#### Organisation of gene programs revealed by unsupervised analysis of diverse

2	gene-trait	associations
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# 35 ABSTRACT

36 Genome wide association studies provide statistical measures of gene-trait associations that reveal how genetic variation influences phenotypes. This study develops an unsupervised 37 38 dimensionality reduction method called UnTANGLeD (Unsupervised Trait Analysis of 39 Networks from Gene Level Data) which organises 16,849 genes into discrete gene programs 40 by measuring the statistical association between genetic variants and 1,393 diverse complex 41 traits. UnTANGLeD reveals 173 gene clusters enriched for protein-protein interactions and highly distinct biological processes governing development, signalling, disease, and 42 43 homeostasis. We identify diverse gene networks with robust interactions but not associated with known biological processes. Analysis of independent disease traits shows that 44 45 UnTANGLeD gene clusters are conserved across all complex traits, providing a simple and powerful framework to predict novel gene candidates and programs influencing orthogonal 46 47 disease phenotypes. Collectively, this study demonstrates that gene programs co-ordinately 48 orchestrating cell functions can be identified without reliance on prior knowledge, providing a 49 method for use in functional annotation, hypothesis generation, machine learning and 50 prediction algorithms, and the interpretation of diverse genomic data.

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#### 59 INTRODUCTION

60 Generation of consortium-scale data such as ENCODE (1), the Human Cell Atlas (2) and the 61 UKBiobank (3) coupled with the development of advanced computational methods is enabling 62 the creation of transformative models that harness the natural diversity of biological systems. 63 These models draw on the relationships and patterns derived from biological data to establish 64 quantitative frameworks that can make highly accurate predictions, with implications for nearly 65 every field of biology. For example, in the field of structural biology, patterns in the sequences 66 and structures of proteins' evolutionary homologs reveal how amino acids interact, enabling 67 prediction of protein structure with atomic accuracy (4). Similarly, patterns of repressive 68 histone methylation (H3K27me3) across hundreds of human cell types enable identification of 69 genes governing cell decisions and functions for any cell type and organism (5). 70

Genome wide association studies (GWAS) characterise the genomic variation underlying complex traits and diseases, providing insights into how genes affect biological processes (6). Despite the wealth of variant-trait association information, GWAS studies predominantly focus on elucidating the genetic basis of a single trait or a group of highly related traits (6, 7). Here, we utilize patterns of genomic variation across hundreds of diverse phenotypes as the basis for an unsupervised method to parse the organisation of gene programs in cells.

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We hypothesised that complex traits are underpinned by conserved gene programs that can be identified by studying associations between genetic variation and phenotypic variation. To test this, we developed UnTANGLeD (Unsupervised Trait Analysis of Networks from Gene Level Data), which identifies patterns of association between genes and hundreds of diverse phenotypes. UnTANGLeD creates a phenotypic signature to cluster genes with similar associations across many traits in an unsupervised manner into gene programs controlling cell biological processes (Figure 1).

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We used a gene-trait association matrix derived from GWAS data for 1,393 complex traits to infer co-ordinately acting gene programs that represent both known and novel biological processes. While the scale of associations available from public GWAS data is underpowered to saturate the accuracy of our model, we demonstrate that UnTANGLeD can be applied to any orthogonal GWAS data to predict the genetic basis of disease including in underpowered and transethnic GWAS data. UnTANGLeD provides a powerful analytical framework for studies in population genetics, cell biology, and genomics, that will improve as more data emerges.

93 Collectively, this study provides a statistical framework for defining genes orchestrating

- 94 biological processes by evaluating genetic signatures across diverse complex traits.
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# 96 MATERIALS AND METHODS

# 97 Data Collection

98 S-MultiXcan results for 1,393 phenotypes with statistically significant SNP-based heritability 99 (p < 0.05) were downloaded from CTG-VL (<u>http://vl.genoma.io</u>). Phenotypes are listed in 100 **Table S2**. SNP-based heritability was estimated using linkage disequilibrium score regression 101 (LDSR). The significance values reflecting the strength of the association between each gene 102 and trait across all tissues were compiled into a gene-trait association matrix.

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# 104 Dimensionality Reduction Analysis Pipeline

All genes with fewer than 2 significant associations across all phenotypes ( $p < 10^{-4}$ ) were 105 106 removed, leaving 16849 genes. Following this, all values in the gene-trait association matrix 107 were chi-squared transformed. Infinite values produced when transforming very small p-value 108 (<1e-300) due to floating point precision were replaced with 1,415, which was 5 greater than 109 the largest non-infinite value. The data was then normalised by the sum of chi-squared values 110 per phenotype and scaled by a factor of 10,000. 10 principal components were retained from 111 the principal component analysis (PCA). Clustering of genes was performed using the native 112 Seurat shared-nearest neighbour algorithm. Clustering iterations were performed at increasing 113 resolutions from 0.2 to 20 in increments of 0.2. The resolution is a parameter from Seurat where 114 increased values lead to a greater number of clusters. Cluster assignments were compiled into a consensus distance matrix, where each gene pair had a value representing how often they 115 116 were grouped together out of 100 potential matches. 100 was then subtracted from the values 117 and they were made absolute to transform the matrix into a dissimilarity matrix. Agglomerative clustering using Ward's minimum variance method, as implemented in the stats package, was 118 119 applied to the consensus matrix directly. The average silhouette score (a metric used to 120 calculate how well a data point relates to its cluster) across all genes was calculated using the 121 *cluster* package from 2 to 300 clusters. The *inflection* package was used to calculate the plateau 122 point, which was determined to be the optimal number of clusters. Pearson's correlation was 123 used to determine the correlation of a gene with the other genes in the same cluster based on 124 chi-squared association values.

#### 126 Enrichment Analyses

127 GO, DO, KEGG enrichment, colocalization and tissue specificity enrichment were performed using *ClusterProfiler* (8). An FDR corrected significance value of p < 0.01 was used. 128 129 Colocalization was determined using *ClusterProfiler* enrichment for the Molecular Signatures 130 Database collection 3: positional gene sets (9). The largest proportion of genes within a cluster belonging to a single genomic region was divided by the total number of genes within the 131 132 cluster to identify the maximum degree of colocalization. STRING enrichment analysis was performed using the STRINGdb package, with a significance threshold of p < 0.001 and a 133 134 confidence threshold of 0.400. STRING enrichment analysis without the text-mining 135 component was performed using the online STRING interface (https://string-db.org/) for 136 clusters found to have PPI enrichment in the prior analysis with a confidence threshold of 0.150 137 to preserve predicted interactions reinforced by other components. For the calculation of the 138 correlation between the loss of enrichment and the degree of colocalization, clusters 111 and 173 were removed due to having well established biological functions despite being highly 139 140 colocalised. Broad enrichment analysis for more specialised gene sets was performed using 141 EnrichR (https://maayanlab.cloud/Enrichr/) across all 192 libraries. Redundant libraries, including GO, KEGG, chromosomal location and NIH-grant associated libraries were 142 excluded. The top significant term from each library for each cluster are reported in Table S9. 143 144 A significance value threshold of 0.01, after correction for multiple testing, was used. For identification of genes possessing the same protein domains or belonging to the same family, 145 the EnrichR library 'Pfam Domains 2019' was used. A distinct protein family or domain was 146 147 defined by collating the family or domain terms together that shared genes until there was no overlap between them. Protein terms did not need to be significantly enriched, but two or more 148 149 members of a protein family had to be present in a single cluster.

