# **PGRNseq Analysis**

Vassily Trubetskoy March 18, 2014

# 1 Overview

Notes on the on going analysis of genetic data in patients undergoing treament with Irinotecan. The current iteration of the study contains data from samples collected across three different sites. In all, there are currently 253 individuals with targeted sequence and Illumina exome chip data. Samples were collected from patients undergoing a chemotherapeutic regimen with Irinotecan.

## 2 DATA

#### 2.1 GENOTYPES

We currently have access to two different sets of genotypes for this population:

- 1. PGRNseq sequence data on VIP pharmacogenes.
- 2. Illumina Exome Chip data. Run as an internal QC measure at the UW.

We have the alignment files for the sequence data. These were run through the consensus calling pipeline on Amazon.

#### 2.2 PHENOTYPES

The primary phenotype for this dataset is pharmacokinetic and pharmacodynamic data on patients. These PK/PD models are being generated at the University of North Carolina Chapel Hill.

In addition to PK/PD phenotypes, we are interested in looking at Neutropenia in patients. This is currently coded as the Neutrophil Count Nadir (NCN). This is defined as the lowest Neutrophil count observed in a patient during their course of therapy.

There are several relevant covariates available:

- · site of collection
- sex
- · drug dose
- self-reported ancestry

## 2.3 QUALITY CONTROL

Consensus calling in the sequence data. The results look very nice for the metrics that we have available (reference separate report, or put figures here).

- Exclude rare SNPs in single marker analyses
- Exclude SNPs out of HWE (has not yet been done)
- Exclude SNPs with high rates of genotype missingness
- Exclude SNPs with low QUAL
- Check PGRNseq and Exome chip genotype concordance

# 2.3.1 Genotype Concordance

Checking concordance between PGRNseq and Exome chip genotypes.

Number of overlapping variant sites: 883 Number of non-overlapping variant sites: 7850 Genotype concordance: 0.9494

This seems low, but the vast majority of sites contain perfectly concordant calls. The average is significantly lowered due to 30 sites with near-zero concordance.

# 2.4 PROPOSED ANALYSES

- Linear regression. Using the estimated effect and its eror as a t-statistic. (GenABEL package in R)
- SKAT burden test. (SKAT package in R)

var ID	p-val	chr	loc	UCSC gene
exm899253	1.611e-06	11	33596398	KIAA1549L
exm1065858	2.292e-06	13	42390907	VWA8
exm1362200	8.388e-06	17	77078092	ENGASE
exm1453164	1.120e-05	19	33579109	GPATCH1

Table 3.1: Top four results for the single marker exome scan. These correspond to the QQ plot found in Fig 3.7

# 3 RESULTS

## 3.1 POPULATION STRUCTURE

Sequence data provides too few variants to adequately identify ancestry through PCA.

The Exome chip data *does* provide enough markers to differentiate samples. The spatial axes are well defined (see Fig 3.1). These PC's were used to identify a set of genetically homogenous samples (see Fig 3.2 and Fig 3.3).

#### 3.2 SINGLE MARKER

#### 3.2.1 Notes and Issues

There may be some genomic inflation present in the single marker tests. This needs to be formally reported as a calculation of lambda.

#### 3.2.2 GENABEL LINEAR REGRESSION

No results in sequence data for common variants. The QQ plot without covariates shows some deflation in the tail (see Fig 3.4). Once adjusting with available covarites, the scan looks reasonably null (see Fig 3.5).

One significant marker in Exome chip data, with a couple of suggestive hits. The p-value improves with a full model including covariates.

#### 3.3 GENE-BASED

## 3.3.1 Notes and Issues

I'll have to think some more about which SKAT test is appropriate for this data. There is an excellent article by Lee et. al (2012) that proposes the optimal rare variant test SKAT-O. This statistic is a linear combination of burden and variance component statistics. You can also specify different kernels. The default kernel weights variants based on rareness.

ENTREZ ID	SKAT asymptotic p	Num. Snps	Symbol
3177	8.183411e-08	8	SLC29A2
6799	2.130900e-05	14	SULT1A2
8824	3.287893e-04	78	CES2
6817	5.014952e-02	27	SULT1A1
4881	6.311036e-04	27	NPR1

Table 3.2: Top five results for the burden test. P-values were generated with an theoretical distribution of the test, as well as with  $10^5$  bootstrap iterations in a resampling procedure. The expensive nature of bootstrap means that the smallest possible observable pvalue is  $10^{-5}$ .

The variants need to be cleaned. Right now, a lot of genes are dropped due to high rates of missingness. Additionally, many genotypes are being imputed through the SKAT package based on HWE.

I am not using the finite sample correction that the author's suggest for n < 2000. P-values are computed with a parametric bootstrap with n = 10000 iterations.

