

A High-Resolution Linkage-Disequilibrium Map of the Human Major Histocompatibility Complex and First Generation of Tag Single-Nucleotide Polymorphisms

Marcos M. Miretti,¹ Emily C. Walsh,² Xiayi Ke,³ Marcos Delgado,¹ Mark Griffiths,¹ Sarah Hunt,¹ Jonathan Morrison,¹ Pamela Whittaker,¹ Eric S. Lander,² Lon R. Cardon,³ David R. Bentley,¹ John D. Rioux,² Stephan Beck,¹ and Panos Deloukas¹

¹Wellcome Trust Sanger Institute, Hinxton, United Kingdom; ²Broad Institute of Massachusetts Institute of Technology and Harvard, Cambridge, MA; and ³Wellcome Trust Centre for Human Genetics, University of Oxford, Oxford, United Kingdom

Autoimmune, inflammatory, and infectious diseases present a major burden to human health and are frequently associated with loci in the human major histocompatibility complex (MHC). Here, we report a high-resolution (1.9 kb) linkage-disequilibrium (LD) map of a 4.46-Mb fragment containing the MHC in U.S. pedigrees with northern and western European ancestry collected by the Centre d'Etude du Polymorphisme Humain (CEPH) and the first generation of haplotype tag single-nucleotide polymorphisms (tagSNPs) that provide up to a fivefold increase in genotyping efficiency for all future MHC-linked disease-association studies. The data confirm previously identified recombination hotspots in the class II region and allow the prediction of numerous novel hotspots in the class I and class III regions. The region of longest LD maps outside the classic MHC to the extended class I region spanning the MHC-linked olfactory-receptor gene cluster. The extended haplotype homozygosity analysis for recent positive selection shows that all 14 outlying haplotype variants map to a single extended haplotype, which most commonly bears *HLA-DRB1*1501*. The SNP data, haplotype blocks, and tagSNPs analysis reported here have been entered into a multidimensional Web-based database (GLOVAR), where they can be accessed and viewed in the context of relevant genome annotation. This LD map allowed us to give coordinates for the extremely variable LD structure underlying the MHC.

Introduction

Characterization of patterns of linkage disequilibrium (LD) at a fine scale across the genome is central to the optimal design and execution of association studies of common variants that influence susceptibility of common diseases, as well as the understanding of processes such as recombination, mutation, and selection. LD maps report directly on marker correlation in the population and can thus guide marker selection for association studies. It is well established that regions of high LD display low haplotype diversity; thus, common haplotypes can be efficiently tagged with only a subset of all common variants, known as “haplotype tag SNPs” (tagSNPs) (Daly et al. 2001; Johnson et al. 2001; Chapman et al. 2003; Goldstein et al. 2003; Clayton et al. 2004). However, the use of tagSNPs carries some loss of power for

detection of all other common variants, with the exception of completely correlated markers. Many methods exist to identify an optimal set of tagSNPs that are based on observed correlations among an initial set of markers, to avoid the collection of redundant information (see Goldstein et al. [2003] for review). These methods take advantage of high-resolution LD maps that show increasing efficiency at higher marker density (Ke et al. 2004a).

Several linkage scans and association-mapping studies of most, if not all, autoimmune diseases—such as type 1 diabetes, multiple sclerosis, systemic lupus erythematosus (Davies et al. 1994; Marchini et al. 2003; Harbo et al. 2004; Marrosu et al. 2004; Oksenberg et al. 2004)—and infectious diseases like malaria and AIDS (Hill et al. 1991; Carrington and O'Brien 2003) have given strong signals in the major histocompatibility complex (MHC), a biologically and medically important region on human chromosome 6p that harbors many immune genes. Partly because of these disease studies, this region attracted a lot of attention, particularly with respect to the structure of LD and recombination. Whereas, historically, only the classical HLA loci were studied by genotyping, recent systematic efforts to characterize LD patterns across the whole of

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Address for correspondence and reprints: Dr. Stephan Beck, The Wellcome Trust Sanger Institute, Wellcome Trust Genome Campus, Hinxton, Cambridge, CB10 1SA, United Kingdom. E-mail: beck@sanger.ac.uk

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the MHC have been reported at increasing marker densities by use of SNPs (Walsh et al. 2003; Stenzel et al. 2004). These complement a number of studies of the recombinational rates in the region. In three different studies, Cullen and colleagues (1995, 1997, 2002) have mapped recombination rates across the MHC class II region and have demonstrated the presence of recombination hotspots near *HLA-DNA*, *BRD2* (formerly *RING3*), and *HLA-DQB1* and within the *TAP2* gene. But, perhaps most importantly, examining the relationship between LD and recombination with a combination of SNP genotyping and sperm recombination mapping, Jeffreys et al. (2001) demonstrated directly the correlation between regions of low LD and the presence of recombination hot spots in a discrete 200-kb segment of the MHC. However, all the studies named above have been limited by the unavailability of a dense SNP map of the entire MHC. Recently, the total number of known SNPs across the MHC has risen to >36,000 (dbSNP121), mainly through efforts of the HapMap Project (The International HapMap Consortium 2003) and the complete resequencing of the MHC region in homozygous cell lines, which contributed >50% of the available SNPs (Stewart et al. 2004). This vast resource allowed us to construct a 1.9-kb-resolution LD map across the MHC in U.S. pedigrees with northern and western European ancestry collected by CEPH and to suggest a first-generation set of haplotype tagSNPs for use in association studies. The map shows a high degree of variability in average LD between MHC subregions and reveals that the olfactory-receptor gene cluster in the extended class I region falls within the largest regions of high LD currently known in the human genome. Using Phase 2.1 (Li and Stephens 2003; Crawford et al. 2004) and the method described by McVean et al. (2004), we obtain estimates of recombination rates across the entire MHC region and show that 90% of predicted peaks (hotspots) correlate with LD breaks.