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#### 151 **Permutations**

Five permutations were generated by re-ordering the values within the gene-trait association matrix. These permutations were analysed as described above. A one-way ANOVA with FDR corrected pairwise comparisons was performed to identify significant differences in the number of enriched clusters, total enriched GO terms and the most significant GO enrichment of any cluster.

#### 158 Phenotype Associations

The gene-trait association matrix containing p-values was -log10 transformed. All infinite values generated due to floating point precision were windsorized with 315, which was 5 greater than the maximum finite value. The phenotypic associations for the genes within a cluster were extracted, averaged, normalised for their average associations across the dataset and ranked.

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#### 165 Clustering quality in dimensionality reduction methods

We extracted the UMAP coordinates for all genes as calculated by *Seurat*. Following this, we identified the 10 closest neighbours for each gene and calculated the average correlation of chisquared association values between the gene and its neighbours. The UMAP was re-plotted representing the average correlation with each point colour. We repeated the process, instead colouring by the number of significant associations for each gene.

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# 172 Prediction of novel genes using an underpowered GWAS of the same trait

173 Data collection and S-MultiXcan Analysis

174 We selected 13 phenotypes for which GWAS studies had been performed at differing cohort 175 sizes or ethnicities for the same, or comparable traits. The specific studies and their respective 176 details can be found in Table S1. Summary statistics were downloaded from various sources 177 and harmonised using MetaXcan's in-built harmonization (https://github.com/hakyimlab/MetaXcan) to be compatible with the MASHR models. We then 178 179 performed S-MultiX can analysis of each trait using the MASHR models built off the V8 GTEx 180 release. Associated genes were defined as those found to have a significance of  $p < 10^{-4}$ 181 by S-MultiXcan.

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#### 183 Global Clustering Coefficient Calculation

The genes identified for an independent GWAS were projected onto the 173 identified clusters. Following this, we generated an unweighted adjacency matrix in which genes in the same cluster were represented by a 1, and genes in different clusters by a 0. A comparison between the same gene was represented by 0. Finally, the global clustering coefficient (GCC) for the genes was calculated. To derive a statistical significance, we randomly sampled the same number of genes as there were significant genes for the phenotype and calculated the GCC onehundred times. A Z score was calculated from the curve generated by the sampled values.

#### 192 Gene Prediction

193 We took a simple approach of predicting which clusters were associated with the trait using the S-MultiXcan associations from the smaller GWAS and then checking whether novel gene 194 195 associations identified by the larger GWAS were in those clusters. A chi-squared enrichment 196 test was used where the minimum expected frequency was greater than 5, and a fisher's test if 197 not. Several approaches to predict clusters associated with the trait were trialled. The first was 198 to identify any of the 173 clusters with a significant gene in it. The second was to integrate the 199 additional phenotype into the trait-gene association matrix. Next, clusters were identified which 200 had an overall significance signature > 1.5 times the average or were significantly (p < 0.05) 201 higher than the average signature. Different values were tested for these thresholds, with these 202 providing the best performance. The third approach was to predict associated clusters from the previously established 173 clusters using the thresholds taken in approach two. A one-way 203 204 ANOVA was performed with pairwise comparisons to determine the best approach. Approach 205 three was the most effective, albeit not significantly, while maintaining a low computational 206 burden. In instances where transethnic GWAS were compared, the East-Asian GWAS was 207 used to predict the trait relevant clusters, and the European GWAS was used as the test set.

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#### 209 Gene Prioritization Analysis

The GWAS with the largest sample size for each of the 13 traits listed in **Table S1** was used to determine the potential of our pipeline for prioritizing genes within a locus. Clumping was performed on each summary statistic using PLINK (https://www.cog-genomics.org/plink/) and 1000 genomes phase 1 genotype data with an LD threshold of 0.5. This was followed by clumping for long distance LD, at the same threshold. Next, we identified individual 500kb regions around the lead SNPs and the genes within that region.

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We took a leave one chromosome out (LOCO) approach, where we removed all potential genes on one chromosome. With the remaining genes, we identified which clusters were enriched for genes associated with the trait. To calculate enrichment, we treated all genes associated with one locus as one positive, so that enrichment was for different loci and not genes at the same locus. A fisher's enrichment test was used to determine significance. Finally, we assessed at what proportion of loci the UnTANGLeD clusters identified a gene when that chromosome was left out of the analysis.

#### 225 Normalisation

We trialled relative count, centralised-log ratio and logarithmic normalisation on the chisquared transformed values of the gene-trait matrix across phenotypes. We evaluated their effects on the following metrics: correlation score, silhouette score, GO and STRING enrichment, global clustering coefficient, prediction of GWAS. A Kruskal Wallis one-way analysis of variance was used to evaluate differences. Relative count was used for the final pipeline.

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# 233 Phenotype Filtering Based on Euclidean Distance

A distance matrix between phenotypes using chi-squared transformed, RC-normalised data was generated using the Euclidean distance formula from the package *wordspace*. Phenotypes with a Euclidean distance below a set threshold, which indicated a high degree of relatedness, were removed from the data, leaving the phenotype with the highest number of significant associations. This was performed for thresholds 0 to 62, at which too few phenotypes remained to cluster the genes using the dimensionality reduction methods. GO enrichment was used to evaluate the clustering efficacy at each threshold.

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# 242 Phenotype Subsampling and Sensitivity Analyses

243 Phenotype subsampling was performed on two datasets; MultiXcan results for 1393 phenotypes 244 across 16,849 genes generated in this paper, and another dataset containing MultiXcan results 245 for 4091 phenotypes across 15,734 genes (phenomexcan.org). For the data containing 1393 246 phenotypes, subsampling was performed randomly without replacement from 50 to 1393 247 phenotypes in 20 equal increments across 5 replicates for each number of traits. The full UnTANGLeD clustering pipeline was applied to each subsampled matrix. Adjusted rand index 248 249 (ARI) was calculated for each of the subsampled clustering configurations compared to the full 250 dataset. This analysis was repeated for the data containing 4091 phenotypes; however, 251 subsampling was performed from 50 phenotypes to 4091 phenotypes in 50 equal increments.

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#### 253 Cluster Conservation

To explore the cause for the marked increase in ARI between 1322 phenotypes and 1393 phenotypes, cluster conservation was calculated between them. For each cluster from 1393 phenotypes, the proportion of genes that remained grouped together in each of the clusters from

257 1322 phenotypes was calculated. That proportion was used to assign a conservation score to

each gene, depending on how large the proportion of cluster the specific gene remained with

259 was. The same approach was applied between 4091 phenotypes and 4009 phenotypes.

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# 261 **RESULTS**

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# 263 Unsupervised identification of gene groups with shared complex trait associations

We used MultiXcan results from CTG-VL (10) derived from publicly available GWAS 264 265 (primarily from UK Biobank, on ~400,000 individuals) to create a gene-trait association matrix 266 for 16,849 genes and 1,393 traits (Figure 1, Figure S1, Table S2). For each gene trait pair, 267 MultiXcan predicts whether trait-associated variants alter the gene's expression. The chi-268 squared transformed significance value for each gene-trait association pair was compiled into 269 the gene-trait association matrix (Figure 2A). These values were normalised using relative 270 count normalisation to account for the difference in power between phenotypes. Performance 271 was not significantly different using other normalisation methods including centralised log ratio 272 or log normalised data (Figure S2). The data was then clustered using *Seurat*, a dimensionality 273 reduction method commonly used to analyse single cell RNA sequencing data to cluster cells 274 into related groups (11). Here, we use Seurat to test whether the calculated gene-trait 275 associations could be simplified into biologically enriched gene clusters. Clustering was 276 performed across 100 stepwise increases in resolution, a parameter which increases the number 277 of gene clusters. Repeat iterations provided an opportunity to survey both the broad scope of 278 biological processes that could be identified, as well as the specificity that could be achieved 279 with each biological process (Figure 2B).

