
Bayesian model comparison for rare variant association studies of multiple phenotypes

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Abstract

Whole genome sequencing studies applied to large populations or biobanks with extensive phenotyping raise new analytic challenges. The need to consider many variants at a locus or group of genes simultaneously and the potential to study many correlated phenotypes with shared genetic architecture provide opportunities for discovery and inference that are not addressed by the traditional one variant-one phenotype association study. Here we introduce a model comparison approach we refer to as MRP for rare variant association studies that considers correlation, scale, and location of genetic effects across a group of genetic variants, phenotypes, and studies. We consider the use of summary statistic data to apply univariate and multivariate gene-based meta-analysis models for identifying rare variant associations with an emphasis on protective protein-truncating variants that can expedite drug discovery. Through simulation studies, we demonstrate that the proposed model comparison approach can improve ability to detect rare variant association signals. We also apply the model to two groups of phenotypes from the UK Biobank: 1) asthma diagnosis, eosinophil counts, forced expiratory volume, and forced vital capacity; and 2) glaucoma diagnosis, intra-ocular pressure, and corneal resistance factor. We are able to recover known associations such as the protective association between rs146597587 in *IL33* and asthma. We also find evidence for novel protective associations between rare variants in *ANGPTL7* and glaucoma. Overall, we show that the MRP model comparison approach is able to retain and improve upon useful features from widely-used meta-analysis approaches for rare variant association analyses and prioritize protective modifiers of disease risk.

Author summary

Due to the continually decreasing cost of acquiring genetic data, we are now beginning to see large collections of individuals for which we have both genetic information and trait data such as disease status, physical measurements, biomarker levels, and more. These datasets offer new opportunities to find relationships between inherited genetic variation and disease. While it is known that there are relationships between different traits, typical genetic analyses only focus on analyzing one genetic variant and one phenotype at a time. Additionally, it is difficult to identify rare genetic variants that are associated with disease due to their scarcity, even among large sample sizes. In this work, we present a method for identifying associations between genetic variation and disease that considers multiple rare variants and phenotypes at the same time. By sharing information across rare variant and phenotypes, we improve our ability to identify rare variants associated with disease compared to considering a single rare variant and a single phenotype. The method can be used to identify candidate disease genes as well as genes that might represent attractive drug targets.

Introduction

Sequencing technologies are quickly transforming human genetic studies of complex traits: it is increasingly possible to obtain whole genome sequence data on thousands of samples at manageable costs. As a result, the genome-wide study of rare variants (minor allele frequency [MAF] < 1%) and their contribution to disease susceptibility and phenotype variation is now feasible [14].

In genetic studies of diseases or continuous phenotypes, rare variants are hard to assess individually due to the limited number of copies of each rare variant. Hence, to boost the ability to detect a signal, evidence is usually ‘aggregated’ across variants. When designing an ‘aggregation’ method, there are three questions that are usually considered. First, across which biological units should variants be combined; second, which variants mapping within those units should be included [5]; and third, which statistical model should be used [6]? Given the widespread observations of shared genetic risk factors across distinct diseases, there is also considerable motivation to use gene discovery approaches that leverage the information from multiple phenotypes jointly. In other words, rather than only aggregating variants that may have effects on a single phenotype, we can also bring together sets of phenotypes for which a single variant or sets of variants might have effects.

In this paper, we present a Bayesian multiple rare variants and phenotypes (MRP) model comparison approach for identifying rare variant associations as an alternative to current widely-used statistical tests. The MRP framework exploits correlation, scale, or location (direction) of genetic effects in a broad range of rare variant association study designs including: case-control; multiple diseases and shared controls; single continuous phenotype; multiple continuous phenotypes; or a mixture of case-control and multiple continuous phenotypes (Fig 1). MRP makes use of Bayesian model comparison, whereby we compute a Bayes Factor (BF) defined as the ratio of the marginal likelihoods of the observed data under two models: 1) a pre-specified null where all genetic effects are zero; and 2) an alternative model where factors like correlation, scale, or location of genetic effects are considered. The BF is an alternative to *p*-values from traditional hypothesis testing. For MRP, the BF represents the statistical evidence for a non-zero effect for a particular group of rare variants on the phenotype(s) of interest.

While many large genetic consortia collect both raw genotype and phenotype data, in practice, sharing of individual genotype and phenotype data across groups is difficult to achieve. To address this, MRP can take summary statistics, such as estimates of effect

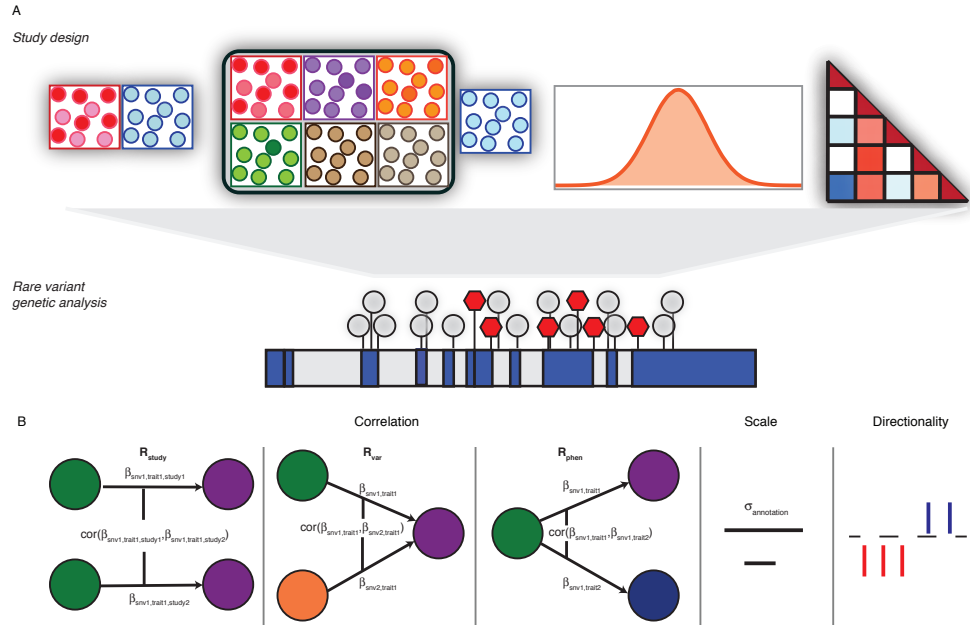


Fig 1. Schematic overview of MRP. A: MRP is suitable for a broad range of rare variant association study designs including (from left to right): i) case-control, ii) multiple diseases with shared controls, iii) single quantitative phenotype, and iv) mixture of case-control and quantitative phenotypes. B: Diagram of factors considered in rare variant association analysis including the correlation matrices: $\mathbf{R}_{\text{study}}$ (expected correlation of genetic effects among a group of studies), \mathbf{R}_{var} (expected correlation of genetic effects among a group of variants), and \mathbf{R}_{phen} (expected correlation of genetic effects among a group of phenotypes); the scale parameter for genetic variant annotation; and the location of genetic effects, which may be used to prioritize or identify protective modifiers of disease risk.

size and the corresponding standard error from typical single variant-single phenotype linear or logistic regressions, as input data. Furthermore, we use insights from Liu et al. [7] and Cichonska et al. [8] who suggest the use of additional summary statistics, like covariance estimates across variants and studies, respectively, that would enable lossless ability to detect gene-based association signals using summary statistics alone.

Aggregation techniques rely on variant annotations to assign variants to groups for analysis. MRP allows for the inclusion of priors on the spread of effect sizes that can be adjusted depending on what type of variants are included in the analysis. For instance, protein truncating variants (PTVs) [9][10] are an important class of variants that are more likely to be functional because they often disrupt the normal function of a gene. This biological knowledge can be reflected in the choices of priors for PTVs in MRP. Since PTVs typically abolish or severely alter gene function, there is particular interest in identifying protective PTV modifiers of human disease risk that may serve as targets for therapeutics [11][13]. We therefore demonstrate how the MRP model comparison approach can improve discovery of such protective signals by modeling the location (direction) of genetic effects which prioritizes variants or genes that are consistent with protecting against disease.

To evaluate the performance of MRP and to study its behavior we use simulations and compare it to other commonly used approaches. Some simple alternatives to MRP

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include univariate approaches for rare variant association studies including the sequence kernel association test (SKAT) [14], and the burden test, which we show are special cases of the MRP model comparison when we assign the prior correlation of genetic effects across different variants to be zero or one.

