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Genome-wide association analysis of chronic lymphocytic leukaemia, Hodgkin lymphoma and multiple myeloma identifies pleiotropic risk loci

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B-cell malignancies (BCM) originate from the same cell of origin, but at different maturation stages and have distinct clinical phenotypes. Although genetic risk variants for individual BCMs have been identified, an agnostic, genome-wide search for shared genetic susceptibility has not been performed. We explored genome-wide association studies of chronic lymphocytic leukaemia (CLL, N = 1,842), Hodgkin lymphoma (HL, N = 1,465) and multiple myeloma (MM, N = 3,790). We identified a novel pleiotropic risk locus at 3q22.2 (*NCK1*, rs11715604, $P = 1.60 \times 10^{-9}$) with opposing effects between CLL ($P = 1.97 \times 10^{-8}$) and HL ($P = 3.31 \times 10^{-3}$). Eight established non-HLA risk loci showed pleiotropic associations. Within the HLA region, Ser37 + Phe37 in HLA-DRB1 ($P = 1.84 \times 10^{-12}$) was associated with increased CLL and HL risk ($P = 4.68 \times 10^{-12}$), and reduced MM risk ($P = 1.12 \times 10^{-2}$), and Gly70 in HLA-DQB1 ($P = 3.15 \times 10^{-10}$) showed opposing effects between CLL ($P = 3.52 \times 10^{-3}$) and HL ($P = 3.41 \times 10^{-9}$). By integrating eQTL, Hi-C and ChIP-seq data, we show that the pleiotropic risk loci are enriched for B-cell regulatory elements, as well as an over-representation of binding of key B-cell

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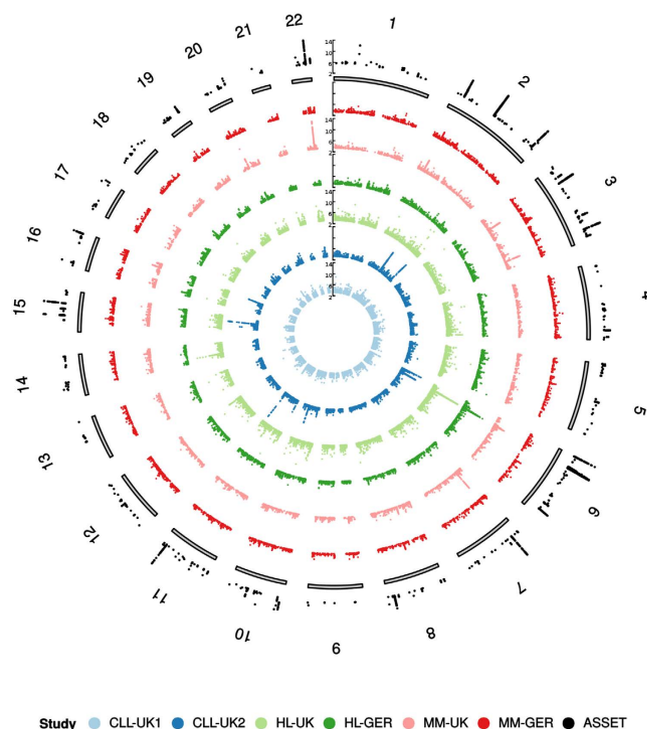


Figure 1. Manhattan plots ($-\log_{10}(P)$) by chromosome. Innermost to outermost ring – chronic lymphocytic leukaemia (CLL)-UK1, CLL-UK2, Hodgkin lymphoma (HL)-UK, HL-GER, multiple myeloma (MM)-UK, MM-GER, and ASSET association test. For clarity, only data with $P < 1 \times 10^{-3}$ are shown.

transcription factors. These data identify shared biological pathways influencing the development of CLL, HL and MM. The identification of these risk loci furthers our understanding of the aetiological basis of BCMs.

Differing in their clinical phenotype, chronic lymphocytic leukaemia (CLL), Hodgkin lymphoma (HL), and multiple myeloma (MM) are all malignancies resulting from the unrestrained clonal expansion of B-cells at different stages of maturation^{1–4}. Evidence for inherited genetic susceptibility to CLL, HL and MM has been provided by studies of familial risk, and more recently from genome-wide association studies (GWAS) which have identified risk SNPs for each tumour type^{5–20}. While the familial risks for CLL, HL and MM are primarily tumour-specific^{21,22}, there is some epidemiological evidence for shared susceptibility^{23–26}. An example of this is provided by the pattern of familial risks associated with B-cell malignancies (BCMs) in Swedish populations²⁵.