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281 To test the biological validity of the derived clusters, we used positive gene sets as defined by 282 gene ontology (GO) (12) and STRING (13) to show that gene clusters have significant 283 enrichment for GO biological processes and STRING protein-protein interactions (Figure 2C 284 and 2D). To demonstrate that the observed enrichment is driven by distinct gene-trait 285 association signatures rather than chance, we performed permutation analyses in which the 286 values in the data matrix were randomly re-ordered. Permutations had significantly fewer 287 enriched clusters, GO terms and a lower strongest significance compared to the real data (p < p4x10<sup>-27</sup>) (Figure 2C, Figure S3A-B). Furthermore, we validated that GO enriched clusters 288 289 were more likely to also have enrichment for protein-protein interactions, suggesting the 290 enrichment is robust (Figure 2D). This analysis revealed that genes possessing similar

associations to complex trait phenotypes cluster meaningfully into biologically enriched groupsand the enrichment is not stochastic.

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294 We next developed an ensemble learning method we call "consensus clustering" that 295 incorporates a measure of clustering robustness and quality. Across each of 100 stepwise 296 increases in clustering resolution we evaluated the robustness of clustering by assessing how 297 often every possible gene combination was clustered together ranging from 100 (always) to 0 298 (never) and compiled these values into a consensus matrix (Figure 2E). Following this, we 299 performed agglomerative hierarchical clustering, evaluating the average silhouette score at 300 each possible number of clusters. The silhouette score quantifies how consistent genes within 301 the same cluster are across Seurat resolutions. To derive the optimal number of gene clusters, 302 we calculated the plateau point of the average silhouette score, which informs the number of 303 clusters at which point further splitting no longer improves the stability of clustering 304 assignments (Figure S3C). Applying this methodology to gene-trait associations for 16,849 305 genes, we identified 173 clusters with an average of 97 genes (Figure 2F, G). Across each 306 cluster, we measured the silhouette score, a metric of cluster robustness and the correlation 307 score, a metric of relation across phenotypes, thereby providing two metrics to quantify the 308 quality of clustering (Figure 2H). Collectively, we call this approach UnTANGLeD: 309 Unsupervised Trait Analysis of Networks from Gene Level Data.

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#### 311 Consensus clustering identifies robust gene groups enriched for known gene sets

312 We analysed each cluster by reference to curated annotations of gene programs (GO, disease 313 ontology (14)), signalling pathways (KEGG (15)), protein-protein interactions (STRING), and 314 tissue specificity (16) to evaluate the ability of UnTANGLeD to identify distinct, biologically 315 established gene programs in an unsupervised manner (Figure 3A, Figure S3D, Tables S3-8). 316 This analysis revealed significant enrichment of cell biological pathways and networks across 317 gene clusters, with stronger enrichment among clusters with higher silhouette and correlation 318 scores (Figure 3A). We further performed enrichment analysis of the UnTANGLeD clusters 319 using the EnrichR database (17) (Figure S4A-C, Table S9), finding considerable enrichment 320 for disease-associated genes, gene-expression perturbations associated with disease states or 321 drugs and protein domains and families. We note that although many clusters contain multiple 322 members of a protein family (18) the proportion of any one protein family in the cluster is 323 minor (Figure S4D, Table S10).

325 We next investigated the relationship between individual gene clusters and the traits most 326 strongly influencing the genes within the clusters, using enriched GO processes as a proxy for the functional profile of a cluster (Figure 3B). Each cluster is defined by a distinct gene-trait 327 328 association 'signature' indicated by the variation and strength of association across 1,393 329 diverse complex traits. In some instances, the enriched biological processes for certain gene clusters are clearly related to the cluster's most significantly associated complex trait 330 phenotypes (e.g., cluster 119: GO enrichment: Cholesterol Homeostasis; Dominant complex 331 332 trait phenotypes: Low-density lipoprotein, Alipoprotein B quantile).

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334 Since UnTANGLeD draws on associations across diverse phenotypes to inform gene-gene 335 relationships, the method can identify gene groups with enriched functions that are apparently 336 biologically independent of the phenotypes most significantly associated with the genes in the cluster. For example, cluster 80, enriched for embryonic morphogenesis (GO:004859), is most 337 338 significantly associated to the phenotype Bone Mineral Density and cluster 111, enriched for 339 nucleosome organisation (GO:0034728), is most significantly associated to the phenotype 340 Mean Corpuscular Haemoglobin. These results support the central hypothesis that genes with 341 shared effects across diverse phenotypes can be clustered into gene groups controlling shared 342 biological functions and processes in an unsupervised manner (Figure 3B).

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Importantly, we show that the GO enriched gene clusters show no overlap in their strongest enriched biological functions, and almost no overlap in their top 5 enriched terms, demonstrating the use of gene-trait association data to parse novel biological gene programs encoded within the genome (**Figure 3C**).

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Stratifying clusters by their silhouette and correlation scores reveals a higher level of GO, STRING, KEGG, DO and tissue specificity enrichment with higher clustering quality, indicating that the metrics provide an accurate representation of cluster quality (**Figure 3A, D**). Furthermore, both the robustness of clustering and the presence and strength of GO and STRING enrichment are correlated with the number of significant associations to phenotypes per gene (Pearson's correlation, r > 0.65), as well as the stability of clustering (Pearson's correlation, r > 0.69) (**Figure 3E-G, Figure S5A-F**).

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Lastly, we note that there is considerable colocalization of genes within clusters, with a stronger
 relationship between the correlation score and the degree of colocalization for the genes in a

359 cluster (Pearson's correlation, r = 0.77), than the cluster robustness (Pearson's correlation, r =360 0.34) (Figure 3H, Table S11). STRING enrichment may also be inflated due to the textmining component, as findings from GWAS may be incorporated into the database, with genes 361 362 in proximity often being reported together. Indeed, we find that the loss of enrichment due to 363 removal of the text-mining component is correlated with the colocalization of the cluster (r =0.60) (Figure S6A-B). However, clusters with a high degree of colocalization are not 364 necessarily artefacts of false-positive associations identified by MultiXcan. For example, 365 clusters 173 and 111 are strongly enriched for immune processes and chromatin organisation 366 367 respectively, despite being highly colocalised (Figure S6C-D).