We applied MRP to summary statistics for two groups of related phenotypes from the UK Biobank. First, we applied MRP to asthma (HC382: the corresponding phenotype label in Global Biobank Engine <https://biobankengine.stanford.edu>), eosinophil count (INI30150), forced expiratory volume in 1-second (FEV₁, INI3063), and forced vital capacity (FVC, INI3062) and recovered the reported association between a rare PTV in *IL33* and asthma [15,16]. We also applied MRP to glaucoma (HC276), intra-ocular pressure (INI5263), and corneal resistance factor (INI5265) and find evidence that rare coding variants in *ANGPTL7* protect against glaucoma. These analyses show that MRP recovers results from typical single variant-single phenotype analyses while identifying new rare variant associations that include protective modifiers of disease risk.

Methods

Description of MRP

In this section, we provide an overview of the MRP model comparison approach. Refer to S1 Appendix for a detailed description. MRP models GWAS summary statistics as being distributed according to one of two models. The null model is that the regression effect sizes obtained across all studies for a group of variants and a group of phenotypes is zero. The alternative model is that summary statistics are distributed according to a multivariate normal distribution with mean zero and covariance matrix described below. MRP compares the evidence for the null and alternative model using a Bayes Factor (BF) that quantifies the amount of evidence for each model as the ratio of the marginal likelihoods of the observed data under two models.

To define the alternative model, we must specify the prior correlation structure, scale, and location (direction) of the effect sizes. Let N be the number of individuals and K the number of phenotype measurements on each individual. Let M be the number of variants in a testing unit \mathbf{G} , where \mathbf{G} can be, for example, a gene, pathway, or a network. Let S be the number of studies where data is obtained from - this data may be in the form of raw genotypes and phenotypes or summary statistics including linkage-disequilibrium, effect sizes (or odds ratio), and standard error of the effect size. When considering multiple studies ($S > 1$), multiple rare variants ($M > 1$), and multiple phenotypes ($K > 1$), we define the prior correlation structure of the effect sizes as an $SMK \times SMK$ matrix \mathbf{U} . In practice, we define \mathbf{U} as a Kronecker product, an operation of matrices of arbitrary size, of three sub-matrices:

- an $S \times S$ matrix $\mathbf{R}_{\text{study}}$ containing the correlations of genetic effects among studies where different values can be used to compare different models of association, such as for identifying heterogeneity of effect sizes between populations [17];
- an $M \times M$ matrix \mathbf{R}_{var} containing the correlations of genetic effects among genetic variants, which may reflect the assumption that all the PTVs in a gene may have the same biological consequence [9,10,18] or prior information obtained through integration of additional data sources, such as functional assay data [5,19], otherwise zero correlation of genetic effects may be assumed, which is used in dispersion tests like C-alpha [20,21] and SKAT [14]; and
- a $K \times K$ matrix \mathbf{R}_{phen} containing the correlations of genetic effects among phenotypes, which may be obtained from common variant data [22,24].

The variance-covariance matrix of the effect sizes may be obtained from readily
available summary statistic data such as in-study LD matrices, effect size estimates (or
log odds ratios), and the standard errors of the effect size estimates (S1 Appendix).
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MRP allows users to specify priors that reflect knowledge of the variants and
phenotypes under study. For instance, we can define an independent effects model
where each variant in the model may have different effect sizes. In this case, \mathbf{R}_{var} is the
identity matrix which reflects the assumption that the effect sizes of the variants are not
correlated. We can also define a similar effects model by setting every value of \mathbf{R}_{var} to
 ~ 1 . This model assumes that all variants under consideration have similar effect sizes
(with possibly differences in scale). This model may be appropriate for PTVs where
each variant completely disrupts the function of the gene, leading to a gene knockout.
The prior on the scale of effect sizes can also be used to denote which variants may have
larger effect sizes. For instance, emerging empirical genetic studies have shown that
within a gene, PTVs may have stronger effects than missense variants [25]. This can be
reflected by adjusting the prior spread of effect sizes (σ) for PTVs (S1 Appendix).
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Similarly, we can utilize a prior on the location (direction) of effects to specify
alternative models where we seek to identify variants with protective effects against
disease. Thus far we have assumed that the prior mean, or location, of genetic effects is
zero which makes it feasible to analyze a large number of phenotypes without
enumerating the prior mean across all phenotypes. To proactively identify genetic
variants that have effects that are consistent with a protective profile for a disease, we
can include a non-zero vector as a prior mean of genetic effect (S1 Appendix). We can
exploit information from Mendelian randomization studies of common variants, such as
recent findings where rare truncating loss-of-function variants in *PCSK9* were found to
decrease LDL and triglyceride levels and decrease CAD risk [11][26][28] to identify
situations where such a prior is warranted.
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Applying MRP to variants from a testing unit \mathbf{G} yields a BF for that testing unit
that describes the evidence that rare variants in that testing unit have a nonzero effect
on the traits used in the model. For instance, consider genes as testing units. By
running MRP, we obtain a BF for each gene that represents the evidence that rare
variants in that gene affect the traits of interest. These BF can be used to identify
specific genes that may be linked to disease. Although we see advantages in adopting a
Bayesian perspective for MRP, our approach could be used in a frequentist context by
calculating a BF and using it as a test statistic to compute p-values (S1 Appendix,
Fig 2).
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HDF5 Tables

Although summary statistics are quicker to read and process than raw data, the number
of studies meta-analyzed in this work is expected to be sufficiently large to require
optimizations in data representation and processing (S1 Fig). Our solution was the use
of the HDF5 (Hierarchical Data Format 5) data representation to enable rapid
processing of effect size, uncertainty, and cross-trait estimate data. HDF5 is a fast and
lightweight file format designed for scientific data. It has bindings for R, Python,
C/C++, Java, and nearly every other population programming language. Reading data
from a table within a HDF5 file can be an order of magnitude faster than reading text
files from a Unix file, and it makes it easier to organize data within an internal structure.
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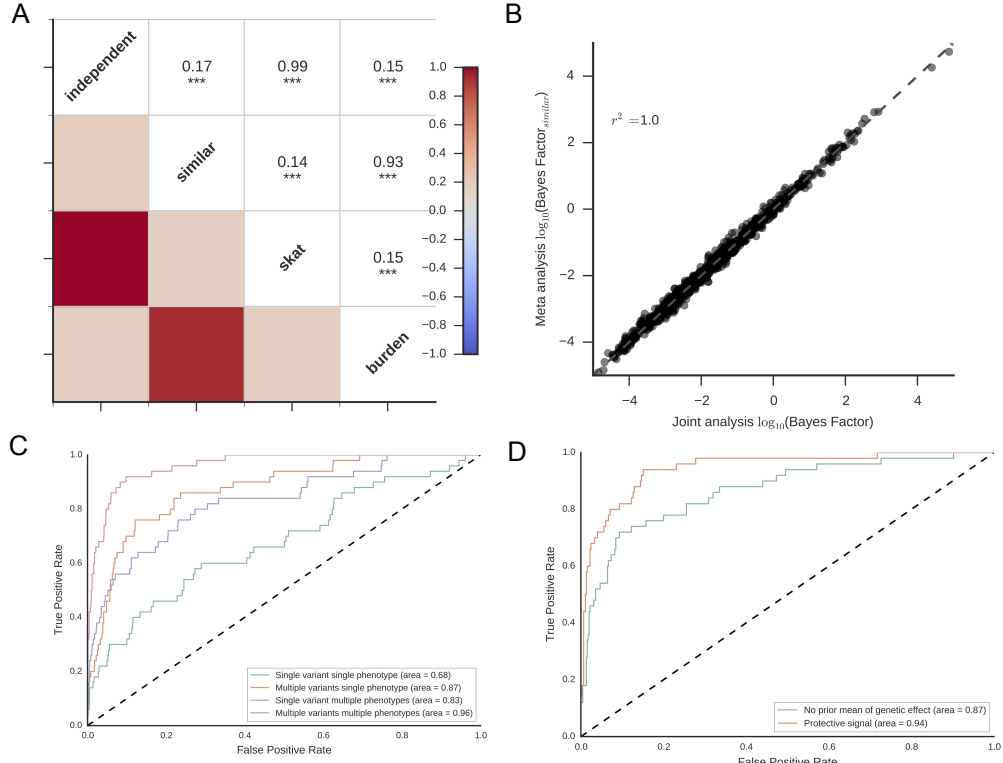


Fig 2. Simulation studies. A: Comparison of $-\log_{10}(p\text{-values})$ from frequentist BF_{MRP} approximation for an independent effects and a similar effects model to commonly used gene-based statistical tests (skat and burden). B: Comparison of $\log_{10}(\text{Bayes Factors})$ obtained when raw genotype and phenotype data is available to a scenario where summary statistics only was available and similar effects across studies is assumed. C: From single variant and single phenotype to multiple variants and multiple phenotypes gene discovery: ROC curves for detecting gene association to any of the phenotypes using single variant/single phenotype association (green) to multiple variants and multiple phenotypes association (purple). D: ROC curves for detecting gene association when incorporating prior mean of genetic effects (orange) to identify protective alleles.

