Jaccard Kernel PCA in genotype and gene-burden data for ALS

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Abstract

This is a summary of important results obtained during my four-and-a-half months long internship at the UMC Utrecht Brain Center under the guidance of Dr. Kevin P. Kenna. It should be readable by non-bioinformaticians, but assumes familiarity with basic statistics and principal component analysis.

The goal of the project was to aid an ongoing GWAS on Amyotrophic lateral sclerosis (ALS) by improving on its population stratification control method. Given the nature of the data (exome genes and rare variants only; gene-burden data instead of genotype data), the usual method (principal component analysis) is unsatisfactory. A good candidate method is Jaccard principal component analysis (jPCA) - a particular case of Kernel principal component analysis.

I developed a set of scripts capable of running jPCA on very large datasets (through parallelization) and another one allowing for arbitrary positive integers burden values - thus working not only on genotype data but also on gene-burden data. My implementation of jPCA was tested by checking that its output is identical to the output of another, less general implementation of jPCA when applied to the 1000 genomes project data [1].

Applying jPCA on exome data from dbGaP led to surprising results: although jPCA captures population stratification on the Human Genome Project data [1], it is not capable of doing the same in the exome data explored here, whether we use common or rare variants. The problem seems to lie either on the specific distribution of the exome variants or the data collection limitations. Exactly how this happens is left as an open question for future work.

1 Introduction

Amyotrophic lateral sclerosis (ALS) is a neurodegenerative disease mostly affecting motor neurons, causing progressive weakness of voluntary muscles and eventually death by respiratory arrest [2]. Its cumulative lifetime risk is approximately $\frac{1}{300}$.

It is of obvious importance to identify what genes¹ are associated with ALS. Nowadays it is possible to sequence the entire genome, and use statistical methods to analyze the data in so-called genome wide association studies (GWAS) (see §1.2. The distribution of disrupting variants can be compared to population controls to identify genes with an excess of potentially pathogenic variants in sick patients. But these datasets are enormous, and the results may be prone to misinterpretation: so many sequences can be considered that by chance alone one could find a certain sequence over-represented in sick patients. Auxiliary genetic information can help reducing the search space, e.g. one may prioritize variants with higher predicted pathogenicity based for example on alterations of the encoded proteins. Furthermore, mutations of the non-exome part of the genome are even harder to interpret since we don't understand many of its functions. For these reasons, and for its higher availability, we use exome sequencing

Some problems arise specifically in a ALS GWAS be-

cause the variants contributing to risk are rare variants. This complicates matters for a few reasons [2]:

- 1. Not all protein-changing mutations found in patients but not in controls indicate pathogenicity: every person has on average 10 major protein-altering mutations with no disease effects.
- 2. When the disease-causing mutations are rare one needs more data.
- 3. Rare variants may be population-specific, making replication difficult.
- 4. The non-presence of rare possibly pathogenic variants in patients does not mean that we can discard it after all, it is rare.
- 5. Rare variants may have different distributions than common variants [3], rendering useless some methods usually applied to common variants.

Some possible solutions are [2]:

- 1. Use extremely large datasets from global efforts like Project MinE.
- 2. Support the results with biochemical functional analysis.
- 3. Use gene-burdens (see § 1.1): lower resolution data may identify pathogenic genes, even if sacrificing a nucleotide-scale view of the data.

¹For the reader unfamiliar with biology lingo, there is a glossary in §5.2 with some of the most important terms used in this text.

Let us briefly discuss the type of data we are dealing with and identify the problem to solve.