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#### 369 Subsampling reveals need for more data to improve accuracy of UnTANGLeD

370 We next sought to determine how the number and diversity of phenotypes influences the 371 accuracy and utility of UnTANGLeD clusters. We show that the number of GO enriched 372 clusters is highly correlated with the number of phenotypes utilised in the analysis (Pearson's 373 correlation, r = 0.85), even when phenotypic diversity is preserved (Figure S7A). To further 374 test this, we performed phenotype subsampling and evaluated clustering accuracy using an 375 adjusted rand index (ARI) analysis. We found that clustering accuracy compared to the full 376 data improved with the addition of more phenotypes, however a marked increase in ARI 377 between 1322 and the full data set suggests that inaccuracy in clustering that isn't determined 378 by phenotypic diversity can be attributed to genes which have weak signatures and few 379 significant associations (Figure S7B). We repeated subsampling in a larger dataset containing 380 MultiXcan analysis of 4091 phenotypes retrieved from Pividori et al. (2020) which resulted in 381 the same outcome (Figure S7C). Comparison of the two data sets revealed that genes already 382 having many significant associations simply had more associations in the larger dataset with 383 both datasets possessing an equal proportion of genes with few to no significant associations 384 (Figure S7D). Further, that genes with higher numbers of significant associations have higher 385 degrees of conservation (Figure S7E-F). It's likely that the effective number of traits is similar 386 between the two datasets, as both mostly draw on the UK Biobank and have many highly 387 correlated phenotypes

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389 Cumulatively, these findings indicate that the quality of gene clustering is dependent on the 390 scale and quality of data needed to derive high silhouette and correlation scores as a basis for

391 efficient enrichment of functional gene clusters. Accordingly, as more data becomes available,

392 the quality and accuracy of UnTANGLeD will improve. However, simply increasing the

number of phenotypes leads to an increase in redundant associations, and therefore strategies
 to increase the number of significant gene-trait associations across the genome should be
 employed, such as diversifying phenotypes and increasing cohort size.

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# 397 UnTANGLeD clusters are conserved across traits and can predict novel trait associated 398 genes

GWAS require collections of large cohorts comprising thousands of individual-level genotype
data to characterise the genetic architecture of a trait. Furthermore, collecting enough samples
can prove challenging for many diseases, and as such they are often underrepresented in
biobanks.

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We hypothesised that UnTANGLeD gene clusters would be conserved across complex traits. 404 405 To test this, we investigated an independent GWAS of ulcerative colitis (UC) (19) (Figure 4A). We show that the 278 genes associated with UC ( $p < 10^{-4}$ ) (Figure 4B) were significantly more 406 clustered within the UnTANGLeD clusters than expected by chance ( $p = 2x10^{-9}$ ) (Figure 4C). 407 408 The result shows that despite not being used to construct the clusters, UC associated genes 409 nevertheless group within the UnTANGLeD clusters, demonstrating that the defined gene programs are conserved. We replicate our findings in 6 additional independent GWAS 410 phenotypes, highlighting that the UnTANGLeD clusters are conserved across a broad 411 412 phenotypic space (3, 20–28) (Figure 4G).

413

414 We next tested whether the gene clusters can be used to predict novel genes and cellular 415 processes underpinning independent complex trait data. To test this hypothesis, we examined two GWAS for UC. The first was performed in 2013 with 6,687 cases and 19,718 controls (29), 416 and the latter in 2017 with 12,366 cases and 33,609 controls (19) (Figure 4A). MultiXcan 417 418 analysis of the summary statistics identified 153 and 278 genes respectively, with an overlap 419 of 53 genes (Figure 4B). We projected the MultiXcan associations for the 2013 GWAS onto 420 the 173 clusters, identifying clusters were statistically associated with UC (Figure S8). Finally, 421 we tested whether the clusters predicted from the 2013 GWAS contained novel genes identified 422 by the 2017 GWAS. Of the 225 novel genes identified by the 2017 GWAS, our approach was 423 able to use the 2013 GWAS to predict 120 with a significant enrichment for predicting UC associated genes compared to other genes ( $p < 3x10^{-121}$ , chi-squared test) (Figure 4D). 424

426 GWAS of the same complex trait conducted in populations of differing ancestries may 427 implicate both shared and distinct loci. We tested whether UnTANGLeD clusters are conserved for genes specific to non-European ancestries, given that the UnTANGLeD gene clusters are 428 429 built from a European cohort. To test this, we examined a GWAS for triglyceride levels in an 430 East Asian population, which identified 34 genes (30) (Figure S9A-B). Mirroring our findings in a GWAS conducted on a European population, we found that the genes associated with 431 432 triglyceride levels in an East Asian population are significantly more clustered than expected 433  $(p = 1x10^{-9})$  and replicate this finding in 4 other GWAS conducted in populations of non-434 European ancestry (30–32). We further tested whether the GWAS conducted in the East Asian 435 cohort could be used to predict novel genes identified in a European cohort. We found that 436 clusters implicated in triglyceride levels using the East Asian GWAS were highly enriched for 437 genes identified by the European GWAS ( $p = 6x10^{-109}$ ) (Figure 4F).

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All together, we show significant enrichment for prediction of novel genes across GWAS
performed for 7 traits in differing cohort sizes in a European population, and 4 traits for which

- 441 GWAS were performed in different ancestries (3, 20-28, 30–33) (Figure 4G, Figure 89C).
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443 We further tested whether the UnTANGLeD clusters could be used to prioritize causal genes 444 at any given locus. It is difficult to accurately identify the causal genes from GWAS identified 445 variants due to linkage disequilibrium and complex regulatory effects of intergenic variants. 446 For each independent trait, we identified potential gene candidates within 500kb of each 447 independent significant SNP then took a leave one chromosome out approach (LOCO) to investigate whether genes on the removed chromosome would be implicated in the clusters 448 449 associated with the remaining genes. (Figure 4H). We are able to identify a major proportion 450 of loci independently across all traits and reduce the potential candidates at each locus 451 considerably, further highlighting the utility of UnTANGLeD (Figure 4I).

#### 453 **DISCUSSION**

This study demonstrates that gene programs governing biological processes can be identified without reliance on prior knowledge, by analysing the association between genetic variation and a large range of diverse complex traits. Several prior studies have constructed small gene networks using a limited number of disease phenotypes and their associated genes from curated GWAS databases and restricted sources of rare genetic variants. Other studies, like PheWAS (34, 35) and PhenomeXcan (36) have collated genomic associations across numerous phenotypes to create resources of variant-trait and gene-trait associations.

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462 Here, we construct a gene-trait association matrix for 16,849 genes across 1,393 complex traits 463 similarly to PhenomeXcan, and further the concept by using UnTANGLeD to identify gene programs. We apply dimensionality reduction methods, which can harness the high 464 465 dimensional, complex gene-trait association data, allowing us to greatly expand on the scale of 466 studies previously attempting to build gene networks. By increasing the scale of data, we not 467 only identify gene programs enriched for biological processes specific to associated phenotypes 468 but also reveal gene programs enriched for central processes governing diverse mechanisms of 469 cellular development and homeostasis.

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471 The UnTANGLeD framework is a powerful approach to identify gene programs orchestrating 472 key biological processes. We implicate novel genes in clusters enriched for known processes 473 and identify numerous novel gene programs with enrichment for protein-protein interactions 474 and no known function. We further highlight the utility of UnTANGLeD for hypothesis 475 generation and functional annotation of genes, which may be particularly valuable for non-476 coding genes, as they are notoriously difficult to annotate in silico (37). Finally, the 477 UnTANGLeD framework reveals relationships between complex traits, linking phenotypes by 478 the gene programs that underpin them.

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We demonstrate the utility of UnTANGLeD for predicting genes associated with complex traits and diseases using a low-powered GWAS of the same trait. Currently, standard methods use gene-set analysis to improve power to identify genes and pathways involved in a phenotype, such as MAGMA, or GIGSEA (38–41). Our method eliminates the need to define gene-sets and instead uses gene-trait association data to learn gene sets governing complex traits (39), enabling us to implicate novel trait associated genes and loci from a much smaller cohort size.

487 We further highlight the use of the UnTANGLeD clusters for gene prioritization, showing that 488 they effectively select gene candidates at different loci related to the same phenotype. Current 489 gene prioritisation approaches use either distance-based metrics or mapping to eQTLs to predict 490 changes in gene expression (42). However, these also suffer from a considerable false positive 491 rate and may not always distinguish between two genes in proximity, as noted in our data (42). 492 Some recent methods have integrated biological data, such as gene sets, RNA sequencing and protein-protein interaction databases to further prioritise genes at a locus (43). Our framework 493 494 can be used independently or integrated with any of these approaches to advance understanding 495 of complex trait biology.