UK Biobank Data

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GWAS Summary Statistics

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We performed genome-wide association analysis using PLINK v2.00a(17 July 2017) as previously described [15]. For asthma, we used the Firth fallback in PLINK, a hybrid algorithm which normally uses the logistic regression code described in [29], but switches to a port of `logistf()` (<https://cran.r-project.org/web/packages/logistf/index.html>) in two cases: (1) one of the cells in the 2x2 allele count by case/control status contingency table is empty (2) logistic regression was attempted since all the contingency table cells were nonzero, but it failed to converge within the usual number of steps. We used the following covariates in our analysis: age, sex, array type, and the first four principal components, where array type is a binary variable that represents whether an individual was genotyped with UK Biobank Axiom Array or UK BiLEVE Axiom Array. For variants that were specific to one array, we did not use array as a covariate.

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Asthma and glaucoma cases were defined using both Hospital Episode Statistics and verbal questionnaire responses. We used the provided values from the UK Biobank for eosinophil counts, forced vital capacity (FVC), forced expiratory volume in 1-second (FEV₁), intra-ocular pressure, and corneal resistance factor. The phenotype codes used throughout (asthma=HC382, eosinophil count=INI30150, FEV₁=INI3063, FVC=INI3062, glaucoma=HC276, intra-ocular pressure=INI5263, and corneal resistance factor=INI5265) correspond to the phenotype codes used in on the Global Biobank Engine <https://biobankengine.stanford.edu>.

Genetic Correlations

We calculated the genetic correlation between the two groups of traits (asthma, eosinophil counts, FVC, FEV₁ and glaucoma, intra-ocular pressure, corneal resistance factor) using the MultiVariate Polygenic Mixture Model (MVPMM) [30]. Briefly, MVPMM estimates genetic correlation given GWAS summary statistics (effect size and standard error of effect size estimate) by modeling GWAS summary statistics as generated from one of two mixture components. Summary statistics from variants in the null component are modeled as being drawn from a multivariate normal distribution with zero mean and covariance matrix that captures correlation in the summary statistics due to the use of shared subjects or other sources of correlation. Summary statistics from variants in the non-null component are modeled as being drawn from a multivariate normal distribution with zero mean, but the covariance matrix for the non-null component combines the covariance matrix from the null component with another covariance matrix that captures the genetic correlation between the phenotypes being considered. We observed similar genetic correlations using LD score regression (S2 Fig) [24].

UK Biobank Asthma and Glaucoma Applications

We applied MRP for both sets of traits assuming both an independent effects model and a similar effects model. We applied Bayesian model averaging to these results by averaging the BF from each model for each gene; these results are reported in the main text. The results for each individual model are included in the Supporting Information. For the Manhattan plots and tables, we removed any genes with non-unique gene symbols. In cases where genes overlapped such that they shared rare variants and therefore the same BF, we removed one gene. *ANGPTL7* protein expression was assessed using the HIPED protein expression database accessed through genecards.org on 2017/1/29 [31]. We identified the protein 1JC9_A as homologous to the *ANGPTL7* protein using the “3D structure mapping” link from dbSNP (https://www.ncbi.nlm.nih.gov/projects/SNP/snp_ref.cgi?rs=28991009). We retrieved the 3D structure image from the iCn3D Structure Viewer (<https://www.ncbi.nlm.nih.gov/Structure/icn3d/icn3d.html>).

Variant Filtering

We used the `variant_filter_table.tsv` file available at <https://github.com/rivas-lab/public-resources> (6f9f726) to filter variants on the UK Biobank array for use with MRP. We first chose variants with minor allele frequency less than 1%. We then filtered out all variants with `all_filters` less than one. This removes variants with missingness greater than 1% (calculated on an array-specific basis for array-specific variants) or Hardy-Weinberg equilibrium $p < 10^{-7}$. This also removes some PTVs for which manual inspection revealed irregular cluster plots [15]. We LD pruned the variants by only using variants with `1d` equal to one. We

included missense variants and PTVs indicated by the following annotations:
missense_variant, stop_gained, frameshift_variant, splice_acceptor_variant,
splice_donor_variant, splice_region_variant, start_lost, stop_lost.

Results

Simulation studies

We first verified the analytical derivations and examined the properties of the approach
under a simulation framework.

Comparison to frequentist gene tests

For the analysis of multiple rare variants and a single phenotype we compared it to the
burden test and the SKAT test, commonly used statistical tests in rare variant
association studies of a single phenotype. We observe concordance between the
frequentist methods and the Bayesian models. To compare the Bayesian models we
compute p-values by using the BF as the test statistic and approximating it using
distribution properties of quadratic forms (S1 Appendix). As expected, an independent
effects model has high correlation with the gene-based test SKAT ($r^2 = 0.99$), whereas
the similar effects model has high correlation with the burden test ($r^2 = 0.93$, Fig 2A).

Summary statistic data

To study the behavior of MRP using summary statistics we simulate two scenarios: first,
the scenario where analysts have access to all the raw genotype and phenotype data;
and second, the scenario where analysts only have access to summary statistics data 7.
We conducted 1000 simulation experiments where we let K (the number of phenotypes)
= 3, M (the number of variants) = 10, S (the number of studies) = 2, N_0 (number of
individuals in study with access to all the data) = 10000, N_1 (meta-analysis study 1)
= 5000, N_2 (meta-analysis study 2) = 5000. We find that, under the scenario where
similar effects are assumed across studies, the Bayes Factors obtained using summary
statistics alone are strongly correlated ($r^2 = 1$) to Bayes Factors obtained by the full
genotype and phenotype data (Fig 2B).

From single variant and single phenotype analysis to multiple variants and multiple phenotypes

To validate the flexibility of the approach we conducted a simulation experiment where
we assumed an allelic architecture consistent to that discovered for *APOC3* in relation
to coronary artery disease (CAD), triglycerides (TG), low-density lipoprotein cholesterol
(LDL-C), and high-density lipoprotein cholesterol (HDL-C) 28,32,34. We simulated
three studies and applied the model comparison unit jointly to summary statistic data
obtained for each study (Supplementary Note). Overall, we observed that considering
the joint effects across multiple studies in a group of variants and phenotypes may
improve ability to detect gene-based signals (Fig 2C), and that considering prior mean
of genetic effects should aid in efforts to identify protective modifiers of disease risk
(Fig 2D).

Applications

We applied the MRP model comparison approach to summary statistic data generated
from single variant logistic regression and linear regression analysis for coding variants

on the UK Biobank array (Methods). We applied MRP separately to asthma and three related traits as well as glaucoma and two related traits.

Asthma, eosinophil counts, forced expiratory volume, and forced vital capacity

We first applied MRP to GWAS summary statistics for asthma, eosinophil count, forced expiratory volume in 1-second (FEV_1), and forced vital capacity (FVC) phenotypes. Recent work has identified associations between the PTV rs146597587 in *IL33* and asthma and eosinophil counts [15,16]. FEV_1 and FVC are measures of pulmonary function that are used to diagnosis and classify pulmonary disease [35]. To demonstrate the advantage of considering the phenotypes jointly, we applied MRP to rare missense variants and PTVs (MAF < 1%) for each phenotype separately (Fig 3A-D) as well as to all phenotypes jointly (Fig 3E,F) and obtained \log_{10} BF for each gene. We applied both independent and similar effects models and used Bayesian model averaging to compute a single BF per gene [36]. In agreement with previous studies, we observed evidence that rare missense variants and/or PTVs in *IL33* affect eosinophil counts and offer protection from asthma from the single-phenotype analyses, though the evidence of association was strongest for the joint analysis (S1 Table) [15,16]. We performed an analysis focused on identifying protective variants which also identified the *IL33* association (Fig 3F). The results were similar using only either the independent effects (S3 Fig) or similar effects models (S4 Fig). We inspected the effect sizes from the marginal GWAS regressions for the rare variants included in the analysis and found that the association identified by MRP is likely driven by the PTV rs146597587 (Fig 3G).