Material and Methods

DNA Samples and Genotyping

We selected a total of 3,892 SNPs, across a 4.46-Mb region spanning the MHC (human chromosome 6, 28,918,812–33,377,873 bp [National Center for Biotechnology Information build 34]), from dbSNP121 and the International HapMap Project. Selection criteria required that SNPs have spacing >500 bp and have an assay design for Illumina's Golden Gate genotyping platform (Oliphant et al. 2002). Markers were typed across a panel of 190 DNA samples from CEPH families (Utah residents with ancestry from northern and western Europe); DNA was obtained from the Coriell Cell Repository. In total, 180 founder chromosomes were used for

the haplotype construction. Note that this panel includes all the samples used both by the International HapMap Project and in the map by Walsh et al. (2003) and hence allowed full integration of data on the ~820 markers genotyped by this effort (see sample list at the Human Chromosome 6 Project Overview Web site). Genotyping was performed at a multiplex level of 1,536 SNPs per well, and data quality was assessed by duplicate DNAs ($n = 8$) and inheritance tests among parent-child trios. SNPs with more than two discrepant calls were removed. PEDCHECK (O'Connell and Weeks 1998) and PEDSTATS (Center for Statistical Genetics) were used to identify and remove genotypes that lead to Mendelian inconsistencies. The genotype confidence score for keeping data was set to 0.25. Finally, we removed loci with <80% of all possible genotypes, out of Hardy-Weinberg equilibrium ($\chi^2 \geq 10$) and zero heterozygosity.

In total, we retained 3,122 SNPs for further analysis. Of those, 787 were either nonpolymorphic or had a minor-allele frequency (MAF) <5% in our sample (5% is the MAF cutoff used by International HapMap Project). The distribution of these 787 markers is not random, which suggests variation deserts in three regions in which the density of such markers is clearly increased (fig. 1, *black arrows*). Thus, we compared the heterozygosity distribution obtained here with that from a genotyping data set of 396 SNPs assayed with the MassEXTEND assay and mass spectrometry (Sequenom platform) in the same sample (Walsh et al. 2003). Data sets from both platforms showed the same trend, which supports the presence of low-heterozygosity regions rather than genotyping problems. We also observed a higher failure rate (fig. 1, *blue line*) around the highly polymorphic genes *HLA-A*, *HLA-B*, *HLA-C*, *HLA-DQB*, and *HLA-DRB1*, which suggests that alternative typing assays may be needed in these regions. The 2,335 loci with MAF $\geq 5\%$ —an average spacing of 1 SNP per 1.9 kb—were used in all further analyses (marker and raw genotypes are available at the Human Chromosome 6 Project Overview Web site and have been submitted to dbSNP).

Estimation of LD

To capture the strength of LD between markers, we estimated two LD measurements on the basis of the pairwise disequilibrium coefficient— D' (Lewontin 1964) and r^2 —by using the HaploXT program in the GOLD package (Abecasis and Cookson 2000). Nonrecombinant haplotypes were derived by employing Merlin (Abecasis et al. 2002) and were then used to estimate haplotype frequencies in founder chromosomes. To compare decay rates among MHC regions, the decay of LD was plotted as a function of physical distance. The decay rates represent the average r^2 and D' values for all markers sep-

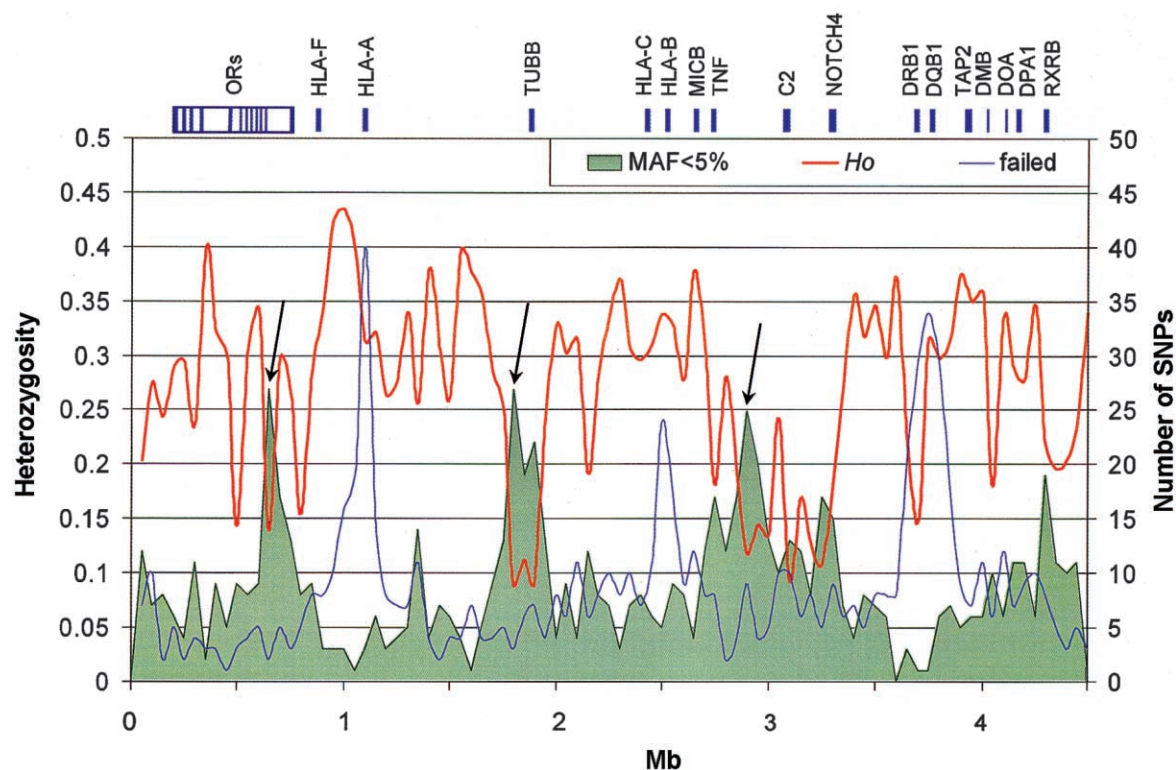


Figure 1 Low-heterozygosity regions in the MHC. The observed heterozygosity (“Ho,” red line) averaged in 50-kb windows is plotted with the number of loci with MAF <5% (green area). SNPs showing MAF <5% are not randomly distributed across the analyzed 4.459-Mb region. Specifically, three regions (black arrows) presented loss of heterozygosity; most SNPs are monomorphic in this population. These fragments contain *OR2H1* and *MAS1L*; *DHX16*, *NRM*, *MDC1* *TUBB*, and *FLOT1*; *BAT5*, *LY6GD*, *C6orf25*, *DDAH2*, *C6orf26*, and *VARS2*, respectively. The uneven distribution of the 770 SNPs that failed to generate genotyping data is represented by the blue line (“failed,” number of SNPs in 50-kb windows). Typing failures are concentrated mainly in three genomic regions (blue peaks), including *HLA-A*, *HLA-B*, *HLA-C*, *HLA-DQB1*, and *HLA-DRB1*, which suggests that the highly polymorphic nature of these genes might be responsible for the failure.