1.1 The genotype matrix and gene-burdens

A gene can be seen as a sequence of nucleotides from {A, T, G, C}. Each person has two copies of each gene, one from the mother and one from the father.

from mother: A T T G A C $C \dots$ from father: A A T G G C $C \dots$

Single nucleotide polymorphisms (SNPs) are positions in the genome where there are variations within the sampled population.

person 1:
$$G$$
 T C A A C C person 2: G T A A A G C person 3: G T C A A G C person 4: G T C A A C C

A SNP has at least two variants, but can have up to four: one for each distinct nucleotide. The variant with the highest frequency in the population is called the *major allele*. All other variants are *minor alleles*.

for the individual i and SNP j, the genotype value (or SNP-burden) $b_{i,j}$ is:

$$b_{i,j} = \begin{cases} 0, \text{ no minor alleles at } j \text{ on father and mother genes.} \\ 1, \text{ a minor allele at } j \text{ on one of the parents' genes.} \\ 2, \text{ minor alleles at } j \text{ on the genes of both parents.} \end{cases}$$

the matrix $[b_{i,j}]$ is called the *genotype matrix*.

As discussed in §1, when dealing with rare variants one may want to consider *gene-burdens*, which are simply sums of the SNP-burdens in a gene.

1.2 GWAS and controlling for population stratification

In a genome-wide association study (GWAS), one tries to assess the pathogenicity of the SNP variants w.r.t. a certain disease, using all the SNPs of the genome.

In the language of statistics, the predictors are the genotype values $b_{\cdot,j}$ and the response is a "is sick?" boolean, called the *phenotype*.

Since the number of SNPs in the human population is enormous, very large quantities of data are needed, usually from individuals around the world. This is true even when one restricts oneself to the exome, as we do. In such a scenario, one cannot simply run a classification modelsay, multivariate logistic regression - and expect good results: the ancestry differences between individuals, which are highly correlated with their geographical differences, is a confounding variable that we must account for. Indeed, correcting for population stratification is now common practice in GWASs.

In [4], it is shown that the first two principal components of the data obtained from principal component analysis (PCA) on genotype data from the 1000 genome project (OTGP) are highly correlated with geographical axes. This means that one can use the principal components of a population's genotype data to control for ancestral differences - whose effect on the population genome is known as *population stratification* - by simply taking those principal components as covariates in whatever classification model we choose to use.

1.3 The gwas pipeline

Given what we said above, the pipeline for a GWAS can be separated in four parts: The input data; a method for stratification control (e.g. PCA); a supervised classification model (e.g. logistic regression; and some methods for analysing the resulting p-values (e.g. manhattan plots and applots, which we will not describe here).



Each of these parts can be modified. In this work, we only tinker with the first two:

- Instead of a genotype matrix, my supervisor proposed the use of a matrix of gene-burdens for the reasons stated in §1. Thus I tested my methods on both genotype and gene-burden data.
- Instead of PCA, one must use methods better suited for rare variants. As we will see shortly, I ended up using two types of so-called jaccard PCA.

2 Data and methods

As explained in §1, I tested my methods to capture population structure in both exome genotype and gene-burden data. The gene-burden matrix was obtained from the genotype data by adding the SNP-burdens of each gene, obtaining as many rows as there are genes in the human exome. Instead of this matrix, I used filtered versions of it, containing only mutations that affect the protein encodings.

Concretely, this data is divided in three datasets:

- 1. Genotype data from the database of genotypes and phenotypes (dbGaP), corresponding to 1343816 exome SNPs and 30820 individuals, approximately 23% of which are ALS patients (phenotype = 1). For the parts where we only wanted rare variants, we extracted them from the genotype matrix.
- 2. The LOF dataset: a gene-burden matrix from 30820 individuals and 12802 exome genes. It only takes

into account rare and severe mutations that are known to cause a loss of function of the encoded protein. Also, this data has been mean-imputated².

3. The *moderate* dataset: a gene-burden matrix from 30820 individuals and 17881 exome genes. It takes into account moderate (mostly missense) rare mutations. Like LOF, this data has been mean-imputated.

2.1 Alternatives to PCA

PCA selects the directions for which the variance is maximized. Those directions are called the *principal components* (PCs) of the data.