We also found moderate evidence for association between rare coding variants in *CCR3* and asthma. The \log_{10} BFs for *CCR3* was 3.3 in the joint model compared to only -0.5 in the asthma-only analysis (Fig 3, S1 Table). *CCR3* is a chemokine receptor that is highly expressed on eosinophils and has been a therapeutic focus for asthma [37,38]. *CCR3* was not reported in a large GWAS for allergic disease including asthma [39] though *CCR3* is near a locus associated with atopy in a previous meta-analysis [40]. These results demonstrate that MRP can identify biologically meaningful therapeutic targets that may be missed by standard GWAS approaches.

Considering multiple phenotypes jointly allows for the efficient prioritization of disease genes. For instance, some genes like *IL18RAP*, *ATP2A3*, and *FLG* had \log_{10} BFs greater than 4 in the asthma-only analysis but much smaller BFs in the joint analyses indicating that rare variants in these genes are less likely to affect this group of traits. Similarly, there were other genes like *RP11-39K24.9* and *IL17RA* that had larger BFs in the eosinophil count-only analysis but small BFs for the joint analyses demonstrating MRP's ability to integrate information across all phenotypes considered.

Glaucoma, intra-ocular pressure, and corneal resistance factor

We also applied MRP to missense variants and PTVs for glaucoma, intra-ocular pressure, and corneal resistance factor as well as performing joint analyses. Intra-ocular pressure is a measure of the fluid pressure in the eye, is associated with glaucoma risk, and has been linked to genetic variants associated with glaucoma [41]. Corneal resistance factor is a measure of the cornea's ability to resist mechanical stress and has been associated with glaucoma presence and severity [42-44]. While the individual glaucoma analysis did not yield any associations with \log_{10} BF greater than three, the joint analysis identified rare coding variants in both *ANGPTL7* and *WNT10A* as associated with glaucoma (Fig 4A-D, S2 Table). Applying the MRP model identified *ANGPTL7* with effects consistent with protection to glaucoma, and additional

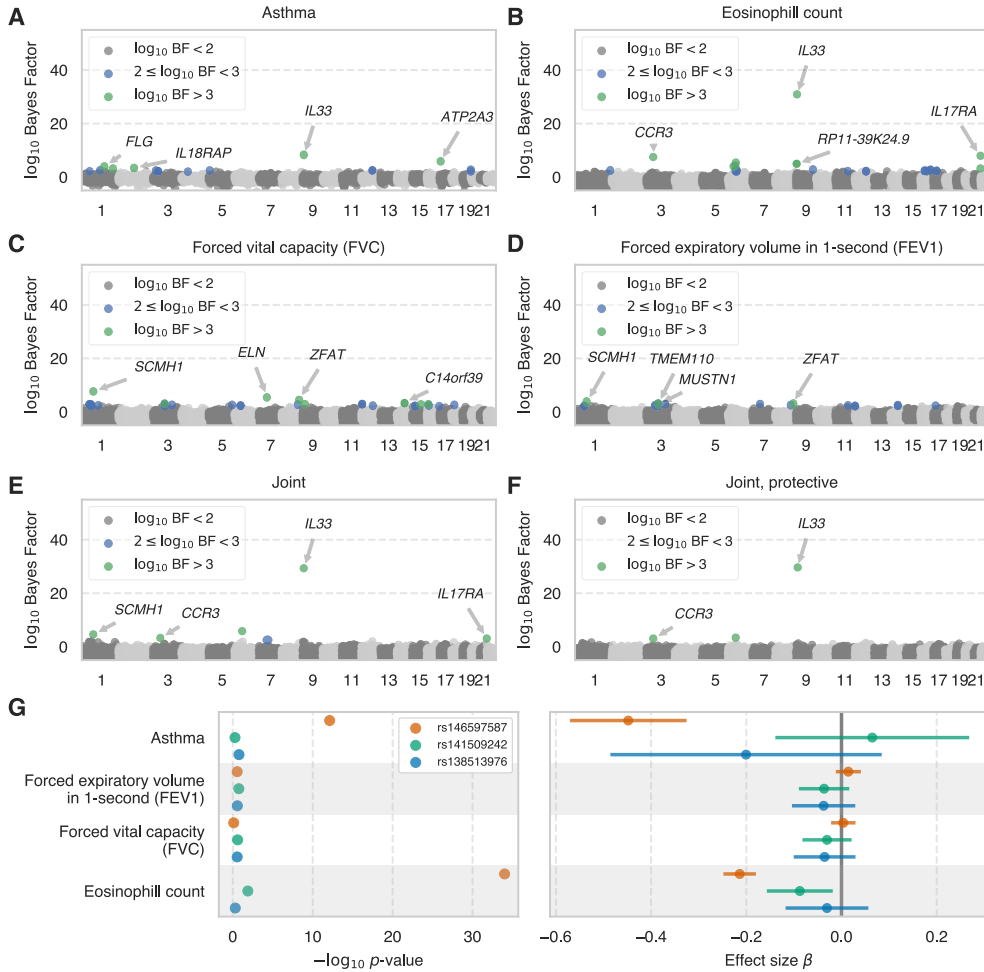


Fig 3. Results for asthma application. log₁₀ Bayes Factors from applying MRP and Bayesian model averaging to summary statistics for missense and protein-truncating variants from (A) asthma (HC382), (B) eosinophil counts (INI30150), (C) forced vital capacity (FVC, INI3062), (D) forced expiratory volume in 1-second (FEV₁, INI3063), (E) all four traits jointly, and (F) all four traits jointly with focus on protective effects. The four genes outside of chromosome 6 with the largest Bayes Factors greater than three are labeled in each plot. Only log₁₀ Bayes Factors greater than -5 are plotted. (F) – log₁₀ p-values (left panel) and estimated effect sizes with 95% confidence intervals (right panel) for missense variants and PTVs in *IL33* for each phenotype

association to *WNT10A* (Fig 4E). We obtained similar results using the independent effects (S5 Fig) or similar effects models (S6 Fig).

Expression of *ANGPTL7* is upregulated in glaucoma and has been proposed to regulate intra-ocular pressure and glaucoma risk [45,46]. The GWAS summary statistics for the rare variants in *ANGPTL7* suggest that the association with glaucoma is driven by the missense variant rs28991009 that changes residue 175 from glutamine to histidine (Fig 4F, G). According to the HIPED protein expression database, *ANGPTL7* protein is expressed at ~ 0.7 parts per million in vitreous humor, the material between the lens and retina of the eyeball; in contrast, the expression of *ANGPTL7* protein is less than 0.01 parts per million in 68 other normal tissues [31]. Such tissue-specific activity may

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make *ANGPTL7* a useful therapeutic target. *WNT10A* has not been previously associated with glaucoma though an exonic variant rs121908120 in *WNT10A* is associated with central cornea thickness and increased risk of keratoconus, a disease of the cornea, indicating that this gene may play a role in ocular diseases.

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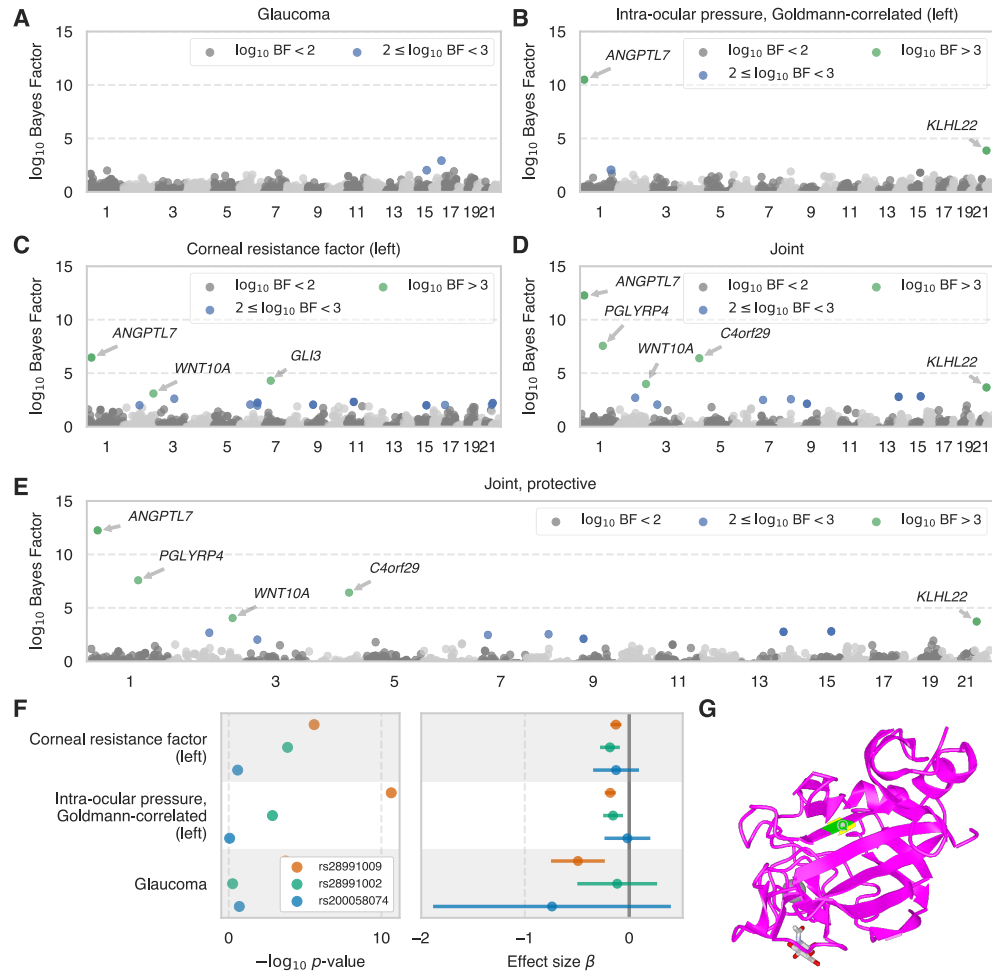