arated by distance S (for $S = 10$ kb, 20 kb, 30 kb, ... 500 kb). Moving average (sliding window [SW]) of pairwise LD coefficients (r^2) was performed at 500-kb windows across the MHC to display general trends of LD. Pairwise r^2 values from all markers spaced between 25 kb and 250 kb were averaged and plotted in 500-kb successive windows. At each step, the 500-kb window moved forward 50 kb. A short increment between windows—or large overlapping—contributes to smoothing the curve and avoids distinct peaks. LD patterns, particularly LD breaks and high-LD long-range regions, were visualized using the GOLDsurfer program (Pettersson et al. 2004) independent of block assessment.

To define haplotype blocks, a set of consecutive sites between which there is little or no evidence of historical recombination, we adopted the D' -based criteria of Gabriel et al. (2002). Haplotype-block-structure assessment, as implemented in Haploview 2.05 (Haploview Web site), was performed for the whole data set without segmentations (3,122 SNPs), and the “block-like” features—number, distribution, and average size—were evaluated.

Exact map positions of LD breaks—determined by SNP coordinates at block boundaries—were identified by analyzing the haplotype-block distribution across the region, with consideration of multiallelic D' values between consecutive blocks. Pairwise disequilibrium estimates, regardless of the haplotype-block definition (Haplo.XT), were interpreted using GOLDsurfer, and the resultant LD-break distribution was compared with that from the haplotype-block analysis.

tagSNP Selection

Application of a stringent r^2 threshold ($r^2 > 0.8$) between SNPs would allow the selected tagSNPs to resolve >80% of all existing haplotypes (Chapman et al. 2003). The LDselect algorithm (Carlson et al. 2004) groups tagSNPs in bins in which all pairwise r^2 values exceed the threshold, so that only one tagSNP would need to be genotyped per bin. The binning process takes place, regardless of the strength of the underlying LD (low or high), for a given region, and behaves independent of any block requirements. Tagging efficiency was defined

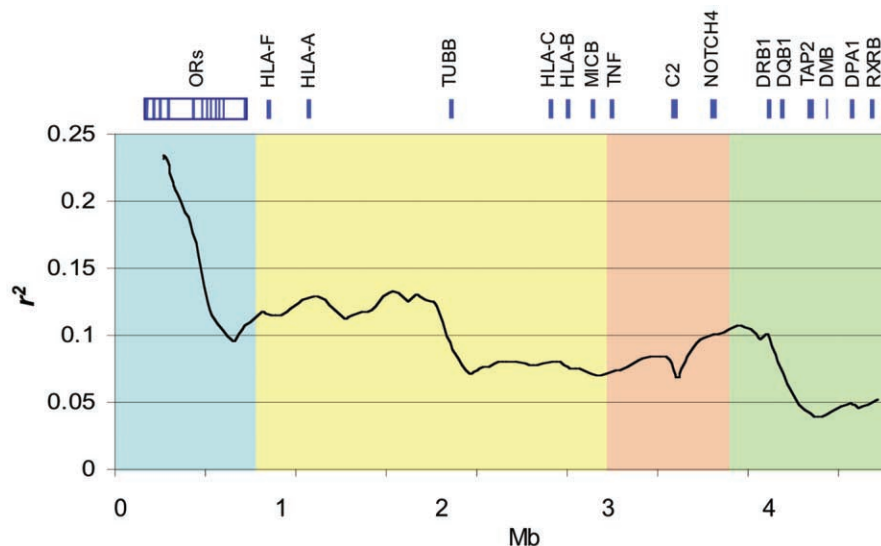


Figure 2 SW plot of average r^2 across the MHC. Average r^2 was calculated from 25 kb to 250 kb in 500-kb SWs, with 50-kb increments between windows. The SW plot captures trends of LD (r^2) by averaging r^2 values between a given marker and all the SNPs up to 250 kb. Avoiding pairwise comparisons with surrounding markers (25 kb each side) excludes the raising effect of closely linked loci on LD. MHC extended class I, class I, class III, and class II regions (*blue, yellow, orange, and green*, respectively) present comparatively distinct variation patterns of long-range LD, which is reflected in the haplotype-block analysis and interferes in the SNP-tagging process.

as n/n_b , where n_b is the number of tagSNPs selected to cover the region and n the total number of genotyped SNPs in the region (Ke et al. 2004a). The first SNP from each bin SNP list was selected for the final list, which was loaded into the GLOVAR database (Human Genome Server).

Variation of Recombination Rate across the MHC Region

Phase 2.1 was used for estimation of recombination rate and hotspot evidence (Li and Stephens 2003; Crawford et al. 2004). The region was separated into windows of 50 SNPs, with 5 SNPs overlapped between neighboring windows. Phased founder haplotypes, which were obtained by running Merlin, were fed into the program (Abecasis et al. 2002). To rescale population-genetic estimates, N_e was estimated from a comparison of the genetic map distance across the MHC region (Cullen et al. 2002) and the estimated population-recombination rate for the same region (McVean et al. 2004).

Coordinates of recombination hotspots were mapped to the current genome assembly to determine the context in which hotspots take place (intra- or intergenic), and consistency was verified by comparison with the haplotype-block (boundaries) distribution

Extended-Haplotype Homozygosity Analysis

Initial SNP chromosomal phasing was done using the Genehunter program (Kruglyak et al. 1996). Further haplotype inference of SNP genotyping data was per-

formed using a refinement of the model used in PHASE (Stephens et al. 2001), with allowance for recombination (Fearnhead and Donnelly 2001). Both unphased and missing SNP data were inferred in this manner. Since we have a dense set of markers and since most markers are in strong LD with several other markers, we do not believe that the phasing has introduced serious bias into our results. Only independent grandparental chromosomes from all families were analyzed.

