Genetic susceptibility to multiple sclerosis: interactions between conserved extended haplotypes of the MHC and other susceptibility regions

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Abstract
Background: To study the accumulation of MS-risk resulting from different combinations of MS-associated conserved-extended-haplotypes (CEHs) of the MHC and three non-MHC “risk-haplotypes” nearby genes EOMES, ZFP36L1, and CLEC16A. Many haplotypes are MS-associated despite having population-frequencies exceeding the percentage of genetically-susceptible individuals. The basis of this frequency-disparity requires explanation.

Methods: The SNP-data from the WTCCC was phased at the MHC and three non-MHC susceptibility-regions. CEHs at the MHC were classified into five haplotype-groups: (HLA-DRB1*15:01 ~ DQB1*06:02 ~ a1)-containing (H+); extended-risk (ER); all-protective (AP); neutral (0); and the single-CEH (c1). MS-associations for different “risk-combinations” at the MHC and other non-MHC “risk-loci” and the appropriateness of additive and multiplicative risk-accumulation models were assessed.

Results: Different combinations of “risk-haplotypes” produce a final MS-risk closer to additive rather than multiplicative risk-models but neither model was consistent. Thus, (H+)-haplotypes had greater impact when combined with (0)-haplotypes than with (H+)-haplotypes, whereas, (H+)-haplotypes had greater impact when combined with a (c1)-haplotypes than with (0)-haplotypes. Similarly, risk-genotypes (0,H+), (c1,H+), (H+,H+) and (0,c1) were additive with risks from non-MHC risk-loci, whereas risk-genotypes (ER,H+) and (AP,c1) were unaffected.

Conclusions: Genetic-susceptibility to MS is essential for MS to develop but actually developing MS depends heavily upon both an individual’s particular combination of “risk-haplotypes” and how these loci interact.

Keywords: Genetic, Susceptibility, Multiple sclerosis, Risk, Additive, Multiplicative, Combination, Epidemiology

Background
The nature of susceptibility to multiple sclerosis (MS) is quite complex and involves both environmental and genetic factors [1–4]. Recently, considerable progress has been made in our understanding of the basis for “genetic susceptibility” in MS. Thus, to date, over 200 common risk variants (located in diverse, largely autosomal, genomic regions) have been identified as being MS-associated by genome-wide association screens (GWAS) using large arrays of single nucleotide polymorphisms (SNPs) scattered throughout the genome [5–14]. Despite this recent explosion in the number of identified MS-associated regions, however, certain alleles of the human leukocyte antigens (HLA) inside the major histocompatibility complex (MHC), known for decades, are
still the main genetic contributors to this susceptibility [11, 15–22]. The importance of these new observations to our understanding of “genetic-susceptibility” in MS is tempered by the fact that any single SNP is generally the marker of a genomic region. Indeed, the presumptively associated (i.e., “candidate”) genes can be at a considerable genetic distance from the location of the SNP itself [12–14].

For example, we described an 11-SNP haplotype (a1), which spans 0.25 megabases (Mb) of DNA surrounding HLA-DRB1, and which has the most significant association with MS of any SNP haplotype in the genome [23, 24]. Moreover, 99% of a1 SNP haplotypes carry the HLA-DRB1*15:01~HLA-DQB1*06:02 haplotype and, conversely, 99% of these HLA-haplotypes carry the a1 SNP haplotype. Individually, each SNP is highly associated both with this particular HLA-haplotype and with MS, but for none of them is this association exclusive [26]. Thus, each of these SNPs is also found in association with other HLA-haplotypes [24, 26]. Consequently, even with the large number of SNPs now identified as being MS-associated [13, 14], any such association can only be viewed as simply tagging a relatively large genomic region; it cannot be used with confidence to identify any specific gene or to implicate any specific allele with respect to its role in causing, or contributing to, a genetic-susceptibility for MS.

We and others have reported that the MHC region is composed of a relatively small collection of highly conserved extended haplotypes (CEHs), stretching across all of the “classical” HLA genes (HLA-A, HLA-C, HLA-B, HLA-DRB1, and HLA-DQB1)—a distance spanning more than 2.7 Mb of DNA [25, 26]. As shown in Additional file 1: Figure S4, this same basic population structure is also found in numerous other widely separated human populations around the world [25]. These CEHs seem to be under a strong selection pressure, presumably based upon favorable biological properties of the complete haplotype [26]. Lastly, this population structure is unlikely to be the result of a linkage disequilibrium caused by the founder effects of a small population migrating out of Africa and radiating throughout Eurasia and the Americas. Rather, the marked divergence of the CEH composition both among and between these different human groups, including Africans (Additional file 1: Tables S4a & S4b), indicates that this population structure must be due to local selection [26]. Consequently, “genetic-susceptibility” to MS is not likely to be attributable to any specific HLA allele but, rather, seems to depend upon the nature of each CEH [26]. For example, we have described a collection of SNP-haplotypes that are composed of unique combinations of the 11 SNPs (rs2395173; rs2395174; rs3129871; rs7192; rs3129890; rs9268832; rs532098; rs17533090; rs2187668; rs1063355; and rs9275141), and which span 0.25Mb of DNA surrounding the HLA-DRB1 locus [23, 24]. Ten of these SNPs are within intergenic regions whereas rs1063355 is within exon 5 of the DQB1 gene. One of these SNP-haplotypes (a1) is strongly linked to HLA-DRB1*15:01~HLA-DQB1*06:02. Thus, almost all of the CEHs, which contain the HLA-DRB1*15:01~HLA-DQB1*06:02 haplotype, are linked to the a1 SNP-haplotype, and all of these are associated with an increased MS-risk, although the magnitude of the association varies significantly among the different CEHs [25]. Nevertheless, some rare haplotypes, which include the Class II motif of HLA-DRB1*15:01~HLA-DQB1*06:02 but are not linked to (a1), seem not to carry any risk [26]. By contrast, haplotypes containing (a1), but not this Class II HLA-motif, still carry substantial risk [26]. Similarly, the disease risk for CEHs carrying HLA-DRB1*03:01~HLA-DQB1*02:01~a6 and, for the latter group of CEHs, the disease association varied widely depending upon the exact CEH composition (Additional file 1 Table S2). Finally, the MHC allele HLA-A*02:01 has been previously reported to be protective [27]. In the WTCCC, this allele was also found to be “protective” (OR = 0.69; p < 10−29), although, again, the association depends upon which CEH this allele resides rather than upon the presence of the HLA-A*02:01 allele itself (Additional file 1 Tables S1 & S2). These examples underscore the complex interactions that take place between the various MHC alleles/ haplotypes and MS-risk.