Fig 4. Results for glaucoma application. \log_{10} Bayes Factors from applying MRP and Bayesian model averaging to summary statistics for missense and protein-truncating variants from (A) glaucoma (HC276), (B) intra-ocular pressure (INI5263), (C) corneal resistance factor (INI5265), and (D) all three traits jointly. (E) shows the results of a joint analysis focused on finding rare variants that protect against glaucoma. The genes outside of chromosome 6 with Bayes Factor greater than three are indicated by arrows. Only \log_{10} Bayes Factors greater than zero are plotted. F: $-\log_{10} p$ -values (left panel) and estimated effect sizes with 95% confidence intervals (right panel) for missense variants and PTVs in *ANGPTL7* for all three phenotypes. G: Location of rs28991009 variant (green) for the protein 1JC9_A homologous to *ANGPTL7*.

Discussion

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In this study, we developed a Bayesian model comparison approach MRP that shares information across both variants and phenotypes to identify rare variant associations. We used simulations to compare MRP to the widely used burden and SKAT tests for identifying rare variant associations and found that jointly considering both variants and phenotypes can improve the ability to detect associations. We also applied the MRP model comparison framework to summary statistic data from two groups of traits from the UK Biobank: asthma diagnosis, eosinophil counts, FEV₁, and FVC; and glaucoma diagnosis, intra-ocular pressure, and corneal resistance factor. We identified strong evidence for the previously described association between the PTV rs146597587 in *IL33* and asthma [15][16]. We also found evidence for a link between rare variants in *ANGPTL7* and glaucoma, consistent with previous experiments that suggested a role for *ANGPTL7* in glaucoma [45][46]. These results demonstrate the ability of the MRP model comparison approach to leverage information across multiple phenotypes and variants to discover rare variant associations.

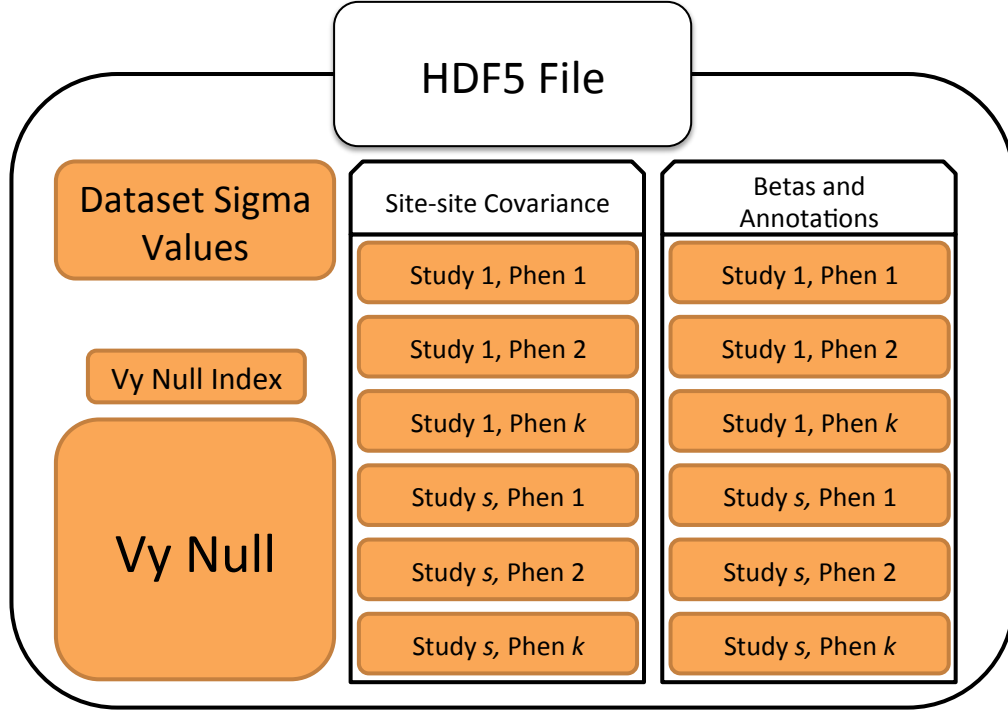
As genetic data linked to high-dimensional phenotype data continues to be made available through biobanks, health systems, and research programs, there is a large need for statistical approaches that can leverage information across different genetic variants, phenotypes, and studies to make strong inferences about disease-associated genes. The approach presented here relies only on summary statistics from marginal association analyses which can be shared with less privacy concerns compared to raw genotype and phenotype data. Combining joint analysis of variants and phenotypes with meta-analysis across studies offers new opportunities to identify gene-disease associations.

Supporting information

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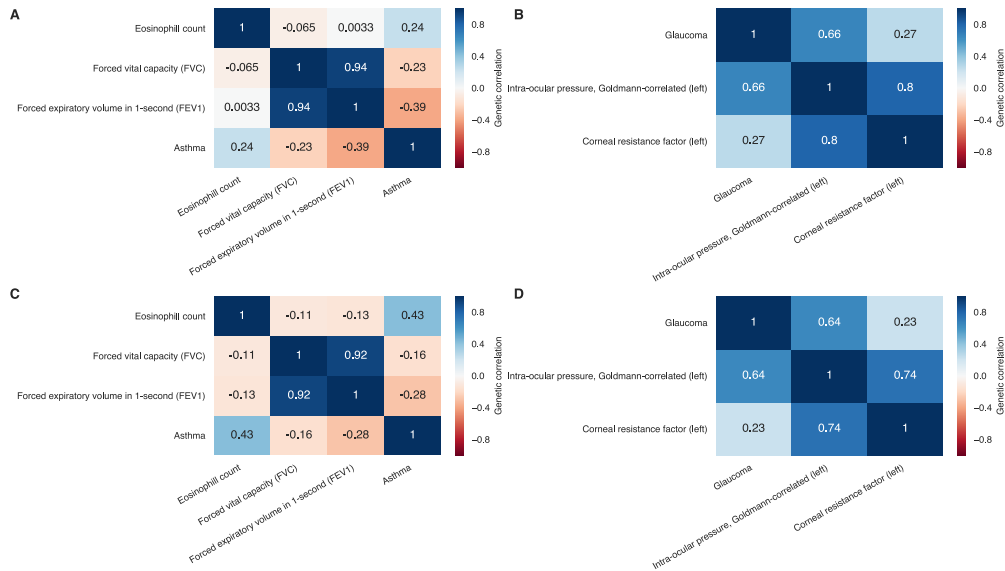
S1 Appendix. MRP model details. Specification of the MRP model including the likelihood function, priors, and Bayes factor calculation.