Extended haplotype homozygosity (EHH) analysis (Sabeti et al. 2002) was performed for each haplotype-block allele (Gabriel et al. 2002), by use of Finch software (P. C. Sabeti, P. Varilly, B. Fry, E. Lander, unpublished material) and with centimorgan estimations as distance. Genetic distances were estimated using the empirical sperm recombination map of Cullen et al. (2002). Since the mapping efforts of Cullen et al. (2002) covered only a subset of the region analyzed in our study, genetic distances from marker *rs422331* to marker *rs1233384* (749 kb) were estimated by examination of ancestral recombination by use of methods of McVean et al. (2004). Outlying variants were chosen on the basis of two criteria designed to pick alleles with high EHH values for their frequency class. First, as a simple approx-

The figure is available in its entirety in the online edition of *The American Journal of Human Genetics*.

Figure 3 Decay of LD as a function of distance

imation of the distribution, we ranked scores by EHH value \times allele frequency. Values >4.5 SD above the mean were considered outliers. Second, all variants were sorted by frequency into 5% bins. Outliers had EHH values ≥ 4.5 SD above the mean for the remaining values in that bin.

Results

High-Resolution LD Map of the MHC

We considered 2,335 loci with an MAF $\geq 5\%$ for construction of an LD map across a 4.46-Mb region of chromosome 6 that contains the MHC. The average marker density is one SNP per 1.9 kb (median 1.2 kb), with only two gaps >25 kb; the largest was 63 kb. Of all gaps, 85% were <3 kb.

We first examined LD at a broad level across the region using an SW plot of average r^2 . Figure 2 shows that LD fluctuates greatly along the MHC, consistent with the findings of extreme LD variability throughout the human genome. It is interesting, though, to note the striking differences in long-range LD trends between MHC subregions. For example, the extended class I region, which harbors an olfactory-receptor gene cluster (Younger et al. 2001), is a contiguous block of high LD that extends ~ 540 kb. The class I region shows lower overall LD than does the extended class I region, with the telomeric half having higher LD than the centromeric half. LD range is lowest within the MHC class II and extended class II regions, consistent with the presence of multiple known recombination hotspots (Cullen et al. 1995, 1997, 2002; Jeffreys et al. 2001; Jeffreys and May 2004). We also examined the decay of LD as a function of physical distance for each MHC domain separately. The decay plots show fairly distinctive slopes (fig. 3), consistent with the patterns emerging from the SW analysis (fig. 2). The LD-decay pattern of the extended class I and class II regions are likely to fall within the extremes of genome distribution. For example, we compared LD decay between the MHC and a 10-Mb region of chromosome 20 (Ke et al. 2004b), which represents the long-

est genomic fragment with available typing information close to the SNP density reported here. LD decay between the two regions is comparable only when values are averaged across the MHC (fig. 3).

To examine the microstructure of LD obtained with the SW analysis, we used the method described by Gabriel et al. (2002) to define haplotype blocks. This method was previously employed for another data set from this region by Walsh et al. (2003), who found 17 haplotype blocks covering only 25% of the MHC sequence; additional block-like regions have been reported by Stenzel et al. (2004). Using the same criteria (Gabriel et al. 2002), we obtained 202 blocks covering 81.95% of the MHC region as a direct result of the much higher marker density employed in the present study. Haplotype blocks have an average size of 18 kb and 6.4 haplotypes per block. To visualize the distribution of haplotype blocks across the MHC in the context of other genomic features such as gene structures and known SNPs, we used the GLOVAR genome browser maintained at the Sanger Institute. Figure 4A provides an overview of the whole region and shows the SNP density, manually annotated protein-coding genes (Mungall et al. 2003), and haplotype blocks.

As expected from the SW analysis, a large portion of the extended class I region (540 kb) that extends over the olfactory-receptor cluster is represented in just two LD blocks interrupted by a single recombination hotspot spanning ~ 750 bp (fig. 4B). In general, large blocks cover most of the classical class I region, except for areas surrounding *HLA-A*, *HLA-B*, *HLA-C*, and *MICA*, in which the LD pattern is disrupted and blocks are shorter. Several studies have shown experimentally that most recombination takes place in narrow hotspots, delimiting regions of high LD (Cullen et al. 1997, 2002; Jeffreys et al. 2001; Jeffreys and May 2004). There are at least four prominent LD breaks intermingled among relatively long blocks in the telomeric portion of the class I region. On its proximal side, recombination appears to take place more uniformly, which results in a number of short stretches of high LD. Three additional hotspots are ap-

Figure 4 LD structure across the MHC. *A*, Distribution of haplotype blocks across the MHC region, as viewed in the GLOVAR genome browser. Haplotype blocks, according to criteria of Gabriel et al. (2002) implemented by Haploview 2.05 (Haploview Web site), are represented by red bars. Each bar corresponds to an individual haplotype block comprising a number of SNPs (red marks), which are located according their map position. This enables an accurate interpretation of the LD-block distribution, size, and gaps in the context of additional genomic features, such as gene annotation, SNP density, and physical distance. The distribution of tagSNPs selected in this work is generated in the GLOVAR genome browser and is indicated by a green track under the haplotype blocks. *B*, High-resolution view of 720 kb of the extended MHC class I region, as represented by GOLDsurfer 3D view of D' values (Pettersson et al. 2004). This region contains a large cluster of olfactory-receptor genes in high LD (540 kb), interrupted by a single recombination hotspot between *OR12D3* and *OR12D2*. This long-range LD region includes 13 contiguous haplotype blocks, according to the criteria of Gabriel et al. (2002) (see corresponding inset in panel A). *C*, View of the LD structure (D' values) within the MHC class II region for which experimental evidence for recombination hotspots have been described elsewhere (Cullen et al. 1997; Jeffreys et al. 2001). High-LD areas (red blocks) are separated by recombination hotspots. The first three LD breaks correspond to recombination hotspots mapped at *TAP2* and *HLA-DMB* and between *BRD2* and *HLA-DOA* (Cullen et al. 1997; Jeffreys et al. 2001). Another LD break is visualized between *HLA-DOA* and *HLA-DPA1*.

parent in the class III region; the most centromeric one resides after *NOTCH4* before the class II border, 70 kb 5' of the hotspot mapped by Stenzel et al. (2004). Within the class II region, those recombination hotspots described elsewhere (Cullen et al. 1997; Jeffrey et al. 2001) at *TAP2*, *HLA-DMB*, and between *BRD2* and *HLA-DOA* were clearly identified (fig. 4C). Two additional hotspots are evident within the centromeric portion of the class II region, the stronger one between *HLA-DOA* and *HLA-DPA1* and the other between *HLA-DPB1* and *HLA-DPB2*.