However, in addition to the MHC, other “risk” loci are worth considering. Here we focus on three non-MHC “risk-regions” of interest, which are nearby genes: EOMES, (Region d1), a transcription factor specific related to T-cell differentiation; ZFP36L1, (Region d2), a transcription factor involved in cell activation; and CLEC16A, (Region d3), a cell surface protein whose family is involved in cell activation.

In the present manuscript, we explore these relationships and interactions between the different disease-associated CEHs in the MHC region and the other “risk” haplotypes elsewhere in the genome, in order to shed light on the nature of genetic susceptibility to MS. In addition, we evaluated the manner in which disease-risk is accumulated by the combination of one or more MS-risk factors in the same individual. There are two basic epidemiological models for this accumulation of disease-risk (see Additional file 1), which have been widely utilized—the so-called additive and multiplicative relative-risk (RR) models [28–32]. Often, however, actual epidemiological observations don’t fall neatly into one model or the other. In studies of the genetic susceptibility
to MS, multiplicative risk models have generally been utilized [33–35], although this choice may not be appropriate in all circumstances (see Additional file 1).

Although, in this manuscript, we focus on the genetic aspects of MS pathogenesis, MS susceptibility is complex and involves genetic factors, environmental factors, and their interaction [1–4]. In fact, we have recently published an in-depth consideration of these issues as well as the important implications that studies of monozygotic, dizygotic twins, and non-twin siblings have regarding both genetic and environmental aspects susceptibility to MS [36]. Here, however, we will not consider further the important environmental aspects of MS pathogenesis.

Methods

Ethics statement

This research has been approved by the University of California, San Francisco’s Institutional Review Board (IRB) has been conducted according to the principles expressed in the Declaration of Helsinki.

Study participants

Wellcome trust case control consortium (WTCCC)

The WTCCC cohort included 18,872 controls and 11,376 cases with MS. The patients enrolled in the multinational WTCCC were of European or European-American ancestry. This cohort has been described previously [12–14]. For 380 controls and 232 cases the SNP data was incomplete. All of the SNP data was post-quality control [12]. Of the cases, the average age-of-onset was 32.3 years, 72.9% were women, and the mean Extended Disability Status Score (EDSS) was 3.9 [12]. The large majority (89%) of the cases had a relapsing–remitting onset [12]. The diagnosis of clinically definite MS was based upon international criteria [37–39]. Control subjects were composed of healthy individuals with European ancestry [12]. The protocol was approved by the ethical committees or institutional review boards at each of the participating centers. Informed consent was obtained from each study participant. The WTCCC granted data access for this study.

Genotyping, and quality control

The WTCCC methods for genotyping and quality control have been described previously [12–14, 16, 18, 19]. Genotyping was performed at the Wellcome Trust Sanger Institute on the Illumina Infinium platform and case samples were genotyped using a customized Humangen60-Quad chip. Common controls were genotyped on a second customized HumanLM-Duo chip (utilizing the same probes). This provided data on 441,547 autosomal SNPs scattered throughout the genome in both MS patients and control subjects after quality control. We used the HIBAG method [40] to impute the identities of the five HLA alleles in the MHC region (A, C, B, DRB1 and DQB1). Imputation methods, apart from HIBAG, such as the so-called ‘SNP2HLA’ method [41], have also been proposed. However, in a comparative study in European Americans, the HIBAG method had slightly better concordance rates and had slightly worse call rates compared to SNP2HLA [42]. Nevertheless, these differences were minimal, and both imputation methods seem equally good and the use of either is appropriate [42].

Statistical methods

Phasing

Alleles at each of five HLA loci (HLA-A, HLA-C, HLA-B, HLA-DRB1 and HLA-DQB1) and the SNPs in the Class II region of the DRB1 gene were phased using previously published probabilistic algorithms [23, 24, 43, 44]. SNP-haplotypes from 3 of the 102 non-MHC genomic regions, which had been identified previously as being significantly MS-associated, were also included in our analysis [24]. In our previous report, the MS-associated SNP haplotypes were numbered (arbitrarily) from 1 to 932. These three particular regions (arbitrarily labeled d1, d2, and d3) were selected based on their having a “risk” SNP-haplotype with 500 or more representations in the WTCCC dataset and also having the largest ORs for disease-association of any haplotype meeting this specification. The reason for choosing only three regions was that, when more regions were added, there was an insufficient number of observations to estimate the ORs for any of the possible higher order combinations [36]. Moreover, we only analyzed combinations having 15 or more representations in case or controls (combined) and of the 83 observations presented in Fig. 1, 18% had a marginal total of less than 25, 17% had a marginal total from 25 to 49 and 65% had a marginal total of 50 or more. Because the effect sizes were not able to be estimated in advance, it is possible that this approach may have considered some ORs to be non-significant because the sample sizes were too small. Nevertheless, for context, the percentage of comparisons reaching a nominal level of statistical significance (p < 0.05) within each of these three groups was similar (27%, 29%, and 39%, respectively).

The three non-MHC regions that we analyzed were located at chromosomal locations 3p24.2, 14q24.1, and 16p13.13 and in the vicinity, respectively, of the genes EOMES, ZFP36L1, CLEC16A [13, 14]. Chromosome 3; Region 22 (d1) spanned 0.65 mb of DNA and the 11-SNP-haplotype (number 234) was used [24]. This region consists of 185 different SNP combinations, of which, 1,243 (2%) are the “risk” haplotype (01,100,000,100). Chromosome 14; Region 78 (d2) spanned 0.68 mb of DNA and the 3-SNP-haplotype (number 734) was used [24].
This region consists of 7 different SNP combinations, of which, 14,091 (23%) were the “risk” haplotype (111) Chromosome 16; Region 85 (d3) spanned 0.20 mb of DNA and the SNP-haplotypes (numbers 814, 818, and 822) were combined into a single 15-SNP haplotype [24]. This was done because each of these risk-haplotypes were adjacent to each other and because the individual risk SNP haplotypes were part of the same extended 15-SNP-haplotype. This region consists of 210 different SNP combinations, of which, 24,709 (41%) are the “risk” haplotype (000010000000000).