## 3.3.2 SKAT BURDEN

Currently running for the model: ANC = Geno + sex + site + dose. Eleven samples are excluded for having missing covariate information. Many sites contain high rates of missingess, and imputation is performed by the SKAT package. Fig 3.4 shows the QQ-plot that contains a number of significant associations in the tail. These top hits are summarized in the Table 3.1.

# 3.3.3 SKAT RARE + COMMON

Many of the variants are rare in this dataset. Our top hits are nearly identical to the burden top hits since the common test is being down weighted.

## 4 REFERENCES

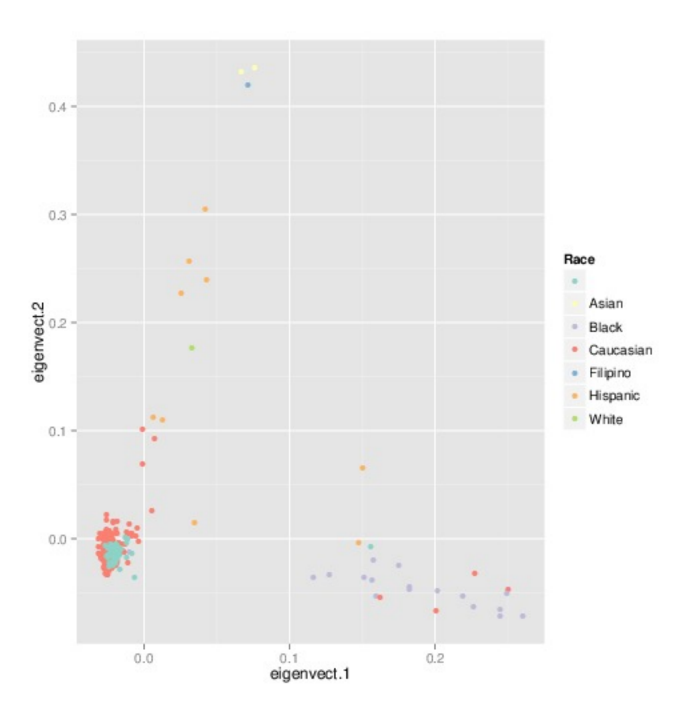


Figure 3.1: First two principal components of exome chip genotype data. The samples segregate fairly well into their self-reported ancestries.

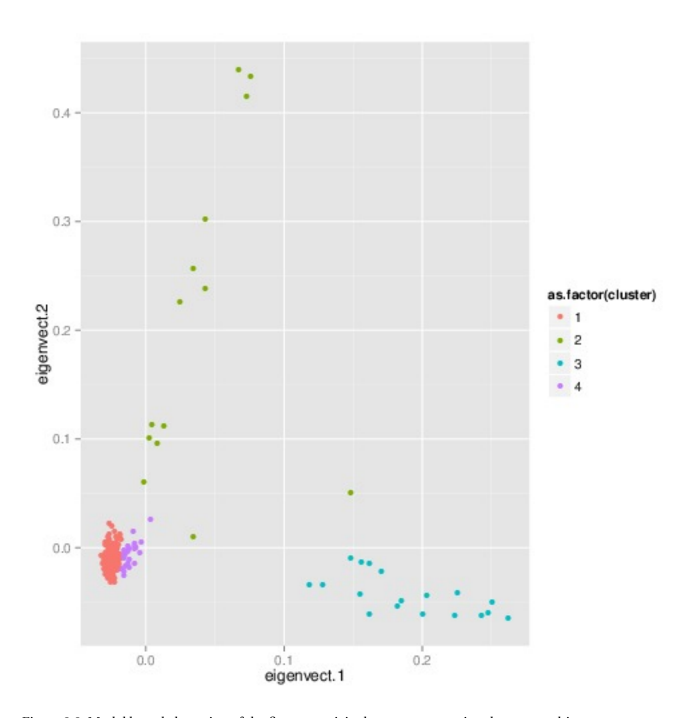


Figure 3.2: Model based clustering of the first two pricipal components using the exome chip genotypes. These clusters were used to select a homogenous set of samples. In this case, we subset our data to the 161 samples contained in cluster 1.

