Significance and Background

A large proportion of common diseases (i.e. prevalence > (XX) % are genetically complex. In contrast to Mendelian diseases, genetic causation of complex diseases is not straightforward, but rather genetic risk for any given disease is spread across a large number of genes, each individually having a relatively minor effect. While successful efforts to map the genetics of Mendelian diseases date well back into the 20^{th} century, only in the past decade with the maturation of the genome wide association study (GWAS) approach has some understanding of the genetic basis of complex disease begun to emerge. 1,? It's become clear now that estimates of heritability based on the twin study design were broadly accurate (if slightly overestimated in some cases), and that the heritabilities of common complex diseases are generally high. GWAS and related approaches have also shown that a substantial proportion of the variance in risk can be attributed to a very large number of common alleles (generally each with small effects), 2,3,4,5 while sequencing and exome studies indicate a potential role for rare variants of large effect as well. 6,7,8 It seems clear now that thousands or perhaps even tens of thousands of individual genetic variants play a role in determining susceptibility to any given complex disease. Less clear however is the issue of why complex diseases are as common as they are, and what forces are responsible for the particular relationships observed between effect size and allele frequency (i.e. the genetic architecture). In contrast to Mendelian diseases, where we have a fairly good idea of how factors such as mutation rate, recessivity, selection⁹ and demography? combine to make these diseases common, we remain for the most part in the dark with respect to the processes that underlying varition in complex disease burden. Certainly these factors are at play here as well, but at present we lack any quantitative framework within which to address their relative contributions.

(need to introduce idea of variation in prevalence and architecture as tool to greater understanding)

Nonetheless, several qualitative mechanisms have been proposed which may potentially explain the observed patterns. Perhaps the most straightforward is that the present distribution of genetic risk reflects a simple balance between mutation and selection.¹⁰ The prevalence of common disease may therefore be ascribed to relatively large mutational target sizes for many diseases. Variation in the prevalence of diseases with similar fitness costs may then simply be down to difference in their mutational target sizes, and variation in genetic architectures may arise chiefly from differences in the distribution of mutational effect sizes. Conversely, variation in the fitness costs of specific diseases will lead to differences in the strength of direct selection against risk incresing variants associated with those diseases, which should account for some amount of variation in both architecture and prevalence. The direct selection experienced by a given allele is likely only part of the story, however, as the extreme polygenicity of complex diseases indicate that pleiotropy must necessarily be extensive, 11, 12 and indirect selection due to these pleiotropic relationships may therefore by play a major role in determining architecture and prevalence. This may be true in steady-state scenarios (wondering if problematic to drop "steady state" here without any background on what I'm talking about), where mutations which increase disease risk may have protective effects on other diseases influence other beneficial phenotypes, thus raising the frequency of the allele (i.e. altering architecture) and increasing prevalence, or under more dynamic hypotheses, where recent positive selection on a trait genetically correlated to disease has resulted in an increase in prevalence above what one would expect under an equilibrium mutation-selection balance scenario, as has been argued for multiple diseases with limited convincing conclusion. 13, 14, 15, 16, 17, 18 Alternative explanations invoke the recent and profound changes in the human environment (diet, lifestyle, etc.) and the possibility that the mismatch between the ancestral human environment and present conditions that have resulted in a large increase in disease incidence. 19,20 Such was the original basis for the "thrifty genotype" hypothesis, 21,22 and subsequent observation of large scale oscilations in type 2 diabetes incidence in response to food shortage and economic crisis and subsequent recovery in Cuba in the 1990s²³ indicate that this effect can be profound indeed. While some of these ideas likely apply to some diseases (and they are also not mutually exclusive), more often than not they lack quantitative grounding and predictions that allow for explicit inference and model comparisson.

