Bayesian analysis of genetic association across tree-structured routine healthcare data in the UK Biobank

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Genetic discovery from the multitude of phenotypes extractable from routine healthcare data can transform understanding of the human phenotype and accelerate progress toward precision medicine. However, a critical question when analyzing high-dimensional and heterogeneous data is how best to interrogate increasingly specific subphenotypes while retaining statistical power to detect genetic associations. Here we develop and employ a new Bayesian analysis framework that exploits the hierarchical structure of diagnosis classifications to analyze genetic variants against UK Biobank disease phenotypes derived from self-reporting and hospital episode statistics. Our method displays a more than 20% increase in power to detect genetic effects over other approaches and identifies new associations between classical human leukocyte antigen (HLA) alleles and common immunemediated diseases (IMDs). By applying the approach to genetic risk scores (GRSs), we show the extent of genetic sharing among IMDs and expose differences in disease perception or diagnosis with potential clinical implications.

Large-scale, hypothesis-free approaches for identifying genetic risk variants, including genome-wide association studies (GWAS) and next-generation sequencing analyses, have greatly advanced understanding of complex traits, with implications for drug development and clinical practice1–3. These approaches typically involve genetic discovery from case–control cohorts where clinically derived phenotypes are considered one at a time. By contrast, resources such as the UK Biobank6,7, which has prospectively collected extensive health-relevant phenotypic and genotypic information from 500,000 participants, allow for simultaneous investigation of multiple traits and are set to lead to a step change in the rate of genetic discovery8,9.

However, capitalizing on the availability of population-based cohorts for biomedical research is complicated by the scale and nature of the data: the phenotypic space is multidimensional and heterogeneous, as data can be subject to observational predilections, non-uniform recording practices and longitudinal biases while phenotype prevalence is variable10–16. This creates new challenges that are not addressed by existing analytical methods for GWAS and phenotype-wide association studies (PheWAS). An open question is how to interrogate the many precise phenotypes obtainable from routine healthcare data at a resolution that identifies associations above and beyond those identified through GWAS, but without sacrificing statistical power. Making use of disease classification hierarchies, such as the tree of International Classification of Diseases, Tenth Revision (ICD-10) codes, provides a tractable solution. Here we have developed a new Bayesian analysis framework for identifying genetic associations across the entire health phenotype space by taking advantage of the relative topology of nodes within two tree-structured phenotypic data sets from the UK Biobank—the self-reported (SR) diagnoses that are organized using the UK Biobank classification tree, which includes 531 diagnostic terms, and the hospitalization episode statistics (HES) data that utilize ICD-10 codes and contain 16,310 diagnostic terms.

RESULTS
Tree analysis approach
To test the association of genetic variation with any given UK Biobank clinical phenotype, we want to construct a statistical framework that meets a set of fundamental requirements. First, the method must accommodate different types of genetic variation, such as (i) SNPs, (ii) haplotypes in a highly polymorphic region like the HLA gene region, or (iii) GRSs constructed using multiple SNPs or haplotypes known to be associated with a quantitative trait or complex disease. Second, for single-locus variation, any genetic model (for example, additive, dominant or full) must be accommodated. Third, the method must allow for joint analysis and quantification of evidence for association at each clinical phenotype and must estimate the genetic coefficients of effects. Next, the method must allow identification of independent genetic effects through conditional analysis. Lastly, the method must model the correlation structure of genetic effects across...
observed clinical phenotypes using a priori knowledge of phenotype relationships obtained from a diagnosis classification tree.