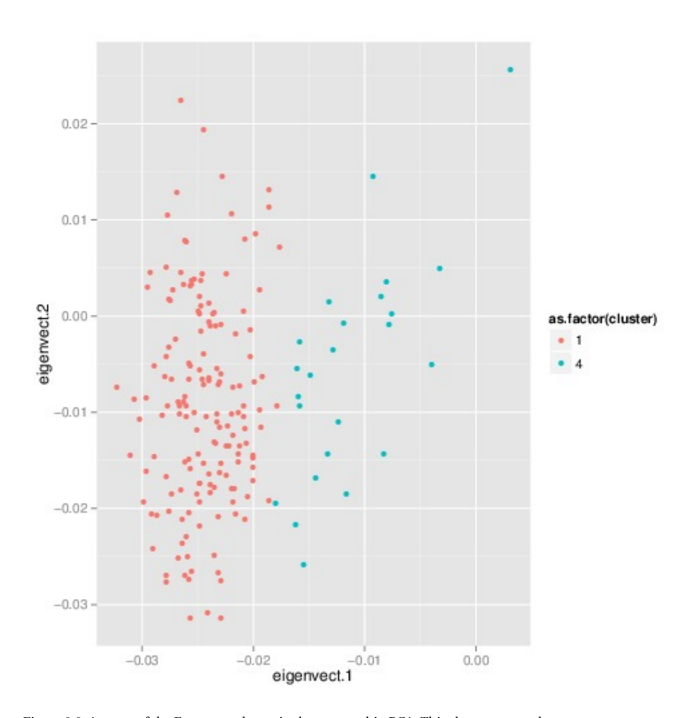


Figure 3.3: A zoom of the European cluster in the exome chip PCA. This shows some substructure, and we ultimately subset the samples contained in cluster 1.

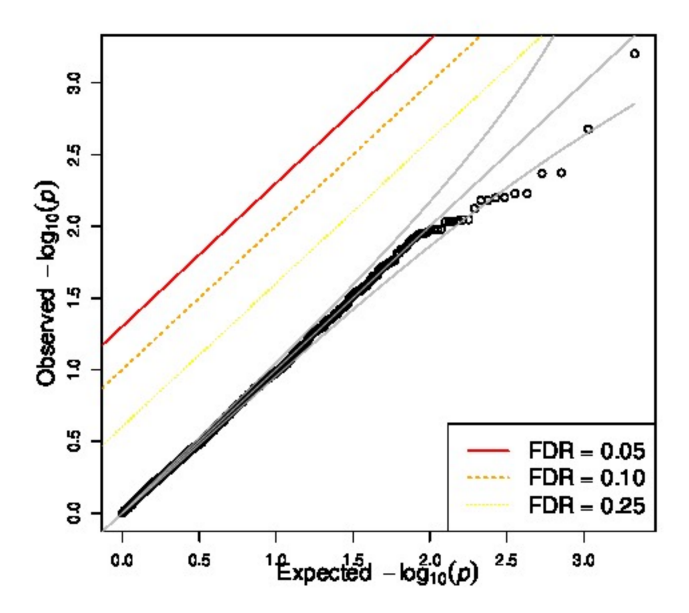


Figure 3.4: GWA scan for sequence data using the basic model: ANC = Geno. SNPs were excluded based on the following criteria: call rate < 0.95, MAF < 0.05, and HWE. Individuals were excluded based on missing phenotype and PCA outliers. This result was produced with 160 individuals with 2,140 markers. There is some deflation seen in extremely small p-values.

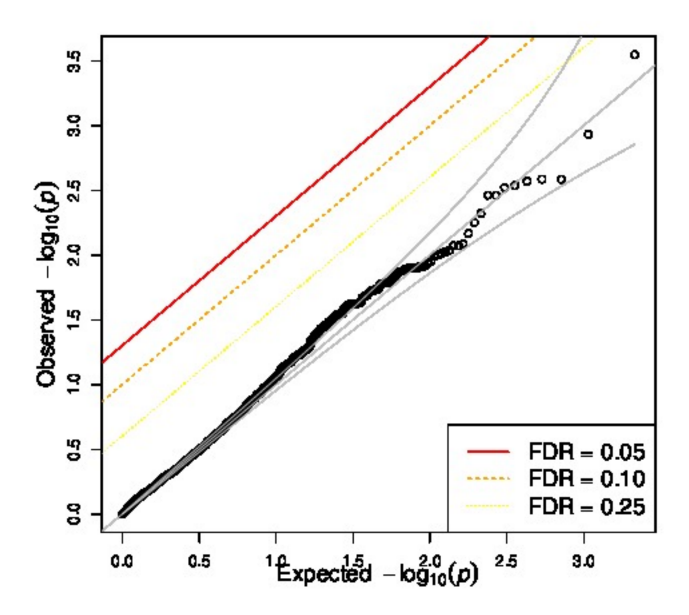


Figure 3.5: GWA scan for sequence data using the full covariate model: ANC = Geno + sex + site + dose. SNPs were excluded based on the following criteria: call rate < 0.95, MAF < 0.05, and HWE. Individuals were excluded based on missing phenotype and PCA outliers. This result was produced with 160 individuals with 2,140 markers. The covariate correction reduces the deflation observed at small p-values.