Each of these explanations are ultimately statements about population genetic processes such as the rates at which disease causing mutations arise and their ultimate effect on fitness, and how these processes have interacted with one another over the course of human evolutionary history to yield the currently observed genetic architectures and disease prevalences. While some models relating population genetic processes with the kind of data emerging from GWAS has been proposed and have been useful in grounding discussions surrounding complex disease architecture, they generally rely on *ad hoc* assumptions that severely limit our ability to draw

reliable inferences about complex disease that are easily interpretable. Nonetheless, two studies in particular have been especially influential. Pritchard, in 2001,²⁴ considered a model in which the effect on disease risk is completely uncoupled from their fitness cost, i.e. an extreme pleiotropy limit. While this paper was enormously influential in grounding the debate surrounding the common disease-common variant hypothesis in population genetic theory, 25 it seems unrealistic that a mutation's effect on disease risk should not have any consequences for its fitness. A second study which has been influential is that of Evre-Walker in 2010.²⁶ who posited that all mutations are deleterious a priori, but arbitrarily assumed a relationship between effect size and selection coefficient of the form $\alpha = \delta S^{\tau}(1+\epsilon)$, where S is the selection coefficient, δ is a randomly chosen sign (i.e. 1 or -1), ϵ is a normally distributed noise term, and τ is a "coupling parameter" meant to capture the effect of pleiotropy: when equal to 1 the effect on the disease and fitness are closely linked, and when equal to zero the model reduces to the extreme pleiotropy limit of Pritchard.²⁴ While this model has seen empirical application (see Specific Aim 2 below for more background on the Eyre-Walker model in this context), it does not have any obvious theoretical justification. While these studies have both been extremely influential, it seems worth reemphasizing the fact that neither posesses any concept of fitness surface relating an explicit disease phenotype to natural selection, and in light of our rapidly accumulating knowledge of the genetic architecture of complex disease, a fresh take on the problem seems due.

Here I propose to develope generative models for the way that genetic architecture and disease prevalence will be affected by evolutionary parameters such as mutation, natural selection, pleiotropy and demography. I will develop these models into statistical inference approaches which take advantage of the rich information present in GWAS data to infer the underlying parameters which govern the evolution of complex disease genetic architecture. Lastly, I will apply the results of my inferences to a problem currently of particular importance in human complex disease genetics, which is the issue of the extent to which polygenic predictors or disease risk trained in one population will be effective in predicting risk in evolutionarily diverge populations, with a particular focus on how the genetic architecture of the disease will affect prediction accuracy.

Approach

Preliminary Results (possibly will take your suggestion to ease into this section even more gently, but leaving for now in order to get more work done elsewhere) Our simplest model and point of departure considers the impact of mutation and selection on a single disease in a constant environment. In this model, each individual's risk of developing disease is a non-linear transform of an underlying (and generally unobserved) disease liability trait, such that

$$R = \ell(Z), \qquad Z = \sum_{i} \alpha_{i} g_{i} + \epsilon$$
 (1)

where R is risk, Z is liability, α_i and g_i are liability scale effect size and genotype at site i, and ϵ is a normally distributed deviate which captures the contribution of stochastic variation in risk among genotypes with the same liability, (i.e. the "environment" of classical quantitative genetics). ℓ is the non-linear link which transforms liability onto the risk scale. Previous work^{27,28} suggests that the exact choice of ℓ is unlikely to be particularly important, so long as it meets a few simple constraints such as monotonicity and a transition from low to high risk that is relatively sharp compared to the variance of liability in the population. Our preliminary investigations (omitted due to space constraints) support this conclusion, and as such I will let ℓ be the probit link function of the liability threshold model for the remainder of this proposal, although other popular choices such as the logit link of the logistic odds model or the truncated exponential of the constrained version of Risch's mulitiplicative model^{28,29} would be equally suitable, and both of these models will be explored alongisde the liability threshold model in the proposed work.

The number, frequencies and effect sizes of the sites contributing to variation in liability arise from population genetics processes. Liability increasing and decreasing mutations arise according to an infinite sites model with free recombination among sites at genome wide rates μ^+ and μ^- , with effect size distributions $f^+(\alpha)$ and $f^-(\alpha)$ respectively. Individuals with the disease have a reduced fitness of 1-S (disease free individuals have fitness 1).