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497 Outside of its utility in GWAS analyses, UnTANGLeD may provide key mechanisms for data 498 analysis in medical and industry pipelines including genetic testing and drug discovery. For 499 example, polygenic risk scores (PRS) are an emerging method that evaluate an individual's 500 disease risk from genetic variants (44). Methods such as UnTANGLeD may help reveal genes 501 and hence genetic variants governing cell programs underlying disease risk and hence improve 502 prediction accuracy. In the context of pharmacogenomics, studies have shown that drug targets 503 with genetic support from either rare or common diseases are more than twice as likely to pass 504 through clinical trials (45, 46). Since UnTANGLeD captures gene programs associated with all 505 complex traits and diseases, its predictive power may help de-risk candidates and thereby 506 decrease cost associated with the drug discovery pipeline. Overall, UnTANGLeD represents a 507 powerful and versatile framework for studying cellular gene programs to interpret diverse 508 sources of orthogonal genetic data.

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510 We note several limitations in our method. Primarily, that the current GWAS data does not 511 represent the whole phenome. Furthermore, many traits are highly correlated, and disease traits 512 are underrepresented in the UK Biobank, the main source of data in this study. Secondly, 513 UnTANGLeD relies on S-MultiXcan to construct the gene-trait association matrix. While S-514 MultiXcan is powered to detect associations across all tissues, it suffers from a high false 515 discovery rate and may perform poorly in tissues with small sample sizes. Moreover, S-516 MultiXcan can identify genes colocalised with a causal gene as significant, which can obscure 517 biological signatures. Other approaches such as SMR MR-JTI may remedy this issue (47). 518 Additionally, UnTANGLeD does not account for the predicted directionality of effect or tissue-519 specific effects, which may help to further increase the quality and biological specificity of the 520 clusters. Biological validation of the method using established gene sets may be inflated due to

521 GWAS data being included in the definition of the gene sets. Finally, we note that although

522 UnTANGLeD is a powerful tool for identifying clusters in an unsupervised manner, the overall

- 523 function of the cluster may be difficult to determine. The development of improved gene-based
- 524 tests and emergence of larger GWAS data spanning the whole phenome will improve the
- 525 accuracy and utility of UnTANGLeD.
- 526

527 This study provides a powerful framework for the identification of gene programs governing 528 biological processes conserved across all complex traits and diseases, with important 529 applications for functional annotation, hypothesis generation, machine learning and prediction 530 algorithms and interpretation of GWAS and diverse other genomic data types. Our approach 531 can be applied to any collection of gene-trait information, harnessing the power of biological 532 patterns in a diverse landscape of phenotypic variation.

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# 540 DATA AND MATERIALS AVAILABILITY

- 541 All source code available on GitHub (<u>https://github.com/palpant-comp/UnTANGLeD</u>) and all
- data available on Zenodo (https://doi.org/10.5281/zenodo.6572617).

# 543 SUPPLEMENTARY MATERIALS

- 544 Supplementary Data are available at NAR online.
- 545

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- 550

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- 553

# 554 AUTHOR CONTRIBUTIONS

- 555
- 556 **DM:** Developed the study, performed all analyses, and wrote the manuscript
- 557 MS and CN: Helped supervise bioinformatics analysis
- 558 GCP: Helped supervise and design GWAS data selection and analysis, interpreted data, and
- 559 wrote the manuscript
- 560 NP: Conceived and supervised the project, raised funding, and wrote the manuscript
- 561

# 562 CONFLICT OF INTEREST STATEMENT

563 GCP is currently an employee of 23andMe Inc. and holds stock options for the company.

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#### 567 **REFERENCE**

- ENCODE Project Consortium (2012) An integrated encyclopedia of DNA elements in the
   human genome. *Nature*, 489, 57–74.
- 2. Rozenblatt-Rosen,O., Stubbington,M.J.T., Regev,A. and Teichmann,S.A. (2017) The
  Human Cell Atlas: from vision to reality. *Nat. 2017 5507677*, **550**, 451–453.
- Sudlow,C., Gallacher,J., Allen,N., Beral,V., Burton,P., Danesh,J., Downey,P., Elliott,P.,
   Green,J., Landray,M., *et al.* (2015) UK Biobank: An Open Access Resource for
   Identifying the Causes of a Wide Range of Complex Diseases of Middle and Old Age.
- 575 *PLoS Med.*, **12**, 1–10.
- 576 4. Jumper, J., Evans, R., Pritzel, A., Green, T., Figurnov, M., Ronneberger, O.,
- 577 Tunyasuvunakool,K., Bates,R., Žídek,A., Potapenko,A., *et al.* (2021) Highly accurate 578 protein structure prediction with AlphaFold. *Nature*, **596**, 583–589.
- 5. Shim,W.J., Sinniah,E., Xu,J., Vitrinel,B., Alexanian,M., Andreoletti,G., Shen,S., Sun,Y.,
  Balderson,B., Boix,C., *et al.* (2020) Conserved Epigenetic Regulatory Logic Infers Genes
  Governing Cell Identity. *Cell Syst.*, 11, 625-639.e13.
- Visscher, P.M., Wray, N.R., Zhang, Q., Sklar, P., McCarthy, M.I., Brown, M.A. and Yang, J.
   (2017) 10 Years of GWAS Discovery: Biology, Function, and Translation. *Am. J. Hum. Genet.*, 101, 5–22.
- 585 7. Bellomo,T.R., Bone,W.P., Chen,B.Y., Gawronski,K.A.B., Zhang,D., Park,J., Levin,M.,
  586 Tsao,N., Klarin,D., Lynch,J., *et al.* (2021) Multi-trait GWAS of atherosclerosis detects
  587 novel pleiotropic loci. *medRxiv*, 10.1101/2021.05.21.21257493.
- Yu,G., Wang,L.-G., Han,Y., He,Q.-Y. (2012) clusterProfiler: an R Package for Comparing
   Biological Themes Among Gene Clusters. *Omi. A J. Integr. Biol.*, 16, 284–287.
- 590 9. Liberzon, A., Subramanian, A., Pinchback, R., Thorvaldsdóttir, H., Tamayo, P. and
  591 Mesirov, J.P. (2011) Molecular signatures database (MSigDB) 3.0. *Bioinformatics*, 27,
  592 1739.
- 10. Cuellar-Partida,G., Lundberg,M., Kho,P.F., D'Urso,S., Gutierrez-Mondragon,L.F. and
  Hwang,L.-D. (2019) Complex-Traits Genetics Virtual Lab: A community-driven web
  platform for post-GWAS analyses. *bioRxiv*, 10.1101/518027.
- 596 11. Butler, A., Hoffman, P., Smibert, P., Papalexi, E. and Satija, R. (2018) Integrating single-cell
   597 transcriptomic data across different conditions, technologies, and species. *Nat.*
- 598 *Biotechnol.*, **36**, 411–420.

