Pervasive additive and non-additive effects within the HLA region contribute to disease risk in the UK Biobank

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# Abstract

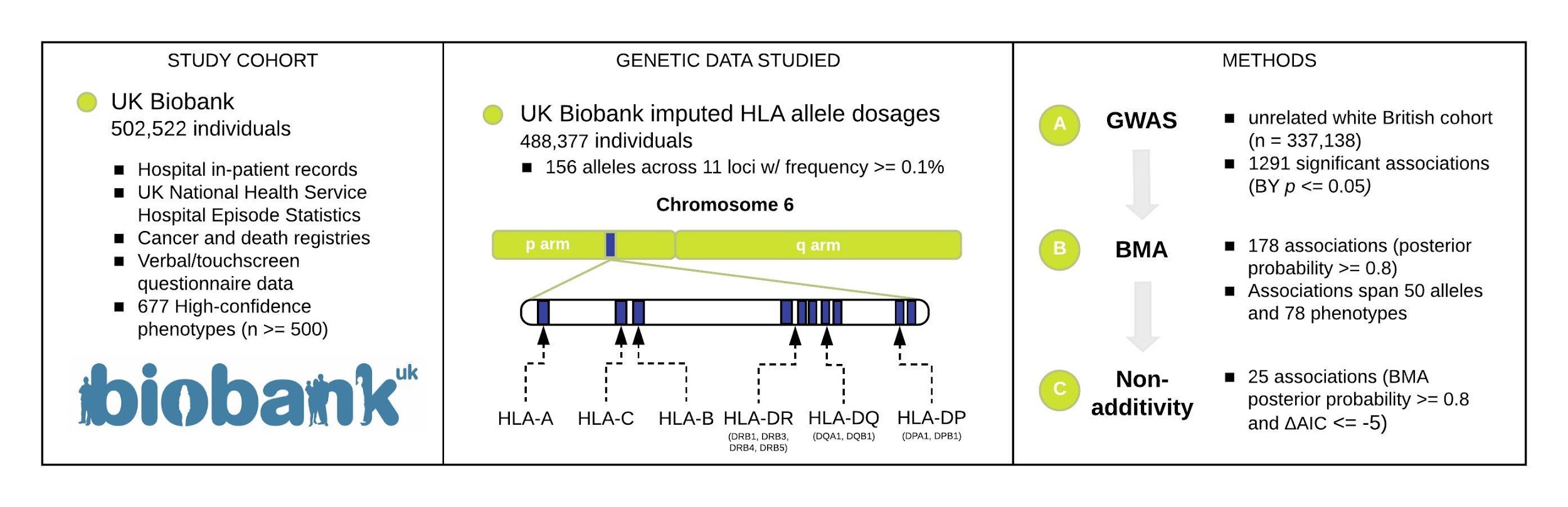
# Introduction

The human leukocyte antigen (HLA) region of the genome is one of the most disease-associated but gene-dense and polymorphic regions of the human genome[1](https://paperpile.com/c/wFd6d6/WpCF). The downstream products of HLA genes generate and present peptides on the cell surface that can be recognized by T cell receptors, making these genes relevant to many disorders of the immune system[2](https://paperpile.com/c/wFd6d6/QjRz). Before the advent of high-throughput genome-wide association studies (GWAS), HLA polymorphisms were associated only with autoimmune disorders and cancers[3,4](https://paperpile.com/c/wFd6d6/DDH8+RgAV). Since, GWAS has uncovered associations between HLA and common infections[5](https://paperpile.com/c/wFd6d6/CcIi), shingles[6](https://paperpile.com/c/wFd6d6/1YBh), chronic Hepatitis B[7](https://paperpile.com/c/wFd6d6/wVWz), Epstein-Barr virus[8](https://paperpile.com/c/wFd6d6/0QPR), and other diseases. The HLA region contains 1.5% of the genes currently in Online Mendelian Inheritance in Man (OMIM, a database of genetic disorders and traits focusing on gene-phenotype relationships) and accounts for roughly 1% (1,827/185,864) of the genome-wide significant SNP associations in the NHGRI/EBI GWAS catalog[9](https://paperpile.com/c/wFd6d6/28Kh), underscoring its importance to disease. Despite being an attractive target for comprehensive GWAS, the HLA region presents many challenges that obscure its roles in disease pathogenesis; complex linkage disequilibrium (LD) structures, structural variation, closely-related genes, paralogs, segmental duplications, and violations of Hardy-Weinberg equilibrium make high-throughput genotyping methods such as fine-mapping and imputation challenging[10,11](https://paperpile.com/c/wFd6d6/U2F6+O6wn). Despite custom-designed chips (e.g. the Illumina MHC SNP Panel[12](https://paperpile.com/c/wFd6d6/wpp7)) and robust reference panels built to study the region[13,14](https://paperpile.com/c/wFd6d6/YtiO+tv8H), there exist well-studied alleles in the HLA region whose impact on disease are unknown. Thus, understanding the role of the HLA region is critical to assessing disease risk.