The genetic architecture of the disease under this simple model is shaped by mutation-selection-drift balance,

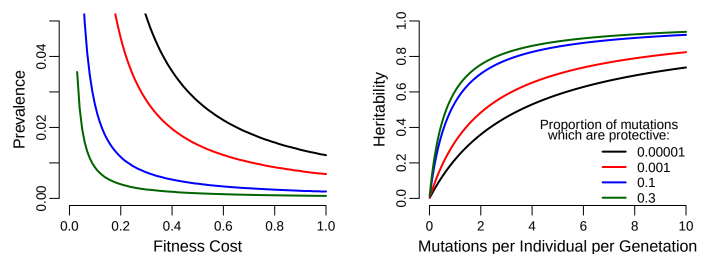


Figure 1: Caption goes here

and can be related to standard results from quantitative genetics. The steady state is reached when

$$U^{+} = U^{-} + V_{A} \underbrace{S\phi\left(\Phi^{-1}\left(1 - P\right)\right)}_{\text{selection gradient}} \tag{2}$$

where $U^+ = \mu^+ \int_0^\infty \alpha f^+ (\alpha) \, \mathrm{d}\alpha$ is the total per generation mutational increase in liability (with U^- defined similarly as mutational pressure toward decreased liability). The final term accounts for the selection pressure toward lower liability; V_A is the additive genetic variance of liability, P is the disease prevalence, and ϕ and Φ are the Gaussian pdf and cdf repsectively. The compound term multiplying V_A is a selection gradient in the standard quantitative genetics sense, and is a simple generalization of the gradient exerted by truncation selection to cases where the "truncated" individuals have fitness greater than zero.?

The steady state requirement allows us to derive expressions for any summary of genetic architecture. While the mutation rates, effect size distributions, and fitness cost of the disease are biological inputs, disease prevalence (P) and genetic variance (V_A) are dependent variables which evolve as part of the system. The additive genetic variance, as a product of the genetic architecture, depends on how the frequencies of individual alleles evolve, which depends on the individual selection coefficients they experience. An individual allele with effect size α on disease liability will experience a selection coefficient

$$s = -2\alpha S\phi\left(\Phi^{-1}\left(1 - P\right)\right) \tag{3}$$

against the liability increasing homozygote, and will evolve under fitness additivity so long as α is small. We can then apply results from diffusion theory on the frequency spectrum of an allele conditional on its selection coefficient^{30,31,?} to find the genetic architecture and disease prevalence at equilibrium. Given a solution to the model, we can derive

Specific Aim 1: Relating Population Genetic Processes with the Architecture of Complex Diseases

I will generalize the basic model from our preliminary results to include the impacts of pleiotropy and environmental and demographic change. In each case, I will solve these generalized models to obtain analytical expressions for the genetic architecture an disease prevalence. In each case, my results will be checked by comparisson to simulations, and in cases where analytical solutions are not tractable, results will be pursued directly via numerical and simulation based methods.

Environmental Change

The most straightforward way to incorporate environmental change may also be the most relevant to human disease. The simplest environmental change models are those in which the mean or the variance of the environmental component of the phenotype shifts or increases suddenly. Given the recent and rapid changes to human

environments (e.g. diet and lifestyle on type 2 diabetes prevalence), we are most interested in the scenario where an environmental shift has just occurred, but there has not been sufficient time for allele frequencies to evolve away from their previous equilibria. In the case of a shift in the enironmental mean, the effect is simply an increase in prevalence such that $P_{new} = 1 - \Phi\left(\Phi^{-1}\left(1 - P_{old}\right) - \delta\right)$, where δ is the shift in the environmental contribution to liability measured in units of the phenotypic standard deviation (Φ , again, is the Gaussian cdf).