For each MHC subregion and for the region as a whole, table 1 summarizes the number of blocks and block characteristics, average size, sequence coverage, and number of markers per block. At an average marker density of one SNP per 1.9 kb, sequence coverage in blocks is >80% across the MHC, with a slight drop in the class II region. The corresponding figures at one SNP per 5 kb (table 1), which are indicative of the first phase of the International HapMap Project, show a net gain of ~15% in coverage at the higher marker density. The corresponding figures from the chromosome 20q study (Ke et al. 2004b) show the same trend and suggest that, because of the extreme variability of LD across the genome, sequence coverage will differ between genomic regions. The average size of blocks in CEPH individuals is 18 kb in both data sets, suggesting that this figure may represent the genome average.

tagSNP Selection

To exploit the dense LD map described above, we proceeded to select a list of maximally informative SNPs for each MHC region that can be used for disease-association studies. We chose to define tagSNPs through a

process that disregards the strength of the underlying LD (low or high) in a given region and hence behaves independent of any block requirements. The caveat in this analysis is that there is an abundance of methods for selecting tagSNPs and that different methods result in different lists of tagSNPs. The results in table 1 show that, at the same marker density (one SNP per 5 kb), the tagging efficiency of the whole MHC region (2.15) is slightly higher than that of chromosome 20 (1.99). There is an overall gain in tagging efficiency (40.9%) when the whole MHC data set is used (one SNP per 1.9 kb), probably as a consequence of the extensive LD in the extended class I region. We observed very good correlation between tagging efficiency and the strength of LD in MHC subregions. Specifically, the extended class I and class III regions appear to approach saturation with only five and three additional tagSNPs necessary to capture the information provided by the extra 113 and 45 assayed SNPs, respectively, in the progression from 1 SNP per 3 kb (70 and 107 tagSNPs, respectively [data not shown]) to full density (table 1). As suggested elsewhere (Ke et al. 2004a), selection of tagSNPs in high-LD regions is largely robust to variability in marker density. Given the high LD underlying the extended class I region, it appears that many features—number of tagSNPs, block size, sequence coverage, and number of blocks—come close to steady state at one SNP per 3 kb (data not shown) where increasing SNP density does not result in further efficiency. A different trend is seen in the class III region, in which sequence coverage is 60% at one SNP per 5 kb, increases to 73% at one SNP per 3 kb, and reaches 81% at full density.

As marker density increases, tagging efficiency also increases in the class I region, but the overall number of

Table 1
LD-Structure Features and tagSNPs in the MHC, at Different Marker Densities

REGION	HAPLOTYPE BLOCK STRUCTURE										SNP TAGGING			
	NO. OF SNPs		No. of Blocks		Block Size (kb)		Sequence Coverage (%)		SNPs per Block		Total No. of tagSNPs		Tagging Efficiency	
	Full	5	Full	5	Full	5	Full	5	Full	5	Full	5	Full	5
Extended class I	408	161	22	18	34	35	90	77	17.9	8.3	75	52	5.44	3.10
Class I	1106	368	99	46	15	18	82	71	10.1	7	356	180	3.11	2.04
Class III	280	142	27	24	23	19	81	60	9.6	5	110	82	2.54	1.73
Class II	466	215	48	35	12	19	73	64	8.8	5.3	204	109	2.28	1.97
MHC ^a	2,260	896	202	116	18	26	82	67	10.7	6.7	745	416	3.03	2.15
Chromosome 20 ^b	3,372 ^c	2,020	367 ^c	278	18 ^c	20	64 ^c	55	7.8 ^c	5.9	1,338 ^c	1013	2.52 ^c	1.99

NOTE.—“Full” density indicates that all genotyped markers were included, whereas 5-kb density means there was, on average, one marker per 5 kb. Haplotype-block analysis was performed using Haploview. tagSNPs were selected using LDSelect (Carlson et al. 2004), with r^2 threshold = .80. Tagging efficiency was defined as n/n_b , where n_b is the number of tagSNPs selected to cover the region, and n is the total number of genotyped SNPs in the region (Ke et al. 2004a).

^a Includes data averaged across the 4.459-kb region covering the complete MHC.

^b Data from chromosome 20 include a 10-Mb region (Ke et al. 2004a)

^c Chromosome 20 data at density of one SNP per 3 kb.

tagSNPs is much higher at maximum density here. This latter trend is depicted in figure 4A (“tag SNPs”) and is particularly evident in regions in which LD is low and sequence coverage needs to be improved. In contrast, sequence coverage at full density (90%) in the extended class I region experienced only a slight increment compared with that at one SNP per 3 kb (89% [data not shown]). Since class II has the shortest LD range and hence the lowest sequence coverage in blocks, the number of tagSNPs increases substantially at maximum marker density (table 1). The relationship between marker density and tagSNPs remain constant, which indicates that LD is very diffuse across the class II region and that more markers will need to be typed. The significantly larger number of blocks (shorter in average) at maximum marker density is consistent with low LD patterns in class II, in which approximately half of the haplotype blocks contain >4 SNPs. Differences in the extended class I and class III regions can be explained by dissimilarities in the underlying LD and LD-block sequence coverage in each region (table 1). The complete list of tagSNPs generated for each MHC region, as well as their distribution, is available at GLOVAR.