Genetic variation at a number of loci, such as 5p15 and 8q24, have been shown to influence the risk of a number of BCM and non-haematological cancers^{27–34}. Identifying risk loci that can have such pleiotropic effects is important for gaining insight into shared and divergent molecular basis of different tumour types.

While conventional meta-analysis provides a powerful tool for combining distinct GWAS, this approach is suboptimal in the presence of disease heterogeneity, such as when SNP associations are only manifest in a specific subset of the diseases, or have opposing effects for different diseases. To address such shortcomings in searching for pleiotropic risk SNPs for BCM, we adopted the previously validated association analysis based on subsets (ASSET) meta-analytic approach^{35,36}. ASSET implements an agnostic analysis exploring all possible subsets of studies to identify the strongest association signal, while accounting for the multiple tests required by the subset search, as well as any shared controls between studies. In doing so, ASSET is able to identify variants that are positively and negatively associated with different diseases.

Applying this statistical procedure to six BCM GWAS (two each of CLL, HL and MM) we report the identification of a novel pleiotropic region influencing BCM risk, as well as eight non-HLA linked pleiotropic loci that have only previously been described in single GWAS. Within the HLA region, we report two novel coding variants in class II HLA proteins which have pleiotropic effects on BCM risk.

Results

Characteristics of the six GWAS are summarised in Supplementary Table 1. After applying quality control filters and imputation of GWAS data (see methods) we analysed over 10 million variants for pleiotropic associations in 7,097 BCM cases and 7,324 controls of European ancestry. Figure 1 shows a Manhattan plot of the association test results for CLL, HL and MM.

To determine whether the global pleiotropic regions of association for pairs of BCMs occurred more often than expected by chance, we generated stratified quantile-quantile (Q-Q) plots to assess enrichment of

associations for a given tumour type conditioned on the P -value for another tumour (Supplementary Fig. 1). The greater departure from the expected line associated with smaller P -values observed in the Q-Q plots provides evidence of pleiotropic effects between CLL, HL and MM³⁷.

To identify the specific regions across the genome that demonstrate pleiotropic effects on risk of BCM we used ASSET^{35,36}. In order for candidate SNPs to be considered, they were required to meet the following criteria: (1) variant associations at $P \leq 5.0 \times 10^{-8}$ for the ASSET test; (2) at least one other variant in the same region (within $r^2 > 0.2$) with the same pleiotropic association at $P \leq 1.0 \times 10^{-6}$; (3) the individual one-sided ASSET subset tests were significant at $P < 0.01$; (4) the variant is not driven by a single study; (5) the variant cannot be both positively and negatively associated in different datasets of the same BCM; and (6) if a variant is positively and negatively associated with different BCM, the 2-sided P -value must be lower than both individual 1-sided P -values. Using these criteria, we identified nine non-HLA regions (607 variants).

Newly identified pleiotropic risk loci. We identified a novel pleiotropic association at 3q22.2 (rs11715604, $P = 1.60 \times 10^{-9}$, Fig. 2) with opposing associations in CLL ($P_{1\text{-tailed}} = 1.97 \times 10^{-8}$) and HL ($P_{1\text{-tailed}} = 3.31 \times 10^{-3}$). rs11715604 maps to intron 1 of *NCK1*, which is integral to T-cell activation^{38,39} and regulates the PI3K/Akt pathway⁴⁰. We also identified a number of promising associations that did not reach genome-wide significance, but exhibited moderate effects in the different BCM (Supplementary Table 2). These included associations at 22q13.33 (rs131821, $P = 7.49 \times 10^{-8}$) and 18p11.31 (rs634212, $P = 5.11 \times 10^{-5}$). rs131821 is intronic of *NCAPH2*, which is important in mitotic chromosome architecture⁴¹, while rs634212 is intronic of *L3MBTL4*, which has been implicated as a tumour suppressor gene for breast cancer⁴².