The PCA algorithm always gives us the PCs of the data, but they are not always meaningful or useful.

PCA is useful when the PCs include most or all of the interesting information in the data. More precisely, this means that the reconstruction error is low enough for our purposes.

For PCA to be useful, the following must be true: [5]

- 1. Linearity: expressing the data in terms of the PCs corresponds to a change of basis.
- 2. PCs are orthogonal: this helps with the linear algebra problem of finding the PCs.
- 3. Directions of largest variance are the important³ ones. This means that signal to noise ratio is high directions with lower variance correspond to noise.

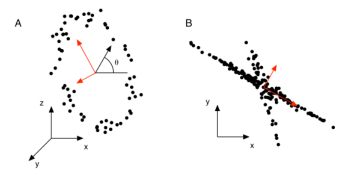


Figure 1: Examples of when PCA fails; taken from [5]. The red arrows represent the two first PCs. A: circular data violates the linearity assumption; the angle θ (from polar coordinates) would be a much better coordinate to use as PC than the PCs from PCA. B: Non-orthogonal PCs - PCA cannot recognize that the PCs are not orthogonal, since it only seaches for othogonal PCs.

Remark: The first and third assumptions are automatically satisfied for (multivariate) normal distribution

of the data, and the second assumption is usually true for not-too-noisy data. So this justifies the common claim that PCA works for Gaussian data.

Given the good results in [4] (mentioned in §1.2), it seems that the three assumptions above are at least approximately satisfied. But there was no 'a priori' reason to think that this would be the case. For instance, the genotype data is clearly not Gaussian distributed, but Binomially distributed⁴.

When dealing with rare variants only, things get worse: it was shown in [3] that, if the genotype data uses rarer alleles, then: the ratio inter-population-variance diminishes; the distance between populations decreases; and the variance explained by the first PCs also decreases.

Hence, there is no reason to think PCA is useful when dealing with the exome data that we use for the study of ALS (and indeed my supervisor's research confirms that things are not so easy in this case).

Possible alternatives

One may start exploring PCA alternatives that fit our needs with the question: how to extract the population stratification in the OTGP data with rare variants only? The answer may suggest methods to use in our own rare variant data.

Fortunately this was already answered in [6]. Two methods stand out: Logistic PCA (LPCA) and Jaccard PCA (jPCA), with LPCA giving slightly better results.

LPCA rests on the assumption that the data follows a binomial distribution. Since at the start of this project the main goal was to control for population stratification in the gene-burden version of the data, which is not binomially distributed, then I decided to explore jPCA, and apply it both to the gene-burden and the genotype versions of the data.

2.2 Jaccard (Kernel) PCA

Before discussing jPCA one must introduce the Jaccard index.

Jaccard index

Given two genotype vectors (two rows of a genotype matrix) a and b, their $Jaccard\ index\ j(a,b)$ is schematically given by

$$j(a,b) = \frac{|(a_{>0} \cap b_{>0})|}{|(a_{>0} \cup b_{>0})|} \tag{1}$$

where $|(a_{>0} \cap b_{>0})|$ is the number of SNPs where a and b have equal, non-zero genotype values, and $|(a_{>0} \cup b_{>0})|$

²Missing values were replaced with the average of their columns.

³In the current context of controlling for population stratification, "important directions" are the ones which capture the ancestry differences.

⁴The other type of data that we use (gene-burdens) has all the same issues. In particular, it is straightforward to check the distribution of the gene-burden values is not Gaussian (although it is also not Binomial).

is the number of SNPs where either a or b (or both) have non-zero genotype values.

The motivation for defining such an index is the following: when comparing two genotype vectors, a first naive idea is to define their similarity index as the ratio of entries on which they agree. Then, a = (0,0,0,1,0) and b = (2,0,0,0,0) would have a similarity index of $\frac{3}{5}$.