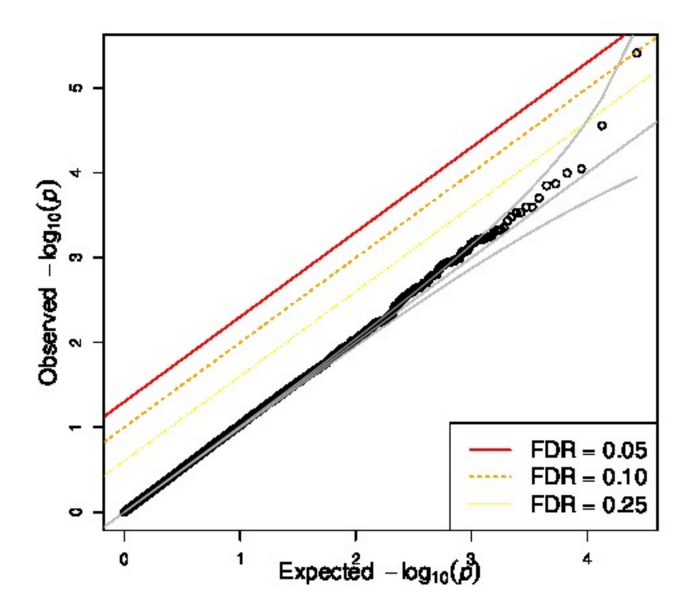


Figure 3.6: GWA scan for exome chip data using the basic model: ANC = Geno. SNPs were excluded based on the following criteria: call rate < 0.95, MAF < 0.05, and HWE. Individuals were excluded based on missing phenotype, missing covariate and PCA outliers. The final dataset consisted of 159 individuals and 26,728 markers.

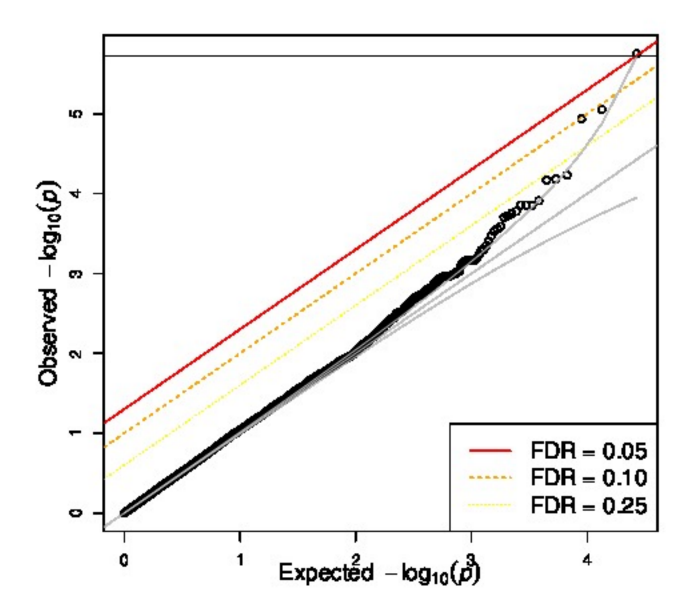


Figure 3.7: GWA scan for exome chip data using the full covariate model: ANC = Geno + sex + site + dose. SNPs were excluded based on the following criteria: call rate < 0.95, MAF < 0.05, and HWE. Individuals were excluded based on missing phenotype, missing covariate and PCA outliers. The final dataset consisted of 159 individuals and 26,728 markers.

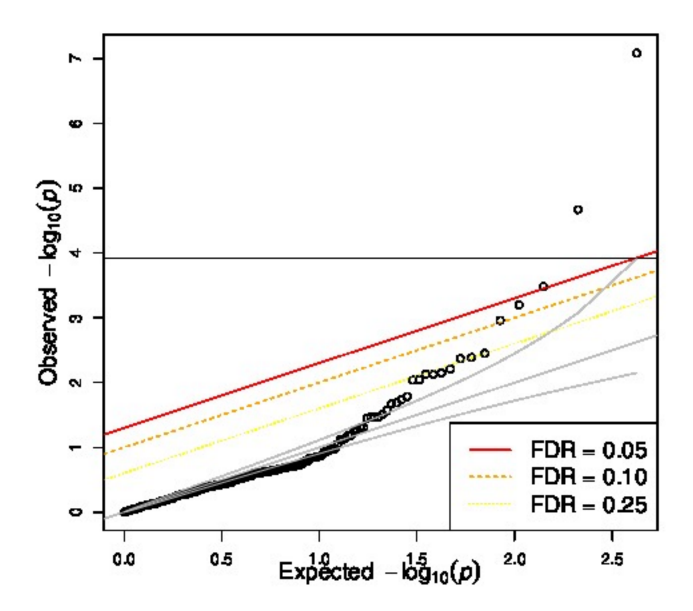


Figure 3.8: QQ plot from a set of 421 gene-based units tested with SKAT's burden test in 150 individuals. P-values were generated with a theoretical distribution (though the top hit is maintained with a resampling based p-value).