The UK Biobank dataset offers researchers genetic and phenotypic data for over 500,000 individuals, and additionally provides the opportunity to analyze HLA region data for 488,377 individuals, generated via imputation strategies that address the aforementioned challenges[13](https://paperpile.com/c/wFd6d6/YtiO). These data can be used to perform association analyses across the rich diversity of phenotypes in the UK Biobank, which are derived from cancer registry, hospital in-patient, primary care, and self-reported questionnaire data. Assessing association between HLA alleles with individual level data can help pinpoint culprit alleles for disease associations via well-powered conditional analysis. We can additionally identify settings where HLA alleles exhibit non-additive effects on phenotypes improving our ability to accurately integrate them in disease risk models.

Here, we performed a single-allele association analysis for 156 HLA alleles across 677 phenotypes in the UK Biobank and identified 1291 associations at a false discovery rate (FDR) of 5% after Benjamini-Yekutieli multiple testing correction. The associations spanned 113 HLA alleles (across all 11 Biobank-provided HLA loci) and 128 binary phenotypes. We performed Bayesian Model Averaging to narrow down to conditionally independent alleles and uncovered 178 high-confidence (posterior probability >= 80%) associations spanning 50 alleles (across all 11 HLA loci) and 78 binary phenotypes (88 [49%] of which were supported by the literature, **Supplementary Table 1**). We assessed these associations for non-additive effects, i.e. “disease contributions beyond the cumulative effect of individual alleles”[15](https://paperpile.com/c/wFd6d6/wKtR), finding 25 associations with significant deviations from additivity (8 [32%] of which were supported by the literature, **Supplementary Table 1**). 9 of these 25 non-additive effects drive intestinal malabsorption and/or celiac disease, but other associations affect a multitude of other autoimmune and endocrine disorders. For example, although HLA-B\*27:05 is known to have strong effects on ankylosing spondylitis and iridicyclitis, and HLA-B\*57:01 is likewise known to have strong effects on psoriasis, we find that these effects are also non-additive. This study links HLA alleles to a variety of phenotypes in the UK Biobank through association analyses and model selection techniques, providing important insight into disease pathogenesis.

# Materials and Methods



**Figure 1. Overview of the study design.**

## Data

For unrelated white British individuals (n = 337,138) in the UK Biobank as determined previously[16](https://paperpile.com/c/wFd6d6/ECdo), three sets of phenotypes were defined: “high-confidence”, “time-to-event”, and “algorithmically-defined”. High-confidence phenotypes were defined by combining disease diagnoses (UK Biobank Field ID 41202) from the UK National Health Service Hospital Episode Statistics (ICD-10 codes H40.[0-6,8,9], H42.8, and Q15.0), the self-reported diagnosis questionnaire (UK Biobank Field ID 20002), and the UK Cancer Registry. Time-to-event phenotypes were derived from First Occurrence of Health Outcomes data as defined by 3-character ICD-10 codes in UK Biobank’s Category 1712. The First Occurrence data-fields were generated by mapping: read code information in the primary care data (Category 3000); ICD-9 and ICD-10 codes in hospital inpatient data (Category 2000); ICD-10 codes in death registry records (Field 40001, Field 40002); and self-reported medical condition codes (Field 20002), reported at baseline or subsequent UK Biobank assessment center visits as 3-character ICD-10 codes. Algorithmically-defined outcomes (based on data from Category 42) include phenotypes of select health-related events obtained through algorithmic combinations of coded information from the UK Biobank’s baseline assessment data collection. The data were derived from self-reported medical conditions, operations and medications together with linked data from hospital admissions and death registries. We included phenotypes with at least 500 cases among the white British cohort (**Supplementary Table 2**) and manually deduplicated several using the FuzzyWuzzy python package’s partial ratio function, resulting in the 677 phenotypes used in the analysis.