To meet these requirements, we have developed a new Bayesian analysis framework, termed TreeWAS, which models genetic coefficients across all phenotypes as a set of random variables. To model the correlation structure, we allow coefficients to evolve down a tree in a Markov process (Fig. 1). A known classification hierarchy determines the tree structure, where each node is a clinical term in the classification, and observations can be made at terminal and internal nodes. The prior \( \theta \) determines the expected correlation between genetic coefficients across phenotypes. The coefficient at a parent node can be inherited by a child node with probability \( \epsilon^{-\theta} \) or can transition to a new uncorrelated value, with probability \( 1 - \epsilon^{-\theta} \). This new value will be zero with a probability of \( 1 - \pi_1 \) or nonzero with a probability of \( \pi_1 \). Thus, parameters \( \theta \) and \( \pi_1 \) define transition probabilities controlling the Markov process. Given the model structure and the Markov process assumption, we can calculate the likelihood over genetic coefficients across all clinical phenotypes using dynamic programming (details are provided in the Supplementary Note), and we estimate a Bayes factor statistic (BFtree) for the evidence that genetic coefficients across all clinical phenotypes, using a priori knowledge of phenotype relationships obtained under these two scenarios when considering a range of allele pairs; it can transition back to the null state (as in the B→A and B→C pairs); or it can transition to a new, uncorrelated nonzero state (as in the B→D pair). An in-depth description of the method is provided in the Supplementary Note.

**HLA-B*27:05 TreeWAS and PheWAS comparison**

We illustrate the advantages of the TreeWAS approach as compared to existing PheWAS tests by analyzing the association of the HLA-B*27:05 allele against the UK Biobank HES data set. The HLA-B*27:05 association with ankylosing spondylitis is one of the strongest genetic effects observed in human complex diseases, with an odds ratio of 17, and was likely driven by associations with M45 (PP = 1.0) (Fig. 1a). As a result of age-related disk degeneration, lack of an HLA-B*27:05 association with M47 is consistent with its non-immunological etiology.

Associations with reactive arthritis (for example, M02.39 Reiter’s disease, PP = 0.78) and anterior uveitis (H20.9 iridocyclitis, unspecified, PP = 0.98) were also observed (Fig. 1b, d), and we detected a previously unreported HLA-B*27:05 association with H40 glaucoma (PP = 0.84) (Fig. 1d). As glaucoma is a common complication of chronic uveitis3, comorbidity may explain this association. Lastly, we observed a weak effect on L40.5 arthropathic psoriasis susceptibility (PP = 0.60), but not non-arthropathic psoriasis (PP ≤ 0.25 for L40 child nodes except L40.5), consistent with previous studies (Fig. 1e). Therefore, our TreeWAS analysis of HLA-B*27:05 in the HES data set recapitulates known associations and demonstrates that our method can identify additional genuine associations in comparison to PheWAS.

**Sensitivity and specificity analysis of TreeWAS approach using simulated data**

Given the capability of TreeWAS to identify multiple associations with HLA-B*27:05, we wanted to further investigate the method's sensitivity and specificity. To assess the relative power of TreeWAS, and to explore its robustness and accuracy, we performed two sets of simulations. In the first set, we assessed power by simulating data from a simple scenario where genetic coefficients are nonzero for a set of five clinical annotations in the tree. These annotations were chosen to occur within a single branch of the tree (clustered nodes) or across distant branches (distributed nodes). We compared the power obtained under these two scenarios when considering a range of allele frequencies. We fitted the TreeWAS model under a two-parameter setting with default parameters \( \theta = 1/3 \) and \( \pi_1 = 0.001 \). For the alternative PheWAS model, we assumed complete independence across annotations, equivalent to setting \( \theta \rightarrow \infty \). Under the simulations with clustered nodes, the relative gain in power for identifying active nodes, where genetic coefficients are nonzero, of TreeWAS as compared to PheWAS was 20–25% across the allele frequencies tested (Fig. 3a). This gain in power was not associated with an increased false-positive rate.
Effects of HLA allelic variation in the phenome

HLA region genetic variation is associated with numerous human disorders, in particular autoimmune and autoinflammatory diseases. Hence, we sought to interrogate HLA effects on the full range of SR and HES phenotypes using TreeWAS. Through conditional analysis (Online Methods and Supplementary Note), we identified independent associations for ten HLA alleles in the SR data ($\log_{10}(BF_{\text{Tree}}) \geq 10$).
and eight HLA alleles in the HES data ($\log_{10}(\text{BF}_{\text{tree}}) \geq 20$) (Fig. 4 and Supplementary Tables 2 and 3). Seven of these alleles or alleles in high linkage disequilibrium (LD; $r > 0.98$) were associated in both data sets (Supplementary Fig. 5).