- 599 12. Ashburner, M., Ball, C.A., Blake, J.A., Botstein, D., Butler, H., Cherry, J.M., Davis, A.P.,
- 600 Dolinski,K., Dwight,S.S., Eppig,J.T., et al. (2000) Gene ontology: tool for the unification
- 601 of biology. The Gene Ontology Consortium. *Nat. Genet.*, **25**, 25–29.
- 13. Szklarczyk, D., Gable, A.L., Lyon, D., Junge, A., Wyder, S., Huerta-Cepas, J., Simonovic, M.,
  Doncheva, N.T., Morris, J.H., Bork, P., *et al.* (2019) STRING v11: protein–protein
  association networks with increased coverage, supporting functional discovery in genomewide experimental datasets. *Nucleic Acids Res.*, 47, D607–D613.
- 606 14. Schrimi, L.M., Arze, C., Nadendla, S., Wayne Chang, Y.-W., Mazaitis, M., Felix, V., Feng, G.
- and Kibbe,W.A. (2012) Disease Ontology: a backbone for disease semantic integration. *Nucleic Acids Res.*, 40.
- 609 15. Kanehisa, M. (2000) KEGG: Kyoto Encyclopedia of Genes and Genomes. *Nucleic Acids*610 *Res.*, 28, 27–30.
- 611 16. Jain, A. and Tuteja, G. (2019) TissueEnrich: Tissue-specific gene enrichment analysis.

*Bioinformatics*, **35**, 1966–1967.

- 613 17. Kuleshov,M. V, Jones,M.R., Rouillard,A.D., Fernandez,N.F., Duan,Q., Wang,Z.,
  614 Koplev,S., Jenkins,S.L., Jagodnik,K.M., Lachmann,A., *et al.* (2016) Enrichr: a
  615 comprehensive gene set enrichment analysis web server 2016 update. *Nucleic Acids Res.*,
  616 44, W90–W97.
- 617 18. El-Gebali,S., Mistry,J., Bateman,A., Eddy,S.R., Luciani,A., Potter,S.C., Qureshi,M.,
  618 Richardson,L.J., Salazar,G.A., Smart,A., *et al.* (2019) The Pfam protein families database
  619 in 2019. *Nucleic Acids Res.*, 47, D427–D432.
- 620 19. de Lange,K.M., Moutsianas,L., Lee,J.C., Lamb,C.A., Luo,Y., Kennedy,N.A., Jostins,L.,
  621 Rice,D.L., Gutierrez-Achury,J., Ji,S.G., *et al.* (2017) Genome-wide association study
  622 implicates immune activation of multiple integrin genes in inflammatory bowel disease.
  623 *Nat Genet*, 49, 256–261.
- 20. Köttgen,A., Albrecht,E., Teumer,A., Vitart,V., Krumsiek,J., Hundertmark,C., Pistis,G.,
  Ruggiero,D., O'Seaghdha,C.M., Haller,T., *et al.* (2013) Genome-wide association analyses
  identify 18 new loci associated with serum urate concentrations. *Nat Genet*, 45, 145–154.
- 627 21. Tin, A., Marten, J., Halperin Kuhns, V.L., Li, Y., Wuttke, M., Kirsten, H., Sieber, K.B., Qiu, C.,
- Gorski, M., Yu, Z., *et al.* (2019) Target genes, variants, tissues and transcriptional pathways
  influencing human serum urate levels. *Nat. Genet.*, **51**, 1459–1474.
- 630 22. Shah, S., Henry, A., Roselli, C., Lin, H., Sveinbjörnsson, G., Fatemifar, G., Hedman, Å.K.,
- 631 Wilk, J.B., Morley, M.P., Chaffin, M.D., et al. (2020) Genome-wide association and
- 632 Mendelian randomisation analysis provide insights into the pathogenesis of heart failure.

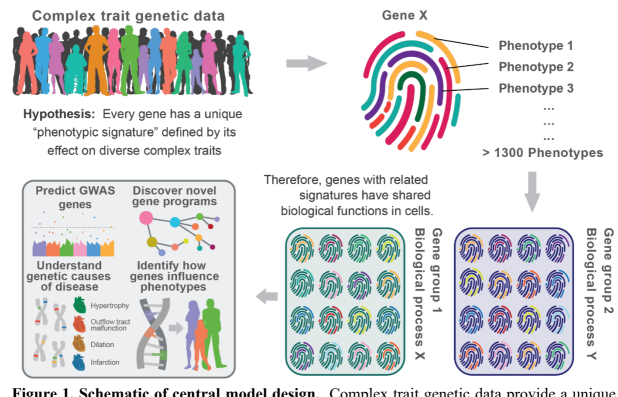
633 *Nat. Commun.*, **11**, 163.

- 634 23. Kathiresan, S., Melander, O., Guiducci, C., Surti, A., Burtt, N.P., Rieder, M.J., Cooper, G.M.,
- 635 Roos, C., Voight, B.F., Havulinna, A.S., et al. (2008) Six new loci associated with blood low-
- density lipoprotein cholesterol, high-density lipoprotein cholesterol or triglycerides in
  humans. *Nat Genet*, 40, 189–197.
- 638 24. Willer, C.J., Schmidt, E.M., Sengupta, S., Peloso, G.M., Gustafsson, S., Kanoni, S., Ganna, A.,
  639 Chen, J., Buchkovich, M.L., Mora, S., *et al.* (2013) Discovery and refinement of loci
  640 associated with lipid levels. *Nat. Genet.*, 45, 1274–1283.
- 641 25. Stahl,E.A., Raychaudhuri,S., Remmers,E.F., Xie,G., Eyre,S., Thomson,B.P., Li,Y.,
  642 Kurreeman,F.A., Zhernakova,A., Hinks,A., *et al.* (2010) Genome-wide association study
  643 meta-analysis identifies seven new rheumatoid arthritis risk loci. *Nat Genet*, 42, 508–514.
- 644 26. Okada, Y., Wu, D., Trynka, G., Raj, T., Terao, C., Ikari, K., Kochi, Y., Ohmura, K., Suzuki, A.,
- 645 Yoshida, S., *et al.* (2014) Genetics of rheumatoid arthritis contributes to biology and drug
  646 discovery. *Nature*, **506**, 376–381.
- 647 27. Schizophrenia Psychiatric Genome-Wide Association Study (GWAS)Consortium (2011)
  648 Genome-wide association study identifies five new schizophrenia loci. *Nat Genet*, 43, 969–
  649 976.
- 28. Schizophrenia Working Group of the Psychiatric Genomics Consortium (2014) Biological
  insights from 108 schizophrenia-associated genetic loci. *Nature*, **511**, 421–427.
- 652 29. Anderson, C.A., Boucher, G., Lees, C.W., Franke, A., D'Amato, M., Taylor, K.D., Lee, J.C.,
  653 Goyette, P., Imielinski, M., Latiano, A., *et al.* (2011) Meta-analysis identifies 29 additional
  654 ulcerative colitis risk loci, increasing the number of confirmed associations to
- 655 47. *Nat Genet*, **43**, 246–252.
- 30. Spracklen, C.N., Chen, P., Kim, Y.J., Wang, X., Cai, H., Li, S., Long, J., Wu, Y., Wang, Y.X.,
  Takeuchi, F., *et al.* (2017) Association analyses of East Asian individuals and transancestry
  analyses with European individuals reveal new loci associated with cholesterol and
  triglyceride levels. *Hum Mol Genet*, 26, 1770–1784.
- 31. Lam,M., Chen,C.-Y., Li,Z., Martin,A.R., Bryois,J., Ma,X., Gaspar,H., Ikeda,M.,
  Benyamin,B., Brown,B.C., *et al.* (2019) Comparative genetic architectures of
  schizophrenia in East Asian and European populations. *Nat. Genet.*, **51**, 1670–1678.
- 663 32. Wang, Y.-F., Zhang, Y., Lin, Z., Zhang, H., Wang, T.-Y., Cao, Y., Morris, D.L., Sheng, Y.,
- 664 Yin,X., Zhong,S.-L., *et al.* (2021) Identification of 38 novel loci for systemic lupus
- 665 erythematosus and genetic heterogeneity between ancestral groups. *Nat. Commun.*, **12**, 772.