HLA alleles HLA-A, -B, -C, -DPA1, -DPB1, -DQA1, -DQB1, -DRB1, -DRB3, -DRB4, and -DRB5 were imputed using the HLA\*IMP:02 program[13](https://paperpile.com/c/wFd6d6/YtiO). HLA-IMP:02 is built on multiple reference panels and a graphical model of the HLA haplotype structure. The UK Biobank provides an imputed dosage file that contains the estimated dosage of 362 HLA alleles across 11 HLA loci from 488,377 individuals (while modeling imputation uncertainty). We included 156 alleles across all 11 loci that had a frequency of 0.1% or greater in our white British cohort (**Supplementary Table 2**). For consistency across models, and because the non-additivity analysis requires integral values for allele dosage, those dosages that were within 0.1 of 0, 1, or 2 for each allele were rounded, and the remaining nonzero entries were excluded. Erroneous total allele counts post-rounding were excluded.

## Association analysis

We performed generalized linear model association analysis (with “Firth-fallback”) as implemented in PLINK v2.00aLM (March 14 2020)[17](https://paperpile.com/c/wFd6d6/xUt9), for the above-described 677 phenotypes and 156 HLA alleles across UK Biobank white British individuals (n = 337,138). Firth-fallback is a hybrid algorithm which normally uses logistic regression but “falls back” to Firth's bias reduction method[18](https://paperpile.com/c/wFd6d6/Te92), equivalent to penalization of the log-likelihood by the Jeffreys prior, in two cases: (1), one of the cells in the 2x2 (allele count-by-case/control status) contingency table is empty; or (2), logistic regression fails to converge within the usual number of steps. We used age, sex, genotyping array, number and length of copy number variants, and the first ten genotype principal components as covariates in our analysis[19](https://paperpile.com/c/wFd6d6/vfuO). We used the Benjamini-Yekutieli (BY) multiple-testing FDR control method[20](https://paperpile.com/c/wFd6d6/fQyV) to adjust the resultant p-values, selecting those associations with BY-corrected p-values less than 0.05 to control FDR at 5%.

## Bayesian Model Averaging

Given the high amount of LD between HLA alleles and the resultant high number of potentially spurious HLA allele-phenotype associations even despite FDR correction, we used a Bayesian Model Averaging (BMA) approach, implemented in the `bma` R package[21](https://paperpile.com/c/wFd6d6/DhjQ), to prioritize which HLA loci were most likely causal for each phenotype. We used BMA to train a model on each possible subset of significantly-associated alleles. The posterior probabilities of each model are reported, and the degree to which an allele is included across all models (the “allele posterior probability”) is then a measure of confidence in the allele-phenotype association.

We identified all of the allele-phenotype pairs that had BY-adjusted p-values less than or equal to 0.05 from the PLINK analysis. Because BMA is exponential in complexity with respect to the number of alleles analyzed (requiring analysis of 2n models for n alleles included in the analysis), we performed BMA only on the 20 alleles with the lowest BY-adjusted p-values from the PLINK analysis for each phenotype. If there were less than two such alleles for a given phenotype, we did not run BMA for that phenotype. Additionally, only the models whose posterior probabilities were within a factor of 1/5 of that of the best model were kept for the final averaging. These requirements filtered our testing base down to 116 phenotypes, with 111 alleles included in at least one analysis. We ran BMA with a binomial error distribution and link function and with age, sex, genotyping array, and the first ten genotype principal components as covariates (copy number variant-related covariates were excluded). We used those allele-phenotype pairs with BMA posterior probabilities >= 0.8 in the subsequent non-additivity analysis.

## Analysis of non-additive genetic effects

To assess whether certain allele-phenotype pairs exhibited non-additive effects on the phenotype, we additionally performed logistic regressions in R using the glm function in R (family = "binomial") and providing dosages as factors (i.e., separate terms indicated whether a subject was heterozygous or homozygous for the HLA allele in question) and included age, sex, genotyping array, number and length of copy number variants, and the first ten genotype principal components as covariates (as in the PLINK single-allele association analysis). Additionally, we computed an additive model in R to generate Akaike Information Criterion (AIC, a measure of goodness-of-fit[22](https://paperpile.com/c/wFd6d6/uifu)), comparing the non-additive to the additive model for model selection. To identify gene-phenotype associations with suspected departures from additivity, we identified allele-phenotype pairs where the BMA posterior probability was greater than 0.8 and the difference between the non-additive model AIC and additive model AIC (ΔAIC) was <= −5.

## Systematic literature review

For each allele-phenotype pair resulting from the BMA analysis, we conducted a systematic literature review to determine whether the association had been previously found. The HLA allele and the phenotype in question were entered into Google Scholar both with and without quotations, and aliases were checked (e.g. B\*08:01, B\*0801, B/08/01, and B/0801). Search results were manually inspected to determine associations (**Supplementary Table 1**).