These associations were fine-mapped, and the majority of the strongest effects were with IMDs, as reported previously through GWAS\textsuperscript{17,25–30} (Fig. 4). For class I alleles, we observed associations with psoriasis (HLA-C*06:02) and ankylosing spondylitis (HLA-B*27:05), and the genetic coefficients for the latter were the largest observed in the SR and HES data sets (Fig. 4a,c). For class II alleles, HLA-DRB1*03:01 and HLA-DQB1*02:02 were observed to be independently associated with celiac disease in both data sets; these alleles tag two of the strongest known celiac disease HLA risk haplotypes, DR3–DQ2 and DR7–DQ2 (ref. 26). In both data sets, HLA-DQA1*03:01 was identified and fine-mapped to rheumatoid arthritis; this allele is in moderate LD with HLA-DRB1*04:01 ($r = 0.71$), which is the likely causal allele driving this association\textsuperscript{27}. Similarly, HLA-DQA1*03:01 was associated with type 1 diabetes (T1D), and we note that this allele is in LD with HLA-DQB1*03:02 ($r = 0.67$), which has been indicated as the most significantly associated T1D class II allele\textsuperscript{26}. In the SR data set, we identified an HLA-DRB1*15:01 association and fine-mapped it to multiple sclerosis (Fig. 4a). In the HES data set, HLA-DQB1*06:02 was identified instead and also fine-mapped to multiple sclerosis (PP = 1; Fig. 4c), but this allele is in strong LD with HLA-DRB1*15:01 ($r = 0.97$) (Supplementary Fig. 5). Lastly, HLA-DRB1*01:03 was fine-mapped to ulcerative colitis and Crohn’s disease in both data sets, and it is the likely causal allele for these two types of inflammatory bowel disease (IBD)\textsuperscript{30}.

Apart from established HLA associations with common IMDs, we also confirmed HLA effects for conditions where GWAS have not been performed, detected associations with clinical annotations linked to disease complications and identified new HLA associations with other IMDs. For example, in the SR data set, we confirmed the association of HLA-DRB1*04:04 with polymyalgia rheumatica and giant cell arteritis, which has previously been identified only through small candidate gene studies\textsuperscript{31,32} (Fig. 4a). The ulcerative colitis– and Crohn’s disease–associated HLA-DRB1*01:03 allele was found also to be associated with surgical procedures linked to complications of IBD, such as Z93.3 colostomy status (PP = 1) and Z93.2 ileostomy status (PP = 1), consistent with findings by the International IBD Genetics Consortium\textsuperscript{33} (Fig. 4c). Of the ten HLA alleles independently associated with clinical phenotypes in the SR data set, five were associated with hypothyroidism/myxoedema, and three of the eight alleles from the HES data were associated with the E03 hypothyroidism code. This disease is thus the phenotype with the largest number of independent HLA associations across both UK Biobank data sets. Associations have been reported with hypothyroidism for both HLA class I and class II loci, but the specific alleles driving these associations are not well resolved\textsuperscript{34,35}, apart from a recently reported HLA-DQA1*05:01–HLA-DQB1*02:01–HLA-DRB1*03:01 (DR3–DQ2 haplotype) association\textsuperscript{36}. In addition to HLA-DRB1*03:01, we refined the HLA associations with hypothyroidism to two further independent risk alleles, HLA-DQA1*03:01 and HLA-DRB1*01:03, and two independent protective alleles, HLA-B*15:01 and HLA-DPB1*01:01 (Fig. 4 and Supplementary Table 4). Our HLA analysis therefore demonstrates the validity of our method as it can identify known genetic associations and can facilitate discovery of new associations for relatively understudied diagnoses.