The result of a change in the environmental variance are slightly more complex, as it impacts both the prevalence and the heritability of the disease. The most straightforward impact is on heritability. If the environmental variance is increased by an amount ψ , then heritability is decreased such that $h_{g,new}^2 = \frac{h_{g,old}^2}{1+\psi}$ (where again, ψ is given here in units of the pre-change phenotypic variance). Prevalence, on the other hand, will increase to $P_{new} = 1 - \Phi\left(\frac{\Phi^{-1}(1-P_{old})}{1+\psi}\right)$. In both of these simple environmental change scenarios, the a given mutation's effect on liability is unchanged. The effect on risk, however, is increased, as a given mutation's contribution to risk conditional on its contribution to liability also depends on the prevalence of the disease (expressions omitted due to space). The result is that the while the additive genetic variance for liability is unchanged under either scenario, we expect that on the *risk* scale it will be increased. A shift in the mean of the environmental contribution should actually cause an increase in heritability on the risk scale, while the impact of an increase in the variance of the environmental contribution on the heritability of risk are not immediately obvious, and may be architecture dependent.

In addition to these simple scenarios, I will also use simulations to study how less recent changes to the environment (e.g. following the Out of Africa event) might have impacted genetic architecture.

Pleiotropy As noted in the background above, mounting evidence suggests that genetic variation is often highly pleiotropic. We will consider pleiotropic effects of two kinds. The first includes pleiotropic effects on other disease traits. Pleiotropy of this form will generally act to modulate the additive selection coefficient felt by a given allele. Liability increasing mutations which have protective effects on other diseases will have less strongly negative selection coefficients than in the preliminary model (or may even be selected for), while those which increase liability for multiple diseases will incur additional selective cost beyond that due to direct selection on the focal disease.

The second form of pleiotropy considered will be due to stabilizing selection on continuously distributed (i.e. non-disease) quantitative traits. In models of stabilizing selection on quantitative traits at equilibrium, there is no directional component to the selection felt by individual alleles. Rather, due to variance reducing selection for individuals to cluster near the optimum, individual alleles experience symetric underdominance with respect to fitness (i.e. the minor allele is always selected against), where the strength of selection depends on the effect size of the mutation on the quantitative trait relative to the strength of stabilizing selection.³²

This observation suggests an interesting relationship which will form the foundation for our modeling of pleiotropy. When a mutation effects only disease, selection is directional and additive, whereas when a mutation effects only quantitative traits, selection is underdominant. Mutations which have large effects on disease and smaller impacts on quantitative traits will experience directional selection with the disease causing mutation being partially dominant for fitness, while mutations which have large effects on quantitative traits and smaller effects on disease will be underdominant for fitness, but asymetrically so, with the liability increasing homozygote being less fit than the liability decreasing homozygote.

The major objective here is to derive an expression for joint distribution of effect size and allele frequency (i.e. the genetic architecture) as a function of the parameters (Θ_A) which describe the nature of pleiotropy impacting the disease architecture. These two quantities are conditionally independent of one another given the specification of the selection coefficient(s), which suggests a tractable approach for theoretical analysis and inference (aim 2). The expression for the genetic architecture can be written as an integral over the selection coefficients

$$p(\alpha, x \mid \Theta_A) = \int \int p(\alpha \mid s, h, \Theta_A) p(x \mid s, h) p(s, h) \, ds dh. \tag{4}$$

The expression for the distribution of allele frequencies $(p(x \mid s, h))$ can be computed from standard diffusion theory,? and this fact can be leveraged in an inference context (aim 2) to learn the distribution of the selection coefficients (p(s, h)) directly from the data (alternately, plausible distribution can be specified in a theoretical context to understand how differences in the distribution of selection coefficients impact architecture). This leaves specifying

the distribution of effect sizes for a given set of selection coefficients and under a given set of parameters governing the effects of pleiotropy $(p(\alpha \mid s, h, \Theta_A))$ as the primary theoretical task.