# Results

## Association analysis

To assess the extent to which HLA alleles affect various phenotypes, we conducted an HLA-wide association study across 337,138 white British individuals in the UK Biobank (Methods). We used 156 HLA alleles (out of 175 total) that were present at greater than 0.1% minor allele frequency in the cohort and 677 de-duplicated binary phenotypes with more than 500 cases in the UK Biobank. These binary phenotypes included autoimmune, lymphatic, cardiovascular, dermal, skeletal, tissue, digestive, respiratory, renal, and endocrine disorders as well as many cancers. To control for false discovery, we corrected for multiple testing across the phenotypes using the Benjamini-Yekutieli (BY) procedure at a FDR of 5%. We found 113 alleles that were associated with at least one of 128 binary phenotypes for a total of 1291 associations.

## Bayesian Model Averaging

Given many HLA alleles are in high LD with one another, we asked whether some of the discovered associations were conditionally independent. We used Bayesian Model Averaging (BMA) on the top 20 BY-significant alleles for each phenotype that had at least 2 BY-significant allele associations. Across 116 test phenotypes and 111 alleles included in at least one analysis, we found 178 putatively causal allele-phenotype pairs among 50 distinct alleles across all 11 loci and 78 distinct phenotypes. Of these 178 putatively causal alleles, 88 have been previously documented as associated allele-disease pairs, and 90 were novelly marked by our analysis as high probability of having causal signal (**Methods, Supplementary Tables 1-2, Figure 2A, Figure 2B**). Of note, 83 putatively causal allele-phenotype pairs with posterior probability 1 were not previously documented in the literature. These 83 allele-phenotype pairs mostly consist of associations that either feature alleles in LD with known associations or associations between alleles and closely-related phenotypes to known associations. For example, we find a link between **non-melanoma skin cancer and DQB1\*03:02**, although the literature supports only a link with an allele in LD with DQB1\*03:02, DQB1\*05:01[23](https://paperpile.com/c/wFd6d6/OyEn). Alleles associated to celiac disease, such as **B\*08:01**[24](https://paperpile.com/c/wFd6d6/ievD), were found to be associated with **intestinal malabsorption and anemias**, which have close ties to celiac disease[25](https://paperpile.com/c/wFd6d6/kbTj). Additionally, alleles associated with Graves' disease and Hashimoto thyroiditis (such as **B\*08:01**[26](https://paperpile.com/c/wFd6d6/KN0C), **B\*39:06**[27](https://paperpile.com/c/wFd6d6/k564), **DQB1\*06:04**[28](https://paperpile.com/c/wFd6d6/WVNA), and **DRB1\*03:01**[29](https://paperpile.com/c/wFd6d6/kbRP)) were found to be associated with **hypo- and hyper-thyroidism** in the UK Biobank and account for 13 of the 84 high-probability associations. Finally, alleles associated with type 1 diabetes (**DQB1\*02:01, DQB1\*03:02, and DRB1\*04:01**) were found to be associated with “**other disorders of pancreatic internal secretion,**” which have obvious links to type 1 diabetes.