**Genetic risk score associations with IMDs**

Outside of the HLA region, over the last decade, understanding of genetic susceptibility to the common IMDs has increased tremendously, with tens to hundreds of risk loci being identified for each disease\textsuperscript{37}. However, given the prevalence of IMDs in the UK Biobank and the typically small effect sizes estimated, we expect low power at individual loci. For example, when considering nine of the most common autoimmune and autoinflammatory diseases (Online Methods), we observed evidence of association ($\log_{10}(\text{BF}_{\text{tree}}) > 0$) for 64 individual

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**Figure 3** Sensitivity and specificity analysis of TreeWAS on simulated data. (a) Rate of active node identification at increasing posterior probability thresholds and different simulated minor allele frequencies (MAFs) of the causal genetic variant, for the TreeWAS method ($\theta = 1/3$ and $r_1 = 0.001$; blue) and for the PhewAS method (a model assuming complete independence among phenotypes with $\theta \to \infty$ and $r_1 = 0.001$; orange). For each simulation replicate ($n = 500$), we simulated five clustered nodes with nonzero genetic coefficients (circles) and, for the remaining nodes, phenotype counts were simulated to match observed disease prevalence and zero genetic coefficients (diamonds). The vertical dashed line denotes the PP = 0.75 threshold used in the analysis. (b,c) Rate of false positives in the $\text{BF}_{\text{tree}}$ statistic (b) and active node identification (c) when genotypes for the HLA-B*27:05 allele were permuted in both phenotypic data sets. Gen., genotype; phen., phenotype.
SNPs (12.96% of GWAS SNPs tested) in the SR data set and 125 individual SNPs (25.30%) in the HES data set. Nevertheless, we can gain power by combining the effects of multiple typed and imputed susceptibility variants in a GRS (Online Methods) and using the TreeWAS approach to assess the relationship of the GRS with the UK Biobank data; we note that simulation analyses estimated an association for each HLA allele with at least one node in the tree in the conditional TreeWAS analysis using Supplementary Table 9. (c) The tree depicts the hierarchical structure of HES-derived clinical phenotypes as determined by ICD-10 classification (showing nodes with PP > 0.75 and their parent nodes). The graph shows estimated effect sizes for the heterozygous genotype of the different HLA alleles on susceptibility to each clinical phenotype. Bars show the 95% credible interval. (b) Evidence of association for each HLA allele with at least one node in the tree (BFtree) in the conditional TreeWAS analysis for the SR data set (Supplementary Table 10). Estimates for heterozygous and homozygous genotype effect sizes and descriptions of all phenotypes shown are available in Supplementary Tables 2 and 3. AS, ankylosing spondylitis; CD, Crohn’s disease; CI, confidence interval; COE, celiac disease; ENT, ear, nose, throat; MAP, maximum a posteriori; MS, multiple sclerosis; PS, psoriasis; RA, rheumatoid arthritis; T1D, type 1 diabetes; UC, ulcerative colitis.

Typically, GRSs best identified those clinical annotations from which they were constructed, with secondary associations detected with greater power by combining the effects of multiple typed and imputed susceptibility variants in a GRS (Online Methods) and using the TreeWAS approach to assess the relationship of the GRS with the UK Biobank phenome (Fig. 5).

For all associations, genetic coefficients were less than 1, demonstrating a degree of dilution in phenotype detection across both the SR and HES data sets; we note that simulation analyses estimated an
expected dilution of ~15% due to winner’s curse (Supplementary Table 5 and Supplementary Note). The least dilution was observed for association of the celiac disease GRS with this disease (β = 0.96 and β = 0.87 in the SR and HES data sets, respectively). The celiac disease phenotypes derived from the UK Biobank healthcare data are thus highly comparable to the clinically ascertained disease phenotype used in the GWAS\textsuperscript{45} from which the variants for the celiac disease GRS were obtained. Across both data sets, the greatest dilution of the association of the celiac disease GRS with this disease (β = 0.43 and β = 0.55 in the SR and HES data sets, respectively), while, in HES data specifically, the ankylosing spondylitis GRS was not associated with the disease (PP = 0.01), potentially owing to the small number of patients with ankyllosing spondylitis in this data set (n = 146), and in SR data the SLE GRS association with SLE had a genetic coefficient of only 0.20 (Fig. 5a,b).