One can see how this reasoning can go terribly wrong when dealing with rare mutations. In fact, one may get vectors a and b with zeros in thousands of entries, and just a couple of non-zero values in different SNPs. Then, the naive similarity index will be extremely high - which for our purposes is nonsensical: a and b should be considered identical only when they have mutations in the same SNPs. The Jaccard index takes this into account by only considering the entries with positive values.

The Jaccard matrix is simply the similarity matrix $J := [j(a_k, a_l)]_{k,l \in \{1...n\}}$, where n is the number of samples/individuals.

Notice that we defined the Jaccard index in a way that generalizes easily to gene-burdens, since nothing stops the vectors' entries of being bigger than 2. This contrasts with e.g. [1], where the author only accounts for burden values in $\{0,1\}$. When clarification is necessary, I will refer to (1) as the generalized Jaccard index.

Kernel PCA

jPCA is sometimes described as "applying PCA on the Jaccard matrix" but this is slightly misleading (see §5). Actually, it is a particular type of the so-called Kernel PCA, which we introduce now.

Kernel PCA (kPCA) is a non-linear dimensionality reduction method where the data is non-linearly mapped to a higher-dimensional space (feature space) where PCA is finnally applied [7][8]. It can be especially useful when trying to separate data that is not separable using standard PCA.

To illustrate kPCA, consider⁵a dataset consisting of two concentric circles (plus noise) - see figure 2. Clearly, PCA would approximately give the original axes as the first two principal components. Hence, because of the non-linearity of the data, PCA is incapable of capturing its structure.

Instead, we can apply a non-linear map $\phi \colon \mathbb{R}^2 \to \mathbb{R}^3$ given by $\phi(x,y) = (x,y,x^2+y^2)$. This captures the structure of the data in the third dimension: the smaller dots from the inner circle will be separated from the bigger dots from the outer circle along the third dimension, since it measures the square of the distance of the original points to the origin.

PCA can now handle the new data and separate the two clusters along one of the first principal components. When applying PCA on a dataset X, one obtains the PCs from X^tX : they are the eigenvectors of X^tX . So in our

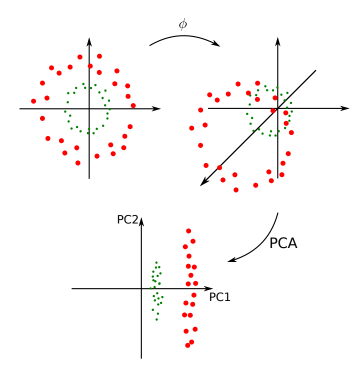


Figure 2: Scheme of kPCA applied to a dataset of concentric circles.

case one must find the eigenvectors of $\phi(X)^t \phi(X)$: they are the kPCA PCs.

But this computation can be expensive. If we instead find a function $k \colon \mathbb{R}^2 \times \mathbb{R}^2 \to \mathbb{R}$ such that ${}^6\forall x,y,\ k(x,y) = \phi(x)^t\phi(y)$, then $\phi(X)^t\phi(X) = [k(x_i,x_j)]_{i,j=1,\dots,n} =: K$, where n is the number of samples in the dataset. Hence the kPCA PCs are the eigenvectors of K, and if we have K from the start we never have to deal with the non-linear function ϕ .

Such a function k is called a kernel. The matrix K is sometimes called the $Gram\ matrix$.

In practice, one rarely chooses an explicit form of ϕ , and instead deals with kernels from the start. One must be careful when choosing a function as a kernel: we cannot just use any function - it must be one for which a ϕ exists such that $\forall x, y, k(x, y) = \phi(x)^t \phi(y)$. Fortunately, one *can* be sure that a certain function k is a kernel if it is "positive definite" as discussed in [9].

The next natural question is then: what kernel should we choose for the genotype and gene-burden datasets?

Jaccard PCA

Jaccard PCA (jPCA) is the particular kPCA method for which the kernel k is taken to be the Jaccard index j [1].