Ke et al. (2004a) showed that tagSNPs selected in western Europeans (to explain 100% of haplotype diversity) can explain 96% of haplotype variation in CEPH founders, who are of western and northern European ancestry, and that ~90% of the tagSNPs selected in one sample would also be selected as tagSNPs in the other. Thus, the MHC list of tagSNPs presented here should be useful for studies involving individuals of western and northern European ancestry.

Ancestral Recombination Estimation across the MHC Region

Previous studies have shown that the rate of recombination varies substantially across certain regions of the MHC (Cullen et al. 1997, 2002; Jeffreys and May 2004). Sperm typing within a 200-kb class II fragment (Jeffreys et al. 2001) has determined the location of recombination hot spots and has shown that they correlate very strongly with LD breaks. The LD map described above has the resolution required to accurately map such events as shown in figure 4C. Thus, we employed the recently described method of McVean et al. (2004) to estimate fine-scale recombination-rate variations based on population-genetic data and to predict the location and intensity of hotspots across the whole MHC. Also, we wanted to assess whether the haplotype-block pattern reflects genuine recombination-rate variation and whether block boundaries are actually delimited by recombination hotspots.

The nature of recombination rates inferred from patterns of genetic variation varied immensely across the

region (fig. 5) and showed patterns mirroring the LD patterns described above. The average recombination rate for the 4.459-Mb region is 0.66714 cM/Mb, with the extended class I region showing a very low rate, 0.195 cM/Mb. The classic MHC has a rate of 0.7852 cM/Mb. More specifically, class I and III recombination rates are somewhat similar to each other (0.443 cM/Mb and 0.469 cM/Mb, respectively), whereas class II has the highest rate (1.712 cM/Mb). It is worth highlighting the extreme variation in recombination rates—a difference of up to 4 orders of magnitude. The data suggest the presence of 29 hotspots—and show >10 times the background recombination rates—corresponding to an average density of one per 150 kb (fig. 5). An additional recombination site falling short of the hotspot threshold, sevenfold higher, was observed between *HLA-C* and *HLA-B*. In total, hotspots cover only 1.6% of the genomic sequence and have an average size of 2.43 kb. Slightly more hotspots were observed between genes than within genes ($n = 19$ vs. $n = 11$, respectively).

All but two of the inferred hotspots coincide with the haplotype-block boundaries we calculated using the method of Gabriel et al. (2002). Interestingly, we find recombination hotspots in *TAP2* and *HLA-DMB* and between *BRD2* and *HLA-DOA* at the same locations as reported by sperm-typing analysis (Jeffreys et al. 2001; Cullen et al. 2002), (fig. 5, *inset*). We observed two recombination coldspots (0.013 and 0.009, 10 times lower than the background recombination), both of which overlap with high-LD fragments (fig. 5). In one case, the lack of recombination perfectly corresponds to one large haplotype block (109 kb) lying in the class III/II border, which contains the *C6orf10* gene and is surrounded by two hotspots and a diffuse pattern of a number of short LD blocks.

Extended-Haplotype Analysis

Positive selection brings rare alleles to higher frequency in relatively few generations, thus affording fewer opportunities for recombination events to separate an allele from its original chromosomal context. Therefore, haplotype-block alleles showing a high EHH score and high frequency in the population may have experienced recent positive selection and could represent functionally important alleles. To identify potential signals of positive selection in the MHC, we used EHH analysis, which determines the length of the chromosomal haplotypes extending from a specific allele at a particular locus (Sabeti et al. 2002). To control for the high positional variation in recombination rates in the MHC that would artificially affect the length of haplotypes, we used the sperm-typing recombination map of the region described by Cullen et al. (2002). Given this map, we scanned the MHC, using each haplotype

block as an independent locus from which to determine EHH values, assessing each of the 1,966 alleles examined. Since no similarly large region of the genome has a direct sperm-based estimate of recombination rate, we compared the EHH values of haplotype-block alleles within the MHC data set with each other and identified allelic variants that are outliers, on the basis of statistical rank of the EHH value relative to allele frequency.

We observed eight alleles at 0.3 cM and six alleles at 0.25 cM that deviate by >4.5 SD from the mean of the $EHH \times$ frequency statistic and also from the mean of the 5% frequency bins (table 2 and fig. 6). All 14 of these alleles map to a single extended haplotype. An example of one of these alleles is shown in figure 7. A mutation on this haplotype may have experienced recent directional positive selection. Interestingly, this haplotype bears allele *HLA-DRB1*1501* 94% of the time (in one case, the allele was *HLA-DRB1*0301*). Although *HLA-DRB1*1501* is an exciting candidate, we cannot rule out other possibly functional variation on the haplotype.

Table 2

EHH Outliers at .3cM and .25cM

The table is available in its entirety in the online edition of *The American Journal of Human Genetics*.

Discussion

We genotyped 2,335 loci across the MHC in CEPH pedigrees. The constructed high-resolution map, one marker per 1.9 kb, provides a comprehensive guide to fine-scale patterns of LD and recombination, as well as the means to select optimal marker sets for disease-association studies. To that end, we suggested one such set of tagSNPs, although we realize that many investigators will use their preferred method.

High and long-range LD has long been interpreted as one of the hallmarks of the MHC (The MHC Sequencing Consortium 1999). The present study revealed that the olfactory-receptor gene cluster falls within a 540-kb region of uninterrupted high LD. Ehlers et al. (2000) showed that these polymorphic olfactory loci contribute