Other associations among these 178 include those between HLA alleles and infectious diseases, asthma, systemic autoimmune disorders, and skin cancers (**Supplementary Table 1**). Among the strongest risk effects are associations between: **B\*27:05 and ankylosing spondylitis** (pBY = 0, BMApostOR = 9.02, 95% CI = [8.19, 9.94])[30](https://paperpile.com/c/wFd6d6/Y7f9), **B\*27:05 and iridocyclitis**, a type of uveitis (pBY = 2.94 x 10-239, BMApostOR = 4.75, 95% CI = [4.32, 5.23])[31](https://paperpile.com/c/wFd6d6/JPqH), **B\*39:01 and psoriatic arthropathy** (pBY = 1.06 x 10-8, BMApostOR = 3.54, 95% CI = [2.51, 5.0])[32](https://paperpile.com/c/wFd6d6/9l72) and **DQA1\*03:01 and seropositive rheumatoid arthritis** (pBY = 6.46 x 10-66, BMApostOR = 3.16, 95% CI = [2.75, 3.63])[33](https://paperpile.com/c/wFd6d6/1Q80). We replicate previously-found strong protective effects of **DQB1\*03:01** (pBY = 1.00 x 10-70, BMApostOR = 0.42, 95% CI = [0.34, 0.51]) and **DQA1\*01:01** (pBY = 2.62 x 10-51, BMApostOR = 0.71, 95% CI = [0.61, 0.81])[15](https://paperpile.com/c/wFd6d6/wKtR) on **celiac disease** and **malabsorption**. We also discover novel associations to these phenotypes: **DQB1\*03:03** (pBY = 2.30 x 10-22, BMApostOR = 0.47, 95% CI = [0.36, 0.62]), DQB1\*03:01 (pBY = 2.00 x 10-53, BMApostOR = 0.42, 95% CI = [0.34, 0.51]), **DRB4\*99:01** (pBY = 4.47 x 10-31, BMApostOR = 0.52, 95% CI = [0.46, 0.58]), and **DRB4\*01:03** (pBY = 1.40 x 10-71, BMApostOR = 0.63, 95% CI = [0.49, 0.81]) are all associated. Of note, we find new HLA associations to anemias, with which HLA alleles have not previously been linked to except for the rare case of aplastic anemia (Latin America, France, Turkey, Brazil, and Barcelona have reported prevalences of only 1.6, 1.5, 0.64, 2.4, and 2.5 per million for the disease)[34](https://paperpile.com/c/wFd6d6/Iw9u). We find **B\*08:01** (pBY = 8.71 x 10-4, BMApostOR = 1.2, 95% CI = [1.11, 1.3]) and **B\*39:06** (pBY = 7.66 x 10-3, BMApostOR = 1.95, 95% CI = [1.49, 2.56]) to be associated with **vitamin B12 deficiency anemia** risk with moderate effects. In contrast, **DQA1\*01:01** protects against **vitamin B12 deficiency anemia** (pBY = 8.41 x 10-7, BMApostOR = 0.76, 95% CI = [0.68, 0.83]) and **pernicious anemia** as well (pBY = 7.20 x 10-3, BMApostOR = 0.73, 95% CI = [0.63, 0.83]).

Several other associations are intriguing. The only previous link between the HLA region and acne is the discovery of the DPB1\*04:02 allele while HLA-typing a patient with acne vulgaris[35](https://paperpile.com/c/wFd6d6/YA0P). Yet, we find a weak protective effect of **B\*08:01 on acne** (pBY = 4.21 x 10-3, BMApostOR = 0.85, 95% CI = [0.79, 0.91]). This allele is additionally found to be associated with multiple systemic autoimmune disorders, like **connective tissue disorder** (pBY = 3.45 x 10-10, BMApostOR = 1.32, 95% CI = [1.23, 1.42]), **sarcoidosis** (pBY = 2.04 x 10-33, BMApostOR = 1.58, 95% CI = [1.36, 1.84]), **Sjogren’s syndrome** (pBY = 1.52 x 10-15, BMApostOR = 1.83, 95% CI = [1.61, 2.09]), and **systemic lupus erythematosus** (pBY = 9.50 x 10-12, BMApostOR = 1.79, 95% CI = [1.55, 2.06]), suggesting diverse and widespread involvement in various immune and bodily functions (**Supplementary Table 1**). Additionally, we find that **B\*57:01** (notably associated with a spike in Alanine aminotransferase levels, among other adverse drug reactions, to the anti-retroviral abacavir[36,37](https://paperpile.com/c/wFd6d6/0FUD+HqvG)) to be associated with **calculus of kidney and ureter**.

HLA links to cancers are well-documented, and here we recover several known associations, including between **DQB1\*05:01** and **DRB3\*99:01** and **cervical cancer** and **DQA1\*03:01** and **female genital tract cancer**. We find **DQB1\*05:01** to be associated with **skin cancer**, **non-melanoma skin cancer**, and **non-hodgkin's lymphoma**. We novelly find several associations between skin cancer, non-melanoma skin cancer, and HLA allelotypes. Statistics on all cancer associations (as well as their novelty) are described in **Table 1**. Taken overall, this body of associations, halfway between novel and previously-discovered, can prove vital in elucidating disease risk across a multitude of complex traits.