One might reasonably question why we used these three regions (chosen based on our WTCCC cohort) to assess the additive nature of the genetic risk when a much larger dataset has now been published and, when using this larger cohort might have led to a different choice for the regions of analysis [14]. However, when we requested access to this larger dataset, we were informed that we were “requesting individual level genotype data which is per definition sensitive data and therefore subject to varying levels of restrictions from all contributing parties within the IMGSC. Your request can therefore not be granted at this time”. Consequently, the current data is all that was available to us and, theoretically, the additive or multiplicative nature of the of the accumulation of genetic risk could be assessed using any 3 non-MHC risk-haplotypes. We chose these three regions because they were the ones that met our criteria and gave us the best statistical power (see above). Adjustment for multiple comparisons [45] only impacted the significance of z-scores less than 2.5.

Also, it is important to note, that this earlier study [14], the authors looked at disease associations for more than 8 million SNPs and, controlling for both population stratification and genomic inflation, they identified (and replicated) disease associations for 233 loci (i.e., SNPs) that were confirmed to be MS-associated. Among these loci identified and, also, among those replicated in the earlier report from the WTCCC [13], were the 4 loci that we studied (i.e., the MHC together with the regions of the EOMES, ZFP36L1, & CLEC16A genes). By contrast, in the present manuscript, we are not looking to identify “true” disease associations. These are already well-established. Rather, we are looking at just four of these well-established and replicated loci, which are known, unequivocally, to be MS-associated to determine whether these loci (i.e., the associated SNP-haplotypes) exhibit either additive or multiplicative properties when forming different combinations of the risk variants.

**Fig. 1** Natural logarithm of the odds ratios (ORs) for the different combinations of MHC and non-MHC genotypes. All ORs were calculated relative to a group consisting of the same MHC genotype and with the genotypes (0,0) at all non-MHC loci involved in the comparison – see text. The MHC genotypes, in order of increasing disease-risk (Fig. S2), are presented on the x-axis (as columns) and the genotypes at non-MHC loci are presented on the y-axis (as rows). The values for ln(OR) and the z-scores for each comparison are represented as numbers at the points of intersection of the column and row for any two haplotypes. Comparisons with a z-score (|z| < 1) are shaded in yellow; comparisons with a z-score (1 ≥|z| < 2) are shaded in either pale blue (negative) or pale red (positive); comparisons with a z-score (2 ≥|z| < 3) are shaded in light red; comparisons with a z-score (3 ≥|z| < 4) are shaded in red; comparisons with a z-score (z ≥4) are maroon. Specific combinations having marginal totals of less than 15 total representations in the WTCCC are indicated by (na).
Haplotype frequencies and association testing

Disease association tests, as measured by ORs and confidence intervals (CIs) comparing cases to controls, were calculated for each of the CEHs and each of the 3 non-MHC risk haplotypes either alone or in different combinations. The WTCCC data was considered in its entirety and not further stratified. MS-associated haplotypes were analyzed by grouping them into five categories of CEHs or haplotype groups (Additional file 1: Table S3), which consisted of: (1) (H+)-carrying CEHs (i.e. those containing the HLA-DRB1*15:01–HLA-DQB1*06:02–a1 haplotype—Additional file 1: Table S1) other increased risk or “extended risk” (ER) CEHs (c23, c27, c34, c46, c68, c81, c85, c96, and c107—Additional file 1: Table S2); (3) decreased risk or “all protective” (AP) CEHs (c5, c15, c18, c24, c30, c32, c51, and c73—Additional file 1: Table S2); (4) all CEHs not in the (H+), ER, or AP groups, which were designated as (0) CEHs; and (5) the (c1) CEH by itself. We also explored “protective” groups, which either included only the (c5) CEH or excluded this CEH. However, these analyses are not presented because the findings were the same as when the AP group was analyzed as a whole. In many circumstances, an individual’s MHC genotype was specified by the haplotype combination that they possessed. For example, by this convention, an individual homozygous for (H+) would be characterized as the (H+, H+) MHC genotype. By contrast, a heterozygous individual would be characterized as having the (H+, 0), the (H+, ER), the (H+, c1), or the (H+, AP) MHC genotypes. In the principal analysis, all MS-associations were assessed compared to a reference group consisting of the (0, 0) MHC genotype. On occasion (e.g., Additional file 1), for notational simplicity, when using the (AP, AP) MHC genotype as a reference, this genotype was referred to as (AP).

Disease associations for the risk SNP-haplotypes on Chromosomes 3, 14, and 16, were assessed compared to a reference group consisting of the (0,0) MHC genotype, and excluded individuals carrying their risk-haplotypes at these chromosomal locations. We designate (collectively) all non-risk-haplotypes at each of these chromosomal locations as the (0) haplotype at each locus.

The significance of the differences in ORs for disease association (comparing cases to controls) for any two haplotypes or genotypes was determined by z-scores calculated from the differences in the natural logarithm of the ORs such that:

$$z = \frac{\ln(OR_1) - \ln(OR_2)}{\sqrt{\text{SE}[\ln(OR_1)]^2 + \text{SE}[\ln(OR_2)]^2}}$$

The Benjamini-Hochberg (BH) method was used to correct for multiple testing of the different possible haplotype combinations both within the MHC and also for combinations of the MHC with the other genetic regions [45]. This method was chosen rather than the Bonferroni method because the BH approach has been proven to be far less stringent than Bonferroni while still controlling the “Type I family-wise error rate” for multiple comparisons to be less than any desired α-level [45]. As discussed in Additional file 1, pair-wise comparisons of ORs are independent of the reference group chosen. The MHC genotype (0, 0) had the largest sample size of any (Additional file 1: Table S3) and, therefore, to maximize the statistical power to detect differences, the ORs used for pair-wise comparisons within the MHC were estimated relative to a reference group consisting of the (0,0) genotype at both the MHC and also at the any non-MHC locus included in the comparison. As noted in the Discussion, such a method eliminates the common reference group disease-risk to yield an estimate of the pairwise RR. Within the WTCCC cohort, we used a principal components (PC) analysis excluding MHC SNPs (Eigensoft) to correct the observations in Additional file 1: Tables S2 & S3 for the possible effects of population stratification, as well as regression analysis to correct for the possible effects of geographic heterogeneity [25]. These adjustments did not significantly alter any of the associations identified (Additional file 1: Tables S1 & S2).