I will approach this problem first by considering simple isotropic models of pleiotropy in which mutations may affect more than one disease/trait, but there is no inherent mutational covariance among diseases or traits. In particulary, I will investigate how the number of pleiotropically related diseases/traits, the strength of selection on them, and the degree of functional overlap among diseases/traits all impact the genetic architecture of disease.

Non-Equilibrium Demography It is clear from population genetic work over the last decade and a half that demographic events such as the Out of Africa bottleneck and recent explosive population growth have had a significant impact on allele frequencies and therefore potentially on genetic architecture. Recent work from Dr Sella's lab,³³ among others (cite others) suggests that both the bottleneck and recent growth may have impacted genetic architecture, though the relative importance of each depends on the (as yet unknown) selection coefficients of disease alleles. However, what most work to date on this problem has been done in the context of a single site in a vaccum with a fixed selection coefficient, with no explicit disease phenotype, and therefore no model of the relationship between effect size and selection coefficient.

I will use simulation based approaches to study how the particular course of human demographic history has impacted the evolution of complex disease architecture and prevalence. Population size changes impact architecture through their modulation of the relationship between genetic drift and natural selection. All else being equal, genetic architectures in larger populations will be composed of more rare alleles, due to the increased efficiency of selection, though the precise impact depends on the details of the model. My investigation of the preliminary model also suggests that disease prevalence at equilibrium is fairly sensitive to population size. This is significant, specifically because the selection coefficient experienced by a particular allele depends on the disease prevalence. If prevalence evolves over time in response to changes in population size, then selection coefficients will change over time as well.

Specific Aim 2: Inference of Model Parameters from Complex Disease GWAS

Background The existing work which comes closest to that proposed in this aim is recent work applying the Eyre-Walker model (discussed in intro above) to investigate the genetic architecture of complex disease. One such line of work, which includes two studies from David Altshuler's group and colleagues, simulates disease architectures under the Eyre-Walker model, and then compares the number causal type 2 diabetes variants discovered by a given study design to that which would be expected under the Eyre-Walker model simulations, with a particular focus on inferring the total mutational target size and the pleiotropy parameter τ , which captures the coupling between fitness and effect size. These studies used the number of genome wide significant variants 34 , 35 as summary statistics in an approximate Bayesian computation (ABC) approach. The initial study, 34 which simply used the total number of variants discovered was able to eliminate the pleiotropic extreme (i.e. the Pritchard model) and a model in which fitness and effect are tightly coupled, but could say little beyond that. A more recent application 35 used the number of significant variants with minor allele frequency (MAF) < 5%, as a summary statistic, and was able to show that relatively small values of τ in the range of 0.1 (i.e. relatively weak coupling between effect size and fitness) are most consistent with the genetic architecture of type 2 diabetes.

Another study in this vain is that of Mancuso et al (2015),? who used the Eyre-Walker model to investigate the genetic architecture of prostate cancer susceptibility. Rather than using signficant hits as a summary statistic, these authors used variance partitioning methods 36 to estimate the proportion of the total genetic variance that is attributable to rare alleles (0.01% > MAF > 1%), and then took this quantity as the ABC summary statistc. The result was support for moderate values of of $\tau \approx 0.5$.

The results of these studies highlight a couple of important facts. One is the difficulty of interpretting the meaning of the τ parameter in the Eyre-Walker model. While it is clear that lower values of τ represent a weaker coupling between fitness and effect size, it remains unclear how to interpret this parameter biologically, as it was not derived from any explicit model of the impact of pleiotropic traits on the focal trait. The Eyre-Walker model is also fundamental not extensible to inference with multiple traits together, which we expect to particularly important in the future as multi-trait GWAS methods continue to mature. $^{?,?}$

One encouraging takeaway is the fact that inference approaches which use information about the frequencies of alleles discovered in GWAS^{2,35} obtained more precise parameter estimates than the one that did not,³⁴ but

it is worth stating that all of these studies leave substantial amounts of the information contained in GWAS data unused, indicating that there should be substantial room for improvement. Finally, it should be noted that all of these studies rely on the assumption that the distribution of selection coefficients experienced by disease associated alleles resembles the gamma distribution often assumed for non-synonymous mutations³⁷ (though perhaps with little real justification³⁸), rather than estimating the distribution of selection coefficients from the data directly. It is therefore clear that there is a need for an inference approach which A) uses the totality of the evidence available in GWAS data, and B) allows for the estimation of parameters which have clear biological interpretations, which is the focus of this aim.