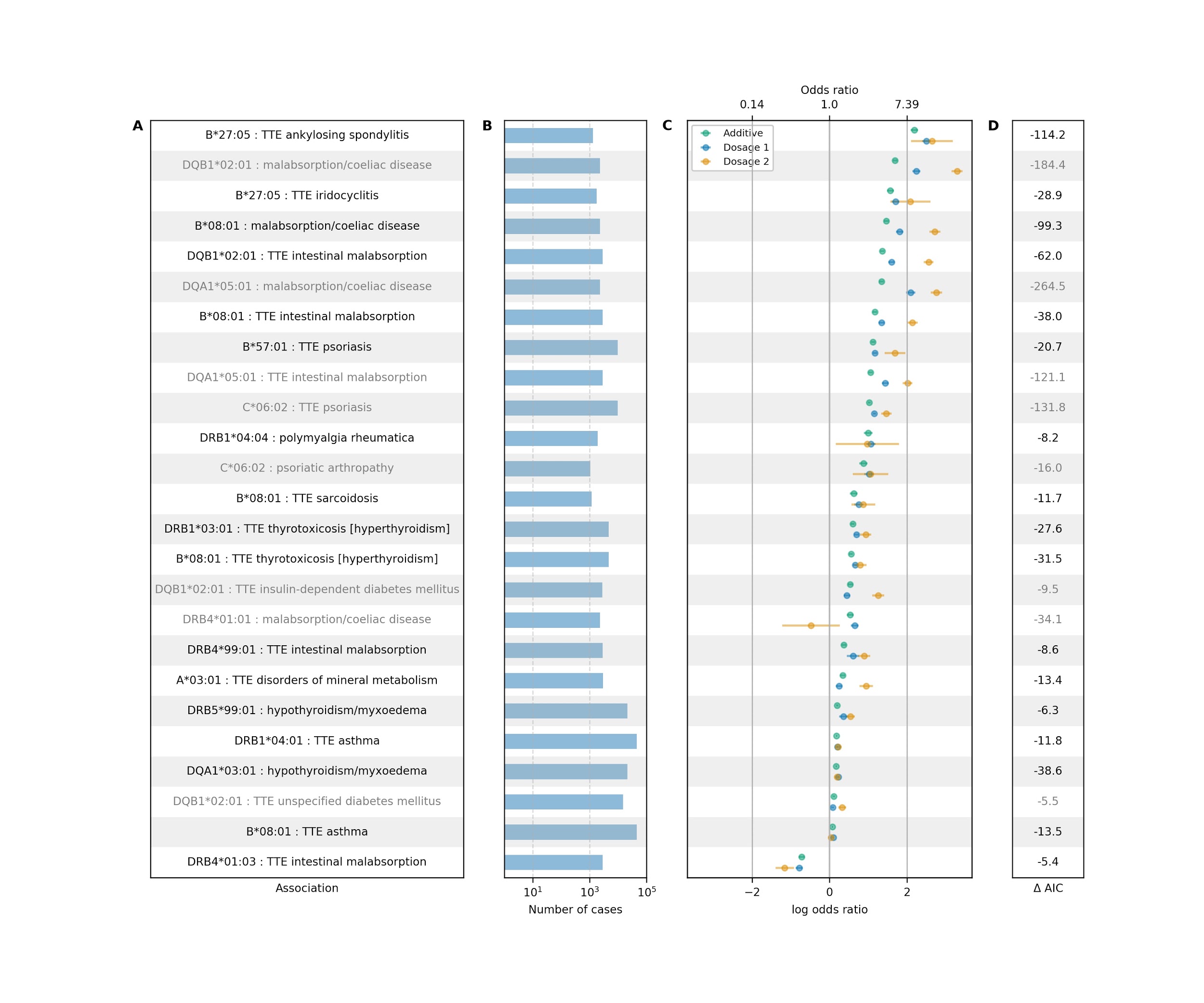
|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **HLA Allele** | **Phenotype** | **BY-adjusted *p-*value** | **BMA OR [95% CI]** | **Novel?** |
| DQB1\*05:01 | cervical cancer | 4.51 x 10-4 | 0.76 [0.69, 0.83] | N[38](https://paperpile.com/c/wFd6d6/99bs) |
| DRB3\*99:01 | cervical cancer | 2.21 x 10-3 | 1.19 [1.13, 1.27] | N[39](https://paperpile.com/c/wFd6d6/JCil) |
| DQA1\*03:01 | female genital tract cancer | 4.77 x 10-5 | 1.16 [1.1, 1.22] | N[40](https://paperpile.com/c/wFd6d6/ELlv) |
| DQB1\*05:01 | skin cancer | 2.12 x 10-15 | 1.11 [1.08, 1.15] | N[23](https://paperpile.com/c/wFd6d6/OyEn) |
| DQB1\*05:01 | non-melanoma skin cancer | 3.27 x 10-17 | 1.13 [1.09, 1.17] | N[23](https://paperpile.com/c/wFd6d6/OyEn) |
| DQB1\*05:01 | non--hodgkin’s lymphoma | 3.68 x 10-3 | 1.31 [1.19, 1.45] | N[41](https://paperpile.com/c/wFd6d6/UqvH) |
| B\*44:03 | skin cancer | 3.65 x 10-5 | 0.89 [0.85, 0.93] | Y |
| DQB1\*03:02 | skin cancer | 2.69 x 10-8 | 0.91 [0.87, 0.95] | Y |
| DQB1\*03:02 | non-melanoma skin cancer | 3.46 x 10-8 | 0.92 [0.88, 0.95] | Y |
| B\*40:01 | non-melanoma skin cancer | 4.11 x 10-5 | 0.89 [0.85, 0.94] | Y |
| B\*44:03 | non-melanoma skin cancer | 5.32 x 10-5 | 0.88 [0.84, 0.93] | Y |