Evaluating additive and multiplicative risk-models

The ORs for the MHC alleles (H+, ER, and 0) were determined relative to the (AP, AP) or (AP) reference group, which was assigned a value of ($R_b = R_{AP} = 1$)—see Additional file. These observed ORs were used to estimate the RR associated with each set of MHC alleles and, in turn, these RRs were used to test the additive and multiplicative risk-models for the accumulation of disease risk considering different allelic combinations at the MHC. Subsequently, using a reference group consisting of the (0,0) MHC genotype, we determined the ORs for susceptibility alleles in the three non-MHC susceptibility regions—(d1), (d2) and (d3). The (0,0) MHC genotype was chosen as the reference both because this was the largest haplotype group in the WTCCC and because there were too few representations of the (AP, AP) MHC genotype in the WTCCC dataset (Additional file 1: Table S3). Nevertheless, these observed ORs were mathematically converted into ORs relative to the (AP, AP) or (AP) MHC genotype and these re-referenced ORs, together with the ORs observed for the different allelic combinations at the MHC, were used to estimate the RRs associated with
each allelic combinations at these four genomic locations (the MHC plus the three non-MHC susceptibility regions). These estimated RRs were then used to test the additive and multiplicative risk-models for accumulation of disease risk considering the different allelic combinations at these four susceptibility regions. In all cases, only ORs estimated from combinations with ≥ 15 representations in the WTCCC were considered.

**Results**

The MHC

In the HLA region, there were 146 CEHs, which had 50 or more representations in the WTCCC dataset, and these accounted for 48% of the total number (59,884) of CEHs present. Information about 45 of these CEHs, which were previously found to have some relationship to MS susceptibility [24], is provided in Additional file 1: Tables S1 & S2. Considering the often low CEH frequencies in the WTCCC, we classified them (Additional file 1: Table S3) into five haplotype-groups: (1) (H+) CEHs (i.e., those containing the HLA-DRB1*15:01 ~ HLA-DQB1*06:02 ~ a1 haplotype.—Additional file 1: Table S1); (2) other increased risk or “extended risk” (ER) CEHs (c23, c27, c34, c46, c68, c81, c85, c96, and c107—Additional file 1: Table S2); (3) decreased risk or “all protective” (AP) CEHs (c5, c15, c18, c24, c30, c32, c51, and c73—Additional file 1: Table S2); (4) the “neutral” group (0) consisting of all those CEHs which did not belong to the (H+), (ER), or (AP) groups; and (5) the (c1) CEH by itself. Each of these groups of CEHs seemed to be segregating independently and, in the control group, frequencies for each of the different combinations were, statistically, at their Hardy–Weinberg expectations. Considering all of the combinations of “risk” CEHs (relative to the (0,0) MHC genotype), the (H+)-haplotypes accounted for 81% of the risk haplotypes in the control population and for approximately the same percentage of this risk in both men and women (80% and 82% respectively). Moreover, the likelihood of men in the control population possessing a “risk” CEH combination (26%) was approximately the same as the likelihood in women (27%). Similarly, the likelihood of men in the control population possessing an AP CEH (9%) was approximately the same as the likelihood in women (8%).

The subset of individuals who don’t carry any (H+), ER, or AP CEHs at the MHC is referred to as the (0,0) MHC genotype. In Additional file 1: Figures S1, S2, & S3, all ORs are presented relative to this group. All of the (H+) CEHs with 50 or more representations were significantly associated with MS-risk (Additional file 1: Table S2), as were, collectively, (H+)–carrying CEHs with fewer than 50 representations in the WTCCC (Additional file 1: Figure S1). Moreover, assessing, collectively, only those (H+)-carrying CEHs that had a single representation in the WTCCC, the disease association is still highly significant and of similar magnitude to other (H+)-carrying CEHs (i.e., OR = 3.0; CI = 2.7 – 3.4; p < 10^{-10}). Consequently, the (H+) haplotype, by itself, seems to contribute to the disease susceptibility in an individual although, as shown in Additional file 1: Table S1, the magnitude of this effect varies among different (H+)-carrying CEHs [25].

In addition, we defined different “risk” CEH combinations as: (1) “single copy risk” [1 copy of any (H+) CEH or any ER CEH] plus one copy of a (0) CEH; and: (2) “double copy risk” [2 copies of any (H+) CEH, the (c1) CEH, or any ER CEH, or combinations of [(H+)+ ER], [(H+)+ (c1)], or {ER+(c1)}]. The different “protective” CEH combinations were defined similarly as: (1) “single copy protective” [1 copy of an AP CEH] plus one copy of a (0) CEH; and: (2) “double copy protective” [2 copies of an AP CEH].

The “single copy risk” of MS for either (H+) or ER CEHs in women (OR = 3.0; CI = 2.8 – 3.2; p < 10^{-20}) was greater (z = 2.4; p = 0.009) than the same risk in men (OR = 2.6; CI = 2.4 – 2.8; p < 10^{-10}). By contrast, the “double copy risk” of MS in men and women was about the same.

The impact on the phenotype of an individual in response to combining two CEHs at the MHC into a single genotype is shown in Additional file 1: Figures S1 & S2. For example, as has been well described previously [11, 15–22], combining two copies of the (H+)-haplotype into a single genotype markedly and significantly increases the disease association Additional file 1: Figure S1. Nevertheless, not all (H+)-carrying haplotypes have the same disease association [26]. For example, the OR for single copy carriers of the (c2) CEH is significantly greater (z = 3.4 – 4.8; p = 10^{-3} – 10^{-6}) than the OR for either single or double-copy carriers of the (c3) CEH.

Similarly, considering the AP group of CEHs (Additional file 1: Figure S2), we found a significant dose-dependent response such that possessing 2 copies of an AP CEH is significantly more “protective” than possessing only a single copy and, in addition, the magnitude of these “protective” effects is similar to the “risk” produced by (H+) CEHs (Additional file 1: Figure S2). Moreover, having an AP CEH significantly and substantially mitigates (z = 5.2; p = 10^{-7}) the disease risk produced by single copies of (c2), (c3), or, more generally, any (H+)-haplotype (Additional file 1: Figure S2). A single copy of an ER CEH adds to the risk of a single copy of (c2), (c3), or any (H+) CEH, although it adds significantly less (z = 2.5; p = 0.006) than does a 2nd copy of an (H+) CEH (Additional file 1: Figure S2). And, finally, the (c1) CEH acts in an apparently recessive manner with little, if
any, disease risk (above the homozygous “neutral” genotype) produced by a single copy of this CEH (Additional file 1: Figure S2). Nevertheless, (and by contrast) a single copy of the (c1) haplotype adds significantly (z = 2.5–6.0; p = 0.006–10^−9) to the disease risk produced by single copies of (c2), (c3), or, more generally, of any (H+) haplotype (Additional file 1: Figure S2).