Previously known risk loci with newly identified pleiotropic effects. We identified genome-wide significant pleiotropic associations (*i.e.* $P \leq 5.0 \times 10^{-8}$) at eight non-HLA linked loci previously identified as risk factors for CLL, HL or MM (Table 1). The CLL risk loci at 6p21.32 (*BAK1*) and 6p25.3 (*IRF4*) were positively associated with HL risk^{11,17}. In contrast the 2q13 (*BCL2L11*) and 11q24.1 (*GRAMD1B*) risk loci for CLL negatively influenced MM risk^{5,11}. The MM risk locus at 3p22.1 (*ULK4*) positively influenced HL⁷, whereas the 2p23.3 (*DTNB*) risk locus for MM negatively associated with CLL⁷. The HL risk locus at 3p24.1 (*EOMES*) was positively associated with CLL¹⁴. The 3q26.2 (*TERC*) a risk factor for MM and CLL, showed a positive association with HL risk^{10,16}. In addition to variation at these eight regions we observed promising pleiotropic associations at 2q37.1 (*SP110*, rs150468793; rs149207840)⁹, 3q27 (*LPP*, rs4459895)⁶, 5q15 (*ELL2*, rs2546191)¹⁸, 8q24.21 (*PVTL1*, rs2720680)¹², 15q15.1 (*BMF*, rs35603048)⁵, and 16q24.2 (*IRF8*, rs4240807)⁹ (Table 1).

Association studies of CLL, HL and MM have demonstrated seemingly different associations between loci within the HLA region and risk^{10,43,44}. To implement an ASSET analysis of the HLA region, we imputed classical alleles, coding variants of HLA proteins, and SNPs using the SNP2HLA software in conjunction with the Type 1 Diabetes Genetics Consortium (T1DGC) HLA reference panel⁴⁵. Figure 3 shows the unconditioned ASSET associations across the 3.7 Mb HLA region. 768 variants demonstrated an association for CLL, HL and MM at $P \leq 5.0 \times 10^{-8}$. To isolate independent pleiotropic associations we performed conditional stepwise logistic regression conditioning on the strongest associated variant from the 2-sided ASSET analysis. We identified Ser37 + Phe37 in HLA-DRB1 ($P_{\text{conditional}} = 1.84 \times 10^{-12}$), positively associated with CLL and HL ($P_{1\text{-tailed-conditional}} = 4.68 \times 10^{-12}$) and negatively associated with MM ($P_{1\text{-tailed-conditional}} = 1.2 \times 10^{-2}$). In addition, Gly70 HLA-DQB1 ($P_{\text{conditional}} = 3.15 \times 10^{-10}$) was positively associated with CLL ($P_{1\text{-tailed-conditional}} = 3.52 \times 10^{-3}$) but negatively associated with HL ($P_{1\text{-tailed-conditional}} = 3.41 \times 10^{-9}$). Additionally, a promising association for Arg62 + Glu62 in HLA-A ($P_{\text{conditional}} = 9.26 \times 10^{-8}$) was found, and was positively associated with CLL ($P_{1\text{-tailed-conditional}} = 8.06 \times 10^{-5}$) but negatively associated with HL ($P_{1\text{-tailed-conditional}} = 5.68 \times 10^{-5}$).

Biological inference of pleiotropic risk loci. To explore whether the identified SNPs are eQTLs, we searched the Blood eQTL browser⁴⁶, and MuTHER⁴⁷ and Geuvadis/1000 Genomes⁴⁸ lymphoblastoid cell line (LCL) datasets. In addition we examined expression data from MM plasma cells⁴⁹. We found evidence for eQTLs (FDR adjusted $P < 0.05$) for nine of the pleiotropic loci in the LCL data, and four loci in the plasma cell data (Supplementary Table 3).

Since spatial proximity between specific genomic regions and chromatin looping interactions are central for regulation of gene expression⁵⁰, we identified patterns of chromatin interactions at candidate pleiotropic SNPs by analysing Hi-C data on GM12878, as a source of B-cell information (Fig. 2 and Supplementary Fig. 2). Looping chromatin interactions were shown at 3q22 (rs11715604), 3p24 (rs9880772), 3q26 (rs12638862), 6p21 (rs210143) and 11q24 (rs4525246). The looping interactions at 3q22, implicates IL-20RB which regulates antigen-specific T-cell responses⁵¹. Furthermore, at 3p24, we observed looping interactions with *AZ12*, which contributes to the activation of NF- κ B⁵².