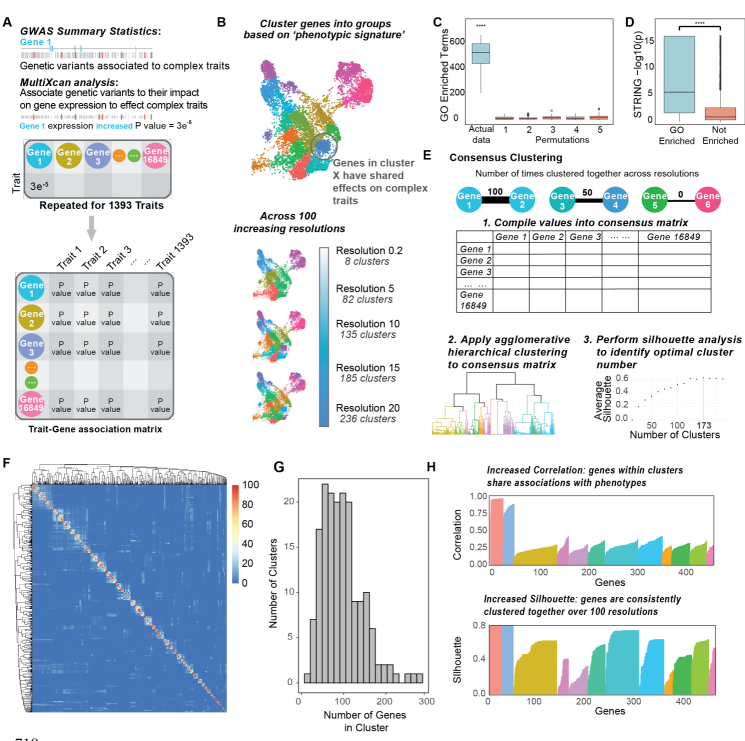
- 666 33. Bentham, J., Morris, D.L., Graham, D.S.C., Pinder, C.L., Tombleson, P., Behrens, T.W.,
- 667 Martín, J., Fairfax, B.P., Knight, J.C., Chen, L., et al. (2015) Genetic association analyses
- 668 implicate aberrant regulation of innate and adaptive immunity genes in the pathogenesis of
  669 systemic lupus erythematosus. *Nat Genet*, 47, 1457–1464.
- 34. Diogo, D., Tian, C., Franklin, C.S., Alanne-Kinnunen, M., March, M., Spencer, C.C.A.,
  Vangjeli, C., Weale, M.E., Mattsson, H., Kilpeläinen, E., *et al.* (2018) Phenome-wide
  association studies across large population cohorts support drug target validation. *Nat.*
- 673 *Commun. 2018 91*, **9**, 1–13.
- 674 35. Pendergrass,S.A., Buyske,S., Jeff,J.M., Frase,A., Dudek,S., Bradford,Y., Ambite,J.-L.,
  675 Avery,C.L., Buzkova,P., Deelman,E., *et al.* (2019) A phenome-wide association study
  676 (PheWAS) in the Population Architecture using Genomics and Epidemiology (PAGE)
  677 study reveals potential pleiotropy in African Americans. *PLoS One*, 14, e0226771.
- 678 36. Pividori,M., Rajagopal,P.S., Barbeira,A., Liang,Y., Melia,O., Bastarache,L., Park,Y.,
  679 Consortium,G., Wen,X. and Im,H.K. (2020) PhenomeXcan: Mapping the genome to the
  680 phenome through the transcriptome. *Sci. Adv.*, 6, eaba2083.
- 37. Perron,U., Provero,P. and Molineris,I. (2017) In silico prediction of lncRNA function using
  tissue specific and evolutionary conserved expression. *BMC Bioinformatics*, 18, 29–39.
- 38. Zhu,S., Qian,T., Hoshida,Y., Shen,Y., Yu,J. and Hao,K. (2019) GIGSEA: genotype
  imputed gene set enrichment analysis using GWAS summary level data. *Bioinformatics*,
  35, 160–163.
- 39. Zhu,X. and Stephens,M. (2018) Large-scale genome-wide enrichment analyses identify
  new trait-associated genes and pathways across 31 human phenotypes. *Nat. Commun.*(2018) *91*, **9**, 1–14.
- 40. de Leeuw,C.A., Mooij,J.M., Heskes,T. and Posthuma,D. (2015) MAGMA: Generalized
  Gene-Set Analysis of GWAS Data. *PLOS Comput. Biol.*, **11**, e1004219.
- 41. Sun,R., Hui,S., Bader,G.D., Lin,X. and Kraft,P. (2019) Powerful gene set analysis in
  GWAS with the Generalized Berk-Jones statistic. *PLoS Genet.*, 15.
- 42. Broekema, R. V., Bakker, O.B. and Jonkers, I.H. (2020) A practical view of fine-mapping
  and gene prioritization in the post-genome-wide association era. *Open Biol.*, 10.
- 43. Schaid, D.J., Chen, W. and Larson, N.B. (2018) From genome-wide associations to
  candidate causal variants by statistical fine-mapping. *Nat. Rev. Genet.*, 19, 491.
- 44. Lewis, C.M. and Vassos, E. (2020) Polygenic risk scores: From research tools to clinical
  instruments. *Genome Med.*, 12, 1–11.

- 699 45. Nelson, M.R., Tipney, H., Painter, J.L., Shen, J., Nicoletti, P., Shen, Y., Floratos, A.,
- Sham, P.C., Li, M.J., Wang, J., et al. (2015) The support of human genetic evidence for
- 701 approved drug indications. *Nat. Genet. 2015 478*, **47**, 856–860.
- 46. King,E.A., Wade Davis,J. and Degner,J.F. (2019) Are drug targets with genetic support
  twice as likely to be approved? Revised estimates of the impact of genetic support for drug
  mechanisms on the probability of drug approval. *PLOS Genet.*, 15, e1008489.
- 47. Zhou,D., Jiang,Y., Zhong,X., Cox,N.J., Liu,C. and Gamazon,E.R. (2020) A unified
   framework for joint-tissue transcriptome-wide association and Mendelian randomization
- 707 analysis. *Nat. Genet.*, **52**, 1239–1246.