Additional file 1: Figure S3 presents the ORs for the various combinations of the non-MHC loci and, in general, as can be appreciated in the Figure, the disease risk for each of these regions seems to be dose dependent. Nevertheless, the increase in disease susceptibility that results from combining susceptibility genotypes at these three non-MHC loci with MHC genotypes is quite different for the different MHC configurations (Fig. 1). Thus, for example, the different combinations of these non-MHC “risk” haplotypes consistently increased the risk for (0, H+), (H+, H+), (0, c1), and (H+, c1) “risk” genotypes (Fig. 1). By contrast, for other “risk” genotypes such as (AP, H+) and (ER, H+) and for “protective” genotypes such as (AP, 0) and (AP, c1), these other non-MHC “risk” haplotypes seemed to contribute essentially nothing to the final risk (Fig. 1).

Figure 2 shows the impact of replacing one MHC haplotype with another in different genotypic contexts. For example, replacing an (0)-haplotype with an (H+)-haplotype has a significantly greater impact when the companion haplotype is an (0)-haplotype compared to when

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**Fig. 2** Lower triangular plots of the natural logarithm of the odds ratios (ORs) for replacing one MHC haplotype by another in different genotypic contexts. For example, at the point of intersection for (0)→(H+) in the 7th column and (c1)→(H+) in the in the 10th row, the value of (0.5) represents the log of the ratio: \( \log(0.5) = -0.6931 \). Only transitions for (H+), “extended risk” (ER), “all protective” (AP), and (0) haplotypes are shown. These transitions are indicated on both the x-axis and y-axis. Positive numbers indicate that OR for the replacement transition (indicated by the column) using the 1st companion haplotype (indicated by 1st haplotype of the transition listed in the row) is greater than the same replacement using the 2nd companion haplotype (indicated by 2nd haplotype of the transition listed in the row). Conversely, a negative number indicates that the OR for the replacement transition using the 2nd companion haplotype is greater than it is using the 1st companion haplotype. Operationally, these replacement transitions are simply ORs comparing cases to controls for the two genotypes (e.g., OR(c1, H+)/OR(c1, 0) = 3.67). Because of symmetry, this describes either of two, mathematically equivalent, replacement scenarios. The first compares the replacement of (0) by (H+) when the companion haplotype is (c1) to the same replacement when the companion is (H+). Therefore, the interpretation for the meaning of the replacement transitions represented by the rows and column are interchangeable. These two, mathematically equivalent, replacement scenarios (i.e., transitions) can be depicted as follows:

1. Only transitions for (H+), “extended risk” (ER), “all protective” (AP), and (0) haplotypes are shown. These transitions are indicated on both the y-axis (as rows) and the x-axis (as columns) and values of ln(OR) for each transition are represented as numbers at the points of intersection of the column and row for any two transitions. Specific combinations having marginal totals of less than 15 total representations in the WTCCC are indicated by (na).
2. Comparisons of the OR for the numerator with the OR for the denominator, and which have an absolute z-score > 3.0, are shaded either dark blue (negative) or dark red (positive); comparisons having an absolute z-score = 2.0–3.0 are shaded either light blue (negative) or light red (positive); comparisons having an absolute z-score = 1.0–2.0 are shaded either pale blue (negative) or pale red (positive) yellow; comparisons having absolute z-scores < 1.0 are shaded in yellow. Adjustment for multiple comparisons [45] only impacted the significance of z-scores less than 2.5.
the companion is an \((H+)-haplotype\) (Fig. 2). Thus, comparing the \((0,H+)\) genotype with the \((0,0)\) genotype had an odds ratio of: \((OR_1 = 3.0)\) whereas, comparing the \((H+,H+)\) genotype with the \((H+,0)\) genotype had an odds ratio of: \((OR_2 = 2.1)\). These two \(ORs\) were significantly different from each other \((z = 4.7)\) and had a ratio of: \(OR_1/\overline{OR_2} = 1.4\); and: \(\ln(1.4) = 0.4\).

By contrast, replacing an \((0)-haplotype\) with an \((H+)-haplotype\) has a significantly smaller impact when the companion is a \((AP)-haplotype\) (Fig. 2). Thus, comparing the \((H+,H+)\) genotype with the \((H+,0)\) genotype had an odds ratio of: \((OR = 2.1)\) whereas, comparing the \((AP,H+)\) genotype with the \((AP,0)\) genotype had an odds ratio of: \((OR = 3.4)\). These two \(ORs\) were significantly different from each other \((z = -3.4)\) and had a ratio of: \(OR_1/\overline{OR_2} = 0.6\); and: \(\ln(0.6) = -0.5\).

As can be appreciated from the figure, the impact of replacing one haplotype with another often depends considerably (and significantly) upon the nature of the companion haplotype, which, together with the haplotype being replaced, constitutes the \(MHC\) genotype (Fig. 2). This reflects the multiple haplotype-haplotype interactions that exist within the \(MHC\). Indeed, if no such interactions were present, each of the comparisons provided in the figure would have an \(OR\) of \(\sim 1.0\)—i.e., \(\ln(OR) = 0\)—i.e., they would be shaded in yellow (Fig. 2).

### Additive versus multiplicative risk

Combinations of the 3 non-\(MHC\) susceptibility regions, together with different genotypes at the \(MHC\) are presented in Figs. 3, 4, 5, 6. In each of these Figures, the \(ORs\) are those derived from a comparison using, as a reference, the group having the \((AP,AP)\) \(MHC\) genotype because this genotype had the lowest MS-risk of any we identified in the WTCCC (see Additional file 1).