Across the BCM pleiotropic risk loci, we confirmed enrichment of regulatory elements in primary haematopoietic stem cells ($P = 2.1 \times 10^{-3}$) and GM12878 cells ($P = 7.4 \times 10^{-3}$, Supplementary Table 4)⁵³. Analysis of ChIP-seq data on 82 transcription factors (TFs) showed an enrichment of binding of key B-cell transcription factors, including CEBPB, RXRA, and POLR3G ($P < 0.05/82 = 6.10 \times 10^{-4}$) (Supplementary Fig. 3). CEBPB is a TF that is involved in immune and inflammatory responses⁵⁴, and can induce reprogramming of B-cells into pluripotent stem cells⁵⁵. RXRA can induce B-cell differentiation⁵⁶, and POLR3G is a DNA-dependent RNA polymerase III⁵⁷.

Pathway analysis and construction of a susceptibility network. We performed a gene-set enrichment analysis to gain insight into the biological pathways perturbed by genetic variation common to CLL, HL and MM, and found eight pathways related to the inflammatory response and antigen processing that showed enrichment (*i.e.* FDR adjusted P -value < 0.05 ; Supplementary Table 5). Following on from this analysis, we

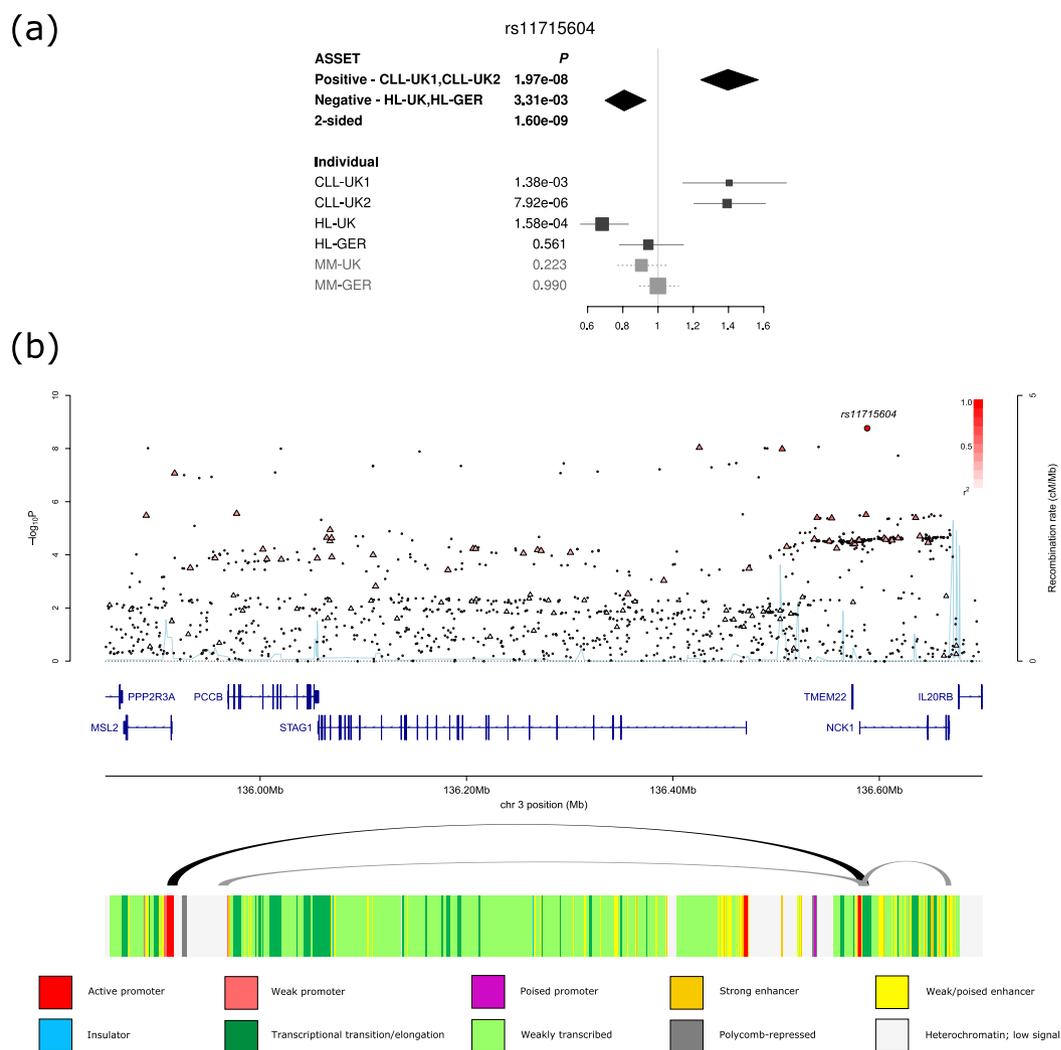


Figure 2. (a) Forest plot of the ORs for the association between rs11715604 and BCM. Studies were weighted according to the inverse of the variance of the log of the OR calculated. Horizontal lines: 95% CI. Box: OR point estimate; box area is proportional to the weight of the study. Diamond: overall summary estimate, with CI given by its width. Unbroken vertical line: null value (OR = 1.0). (b) Regional plot of association and recombination rates. $-\log_{10}(P)$ (y axis) of the SNPs are shown according to their chromosomal positions (x axis). The sentinel SNP is shown as a large circle. The colour intensity of each symbol reflects the extent