Approach I will develop an inference approach which leverages GWAS data to infer the parameters of the models developed in Aim 1 for a number of disease GWAS datasets. The approach is based on the Poisson Random Field and related composite likelihood techniques which have been used extensively in population genetic inference. Given 1) the K genome wide significant variants discovered in a GWAS, 2) estimates of their effect sizes and allele frequencies (α_i, x_i) , 3) an estimate of the heritable variation $(V_{G,b}^*)$ attributable to each of B minor allele frequency bins, and 4) the parameters of the study Θ_S (i.e. the number of cases and controls and the disease prevalence assumed for the GWAS), the likelihood of the evolutionary parameters underlying the architecture (Θ_A) can be written as

$$L\left(\Theta_{A} \mid \{(\alpha_{i}, x_{i})\}_{i=1}^{K}, \{V_{G,b}^{*}\}_{b=1}^{B}, \Theta_{S}\right)$$

$$= Pr\left(K \mid \Theta_{A}, \Theta_{S}\right) \left(\prod_{i=1}^{K} Pr\left(\alpha_{i}, x_{i} \mid \Theta_{A}\right) H\left(\alpha_{i}, x_{i} \mid \Theta_{S}\right)\right) \prod_{b} Pr\left(V_{G,b}^{*} \mid \Theta_{A}, \Theta_{S}\right)$$

The first term gives the probability of observing K genome wide significant variants associated with the disease (which is Poisson distributed). The second term gives the probability density of a variant with a given effect size and frequency, while the third term gives the power to detect such a variant as a function of the study parameters. The final term gives the probability that a given proportion of the heritability not captured by the K genome wide significant variants is apportioned to a given minor allele frequency bin. The first, second, and fourth terms all depend on the evolutionary parameters (Θ_A) and therefore are the major link which connects Aim 1 and Aim 2, as the the theory from Aim 1 will be used to compute these expressions.

Intuitively, the number of genome wide significant variants mostly provides information about the mutational target size, while their frequencies are informative about the strength of selection they experience, and the relationship between frequency and effect size is therefore informative about the nature of the pleiotropic effects of disease associated loci.

The proportion of the "missing heritability" attributable to different minor allele frequency bins essentially contains information about the mean strength of selection experienced by disease variants, and therefore helps constrain the range of possible models which can fit the data. It is a less rich source of information than the individual genome-wide significant variants themselves, as we do not get to observe informative features such as whether it is the derived or ancestral allele which increases disease risk, and we must fold the frequency spectrum, which in particular discards information about protective mutations. Nevertheless, the theory from Aim 1 will make predictions about how variance is distributed across minor allele frequencies (hopefully add a figure here to show how two different choices of evolutionary parameters lead to different distributions of variance among bins), which means it can be used for inference. Intuitively, the stronger seletion on disease variants is, the larger the proportion of variance we expect to be explained by low frequency bins.

The likelihood above is a composite rather than a true likelihood, meaning that it does not account for the non-independence (i.e. linkage disequilibrium) among sites. Such approaches are commonplace in population genetics when full likelihood computation is infeasible. Composite likelihoods have the property that they are unbiased with respect to the maximum likelihood estimate, but understate the uncertainty about that estimate, precisely because non-independence among datapoints is ignored, leading to the appearance of stronger evidence than actually exists, and naive approaches to model comparison therefore erroneously tend to favor more complex models (Gao and Song 2010). This limitation can be overcome by bootstrapping or potentially by adapting recently developed methods from the literature on demographic inference which show promise in circumventing these issues analytically at significantly reduced computational cost.

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