PGRNseq Sample and Phenotype Processing

Format the phenotype data for downstream processing.

Expirical Distributions of Phenotypes

```
anc.plt <- ggplot(pheno.df, aes(x=ANC)) + geom_histogram()
print(anc.plt + ggtitle('ANC'))

## stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this.

loganc.plt <- ggplot(pheno.df, aes(x=log10(ANC))) + geom_histogram()
print(loganc.plt + ggtitle('log10(ANC)'))

## stat_bin: binwidth defaulted to range/30. Use 'binwidth = x' to adjust this.</pre>
```

PCA

```
Prune SNP set for LD, filter on MAF, and run PCA.
```

```
ld.snpset <- snpgdsLDpruning(exome.geno)

## SNP pruning based on LD:

## Sliding window: 500000 basepairs, Inf SNPs

## |LD| threshold: 0.2

## Removing 5465 non-autosomal SNPs

## Removing 154888 SNPs (monomorphic, < MAF, or > missing rate)

## Working space: 253 samples, 82543 SNPs

## Chromosome 1: 18.92%, 4668/24674

## Chromosome 2: 20.38%, 3504/17190

## Chromosome 3: 19.84%, 2882/14528

## Chromosome 4: 23.03%, 2349/10198

## Chromosome 5: 21.60%, 2444/11314

## Chromosome 6: 17.92%, 2718/15171
```

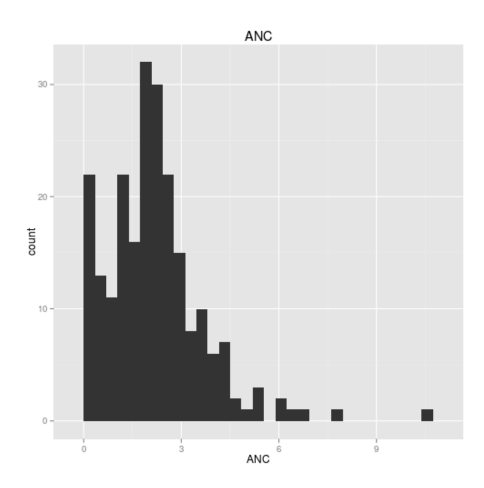


Figure 1: plot of chunk anc_plot

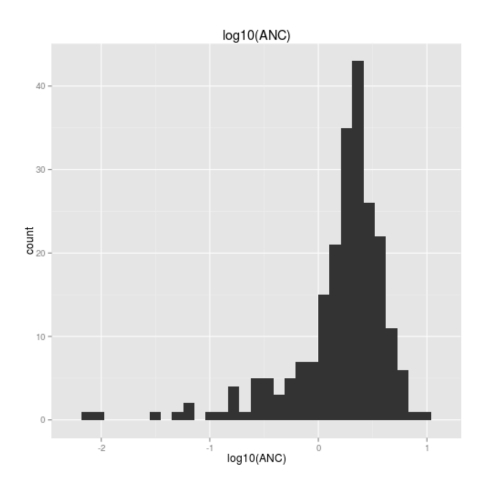


Figure 2: plot of chunk loganc_plt

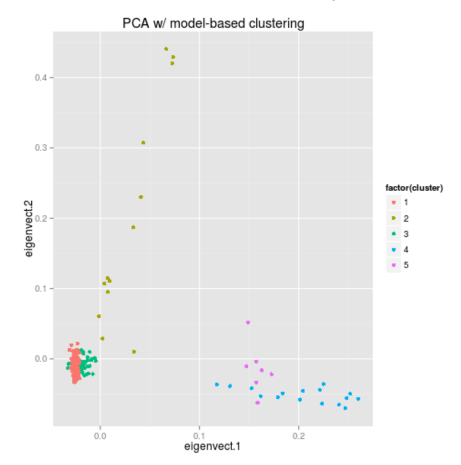
```
## Chromosome 7: 21.33%, 2366/11092
## Chromosome 8: 21.53%, 1921/8921
## Chromosome 9: 20.35%, 2076/10201
## Chromosome 10: 21.80%, 2073/9511
## Chromosome 11: 18.46%, 2866/15528
## Chromosome 12: 19.82%, 2442/12318
## Chromosome 13: 25.00%, 1095/4380
## Chromosome 14: 20.32%, 1569/7721
## Chromosome 15: 19.46%, 1610/8273
## Chromosome 16: 17.75%, 1843/10384
## Chromosome 17: 16.86%, 2185/12963
## Chromosome 18: 24.11%, 904/3749
## Chromosome 19: 15.31%, 2289/14955
## Chromosome 20: 19.42%, 1248/6428
## Chromosome 21: 21.08%, 595/2823
## Chromosome 22: 18.54%, 947/5109
## 46594 SNPs are selected in total.
pca <- snpgdsPCA(exome.geno, snp.id = unlist(ld.snpset), maf = 0.05)</pre>
## Principal Component Analysis (PCA) on SNP genotypes:
## Removing 35251 SNPs (monomorphic, < MAF, or > missing rate)
## Working space: 253 samples, 11343 SNPs
## Using 1 CPU core.
## PCA: the sum of all working genotypes (0, 1 and 2) = 1642261
## PCA: Fri Aug 8 15:36:27 2014
                                     0%
## PCA: Fri Aug 8 15:36:27 2014
                                     100%
                                     Begin (eigenvalues and eigenvectors)
## PCA: Fri Aug 8 15:36:27 2014
## PCA: Fri Aug 8 15:36:27 2014
                                    End (eigenvalues and eigenvectors)
pca.df <- data.frame(sample.id=pca$sample.id, eigenvect=pca$eigenvect)</pre>
Look at the first two PCs, and color the samples by self-reported ancestry.
## Error: object 'eigenvect.1' not found
```

Model-based Clustering

Add principal components to the phenotype data frame, then cluster samples in the first two PCs.

```
pheno.df <- merge(pca.df, pheno.df, by.x = "sample.id", by.y = "id")
names(pheno.df)[which(names(pheno.df) == "sample.id")] <- "id"
cluster <- Mclust(pheno.df[,c("eigenvect.1", "eigenvect.2")])
pheno.df$cluster <- cluster$classification</pre>
```

Examine the results of our PCA and clustering the first two PCs.



Taking a closer look at the clustering of the Europeans. We may be able to explain some genetic variation just by site of ascertainment.

Warning: Removed 35 rows containing missing values (geom_point).

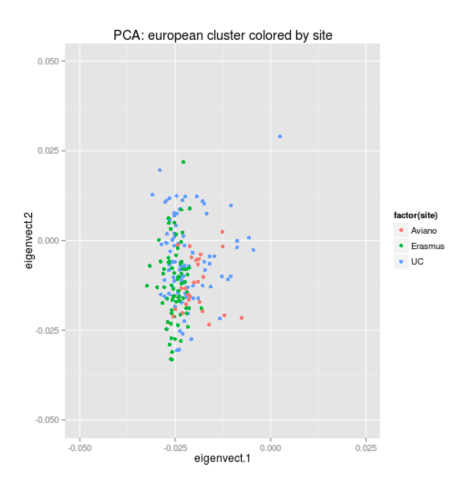


Figure 3: plot of chunk euro_zoom_plot