It turns out that the Jaccard index is positive definite [9], meaning that we can indeed use it as a kernel.

⁵This example is similar to the one given in the excellent video lecture on kPCA by David R. Thompson, from Caltech.

⁶In other words, k acts as a dot product in \mathbb{R}^3 .

3 Results

I ran my own implementation of jPCA on the datasets described in §2, and tested it on the OTGP data as a control, comparing the results with [1].

3.1 jPCA on the 1000 Genomes Project data

In [1], jPCA was successfully applied to the OTGP data to extract population structure, obtaining better results than PCA when applied to rare variants. This paper uses an open-source R library called *jacpop*, which computes the jPCA principal components from a genotype matrix using the function *generate_pw_jaccard*.

Unfortunately, this library was unfit for my work, because:

- 1. The dbGaP dataset was too large to be fed to generate_pw_jaccard: I needed a flexible script that I could parallelize.
- 2. generate_pw_jaccard uses the standard Jaccard index, not the generalized Jaccard index, and is thus unfit for gene-burden data (the LOF and moderate datasets).
- 3. I prefer to work in Python.

So I created a collection of Python scripts that generalized *jacpop* for genotype data by allowing parallelization, and another collection of Python scripts that generalized *jacpop* for gene-burden data by allowing burdens of arbitrary positive value. Clearly, the former should still give the same results as *jacpop* in the OTGP data, so I used both methods using 5e5 randomly chosen variants and verified that they output the identical plots (see figure 3).

3.2 Gene burdens

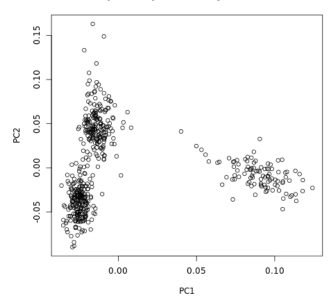
For this part I used the moderate data. I did not round the data to avoid spurious similarities between individuals with the same missing variants (see the appendix §5.1).

The plots obtained by applying jPCA to the geneburden data are not encouraging: in contrast with the results in [1] and [6], the first two PCs do not show a clear separation between the different populations (figure 4).

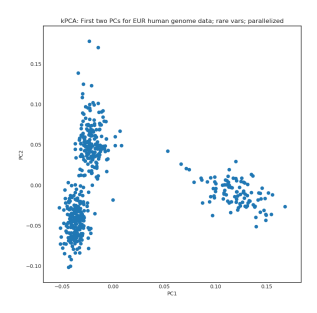
Another surprising result is that the jPCA PCs do not correlate at all with the PCs from PCA applied to the genotype data. Since the latter already capture the population structure fairly well, this confirms our suspicion that jPCA is not able to do it using the gene-burden, rare variants exome data.

We may also be interested in knowing if the PCs separate sick and healthy people. They don't: there's no visible separation (see figure⁷ 5) and, more rigorously, running logistic regression gave terrible results - very large p-values - and the model simply predicted 0 every time (thus

Principal components on Jaccard matrix



(a) Plot from $generate_pw_jaccard$ using 5e5 variants from the OTGP data.



(b) Plot from my (parallelized) jPCA script using 5e5 variants from the OTGP data.

Figure 3: Comparison between plots generated by *generate_pw_jaccard* and by my script when applied to OTGP data.

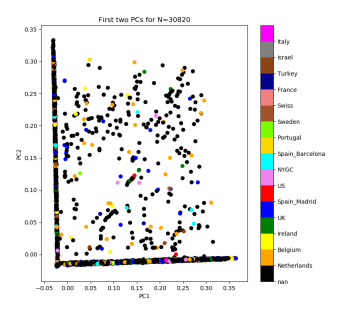


Figure 4: First two jPCA PCs in the Gene-burden data, labelled by cohort.

having an accuracy of $\frac{\text{number of healthy individuals}}{\text{number of individuals}} = 77\%$ and null F1-statistic.