Overall, the GRS associations were largely consistent between the SR and HES data sets, and, for GRSs and their respective diseases, the estimated genetic coefficients were weakly positively correlated (r\textsubscript{corrected} = 0.23, correcting for measurement error) (Fig. 5c). Strikingly, although the capacity of the SLE GRS to identify SLE itself in SR data was so diluted that the SLE GRS was in fact a better predictor of SLE itself in SR data was so diluted that the SLE GRS was in fact a better predictor of SLE itself (β = 0.87 in the SR and HES data sets, respectively). The graph shows the estimated effect size of the GRS on susceptibility to each clinical phenotype. (b) The tree depicts the hierarchical structure of HES-derived clinical phenotypes as determined by ICD-10 classification (showing nodes with PP > 0.75 and their parent nodes). The graph shows estimated effect sizes of the GRS on susceptibility to each clinical phenotype. (c) Comparison of estimated genetic coefficients for each GRS and the respective clinical annotation in both phenotypic data sets. Estimates of effect sizes and a description of all phenotypes shown are available in supplementary Tables 11 and 12.

Figure 5 Association analysis of genetic risk for multiple IMDs derived from clinical phenotypes in the UK Biobank SR diagnosis and HES data sets. (a) The tree depicts the hierarchical structure of SR clinical phenotypes as determined by the UK Biobank classification. Only nodes with a significant association (PP > 0.75) with at least one IMD GRS are shown, along with their parent nodes. The graph shows the estimated effect size of the GRS on susceptibility to each clinical phenotype. (b) The tree depicts the hierarchical structure of HES-derived clinical phenotypes as determined by ICD-10 classification (showing nodes with PP > 0.75 and their parent nodes). The graph shows estimated effect sizes of the GRSs on susceptibility to each clinical phenotype.

Expectations, signs and abnormal clinical and laboratory findings, not elsewhere classified
Factors influencing health status and contact with health services
External causes of morbidity and mortality
Codes for special purposes

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HLA-DRB1*01:03 associations, the ulcerative colitis GRS was associated with colostomy and ileostomy events ($\beta = 0.31$, PP = 0.98 and $\beta = 0.31$, PP = 1, respectively), as was the Crohn’s disease GRS, although the effect size magnitude was lower ($\beta = 0.03$, PP = 0.91 and $\beta = 0.03$, PP = 0.87, respectively). Also paralleling the HLA analysis, hypothyroidism was associated with several GRSs: five and four of the nine GRSs tested were associated with the disease in the SR and HES data sets, respectively, with those for celiac disease, rheumatoid arthritis, SLE and T1D found in both data sets. Hence, hypothyroidism is the single phenotype with the largest number of different GRS associations (Fig. 5a,b and Supplementary Tables 6 and 7).

**DISCUSSION**

By exploiting the inherent hierarchical structure of diagnostic classifications, our Bayesian analysis framework addresses a fundamental challenge for the analysis of high-dimensional, heterogeneous routine healthcare data: how to identify statistically significant genetic associations when interrogating thousands of diagnoses without employing methods that sacrifice phenotypic resolution. When applying TreeWAS to interrogate the effect of the HLA region on the UK Biobank phenotype, associations were identified with 143 and 966 nodes in the SR and HES data sets, respectively. Assessing the impact of IMD GRSs also identified associations with 151 and 810 nodes in the two respective data sets. The total number of nodes identified demonstrates the power of TreeWAS for detecting associations in data sets where numerous weak but correlated effects are present across the classification tree.