In all cases, the disease risk conferred by each genotype at each locus is estimated directly from the WTCCC observations (see “Methods” section). The expectations from the additive and multiplicative risk-models are then compared to the actual observations (Figs. 3, 4, 5, 6). In almost all cases, the additive model fits better with the actual observations than does the multiplicative model, especially as more “risk” loci are included in the combinations (Figs. 4, 5, 6). Nevertheless, neither model fits (or is approximated) consistently. For example, considering only those \(MHC\) “risk” genotypes that have a disease-risk exceeding that of the \((0,0)\) \(MHC\) genotype, the actual disease-risk observed is, in general, greater than predicted by the additive model (Fig. 3). By contrast, considering 2-genotype combinations with only non-\((H+)\) & non-\(MHC\) “risk” genotypes, the observed disease-risk is generally less than predicted by the additive model (Fig. 4). Such a trend becomes more noticeable (but not dramatically so) when more “risk” loci are included in the

| \(MHC\) Genotype | \(RR_1\) | \(RR_2\) | Additive Model | Observed | Multiplicative Model |
|------------------|--------|--------|----------------|---------|---------------------|
| \(AP, AP\)       | 1      | 1      | 1              | 1       | 1                   |
| \(AP, 0\)        | 1      | 4      | 4              | 4       | 4                   |
| \(0, 0\)         | 4      | 4      | 7              | 8       | 16                  |
| \(AP, H+\)       | 1      | 14     | 14             | 14      | 14                  |
| \(0, H+\)        | 4      | 14     | 17             | 23      | 56                  |
| \(H+, H+\)       | 14     | 14     | 27             | 49      | 196                 |
| \(AP, ER\)       | 1      | 8      | 8              | 8       | 8                   |
| \(0, ER\)        | 4      | 8      | 11             | 15      | 32                  |
| \(ER, H+\)       | 8      | 14     | 21             | 33      | 112                 |
| \(AP, c1\)       | 1      | 3      | 3              | 3       | 3                   |
| \(c1, c1\)       | 3      | 3      | 5              | 25      | 9                   |
| \(0, c1\)        | 4      | 3      | 6              | 8       | 12                  |
| \(c1, H+\)       | 3      | 14     | 16             | 31      | 42                  |
| \(c1, ER\)       | 3      | 8      | 10             | 29      | 24                  |

**Fig. 3** Conformity of the observed effect of combining different \(MHC\) haplotypes with an additive and a multiplicative model of combined risk. Yellow bands represent the definitional odds ratios \((ORs)\) relative to a reference group consisting of the \((AP,AP)\) or \((AP*)\) genotype (i.e., as defined in the text: \(R_{ref} = R_{ap,ap} = 1\)). With the exception of \((c1)\), which seems to behave in an unusual fashion, the combination of other risk alleles produced, in almost all \(2\)-genotype combinations with only non-\((H+)\) & non-\(MHC\) “risk” genotypes, the observed disease-risk is generally less than predicted by the additive model (Fig. 4). Such a trend becomes more noticeable (but not dramatically so) when more “risk” loci are included in the
combinations (Figs. 5, 6). By contrast, the marked disparity with the multiplicative model becomes increasingly apparent as more “risk” loci are included in the analysis (Figs. 4, 5, 6).

Discussion
The present findings provide considerable insight to the underpinnings of genetic susceptibility to MS and indicate that this susceptibility is complex as MHC CEHs groups can either be associated with an increased or decreased disease-risk. For example, the combination of 2 “risk” CEHs (H+ or ER) results in an increased disease risk compared to a single copy of “risk” CEH alone (Additional file 1: Figures S1 & S2). Similarly, the combination to 2 “protective” CEHs (AP), results in a decreased disease risk compared to a single copy (Additional file 1: Figure S2). Finally, combining a “risk” CEH together with a “protective” CEH results in an intermediate disease risk compared with having a single copy of either type of CEH alone (Additional file 1: Figure S2). Nevertheless, there are exceptions to this general rule. Notably, a single copy of the (c1) CEH—the highest frequency CEH in both the WTCCC controls and other European populations [25, 26]—is associated with a negligible disease-risk (Additional file 1: Figure S2). By contrast, the disease-risk is significantly increased ($p < 10^{-11}$) in (c1,c1) individuals (Additional file 1: Figure S2), which suggests a recessive model. Nevertheless, a single (c1) CEH increases disease risk when combined with “risk” CEHs but not with “protective” CEHs (Additional file 1: Figure S2).

Our findings also have important implications with respect to additive and multiplicative causal models for the accumulation of genetic risk. In practice, some difficulties are encountered when trying to assess the appropriateness of either model. First, in a case–control studies (such as the WTCCC), because the incidence of the disease is not assessed (as it would be in a prospective cohort study), the actual RRIs cannot be determined [46]. However, for a rare disease such as MS [e.g., where: $P(MS) \approx 0.003$], the ORs and the RRs are almost identical [46] and, thus, can be used interchangeably. Second, selecting an appropriate reference group for calculating the RRs is crucial (see Additional file). This choice will influence how well the observations fit into one or another of these risk models. As discussed further in the Additional file 1, the theoretical underpinnings for both the additive and multiplicative models arise from the same underlying probability assumptions [28–31] and are predicated on the notion that MS-risk for the different potential “risk-factors” is as great or greater than the “risk” in the reference group (Additional file 1). This requires identifying the reference group with the lowest MS-risk of any. In MS, however, because more than 93% of the population has no MS-risk whatsoever [36], the choice of this group is not possible due to the fact that the RRs become infinite or undefined (Additional file 1). In this situation, perhaps, using a reference group having a minimal (but non-zero) MS-risk, could permit the evaluation (approximately) of whether either of these two models fits with the data (Additional file 1). In the WTCCC, the (AP, AP) or (AP*) MHC genotype has the least MS-association [OR=0.13 relative to the (0,0) MHC genotype] of any that we identified (Additional file 1: Figure S2). Therefore, this group was used to assess the appropriateness of the additive and multiplicative disease-risk models. The combination of 2 CEHs into a single MHC genotype, with the exceptions of (c1), produced an effect in between the two causal models (Fig. 3). By contrast, for (c1,c1) and (c1,ER) genotypes, the observations exceed the expectations of both models (Fig. 3).

When the other non-MHC “risk” loci are included in the analysis, the observations are closer to the additive model and even more when additional loci are included (Figs. 3, 4, 5, 6). Thus, the estimates from a multiplicative model ultimately exceed observations by 1–2 orders of magnitude (Figs. 3, 4, 5, 6). This observation is consistent with our previous observation that the distribution of penetrance values within the general population does not follow a lognormal distribution—i.e., the distribution expected for a multiplicative mode [3]. Similarly, as discussed further in Additional file 1, the multiplicative model breaks down as the disease-risk in the reference group becomes progressively lower. Consequently, based upon both theoretical considerations and observation, a multiplicative model for the accumulation of genetic risk in MS seems to be inappropriate.