This is worse than when using the PCs from PCA: the first 5 PCs are significant (with nearly null p-values) and the F1-statistic is nonzero (albeit still unsatisfactory with a value of 0.02).

So it is clear that jPCA does not work with geneburden, rare variant exome data. Let's try it on the genotype data.

3.3 Genotype values - rare variants

We now apply jPCA on the dbGaP data, including only the rare variants.

The results are still unsatisfactory, with no clear separation between cohorts (figures 6 and 7). Furthermore, logistic regression (with the phenotype as the response variable) gives equally bad results as in the gene-burden case, and again the jPCA PCs do not correlate with the standard PCs.

These results are disappointing. They are also surprisingly, since jPCA works so well in the OTGP data, with both common and rare variants. But sure jPCA must at least give good results when using common variants? The answer to this question may help understand why jPCA is failing.

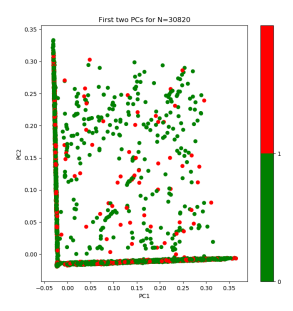


Figure 5: First two jPCA PCs in the Gene-burden data, labelled by phenotype.

3.4 Genotype values - common variants

We again apply jPCA on the dbGaP data, now including the common variants.

The results are more interesting: the first two PCs separate the data in two (see figures 8 and 9). Contrary to our expectations, these two clusters do not have an obvious association with the cohort labels.

On the other hand, logistic regression with phenotype as the response variable is as bad as before, and again the iPCA PCs do not correlate with the standard PCs.

The natural question to ask is: why does jPCA capture the population structure in the OTGP data (for both rare and common variants) but not in the genotype data that we are using (for both rare and common variants)?

To understand this, one may start by looking at the differences between these datasets: although they both consist of genotype data, the OTGP data contains the SNPs of the entire genome of the population, while the db-GaP data only contains the exome SNPs. Furthermore, while the dbGaP data that we are using is an amalgamation of data from different countries and laboratories, the OTGP data comes from a single source sequencing project, meaning that the dbGaP data is a lot less uniform than the OTGP data. This messiness can throw off the results. For example, the two clusters in figure 8 may very well be an artifact of mean-imputation, similar to what happens in §5.1. This is clear by comparing figures 10 and 12a in light of the discussion in §5.1.

From these results alone it is not possible to conclude which of these factors is causing the failure of jPCA.

⁷You may notice that the two plots from figures 4 and 5 are slightly different. This is because in the first plot I removed the points whose cohort label was "nan".

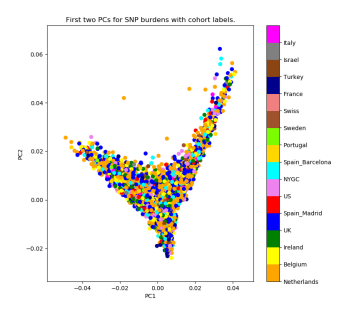


Figure 6: First two jPCA PCs in the genotype data, labelled by cohort.

4 Conclusion

The fact that jPCA fails on moderate (gene-burden) data is not very surprising: it was something quite different from what was tried before. Even the failure of jPCA on the rare variants of the dbGaP genotype data is understandable: as discussed with my supervisor, using all rare variants is probably a bad idea, since there are plenty of super rare variants occurring on only one or two individuals each, which mess up the selection of PCs. So further filtration of the data could solve this.

The true mystery is the failure of jPCA in capturing population structure in the dbGaP genotype data using the common variants. Here, such filtration problems are inexistent, and there seems to be no good reason for jPCA to fails, given that it works on the OTGP data [1][6].

The issue must come from the technical differences between the dbGaP genotype data and the OTGP data, namely: the dbGaP data we used contains exome-sequencing only, which may have a different distribution than the genome-sequencing data; the OTGP data is more uniform - the dbGaP data comes from dozens of different projects, and for example the imputation methods that were used to cope with the missing data may create spurious issues.