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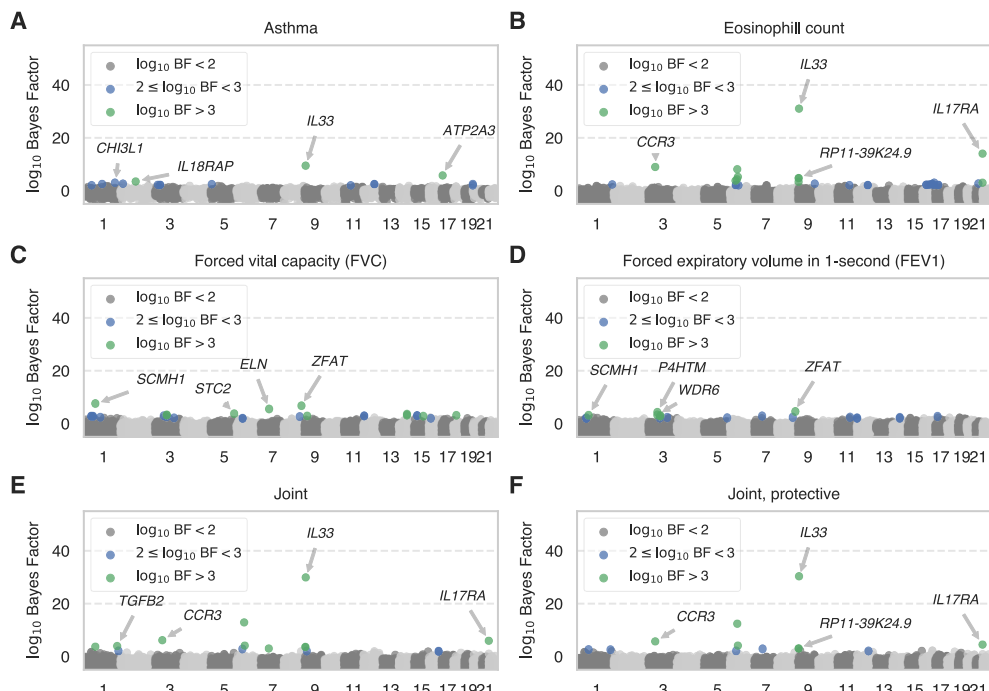
S1 Fig. HDF5 Implementation. Our HDF5 implementation contained the following components: first, a group with one table per annotation file. All effect size (beta) values and study-specific annotations were contained here, and the number of tables is limited by S (the number of studies) $\times K$ (the number of traits). Second, a group with site-site covariance data. While these covariance matrices may have dimension M (the number of variants) $\times M$, we store the data as tables, each row specifying the covariance between two variants. The number of tables should be the same as the previous set, capped by S (the number of studies) $\times K$ (the number of traits). Third, we store one table with sigma values for each study/phenotype combination. In the event that the traits were rank-normal transformation was performed these sigma values are equal to 1. These are used to compute correlation between two datasets. Finally, we store a matrix/table pair for Vy null and its index. The Vy null matrix has dimensions $(S \times K) \times (S \times K)$ each entry specifying the estimated correlation of effect sizes between two datasets. The index table encodes row/column position of each dataset.

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S2 Fig. Genetic correlations. Genetic correlations for (A) asthma and related phenotypes and (B) glaucoma and related phenotypes estimated using MVPMM. Genetic correlations for (C) asthma and related phenotypes and (D) glaucoma and related phenotypes estimated using LD score regression.

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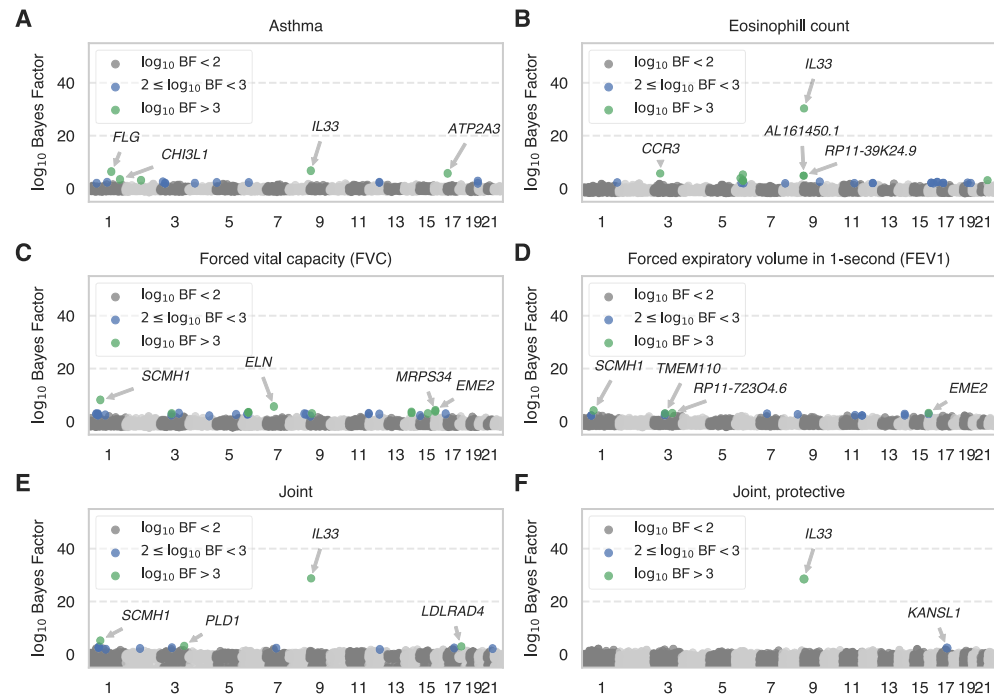


S3 Fig. Results for independent effects model applied to asthma, eosinophil counts, FEV₁, and FVC. \log_{10} Bayes Factors from applying MRP independent effects model to summary statistics for missense and protein-truncating variants from (A) asthma (HC382), (B) eosinophil counts (INI30150), (C) forced vital

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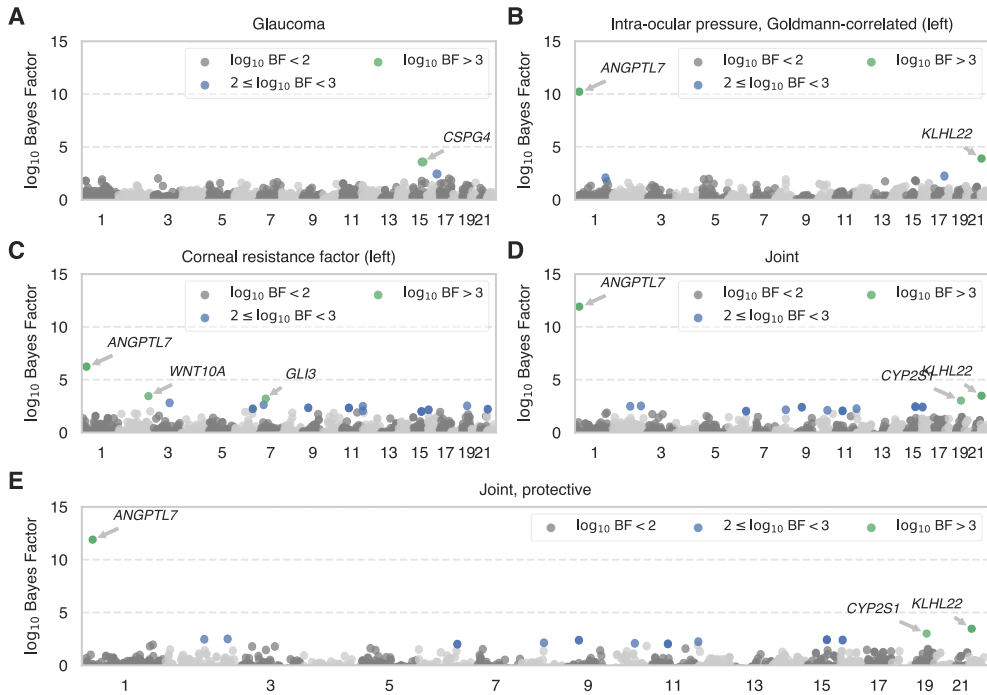
capacity (FVC, INI3062), (D) forced expiratory volume in 1-second (FEV₁, INI3063), (E) all four traits jointly, and (F) all four traits jointly with focus on protective effects. The four genes outside of chromosome 6 with the largest Bayes Factors greater than three are labeled in each plot. Only log₁₀ Bayes Factors greater than -5 are plotted.