**Table 1.** **Cancer associations found by BMA (posterior probability >= 0.8).**

**Figure 2A. Overview of BMA results. a) Direction and magnitude of HLA allele effects on phenotypes.** x-axis indicates allele, y-axis indicates phenotype. Red shows risk effects; blue shows protective effects. A systematic literature search was conducted, yielding a set of previously reported and novel associations (**Supplementary Table 1**). Marker size indicates novelty of discovery - small (previously found) or large (novel). y-axis was clustered by mean effects of alleles on phenotypes.

**Figure 2B. Spider plot showing BMA associations between HLA alleles and binary phenotypes.** The phenotypes were grouped by category as in the legend. A line was drawn between an allele and a phenotype if the allele had a BMA posterior probability ≥ 0.8 for that phenotype. Novel associations (as determined in **Supplementary Table 1**) are shown as solid, opaque lines; previously-found associations are indicated as dashed, faded lines. Line width scales with BMA posterior mean effect size.

## Analysis of non-additive genetic effects

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**Figure 3. Non-additive dosage model associations. (A) Allele-phenotype pair.** Light grey text represents a previously-found non-additive effect (**Supplementary Table 1**). **(B) Number of UK Biobank cases for the phenotype** x-axis is log10 scaled. **(C) Odds ratio and log odds ratio for additive (green) and genotypic model (blue, dosage 1 and yellow, dosage 2).** Graphical measure of model fit. **(D) ΔAIC (AIC additive model - AIC genotype model).** A more negative value represents a larger departure from additivity.

Several analyses have looked at the evidence of non-additive effects within the HLA region of the genome[15](https://paperpile.com/c/wFd6d6/wKtR). As a further model-selection technique, we probed the presence of non-additive effects among the 178 associations resulting from the BMA analysis. Where ΔAIC was defined as the Akaike Information Criterion goodness-of-fit[22](https://paperpile.com/c/wFd6d6/uifu) difference between the additive (standard GWAS) model and the non-additive/”genotypic” model (in which rounded genotype dosage is considered as a factor as opposed to a numerical value), we selected those associations where the ΔAIC was <= -5, i.e. where the genotypic model fit was significantly better than that of the additive model.

We found 25 associations which met these criteria, of which 8 were previously documented[15](https://paperpile.com/c/wFd6d6/wKtR) (**Figure 3**). There is an enrichment for malabsorption (9) thyroid (4), and autoimmune (4) phenotypes, but psoriatic (3), asthma (2), and diabetes (2) phenotypes also feature multiple non-additive associations. We define heterozygote odds ratio (OR) to be the OR associated with Dosage 1 and homozygote OR to be the OR associated with Dosage 2, as in **Figure 3**. We find **C\*06:02** to have a non-additive effect on **psoriasis** (additive OR 2.79, 95% CI [2.69, 2.89]; heterozygote OR 3.19, 95% CI [3.05, 3.33]; homozygote OR 4.35, 95% CI [3.79, 4.99]), as backed by the literature[15](https://paperpile.com/c/wFd6d6/wKtR). We then find a series of novel associations, including that between **B\*27:05** and **ankylosing spondylitis** (additive OR 8.93, 95% CI [8.13, 9.81]; heterozygote OR 12.24, 95% CI [10.94, 13.68]; homozygote OR 14.11, 95% CI [8.24, 24.18]); low homozygote counts result in a large CI for this homozygote OR. In the case of **DRB5\*99:01** and **hypothyroidism/myxedema** (additive OR 1.22, 95% CI [1.19, 1.26]; heterozygote OR 1.44, 95% CI [1.28, 1.62]; homozygote OR 1.72, 95% CI [1.53, 1.93]), the heterozygote and homozygote CIs overlap with each other but not with the additive CI. We see a significantly higher homozygote OR for **B\*57:01** and **psoriasis** (additive OR 3.07, 95% CI [2.93, 3.23]; heterozygote OR 3.22, 95% CI [3.06, 3.39]; homozygote OR 5.42, 95% CI [4.14, 7.09]), representing another type of departure from additivity. There is also the case where all CIs overlap with each other, like for **B\*27:05** and **iridocyclitis** (additive OR 4.83, 95% CI [4.4, 5.29]; heterozygote OR 5.5, 95% CI [4.96, 6.09]; homozygote OR 8.05, 95% CI [4.8, 13.51]).