The additive model, in general, performed better in these circumstances (Figs. 3, 4, 5, 6). Nevertheless, it does not explain perfectly the accumulation of genetic risk in MS. First, (c1)-containing CEH genotypes consistently...
| 2-Genotype Combinations | RR₁ | RR₂ | Additive Model | Observed | Multiplicative Model |
|--------------------------|-----|-----|----------------|----------|---------------------|
| (0, 0) + (0, d₁)        | 15  |     |                | 15       |                     |
| (0, 0) + (d₁, d₁)       | 20  |     |                | 20       |                     |
| (0, 0) + (0, d₂)        | 8   |     |                | 8        |                     |
| (0, 0) + (d₂, d₂)       | 10  |     |                | 10       |                     |
| (0, 0) + (0, d₃)        | 9   |     |                | 9        |                     |
| (0, 0) + (d₃, d₃)       | 11  |     |                | 11       |                     |
| (0, H+) + (0, d₁)       | 23  | 15  | 37             | 42       | 345                 |
| (0, H+) + (d₁, d₁)      | 23  | 20  | 42             | 59       | 460                 |
| (0, H+) + (0, d₂)       | 23  | 8   | 30             | 25       | 184                 |
| (0, H+) + (d₂, d₂)      | 23  | 10  | 32             | 42       | 230                 |
| (0, H+) + (0, d₃)       | 23  | 9   | 31             | 28       | 207                 |
| (0, H+) + (d₃, d₃)      | 23  | 11  | 33             | 31       | 253                 |
| (H+, H+) + (0, d₁)      | 49  | 15  | 63             | 69       | 735                 |
| (H+, H+) + (0, d₂)      | 49  | 8   | 56             | 57       | 392                 |
| (H+, H+) + (d₂, d₂)     | 49  | 10  | 58             | 67       | 490                 |
| (H+, H+) + (0, d₃)      | 49  | 9   | 57             | 59       | 441                 |
| (H+, H+) + (d₃, d₃)     | 49  | 11  | 59             | 54       | 539                 |
| (0, c₁) + (0, d₁)       | 3   | 15  | 17             | 15       | 45                  |
| (0, c₁) + (0, d₂)       | 3   | 8   | 10             | 10       | 24                  |
| (0, c₁) + (d₂, d₂)      | 3   | 10  | 12             | 12       | 30                  |
| (0, c₁) + (0, d₃)       | 3   | 9   | 11             | 10       | 27                  |
| (0, c₁) + (d₃, d₃)      | 3   | 11  | 13             | 11       | 33                  |
| (c₁, c₁) + (0, d₂)      | 25  | 8   | 32             | 36       | 200                 |
| (c₁, c₁) + (0, d₃)      | 25  | 9   | 33             | 30       | 225                 |
| (c₁, c₁) + (d₃, d₃)     | 25  | 11  | 35             | 39       | 275                 |
| (H+, c₁) + (0, d₁)      | 31  | 15  | 45             | 57       | 465                 |
| (H+, c₁) + (0, d₂)      | 31  | 8   | 38             | 31       | 248                 |
| (H+, c₁) + (d₂, d₂)     | 31  | 10  | 40             | 46       | 310                 |
| (H+, c₁) + (0, d₃)      | 31  | 9   | 39             | 35       | 279                 |
| (H+, c₁) + (d₃, d₃)     | 31  | 11  | 41             | 51       | 451                 |
| (Aₚ , 0) + (0, d₁)      | 4   | 15  | 18             | 8        | 60                  |
| (Aₚ , 0) + (0, d₂)      | 4   | 8   | 11             | 4        | 32                  |
| (Aₚ , 0) + (0, d₃)      | 4   | 8   | 13             | 7        | 40                  |
| (Aₚ , 0) + (d₃, d₃)     | 4   | 9   | 12             | 4        | 36                  |
| (Aₚ , 0) + (d₁, d₁)     | 4   | 11  | 14             | 7        | 44                  |
| (Aₚ , H+) + (0, d₃)     | 14  | 9   | 22             | 17       | 126                 |
| (Aₚ , c₁) + (0, d₂)     | 3   | 8   | 10             | 3        | 24                  |
| (Aₚ , c₁) + (0, d₃)     | 3   | 9   | 11             | 3        | 27                  |
| (Aₚ , c₁) + (d₂, d₂)    | 3   | 11  | 13             | 1        | 33                  |
| (0, ER) + (0, d₁)       | 15  | 15  | 29             | 16       | 225                 |
| (0, ER) + (0, d₂)       | 15  | 8   | 22             | 13       | 120                 |
| (0, ER) + (d₂, d₂)      | 15  | 10  | 24             | 30       | 150                 |
| (0, ER) + (0, d₃)       | 15  | 9   | 23             | 20       | 135                 |
| (0, ER) + (d₃, d₃)      | 15  | 11  | 25             | 23       | 165                 |
| (ER, H+) + (0, d₂)      | 33  | 8   | 40             | 33       | 264                 |
| (ER, H+) + (0, d₃)      | 33  | 9   | 41             | 35       | 297                 |
| (ER, H+) + (d₂, d₂)     | 33  | 11  | 43             | 40       | 363                 |
| (ER, c₁) + (0, d₂)      | 33  | 8   | 40             | 29       | 264                 |

Fig. 4 (See legend on previous page.)
exceed the additive expectations (Fig. 3). Second, effect of a given MHC haplotype is dependent on its companion MHC haplotype in a genotype (Fig. 3). Third, the effect of the 3 non-MHC “risk” haplotypes is not consistent across all MHC genotypes (Fig. 4). And fourth, when more loci are included in the analysis, the observations become
increasingly less than what is predicted by the additive model (Figs. 3, 4, 5, 6). Taken together, these observations suggest that the accumulation of genetic risk from these “susceptibility loci” is inconsistent with both models. Rather, the magnitude of any change in disease-risk associated with the inclusion of additional “susceptibility loci” seems to depend upon the exact state at each “risk-locus” and on the interaction across all loci. Such a conclusion is also consistent with theoretical considerations [36].

The MHC is known to have a remarkable diversity [27] although the individual HLA-alleles occur as part of linked haplotypes or CEHs (Additional file 1). Even though some CEHs share common features, such as carrying the (H+)-haplotype, the degree of association with MS varies depending upon the exact CEH considered (Additional file 1: Tables S1 & S2). For example, both (c2) and (c3) CEHs carry the (H+)-haplotype, but their MS-association differed significantly ($z = 4.8; p < 10^{-6}$). It might be tempting to attribute this difference to (c3) carrying the potentially “protective” HLA-A*02:01 allele (S2 File; Table S2). However, other HLA-A*02:01 and (H+) carrying CEHs (e.g., c50, c58, and c139) do not seem to be similarly protected (S1 File; Table S2). Finally, each identified CEH probably represents a diverse set of CEHs. Thus, because the genomic region from HLA-A to HLA-DQB1 (spanning ~3 mb of DNA) is quite “gene-dense”, each of the CEHs that we defined, almost certainly, represent groups of CEHs, which carry many other linked variants of other genes.