Among the many active nodes for which genetic associations were observed, previously established effects of HLA alleles on specific IMDs were detectable, as were effects for relatively understudied conditions. Notably, multiple new associations with HLA alleles were discovered for hypothyroidism. Although not all previously reported HLA associations could be detected for any single IMD (such as ankylosing spondylitis or multiple sclerosis) owing to limited power with the current UK Biobank data sets, the capacity for genetic discovery will improve with increasing cohort size, and associations with nodes displaying substantial granularity of clinical description were already identifiable.

In the GRS analysis, associations between GWAS-derived GRSs and their respective diseases were typically the strongest effects observed, even without HLA allele inclusion, demonstrating that non-HLA variants can provide precision for detecting specific IMDs. Cross-disease associations of GRSs were also identified, particularly for hypothyroidism, and this previously unappreciated extent of genetic sharing indicates a common, genetically determined pathogenesis. For all GRS associations, dilution of the capacity for phenotype detection was observed but was largely comparable between the SR and HES data sets. An intriguing exception was the differential association of the SLE GRS with the respective SLE terms in the two data sets: this GRS could not precisely predict the self-reported disease but could accurately detect the hospitalization-record-derived phenotype. In comparison to the other IMDs investigated, SLE is a more heterogeneous, systemic condition, which consequently presents a substantial diagnostic challenge. Therefore, this discrepancy in the magnitude of SLE GRS association could reflect incorrect reporting of the disease, disease overdiagnosis not discernible in the HES data if hospitalization is associated with more clear-cut diagnosis, or greater disease heterogeneity whereby SLE as defined in GWAS and in the HES data represents only a subset of a more genetically variable syndrome.

Identifying miscategorization, misdiagnosis and miscoding in routine healthcare data is an ongoing challenge, although there are recognized instances, such as inaccuracy in T1D and type 2 diabetes (T2D) differentiation. In the UK Biobank, the T1D GRS is not associated with T2D terms in the SR data (PP = 0.0002) and shows weak evidence of association in the HES data (PP = 0.52). However, the T2D GRS, which can accurately detect T2D terms ($\beta = 0.80$, PP = 1.00 and $\beta = 0.71$, PP = 1.00 in the SR and HES data sets, respectively), is also associated with T1D in the HES data ($\beta = 0.71$, PP = 1.00) but not the SR data (PP = 0.30; Supplementary Table 8 and Supplementary Note). These cross-disease associations may be attributable to T1D/T2D misclassification, misdiagnosis and miscoding (Supplementary Figs. 6 and 7, and Supplementary Note) but also to genetic sharing and poor distinction of patients with latent autoimmune diabetes of adulthood, whose genetic profiles comprise a mixture of T1D and T2D risk loci. Thus, the SLE and diabetes examples demonstrate how exploring the genetic basis of the healthcare phenotype can expose disease areas where improvements are required to ameliorate disease perception or strengthen diagnostic practices. Digital phenotyping using genetic data in combination with longitudinal clinical information, physical measures and biomarkers could help to rectify miscategorization, misdiagnosis and miscoding present in healthcare data and to infer missing phenotypes. This could in turn facilitate patient management, particularly if it enables correction of treatment strategies within an actionable time frame.

Integration of genomic data with routine healthcare information offers much potential to learn about differences in disease risk, diagnosis and reporting within and between healthcare systems, including between countries. Moreover, increased incorporation of correlated, high-dimensional phenotypes (for example, from molecular, cytometry and imaging readouts), including measures of temporal disease progression, may lead to a genetically driven understanding of the architecture of the human phenotype and of causal relationships. The value of TreeWAS lies in enhancing power to identify groups of endpoints affected by specific genetic risk factors, by exploiting the encoding of medical ontologies. A corollary is that structures that better capture the underlying biological process affecting the origin and progression of disease should be better correlated with genetic risk factors. Although generalizing the TreeWAS method to structures reflecting temporal progression and associated quantitative data modalities requires future development, we believe that it is an important step toward the goal of learning a genetically motivated classification of disease and associated phenotypes.