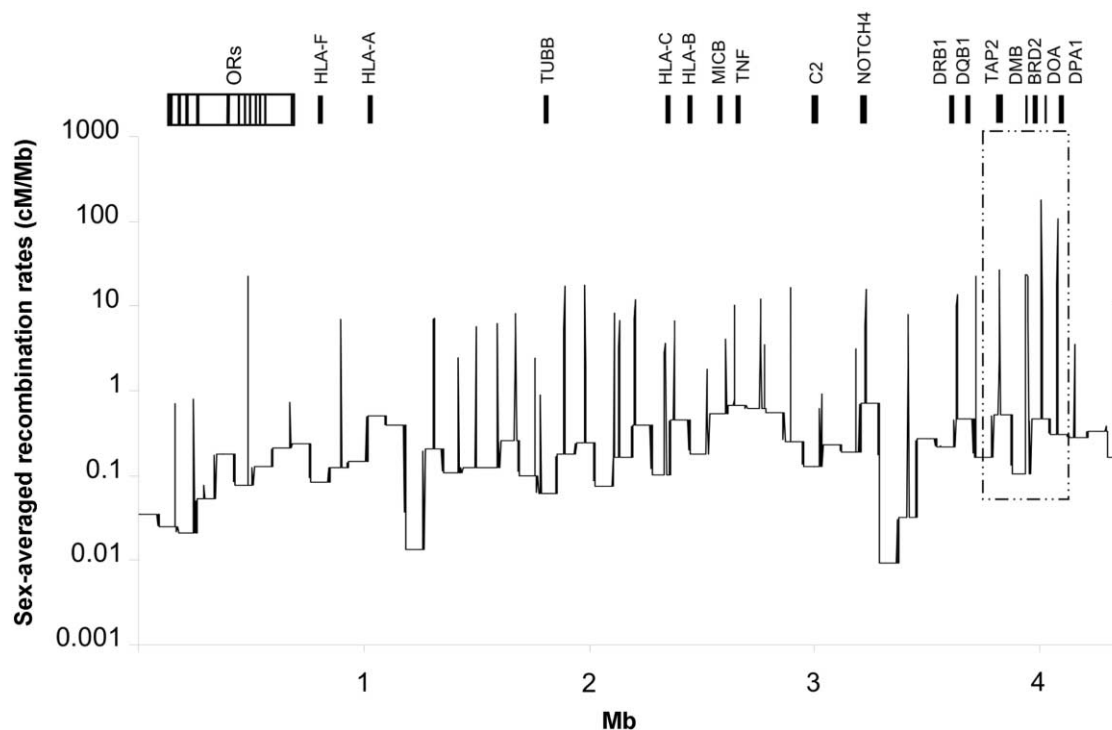


Figure 5 Recombination-rate variation across the MHC. Recombination rates estimated from population-genetic data are far from being uniform; their distribution fluctuates considerably in both, by scale (cM/Mb) and by map position. Recombination hotspots are represented by peaks 10 times higher than the local background level of recombination. Peaks enclosed in the inset correspond to hotspots identified by sperm typing (Jeffreys et al. 2001) located at or near *TAP2*, *HLA-DMB*, *BRD2*, and *HLA-DOA*, observable as LD breaks in figure 4C. The recombination hotspot between *OR12D3* and *OR12D2* in the olfactory-receptor gene cluster (“ORs”), inferred from population genetic data, correlates perfectly with the LD break visible in figure 4B. Note the presence of two coldspots showing recombination rates 10 times lower than the local background level of recombination.

The figure is available in its entirety in the online edition of *The American Journal of Human Genetics*.

Figure 6 EHH outliers at 0.3 cM and 0.25 cM

to extended HLA/OR haplotypes, and a recent study by Füst et al. (2004) indicates that a specific haplotype associated with smoking behavior may even extend to the central region (class III) of the MHC. The HapMap data (International HapMap Project) suggest that this region of strong LD extends farther toward the telomere for 350 kb. It will be interesting to determine the full extent of what appears to be one of the longest regions of high LD in the genome. It also remains to be investigated whether this LD block—which seems to extend over ~900 kb, according to the HapMap data (International HapMap Project)—is also present in other populations.

EHH analysis compares an allele's frequency "age" to its extended haplotype's recombinational "age." Under neutral evolution, new variants require a long time to reach high frequency; during that time, LD adjacent to the new variant will be decreased because of recombination. Consequently, common alleles will be surrounded by short-range LD. High-frequency variants within a long-range LD context may represent variants under positive selection. Our EHH analysis identified a handful of variants in which haplotypes showed comparatively extended homozygosity, given their population frequencies. These variants map to a single long-range haplotype that bears *HLA-DRB1*1501* 94% of the time. This haplotype also bears other variants from the so-called ancestral *DR2* haplotype; however, the correlation is best with the *HLA-DRB1*1501* allele, which suggests that it or another nearby variant may have been responsible for the hypothesized selective event. Variation on the *DR2* haplotype has been associated with susceptibility to multiple sclerosis and systemic lupus erythematosus (Barcellos et al. 2003; Hegarty et al. 2003; Larsen and Alper 2004), whereas it has been associated with a protective effect for type I diabetes (Cucca et al. 2001)

Population-based recombination-rate estimates (coalescent model) provide an average transmission of recombination rates (female-male) over thousands of generations, whereas pedigree analyses are based on only a few meioses and sperm-typing measures of recombination in few extant males. We find that pedigree analysis, sperm typing, and coalescent approaches identify identical recombination hotspots for the 200-kb MHC class II fragment studied by Jeffreys et al. (2001) (figs. 4C and fig. 5, *inset*) This is in agreement with other studies (McVean et al. 2004; Crawford et al. 2004), and

it provides additional weight to the new recombination hotspots identified in the present study. As a consequence of the number of recombination hotspots predicted for the MHC region, we would expect to find a fairly disrupted LD pattern. However, extended haplotypes exist at relatively high frequency in whites. In addition, according to population frequencies, the MHC can be divided into only a few blocks that contain nonrandomly associated alleles at different loci (Yunis et al. 2003). We also observed regions that showed excessively long haplotypes, given their population frequencies. Selective sweeps and population history (genetic drift) can explain the disproportionately long-range LD in some haplotypes, but there is also evidence indicating a strong influence of recombination activity shaping the LD landscape within the MHC. Specifically, a considerable proportion of recombination hotspots can be resolved as gene-conversion events (Jeffreys and May 2004). Additionally, differential intensity in crossover activity and haplotype-specific recombination patterns (Ahmad et al. 2003) could also affect the extent of LD. However, the relationship between haplotype-specific hotspots and population-specific haplotypes observed in a 75-kb fragment of the MHC class II region (Kauppi et al. 2003) remains to be studied.

Our analysis shows that the number of tagSNPs required to explain variation across large genomic regions fluctuates and depends on the extent of LD. High marker density is required in regions of low LD to allow comprehensive coverage with tagSNPs. Extrapolating to the rest of the genome, the International HapMap Project database will provide efficient tagging in regions of extended LD already in phase I (one SNP per 5 kb) but will require the higher density of phase II to achieve comprehensive coverage of the whole genome.