of LD with the sentinel SNP: white ($r^2 = 0$) through to dark red ($r^2 = 1.0$). Genetic recombination rates, estimated from the 1000 Genomes Project, are shown with a light blue line. Physical positions are based on NCBI build 37 of the human genome. Also shown are the relative positions of genes and transcripts mapping to the region of association. The arcs represent Hi-C promoter contacts in GM12878 cells. The colour intensity of each contact reflects the interaction score. The bottom track represents the chromatin-state segmentation track (ChromHMM) for lymphoblastoid cells using data from the HapMap ENCODE Project.

investigated the inter-connectivity of the associated genes⁵⁸. By constructing a network of published and predicted protein-protein interactions, protein co-localisations and protein domain similarity, we delineated two broad clusters – one related to BCL2, and the other related to HLA (Supplementary Fig. 4).

Discussion

Motivated by the stratified Q-Q plots, which suggested the existence of pleiotropy, we utilised genotype data from six datasets in British and German populations, to conduct an agnostic cross-cancer genome-wide analysis to identify specific pleiotropic associations for CLL, HL and MM for both HLA and non-HLA regions. We identified a number of promising associations that have strong biological plausibility, including *NCK1*, *NCAPH2* and *L3MBTL4*. A contemporaneous analysis also used ASSET across a number of different non-BCM cancers, and discovered a novel risk locus at 1q22 involving breast and lung cancer⁵⁹.

Our analysis also provides evidence for common and opposing effects being responsible for BCM pathogenesis, but is not the first to identify opposing risk associations in different cancers^{60,61}. Given that many of the

