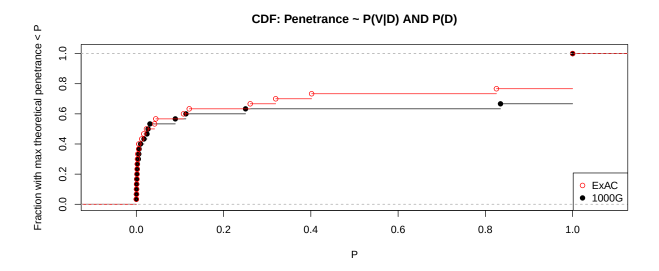
Penetrance by Ancestry (1000 Genomes)



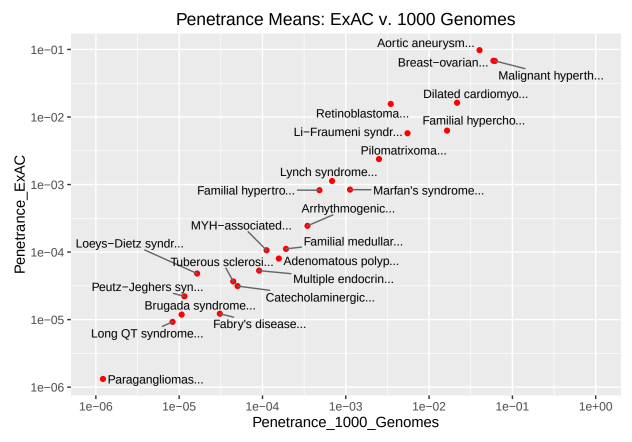
Penetrance by Ancestry (ExAC)



3.3 Empirical CDFs for All Penetrance Plots



3.4 Comparing Mean Penetrance between ExAC and 1000 Genomes



The Pearson correlation is 0.91. Max penetrance values computed using 1000 Genomes are 0.64-fold larger than those computed using ExAC.