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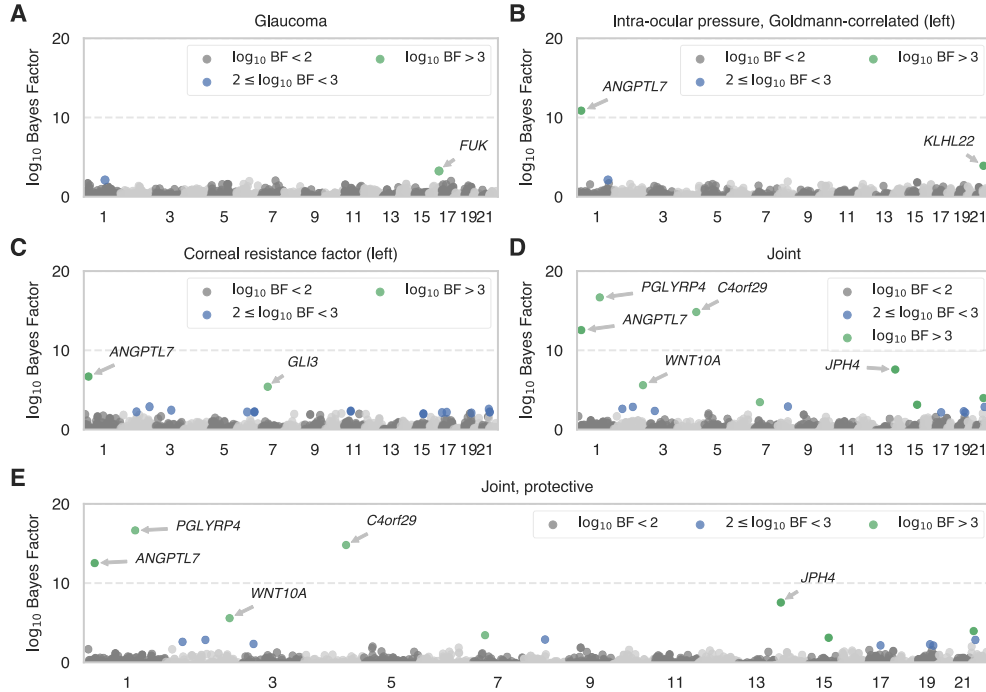
S4 Fig. Results for similar effects model applied to asthma, eosinophil counts, FEV₁, and FVC. log₁₀ Bayes Factors from applying MRP similar effects model to summary statistics for missense and protein-truncating variants from (A) asthma (HC382), (B) eosinophil counts (INI30150), (C) forced vital capacity (FVC, INI3062), (D) forced expiratory volume in 1-second (FEV₁, INI3063), (E) all four traits jointly, and (F) all four traits jointly with focus on protective effects. The four genes outside of chromosome 6 with the largest Bayes Factors greater than three are labeled in each plot. Only log₁₀ Bayes Factors greater than -5 are plotted.

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S5 Fig. Results for independent effects model applied to glaucoma intra-ocular pressure, and corneal resistance factor. \log_{10} Bayes Factors from applying MRP independent effects model to summary statistics for missense and protein-truncating variants from (A) glaucoma (HC276), (B) intra-ocular pressure (INI5263), (C) corneal resistance factor (INI5265), and (D) all three traits jointly. (E) shows the results of a joint analysis focused on finding rare variants that protect against glaucoma. The genes outside of chromosome 6 with Bayes Factor greater than three are indicated by arrows. Only \log_{10} Bayes Factors greater than zero are plotted.

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S6 Fig. Results for similar effects model applied to glaucoma intra-ocular pressure, and corneal resistance factor. log₁₀ Bayes Factors from applying MRP similar effects model to summary statistics for missense and protein-truncating variants from (A) glaucoma (HC276), (B) intra-ocular pressure (INI5263), (C) corneal resistance factor (INI5265), and (D) all three traits jointly. (E) shows the results of a joint analysis focused on finding rare variants that protect against glaucoma. The genes outside of chromosome 6 with Bayes Factor greater than three are indicated by arrows. Only log₁₀ Bayes Factors greater than zero are plotted.

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Gene	Joint, protective	Joint	Eosinophil count	FVC	FEV ₁	Asthma
<i>IL33</i>	29.4	29.3	30.6	-2.3	-2.2	8.1
<i>CCR3</i>	3.1	3.3	7.4	-1.4	-1.6	-0.5
<i>RP11-39K24.9</i>	0.8	1.8	4.9	-0.1	-0.4	0.3
<i>SCMH1</i>	0.5	4.7	-1.5	7.7	3.8	-0.7
<i>MUSTN1</i>	0.4	1.1	-1.2	2.9	2.9	-0.6
<i>ZFAT</i>	0.3	1.3	-2.0	4.7	3.1	-0.4
<i>ELN</i>	0.2	2.5	-1.0	5.6	2.9	-0.6
<i>C14orf39</i>	-0.7	-0.0	-1.1	3.5	2.5	0.0
<i>TMEM110</i>	-0.9	1.1	-1.0	3.3	3.1	-0.6
<i>IL17RA</i>	-4.4	3.1	7.9	-2.7	-2.5	-1.1
<i>IL18RAP</i>	-9.6	-0.9	-1.0	-1.6	-1.7	3.5
<i>ATP2A3</i>	-11.9	-0.8	-1.2	-2.1	-2.2	5.8
<i>FLG</i>	-20.1	-17.2	-6.0	-7.4	-8.5	4.0

S1 Table. Highlighted genes from asthma analysis. log₁₀ Bayes Factors for genes highlighted in Figure 3.

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Gene	Joint, protective	Joint	Glaucoma	Intra-ocular pressure, Goldmann-correlated	Corneal resistance factor
<i>ANGPTL7</i>	11.9	11.9	1.7	10.1	6.2
<i>KLHL22</i>	3.5	3.5	-0.2	3.9	2.2
<i>WNT10A</i>	2.5	2.5	-0.3	-0.5	3.5
<i>GLI3</i>	1.5	1.5	-0.3	-0.7	3.2
<i>RNASE2</i>	-0.5	-0.5	0.4	-1.0	-1.1
<i>FUK</i>	-1.8	-1.8	2.5	-2.1	-2.1

S2 Table. Highlighted genes from glaucoma analysis. \log_{10} Bayes Factors for genes highlighted in Figure 4

Author Contributions

MAR and MP designed the method and derived all analytical calculations. MAR, MP, and CD wrote the manuscript. MAR, MP, CCAS, YT, MA, and CD provided analysis and designed figures. TP designed HDF5 tables and implementation of loaders. MJD provided critical feedback on methodology.

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Appendix S1

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January 28, 2018

1 MRP model comparison for association testing

We consider the multivariate linear regression model

$$\underset{(N \times K)}{Y} = \underset{(N \times K)}{\Psi} + \underset{(N \times M)(M \times K)}{X} \underset{(M \times K)}{B} + \underset{(N \times K)}{E},$$

where the matrices $Y = [y_{ik}]$, $X = [x_{im}]$, $B = [\beta_{mk}]$ and $E = [e_{ik}]$ describe the phenotype values (y_{ik}), copies of minor allele (x_{im}), variant-phenotype effects (β_{mk}), and residual errors (e_{ik}), for individual i , phenotype k , and variant m . We assume that each phenotype has been transformed to a standard normal distribution and that the columns of X have been centered, which means that the estimate for the intercept term Ψ is 0 and independent of the estimate of B . We use vectorized notation where the rows of B form vector $\beta = (\beta_1, \dots, \beta_M)^\top$ of length MK .

We define the MRP model comparison as a Bayes factor (BF) between the alternative model, where at least one variant affects at least one phenotype, and the null model, where all variant-phenotype effects are zero. BF is the ratio of the marginal likelihoods for these two models:

$$BF = \frac{\int_{\beta} p(\text{Data}|\beta) p(\beta|\text{ALT}) d\beta}{\int_{\beta} p(\text{Data}|\beta) p(\beta|\text{NULL}) d\beta},$$

where Data can correspond either to the effect size estimates $\hat{\beta}$ and the estimated variance-covariance matrix of $\hat{\beta}$, \hat{V}_{β} , or to the original phenotypes and genotypes, $\underset{(N \times K)}{Y}$ and $\underset{(N \times M)}{X}$, and any other covariates that we want to regress out from the phenotypes.

The prior distribution for the null model, $p(\beta|\text{NULL})$, is simply the point mass at $\beta = 0$. In section 2 we show how we approximate the likelihood function for β , $p(\text{Data}|\beta)$, in section 3 we define the prior distribution $p(\beta|\text{ALT})$ for the alternative model, and finally, in section 4, we compute the BF.

2 Likelihood function

A maximum likelihood estimator of B is given by the ordinary least-squares method

$$\hat{B} = (X^\top X)^{-1} X^\top Y,$$

that in vectorized form is denoted by $\hat{\beta} = (\hat{\beta}_1, \dots, \hat{\beta}_M)^\top$. An estimator of the variance-covariance of $\hat{\beta}$ is given by

$$\hat{V}_{\beta} = (X^\top X)^{-1} \otimes \hat{V}_Y,$$

where \hat{V}_Y is the estimated residual variance-covariance matrix of Y given X .

Following Band et al. [1], we approximate the likelihood function of β by a multivariate normal distribution with mean $\hat{\beta}$ and variance-covariance matrix \hat{V}_β . Note that by approximating \hat{V}_Y by the trait correlation matrix, this likelihood approximation does not require access to the individual level data X and Y but only to the summary data of effect sizes $\hat{\beta}$, LD-matrix $X^T X$ and a trait correlation estimate.