A diagnose would involve trying jPCA with the entire genotype data (not just the exome), as well as a close look into the origins of the two clusters on figure 8. It may be also interesting to re-run jPCA on the rare-variants, genotype data after filtering out the extremely rare mutations. These are left for future work, since my internship has come to an end.

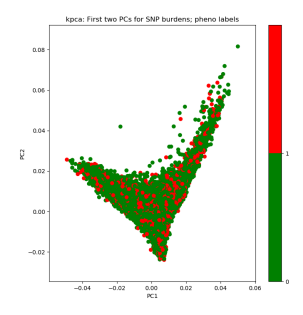


Figure 7: First two jPCA PCs in the genotype data, labelled by phenotype.

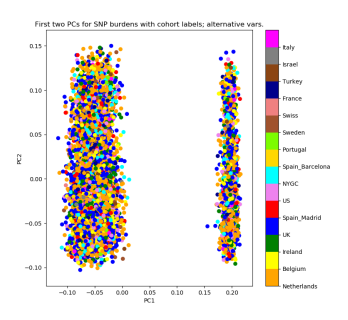


Figure 8: First two jPCA PCs in the genotype data, labelled by cohort.

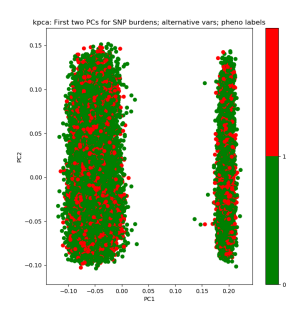


Figure 9: First two jPCA PCs in the genotype data, labelled by phenotype.

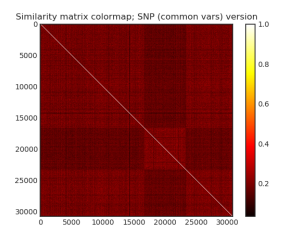


Figure 10: Heatmap for Jaccard scores from the genotype data - common variants.

5 Appendix

5.1 PCA on Jaccard values

In [6] and [1], yhey describe jPCA as applying PCA on the similarity matrix created using the so-called Jaccard index as the similarity value. This is not exactly what jPCA is, as discussed in the main text. But their abuse of terminology suggested a different approach: why not trying PCA on the Jaccard values themselves? After all, the geneburden data may be approximately Gaussian distributed: each gene-burden is the sum of a large number of genotype values (each binomially distributed), and there may be enough consistency between binomail distributions and genes that the central limit theorem suggests an approximate Gaussianity. It was a long shot, but it was worth a try.

I will not write the details here, because they are now irrelevant, but one important lesson came from this exploration: how rounding mean-inputed data can give spurious results. I will illustrate this with the problems that appeared when applying PCA on the Jaccard matrix from the LOF data, but the moderate data case was similar.

Building the Gram matrix from the LOF data one sees that, as we feared, the data Jaccard values are very low and mostly zero or very close to zero (figure 11).

Looking at the LOF data, one may notice that many values are not integers. This is due to the many meaninputed values in the dataset. Because of the way Jaccard values are computed, we will get zeros even if the individuals have similar (but non-integer) gene-burden values.

A natural solution is to round these values. This gives interesting results (figure 12), although still heavily skewed towards zero.

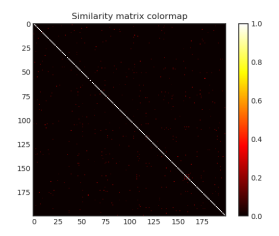
I spent some time zooming in and trying to understand the origins of the non-zero peaks of the histogram.