**URLs.** UK Biobank, http://www.ukbiobank.ac.uk/; UK Biobank genotyping procedure and genotype calling protocols, http://biobank.ctsu.ox.ac.uk/crystal/refer.cgi?id=155580; UK Biobank internal quality control procedures, http://biobank.ctsu.ox.ac.uk/crystal/refer.cgi?id=155580; HLA*IMP, https://oxfordhla.well.ox.ac.uk/hla/; World Health Organization ICD-10 disease classification codes, http://www.who.int/classifications/icd/en/.

**METHODS**

Methods, including statements of data availability and any associated accession codes and references, are available in the online version of the paper.

*Note: Any Supplementary Information and Source Data files are available in the online version of the paper.*

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AUTHOR CONTRIBUTIONS
A.C. and G.M. performed the analyses with contributions from C.A.D. A.C., L.F. and G.M. conceived the study. A.M., D.V., A.D. and S.L. performed HLA imputation. A.C., C.A.D. and G.M. wrote the manuscript, and all other authors reviewed the manuscript.

COMPETING FINANCIAL INTERESTS
The authors declare competing financial interests: details are available in the online version of the paper.

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ONLINE METHODS

UK Biobank data. The UK Biobank is a prospective cohort of over 500,000 men and women aged 40 to 69 years when recruited in 2006–2010. Participants have provided data on lifestyle, environment and medical history through an interview and completion of a questionnaire; physical measures; biological samples for genotyping and biochemical assays; and informed consent to long-term medical follow-up through linkage of national health registries. The UK Biobank has obtained ethical approval covering this study from the National Research Ethics Committee (REC reference 11/NW/0382).

Phenotypic data. We analyzed two phenotypic data sets available through the UK Biobank. The first included SR diagnosis data, ascertained through the completion of questionnaires and interviews with study participants (data field 20002 non-cancer illness code, self-reported); the second data set included the HES registry data set ascertained through linkage of health registries (data fields 41142 and 41078; accessed in September 2016). Clinical diagnoses in these data sets are described with different classification schemes, both of which follow a hierarchical structure. The diagnosis terms used to store the medical history of UK Biobank participants were proposed by the UK Biobank team (data-coding 6), and this classification tree is organized into 11 subclasses with a total of 561 clinical terms, 531 of which are selectable. Diagnosis terms used to store hospitalization events follow the ICD-10 list compiled by the World Health Organization. The ICD-10 classification tree is organized into 22 chapters and contains a total of 19,855 clinical terms, 16,310 of which are selectable. Each hospitalization episode in the data set has a primary diagnosis associated with the event, and an event may be annotated with one or more secondary diagnoses. Disease outcomes for each individual, as a binary trait, were generated for the combined primary and secondary diagnosis annotations. Individuals were considered unaffected for any given diagnostic term unless the diagnosis was reported in the questionnaires and interviews or a hospitalization event with that diagnostic term was observed.