#### 709 FIGURES



- 710
- Figure 1. Schematic of central model design. Complex trait genetic data provide a unique association signature for each gene which can be used to parse the genome into functionally
- 713 related gene sets.
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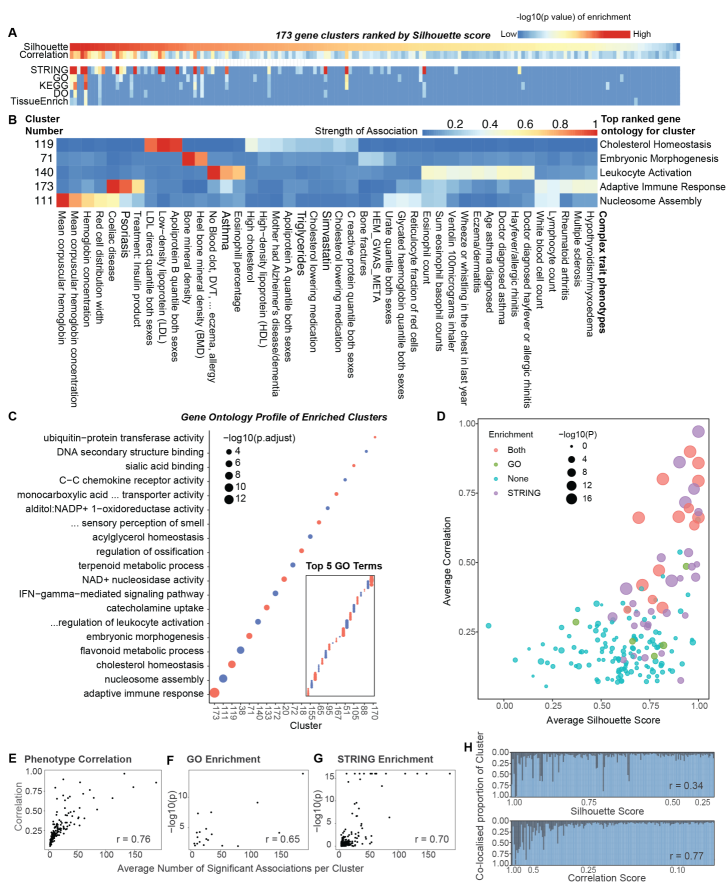
# 719 Figure 2. Consensus clustering method identifies biologically enriched gene clusters.

720 (A) MultiXcan analysis links genetic variants to genes by predicting changes in gene

- expression using eQTLs. The chi-squared values of the associations between each of 1393 traits
- and 25851 genes were compiled into a gene-trait matrix.
- 723 **(B)** Dimensionality reduction clustering of genes based on their phenotypic associations was
- performed using *Seurat* across resolutions 0.2 to 20 in 0.2 increments.

725 (C) Five permutations of the dataset were compared to the real data by the number of enriched

- gene ontology terms per resolution. Enrichment was performed using *ClusterProfiler*, FDR
   corrected p-value < 0.01. Pairwise comparisons between permutations were performed</li>
   with Wilcoxon signed rank test.
- (D) Validation of gene ontology enriched clusters with STRING protein-protein interaction
   enrichment. Wilcoxon signed rank test was used to compare STRING enrichment in gene
   ontology enriched and non-enriched clusters.
- 732 (E) Each gene pair is given a similarity score based on how often they were clustered together
- across 100 resolutions and these values are compiled into a consensus matrix.
  Agglomerative hierarchical clustering is applied to the matrix, with the plateau in the
  average silhouette score defining the optimal number of clusters.
- (F) Heatmap of consensus matrix as clustered using agglomerative hierarchical clustering for173 clusters.
- 738 (G) Histogram of the number of genes in each of the 173 clusters.
- 739 (H) Silhouette scores and correlation scores calculated for each gene to evaluate the clustering
- robustness and quality respectively. Data generated for 450 genes selected from 12 randomclusters.





# Figure 3. Enrichment of identified clusters for known gene sets is dependent on dataquality.

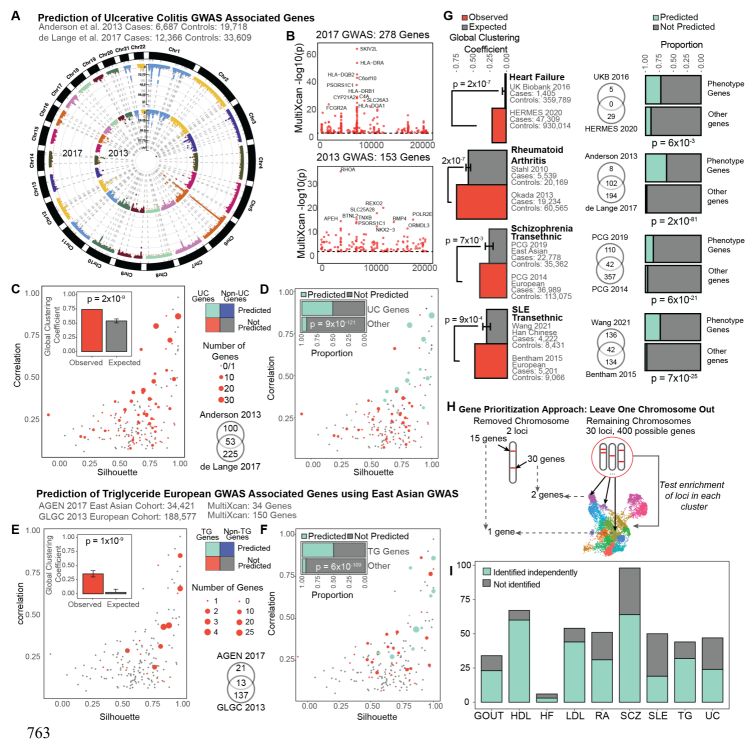
745 (A) Broad enrichment profile of 173 clusters stratified by average silhouette and correlation
 746 scores.

**(B)** 747 Heatmap showing the relationship between the biological profile of five clusters, as 748 proxied by their top gene ontology term, and the unique phenotypic signature. The top 10 749 phenotypes per cluster were selected. Association strength was calculated using negative log 750 transformed significance values, which were then normalised across phenotypes and then per 751 cluster. (C) Top enriched gene ontology biological processes for each cluster have no overlap. 752 Clusters ranked by strength of top enriched term. Specificity of top 5 terms per cluster is also 753 provided. (D) 173 clusters stratified by their average correlation and silhouette scores with 754 their gene ontology and STRING enrichment indicated. The P-value represented is specific to 755 the enriched category, and in the case of both represents the more significant of the two. 756 (E-G) Correlation between the average number of significant gene-trait associations per cluster 757 and E) the average correlation score of each cluster (F), strength of GO enrichment and strength 758 of STRING enrichment per cluster (Pearson's correlation) (G).

759 (H) Presence and degree of colocalization within clusters as stratified by their silhouette and

760 correlation scores (Pearson's correlation).

761



764 Figure 4. Identified clusters are conserved across all phenotypes and can be used for

prediction of genes involved in complex trait biology and prioritization of GWAS genes at
 implicated loci.

- 767 (A) Manhattan plot of loci identified by a 2017 and 2013 GWAS of ulcerative colitis (UC). (B)
- 768 Manhattan plot of S-MultiXcan genes for the two GWAS respectively, genes are ordered
- 769 according to their genomic positions.

770 (C) Distribution of 278 significant genes from the 2017 UC GWAS across 173 clusters.

771 Global clustering coefficient was calculated for the 278 genes. Significance was calculated

- using 100 bootstrap replicates to establish a distribution from which a Z score was calculated.
- (D) Prediction of 2017 UC GWAS genes using 2013 UC GWAS. Chi-squared enrichment
   test was used to determine enrichment for prediction of novel genes compared to non-trait
   associated genes.
- 776 (E) Distribution and global clustering coefficient of 34 significant genes from East Asian
   777 GWAS of Triglyceride levels. Significance was calculated using bootstrapping.
- 778 **(F)** Prediction of 137 novel genes from 2013 European GWAS of triglycerides using 2017
- 779 East Asian GWAS. Enrichment was calculated using chi-squared test.
- (G) Increase in observed global clustering coefficient compared to expected and prediction
   enrichment across four additional traits.
- 782 (H) Schematic of gene prioritization strategy using the leave one chromosome out approach.
- 783 Potential genes at a significant locus were refined using clusters enriched for the trait.
- 784 (I) The proportion of loci at which a gene was successfully identified independently of all785 genes on the same chromosome.