We identified a subset (tagSNPs) of all the typed SNPs with MAF >5% that captures haplotype variation in this sample. The tags are displayed in the context of annotated genomic sequence through GLOVAR. As expected, variation in the underlying LD greatly affects the SNP-tagging process and efficiency in each MHC subregion. The highest SNP tagging efficiency (5.44) could be achieved for the extended class I region, which indicates attainable gains in typing cost for subsequent studies. Higher efficiency can be obtained by using haplotype-based methods (Ke et al. 2004a), although haplotypes and their frequencies must be carefully estimated in regions of complex local LD and recombination profile (e.g., the class II region). The list of tagSNPs generated within the context of LD profiles and distribution of recombination hotspot will allow MHC association studies to be conducted more efficiently. On the basis of data presented here, given a sufficiently large population, it will also be possible to identify rare haplotypes, thus increasing sensitivity for association with

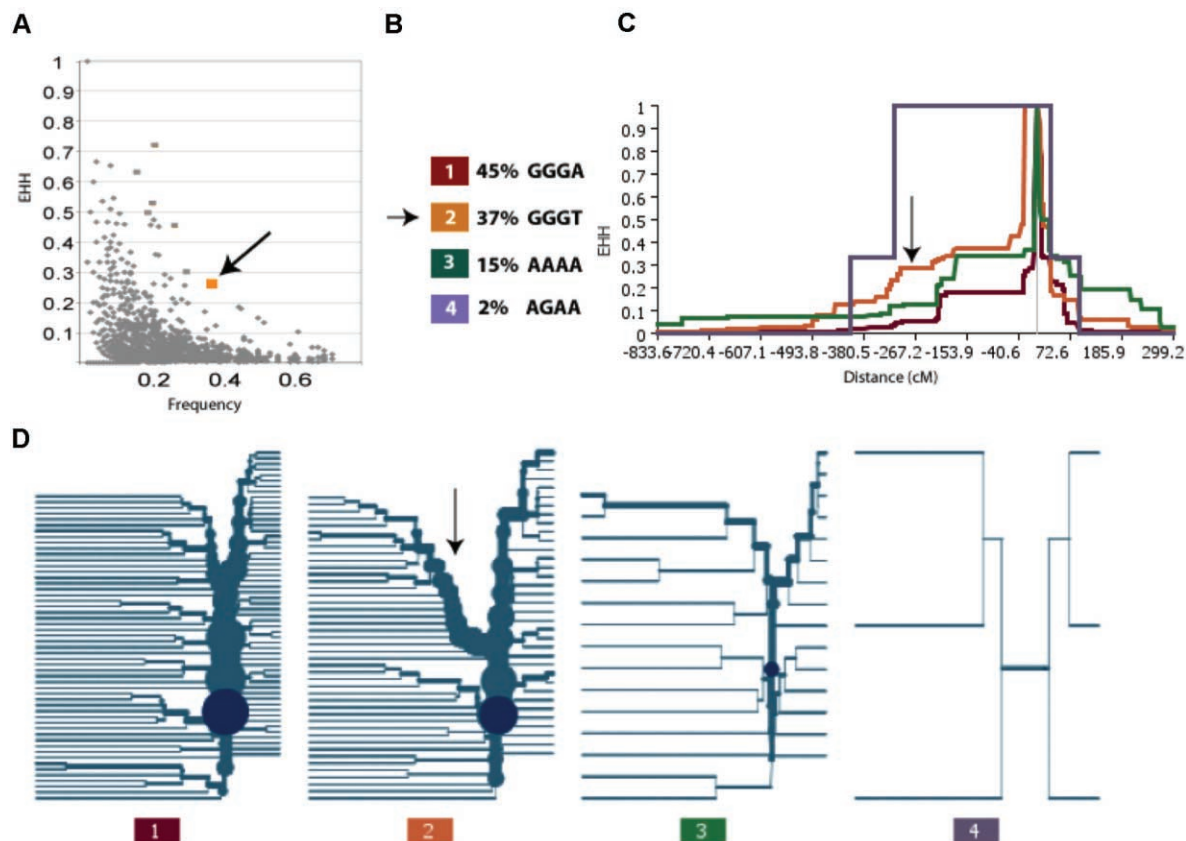


Figure 7 Representative EHH outlier. *A*, EHH by frequency plot at 0.3 cM distance, indicating one of the eight variants at that an outlier with distance that is >4.5 SD in its frequency bin and per its frequency \times EHH statistic. This variant is indicated in figure 6 (*panel 9*) and is part of a block slightly centromeric to the *DQB1* gene. *B*, Haplotype structure of the block containing this haplotype variant. The outlier is the 37% allele (“GGGT”) indicated in orange in all remaining plots (*arrow*). *C*, EHH by distance (cM) plot of all variants in the block. The arrow indicates the region of extended LD for the orange haplotype. *D*, Haplotype bifurcation plots of all variants in the block. The arrow indicates the region of interest for the orange haplotype.

MHC-linked disease. In addition, a reduction in alloreactivity can be envisioned on the basis of the complementary information provided by SNP typing. At least for the extended haplotypes, which represent 25%–30% of all haplotypes in European-derived populations, patient-donor matching can be improved by the identification of haplotype-specific and block-specific variation (Yunis et al. 2003).

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Electronic-Database Information

The URLs for data presented herein are as follows:

Center for Statistical Genetics, <http://www.sph.umich.edu/csg/abecasis/PedStats/> (for PEDSTATS)
 dbSNP Home Page, <http://www.ncbi.nlm.nih.gov/SNP/index.html>
 GLOVAR Genome Browser, http://www.glovar.org/Homo_sapiens/
 Haploview, <http://www.broad.mit.edu/mpg/haploview/index.php>
 Human Chromosome 6 Project Overview, <http://www.sanger.ac.uk/HGP/Chr6/>
 International HapMap Project, <http://www.hapmap.org/>

MHC Haplotype Project, <http://www.sanger.ac.uk/HGP/Chr6/MHC/>

National Center for Biotechnology Information, <http://www.ncbi.nlm.nih.gov/>

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