Perhaps the most dramatic of these is the effect of **DRB4\*01:01 on celiac disease** (additive OR 1.71, 95% CI [1.56, 1.87]; heterozygote OR 1.92, 95% CI [1.74, 2.12]; homozygote OR 0.62, 95% CI [0.3, 1.31]). The heterozygote and additive effect estimates are somewhat opposite the homozygote estimate, albeit the confidence interval of the homozygote OR crossing 0 (**Figure 3**). 12 associations were not found to be non-additive by our analysis, yet showed moderate non-additive effects in the literature[15](https://paperpile.com/c/wFd6d6/wKtR). These range in ΔAIC from -3.368 to +1.99 (**Supplementary Table 2**). Overall, the underlying biology of these associations could be important to disease pathogenesis.

# Discussion

This study is a comprehensive overview of associations of 156 HLA alleles to 677 phenotypes in 337,138 individuals in the UK Biobank. Using single-allele association analysis and subsequent Bayesian Model Averaging, we replicate 88 known associations between HLA alleles and cancer and other binary phenotypes and discover 90 novel associations; many of the novel associations feature phenotypes close to known associations (e.g. intestinal malabsorption and celiac disease, hypo- and hyperthyroidism), but some introduce links that are not known (e.g. anemias and acne). For example, we find that DRB1\*04:04 is associated with a family of autoimmune disorders such as rheumatoid arthritis, polymyalgia rheumatica, and connective tissue disorder, replicating signals found previously in the literature. We add to the body of known HLA associations with strong, novel effects such as that of B\*39:06 on vitamin B12 deficiency anemia. Most importantly, we find several novel associations to skin and reproductive tract cancers that could prove impactful.

These genotype-phenotype associations link the HLA region of the genome to many diverse phenotypes and suggest avenues for uncovering relevant, novel biology and developing therapeutics; however, there are many opportunities for future studies to extend this analysis, and avenues for improvement. Given that BMA is exponential in complexity with respect to the number of alleles analyzed, it is computationally infeasible to run the analysis on all BY-significant alleles for certain phenotypes. As such, it is likely that the 178 allele-phenotype associations we find are actually conservative, underscoring the impact of the HLA on disease risk. Additionally, the non-additivity analysis may be underpowered to determine homozygote odds ratios in some cases where those genotypes are low in number. It is surprising that we do not find any non-additive effects consistent with a recessive-only model, i.e. where the heterozygote OR overlaps 1. It is also unclear whether or not these non-additive effects can be classified as truly dominant, recessive, or co-dominant effects without additional formal model comparisons.

HLA alleles are oftentimes inherited as haplotypes that group several alleles in LD with each other. Better HLA typing could enable accurate haplotype frequency estimation and subsequent haplotype-based tests that enable testing of other targeted hypotheses of HLA diversity. Interactions between HLA allelotypes are also not studied here and would be a logical extension of this work; examples in which interaction effects are responsible for disease risk include the effect of an HLA-B/ERAP1 interaction on ankylosing spondylitis[42](https://paperpile.com/c/wFd6d6/cylk) and HLA-C/ERAP1 on psoriasis[43](https://paperpile.com/c/wFd6d6/mC7G). One of the functions of ERAP1 is to “trim” peptides before they are presented on HLA class I proteins. These studies therefore suggest that the risk effect mediated by certain HLA alleles also depends on peptide pre-processing.As better reference panels, superior imputation techniques, and larger population biobanks are developed, the HLA region will continue to grow in importance in human genetics, and down the line, as effective therapeutic hypotheses are generated, in clinical settings.

# Author Contributions

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# Supplementary Materials

**Supplementary Table 1. Results of systematic literature review.** Allele-phenotype pairs were marked as “novel” if not found in the literature and “non-additive” if the allele was found to have a non-additive effect on the phenotype. PMIDs and links for all references were provided.

<https://bit.ly/hla_sup1>

**Supplementary Table 2. Complete results table.** Results from PLINK additive association analysis, BMA analysis, and non-additivity analysis for each allele-phenotype pair. Number of cases per phenotype and allele frequencies per allele in the UK Biobank white British cohort.

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