One may realize that something odd is going on by looking at the well-defined yellow and red squares in the heatmap. What are the chances that the similar individuals are all in the same region of the dataset? It's much more likely that these apparently interesting results are actually an artifact of mean-imputation followed by rounding the results: many of the individuals with close index numbers are from the same cohort, and thus have missing values in many of the same SNPs, so that mean-imputation + rounding effectively turns two individuals from the same cohort very similar under the Jaccard index.

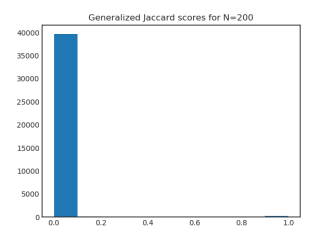
This is the danger of using mean-inputation (with data with many missing values and/or very rare nonzero values) together with the Jaccard index.

5.2 Glossary

Allele: a variant form of a given gene, meaning it is one of two or more versions of a gene. It can also refer to a region of interest in the genome. In this last sense,

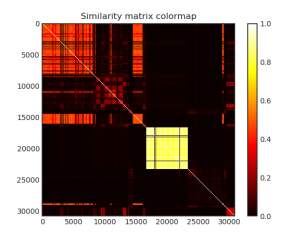


(a) Heatmap for Jaccard scores using 200 individuals.

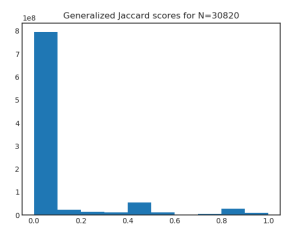


(b) Histogram for Jaccard scores using 200 individuals.

Figure 11: Jaccard scores for the LOF data, using 200 individuals.



(a) Heatmap for Jaccard scores using 30820 individuals.



(b) Histogram for Jaccard scores using 30820 individuals.

Figure 12: Jaccard scores for the LOF rounded data, using 30820 individuals.

alleles can come in different extremes of size. At the lowest possible end one can be the single base choice of a single nucleotide polymorphism (SNP). At the higher end, it can be the sequence variations for the regions of the genome that code for the same protein which can be up to several thousand base-pairs long.

Common variant: SNP variant/allele with an allele frequency smaller than 0.5 in the sample (hence any minor SNP allele is a common variant).

Deoxyribonucleic acid (DNA): is a molecule composed of two polynucleotide chains that coil around each other to form a double helix, being connected by hydrogen bonds at their nucleotides. Its nucleotides are: A, T, C, G.

Gene: a sequence of nucleotides in DNA or RNA that encodes the synthesis of a genetic product, either RNA or protein.

Gene expression: the process by which information from a gene is used in the synthesis of a functional gene product. These products are often proteins, but in non-protein coding genes such as transfer RNA (tRNA) or small nuclear RNA (snRNA) genes, the product is a functional RNA.

Genome: Sum total of an organism's DNA.

Genome-wide association study (GWAS): observational study of a genome-wide set of genetic variants in different individuals to see if any of those variants are associated with a phenotype. Typically, the variants are simply SNPs and the phenotype is the presence or absence of a (human) disease. In more detail: GWA studies compare the DNA of participants with varying phenotypes for a particular trait/disease. The cases are the individuals with the disease, and the controls are the ones without (this is the traditional approach, called phenotype-first). The DNA of every subject is read using snip arrays, letting us know what allele occurs in each person. If one allele is more frequent in cases than in controls, it is said to be associated with the disease.

Nucleotides: molecules consisting of a nucleoside (five-carbon sugar ribose + nitrognous base) and a phosphate group. They are the basic building blocks of DNA and RNA.

Population Structure: A population has structure when there are large-scale systematic differences in ancestry and/or groups of individuals with more recent shared ancestors than one would expect in a randomly mating population.

Rare variant: SNP variant/allele with an allele frequency smaller than 0.01 in the sample.

Variant: An alteration in the most common DNA nucleotide sequence. The term variant can be used to describe an alteration that may be benign, pathogenic, or of unknown significance. The term variant is increasingly being used in place of the term mutation.

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