![Table 1](image)

Table 1: Conformity of the observed effect of combining different genotypes at the MHC and three susceptibility regions with an additive and a multiplicative model of combined risk. The non-MHC susceptibility haplotypes are: (d1), (d2), and (d3)—see “Methods” section. The ORs listed are those actually observed (Figs. 4, 5). Only haplotype combinations with $\geq 15$ or more representations in the WTCCC are shown. Combinations with fewer than 50 representations are shaded in pink; combinations with at least 50 representations are shaded in green.

In the study of human genetics there has been a long-running debate between the so-called “common-disease, common variant” and the “common-disease, rare variant” hypotheses [52]. Nevertheless, with our improved genetic sophistication, it has become increasingly clear that, in different specific circumstances, either (or both, or neither) hypotheses could be operative [52]. In fact, our observations also support this notion. For example, on the one hand, all of the MHC CEH combinations, which impact genetic susceptibility to MS, are quite rare. None
has a population frequency in controls of more than 6.2% and the large majority of them have population frequencies well below 1% (Additional file 1: Tables S1 & S2). On the other hand, considered collectively, those CEH combinations, which include the Class II (H+) haplotype, have a WTCCC Control population frequency of 23%. Indeed, this particular haplotype-group is the most prevalent and, therefore, the most highly selected of all such Class II haplotype combinations in northern American and European populations [26]. Consequently, the most prevalent, and therefore the most highly selected [26], CEHs are also those that are associated with the highest MS-risk (Additional file 1: Tables S1 & S2). Thus, it is clear that these particular CEHs must come with both adaptive and deleterious consequences for the individual. In addition, although the CEH composition differs markedly among long-separated human populations (Additional file 1: Tables S4a & S4b), as shown in Additional file 1: Figure S4, specific CEHs are still being strongly selected in each of them [26]. Consequently, the benefits of the adaptive features of these CEHs must outweigh the risk of any deleterious ones. Obviously, for circumstances, either in which the risk of MS is small or in which MS has little impact on an individual’s eventual number of surviving children, even a modest advantage in favor of a specific CEH might still cause it to be selected. In this regard, a recent French study estimated that women with MS had 31% fewer children than their contemporary controls [33]. If this observation is correct, it suggests that there is a strong selective disadvantage to having MS. Therefore, the explanation for the benefits of these MS-associated CEHs outweighing the risks is likely to lie in an individual’s low risk of MS rather than the disease having little impact on their fertility. Based on our observations, this seems likely to be the case. Thus, because natural selection can only select against those genotypes, which actually carry risk (relative to other genotypes), both the fact that so few individuals are susceptible and the fact that so few of the individuals who are susceptible ever develop MS [36], makes such a favorable tradeoff between adaptive and deleterious features considerably more likely to occur.

Conclusions

Comparing the different combinations of “risk” haplotype-groups to that of the final MS-risk demonstrates that an additive risk model is considerably more likely than a multiplicative risk-model. Nevertheless, even the additive-model is inconsistent for combinations of the four loci considered in this manuscript. For example, (H+) haplotypes have a significantly greater impact when combined with (0)-haplotypes than they have when combined with other (H+)-haplotypes. By contrast, (H+)-haplotypes have greater impact when combined with a (c1)-haplotype than they have when combined with (0)-haplotypes. Similarly, risk-genotypes (0,H+), (c1,H+), (H+,H+) and (0,c1) were additive with risks from non-MHC risk-loci, whereas risk-genotypes (ER,H+) and (AP,c1) were unaffected by similar combinations.

Genetic susceptibility to MS is very uncommon in the population and, yet such susceptibility is essential for MS to develop [36]. How likely MS is to develop depends heavily upon both an individual’s particular combination of genetic “risk-loci” and also how these different loci interact with each other to make genetic-susceptibility more likely in that individual [36]. In addition, a person’s environmental experience, and stochastic processes play important roles in determining whether or not MS actually develops in a susceptible individual [36].

Abbreviations

MHC: Major histocompatibility complex; HLA: Human leukocyte antigen; CEH: Conserved extended haplotype; ER: Extended risk; AP: All protective; SNP: Single nucleotide polymorphism; H+ - The haplotype HLA-DRB1*15:01 – HLA-DQB1*06:02 – a1; EBV: Epstein Barr virus; OR: Odds ratio; WTCCC: Wellcome Trust Case Control Consortium; BH: Benjamini-Hochberg; MS: Multiple sclerosis.

Supplementary Information

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Additional file 1: This Supplemental File describes composition of the CEHs found in the WTCCC dataset as well as their individual relationships to MS susceptibility, the impact on susceptibility of various combinations either of CEHs at the MHC or of the non-MHC loci and, finally, how this CEH composition differs between populations around the world. Furthermore, this file considers the theoretical underpinnings for the commonly used additive and multiplicative Models for the accumulation of disease “risk” with increasing number of “risk haplotypes” being present in an individual’s genotype.

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Authors’ contributions

DSG: Conceptualized and led the project, analyzed and assisted in the interpretation of the data, developed the Susceptibility Model, and wrote the original draft of the manuscript. PK: Assisted in the critical interpretation of the data and review of the manuscript. PA: Assisted in the critical interpretation of the data and review of the manuscript. NV: Assisted in the analysis and critical interpretation of the data and the review of the manuscript. All authors have read and approved the manuscript.

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Availability of data and materials

The Wellcome Trust Case Control Consortium (WTCCC) granted data access for this study. The data is available by request to the authors of the paper from which the WTCCC dataset is taken [12].
Declarations

Ethics and consent to participate
This research has been approved by the University of California, San Francisco's Institutional Review Board (IRB) has been conducted according to the principles expressed in the Declaration of Helsinki. The Ethical Committees or Institutional Review Boards at each of the participating centers approved the protocol and written informed consent was obtained from each study participant.

Consent for publication
Not applicable. This manuscript contains no individual’s data.

Competing interests
There are no competing interests.

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