3 Prior of β in the alternative model

We construct the prior distribution $p(\beta|ALT)$ for the alternative model in three steps allowing user to specify correlations between effects of different variants on different traits across different studies.

In a single study, the prior density for β incorporates the expected correlation of genetic effects among a group of variants (\mathbf{R}_{var}) and among a group of phenotypes (\mathbf{R}_{phen}). In addition, we incorporate an expected spread of the effect size of each variant by scaling \mathbf{R}_{var} as

$$\mathbf{S}_{\text{var}} = \Delta(\sigma_m) \mathbf{R}_{\text{var}} \Delta(\sigma_m),$$

where $\Delta(\sigma_m)$ is a diagonal matrix with entries σ_m determining the spread of the effect size distribution for each variant $m \leq M$. Thus, we can model settings where, e.g., protein-truncating variants have larger effect sizes ($\sigma = 0.5$) than missense variants ($\sigma = 0.2$). Note that when $\sigma_m = 1$ for all m then $\mathbf{S}_{\text{var}} = \mathbf{R}_{\text{var}}$.

All in all, our prior density for β under alternative model is

$$\beta|ALT \sim \mathcal{N}(\mathbf{0}, \mathbf{U}), \text{ where } \mathbf{U} = \mathbf{S}_{\text{var}} \otimes \mathbf{R}_{\text{phen}}.$$

When we have data from multiple studies we allow for possible differences in genetic effects across ethnicities or populations extending the Approximate Bayes Factors of Band et al. [1] and the summary statistics approach of RAREMETAL [2] from univariate to multivariate phenotypes. Let $\hat{\beta} = (\hat{\beta}_{s,m,k}) =$

$(\hat{\beta}_{1,1,1}, \hat{\beta}_{1,1,2}, \dots, \hat{\beta}_{1,1,K}, \hat{\beta}_{1,2,1}, \dots, \hat{\beta}_{1,2,K}, \dots, \hat{\beta}_{1,M,K}, \hat{\beta}_{2,1,1}, \dots, \hat{\beta}_{S,M,K})$, where S is the number of studies, M is the number of variants, and K is the number of phenotypes. As with a single study, we incorporate the expected correlation of genetic effects between a pair of variants and a single phenotype using the matrix \mathbf{S}_{var} , between a variant and a pair of phenotypes using the matrix \mathbf{R}_{phen} , and we introduce the matrix $\mathbf{R}_{\text{study}}$ to specify prior on the similarity in effect sizes across the studies. Thus, the prior is

$$\beta \sim \mathcal{N}(\mathbf{0}, \mathbf{U}), \text{ where } \mathbf{U} = \mathbf{R}_{\text{study}} \otimes (\mathbf{S}_{\text{var}} \otimes \mathbf{R}_{\text{phen}}).$$

It is straightforward to include a non-zero vector μ as a prior mean of genetic effects, in which case the prior is

$$\beta \sim \mathcal{N}(\mu, \mathbf{U}).$$

We use this, for example, when screening for protective rare variants that have a pre-specified beneficial profile on a set of risk factors.

4 BF_{MRP}

The Bayes Factor is the ratio of the marginal likelihoods between the alternative and the null model. The marginal likelihood for the alternative model is

$$\int_{\beta} p(\text{Data}|\beta) p(\beta|ALT) d\beta = c \times \mathcal{N}(\hat{\beta}; \mu, \hat{V}_\beta + \mathbf{U})$$

and the marginal likelihood for the null model is

$$\int_{\beta} p(\text{Data}|\beta) p(\beta|\text{NULL}) d\beta = c \times \mathcal{N}(\hat{\beta}; 0, \hat{V}_{\beta}).$$

The Bayes Factor (derived below in section 4.1) is given by

$$\text{BF}_{\text{MRP}} = \frac{\det(\hat{V}_{\beta} + \mathbf{U})^{-\frac{1}{2}} \exp\left[-\frac{1}{2}(\hat{\beta} - \mu)^T (\hat{V}_{\beta} + \mathbf{U})^{-1} (\hat{\beta} - \mu)\right]}{\det(\hat{V}_{\beta})^{-\frac{1}{2}} \exp\left[-\frac{1}{2}\hat{\beta}^T \hat{V}_{\beta}^{-1} \hat{\beta}\right]}.$$

When $\mu = 0$, BF_{MRP} is an increasing function of the following quadratic form

$$Q(\hat{\beta}; \hat{V}_{\beta}, \mathbf{U}) = \hat{\beta}^T (\hat{V}_{\beta}^{-1} - (\hat{V}_{\beta} + \mathbf{U})^{-1}) \hat{\beta}. \quad (1)$$

Furthermore, this quadratic form is the only part of the BF_{MRP} that depends on $\hat{\beta}$. Thus, by deriving a distribution of $Q(\hat{\beta}; \hat{V}_{\beta}, \mathbf{U})$ under the null model we can compute a p-value when BF_{MRP} is used as a test statistic. According to basic properties of quadratic forms of Gaussian variables, $Q(\hat{\beta}; \hat{V}_{\beta}, \mathbf{U}) \sim \sum_{i=1}^n d_i \chi_i^2$, where χ_i^2 are an independent sample from χ_1^2 distribution (chi-square with one degree of freedom), and d_i are the eigenvalues of matrix $I - (\hat{V}_{\beta} + \mathbf{U})^{-1} \hat{V}_{\beta}$. The distribution function for a mixture of chi-squares can be numerically evaluated by the R-package ‘CompQuadForm’ [3].

4.1 MRP Bayes Factor derivation

To compute the Bayes Factor

$$\text{BF}_{\text{MRP}} = \frac{\det(\hat{V}_{\beta} + \mathbf{U})^{-\frac{1}{2}} \exp\left[-\frac{1}{2}(\hat{\beta} - \mu)^T (\hat{V}_{\beta} + \mathbf{U})^{-1} (\hat{\beta} - \mu)\right]}{\det(\hat{V}_{\beta})^{-\frac{1}{2}} \exp\left[-\frac{1}{2}\hat{\beta}^T \hat{V}_{\beta}^{-1} \hat{\beta}\right]},$$

we first consider the term inside the exponential function:

$$\mathcal{E}(\hat{\beta}, \mu, \hat{V}_{\beta}, \mathbf{U}) = \frac{1}{2} \hat{\beta}^T \hat{V}_{\beta}^{-1} \hat{\beta} - \frac{1}{2} (\hat{\beta} - \mu)^T (\hat{V}_{\beta} + \mathbf{U})^{-1} (\hat{\beta} - \mu).$$

Since \hat{V}_{β} and \mathbf{U} are typically defined through Kronecker products of smaller matrices, their inverses are easier to compute than the inverse of their sum. Hence we use Woodbury matrix identity to write

$$\mathcal{E}(\hat{\beta}, \mu, \hat{V}_{\beta}, \mathbf{U}) = \frac{1}{2} \hat{\beta}^T \hat{V}_{\beta}^{-1} \hat{\beta} - \frac{1}{2} (\hat{\beta} - \mu)^T \left(\hat{V}_{\beta}^{-1} - \hat{V}_{\beta}^{-1} (\mathbf{U}^{-1} + \hat{V}_{\beta}^{-1})^{-1} \hat{V}_{\beta}^{-1} \right) (\hat{\beta} - \mu).$$

To simplify the determinant calculation we write

$$\det(\hat{V}_{\beta} + \mathbf{U}) = \det(\hat{V}_{\beta}) \det(\mathbf{I} + \hat{V}_{\beta}^{-1} \mathbf{U}).$$

The logarithm of the Bayes Factor is then

$$\log(\text{BF}_{\text{MRP}}) = -\frac{1}{2} \log\left(\det(\mathbf{I} + \hat{V}_{\beta}^{-1} \mathbf{U})\right) + \mathcal{E}(\hat{\beta}, \mu, \hat{V}_{\beta}, \mathbf{U}).$$

If studies do not share individuals, \widehat{V}_{β} is a block-diagonal matrix

$$\widehat{V}_{\beta} = \begin{bmatrix} \widehat{V}_{\beta}^1 & 0 & \cdots & 0 \\ 0 & \widehat{V}_{\beta}^2 & \cdots & 0 \\ \vdots & & \ddots & \vdots \\ 0 & 0 & \cdots & \widehat{V}_{\beta}^S \end{bmatrix}.$$

If studies share individuals, e.g., controls, we take the approach of Cichonska et al. [4] to use summary level data to estimate the correlation structure of the non-diagonal blocks caused by overlapping individuals.

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