Genetic data set. The interim release of the UK Biobank genetic data used for this study includes 152,732 individuals, 120,286 of whom were determined to be of British Isles ancestry (Supplementary Fig. 8) and included in the analysis. The initial 50,000 individuals were genotyped on the Affymetrix UK BiLEVE Axiom array as part of a pilot study described elsewhere49, and the remaining 102,732 individuals were genotyped on the Affymetrix UK Biobank Axiom array. Quality control of SNP data and whole-genome SNP imputation was performed by the UK Biobank analysis team and is described on the UK Biobank website (http://www.ukbiobank.ac.uk/scientists-3/genetic-data/). We imputed 356 classical HLA alleles for the HLA-A, HLA-B, HLA-C, HLA-DRB5, HLA-DRB4, HLA-DRB3, HLA-DRB1, HLA-DQB1, HLA-DQA1, HLA-DPB1 and HLA-DPA1 loci at four-digit resolution with the HLA*IMP:02 algorithm50,51 using data from a multi-population reference panel. The imputation panel contained 2,263 SNPs in the MHC region (GRCh37 coordinates chr: 6:29,500,000–33,500,000) that overlapped UK Biobank genotyped SNPs. This SNP set was selected to optimize MHC coverage and imputation performance, and the HLA*IMP:02 algorithm was trained on this SNP set. GRSs, weighted by effect sizes, were generated for nine IMDs using genome-wide associated variants compiled from previous studies: ankylosing spondylitis17, Crohn’s disease39, celiac disease40, multiple sclerosis41, psoriasis42, rheumatoid arthritis43,44, SLE45, T1D39 and ulcerative colitis39. SNP genotypes for the UK Biobank individuals were extracted from the imputed genotype data and maintained if the imputation information score was above 0.85; if a SNP was not typed or imputed successfully, it was not included in the GRS calculation.

Simulated data. To assess the accuracy of the method, we simulated case–control status for 120,000 individuals and the 531 selectable phenotypes in the diagnosis tree used for the self-reported data set and with disease prevalence as observed in the UK Biobank cohort. Simulations were generated under two scenarios. For the first, we assumed a causal relationship between a genetic variant and five clinical terms under the same parent node in the tree (disease prevalence in these nodes ranged between 0.01 and 0.4%). These simulations are referred to as clustered clinical phenotypes. The second set of simulations, termed distributed phenotypes, consisted of five clinical terms with a causal relationship distributed under different branches of the classification tree; these clinical terms were selected with matching disease prevalence, as for the clustered simulations. For each scenario, we simulated genotypes sampled from a multinomial distribution with a fixed allele frequency and genetic coefficients sampled from the prior (Supplementary Fig. 9). Case–control status was determined by using logistic risk with a y intercept matching the observed disease prevalence. Sets of simulations were performed for the allele frequencies 0.005, 0.01, 0.02 and 0.05. For each simulation, we computed the evidence of association in the tree (BFtree) and the evidence of association at each individual node with the parameters $\theta = 1/3$ and $\pi = 0.001$. We compared the power to detect association with at least one node in the tree with an analysis where we assumed no correlation in the genetic coefficients between nodes in the tree, equivalent to setting $\theta \rightarrow \infty$ in the TreeWAS method (Supplementary Note). We performed 500 simulation replicates for each combination of parameters and settings. To assess the robustness of the algorithm to the non-independence between annotations unaccounted for by the tree structure, we performed simulations where we permuted the genotypes whilst leaving the observed phenotypes in the UK Biobank cohort intact. Simulations were performed with the observed self-reported and HES data sets, and we permuted the observed genotypes.

HLA analysis. For each HLA locus, we derived highest-confidence genotypes by taking the allele at each chromosome with the highest imputation posterior probability. Genotypes were used to generate count distributions in affected and unaffected individuals at each terminal node in the tree. To identify independent HLA associations, we performed sequential conditional analysis using an approximation to the likelihood function as described in the Supplementary Note. At each step, BFtree statistics were generated for each allele and the allele with the largest value was selected for conditioning in the next iteration. Conditional analysis was repeated until all observed BFtree statistics were below $10^{-10}$ in the self-reported diagnosis data set and $10^{-20}$ in the HES data set, ensuring a false discovery rate below 0.01, as determined through the simulation analysis. For each significant allele association, we computed the marginal posterior probability for the genetic coefficient being not equal to 0 and the MAP estimate using posterior decoding as described in the Supplementary Note. Association with a clinical annotation was deemed significant if the posterior probability was above 0.75.

Code availability. Code to perform TreeWAS analysis is available from the authors upon request or through the code repository at https://github.com/mcvteamlab.

Data availability. UK Biobank data are available under open access to conduct health-related research after approval of a project proposal46.

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