Locus	SNP ID	Position (bp)	Allele 1	Allele 2	ASSET 2-sided P-value	Disease Group 1			Disease Group 2		
						BCM	Odds Ratio (CI)	P-value	BCM	Odds Ratio (CI)	P-value
2p23.3	rs6546149	25629438	C	G	6.27×10^{-10}	CLL	1.09 (1.01–1.17)	2.14×10^{-2}	MM	0.83 (0.78–0.88)	1.15×10^{-9}
2q13	rs12711846	111856293	A	G	3.48×10^{-14}	CLL	1.44 (1.31–1.58)	6.37×10^{-14}	MM	0.92 (0.86–0.98)	1.53×10^{-2}
3p24.1	rs9880772	27777779	G	A	7.42×10^{-9}	CLL, HL	1.18 (1.11–1.24)	7.42×10^{-9}			
3p22.1	rs6763508	41750989	T	C	7.56×10^{-12}	MM, HL	1.22 (1.16–1.30)	7.56×10^{-12}			
3q26.2	rs12638862	169477506	G	A	1.88×10^{-11}	CLL, MM, HL	1.15 (1.09–1.19)	1.88×10^{-11}			
6p25.3	rs9392017	442357	A	G	6.03×10^{-9}	CLL, HL	1.22 (1.15–1.30)	6.03×10^{-9}			
6p21.32	rs210143	33546837	T	C	6.81×10^{-12}	CLL, HL	1.24 (1.17–1.32)	6.81×10^{-12}			
11q24.1	rs4525246	123395246	G	C	3.37×10^{-14}	CLL	1.40 (1.28–1.53)	6.33×10^{-14}	MM	0.93 (0.87–0.99)	1.50×10^{-2}
2q37.1	rs150468793; rs149207840	231144578	T	TCCTCCTG	9.63×10^{-8}	CLL, MM	1.16 (1.10–1.22)	9.63×10^{-8}			
3q27.3	rs4459895	187954414	A	C	1.70×10^{-7}	CLL	1.12 (1.02–1.23)	1.35×10^{-2}	HL	0.76 (0.68–0.85)	6.43×10^{-7}
5q15	rs2546191	95232541	G	A	4.15×10^{-7}	HL	1.18 (1.07–1.29)	5.38×10^{-4}	MM	0.88 (0.82–0.93)	4.14×10^{-5}
8q24.21	rs2720680	129115217	A	G	6.78×10^{-8}	HL	1.27 (1.16–1.39)	1.13×10^{-7}	CLL	0.92 (0.86–0.99)	2.9×10^{-2}
15q15.1	rs35603048	40391965	C	T	3.64×10^{-7}	HL	1.17 (1.06–1.28)	1.23×10^{-3}	CLL	0.81 (0.74–0.89)	1.58×10^{-5}
16q24.2	rs4240807	85985361	A	C	6.62×10^{-7}	CLL	1.21 (1.11–1.32)	1.35×10^{-5}	HL	0.86 (0.79–0.95)	2.70×10^{-3}

Table 1. Novel pleiotropic associations in genomic regions already identified through single disease genome-wide association studies. Genome-wide significant associations are shown above, and promising associations are below. B-cell malignancies (BCM) in bold indicate the disease the SNP was previously shown to be associated. Odds ratio calculated from allele 2.

identified risk loci harbour genes integral for immune function, it is entirely conceivable that balancing selection may act to ensure immune diversity and thus a selective advantage against temporal environmental risk factors such as infection⁶².

As with standard GWAS analyses ASSET may not identify the causative genetic variant at a locus. Accepting this caveat, many of the identified regions map to eQTL and regulatory elements in B-cells. Moreover, they feature an over-representation of key B-cell TF binding.

The HLA class II region has previously been implicated in multiple BCM including follicular lymphoma^{33,63}, HL¹² and CLL^{17,44}. Here, we additionally show the involvement of this region in the development of MM. By performing a more refined imputation analysis on the HLA region, we found a variant that alters amino acid 37 of HLA-DRB1. This change affects the electrostatic properties of the P9 binding pocket⁶⁴, altering T-cell receptor recognition⁶⁵. The second pleiotropic association at HLA region at amino acid 70 of HLA-DQB1 is located in the P4 binding pocket, which is also a critical residue influencing antigen T-cell receptor binding⁶⁶. A previous study of a number of different B-cell lymphomas using over 7,000 cases also found an association in the HLA region⁶⁷, further highlighting the importance of this region to the development of BCM. In addition to the HLA association, we identified other associations that were independently ascertained in the BCM specific GWAS, including 3p24.1 (*EOMES*) for HL¹⁴ and CLL⁶, and 3q26.2 (*TERC*) for MM¹⁰ and CLL¹⁶, thus adding confidence that ASSET method is able to identify common genetic components.

Although predicated on protein-protein interactions, our pathway analysis provides two core cellular functions influencing BCM susceptibility. Firstly, antigen presentation/T-cell regulation, centred around HLA, and secondly cellular growth and apoptosis, centred around *BCL2*, which are interconnected through the key B-cell regulators, MYB and *GATA3*^{68,69}. *BCL2* is commonly overexpressed in BCM and is relevant to tumour escape apoptosis^{70–72}. It is noteworthy that Venetoclax, a *BCL2* inhibitor used in treatment of CLL⁷³, may also be efficacious in treating other forms of BCM⁷⁴. This exemplifies that targeting pathways identified through GWAS may inform drug discovery initiatives⁷⁵.

In conclusion, using data from six GWAS we have identified associations with multiple BCM. There are likely additional loci that have an effect, but their detection will require additional efforts with larger datasets. Such future analyses should also address the disparity in sample sizes of each of the BCM series that characterises our study.

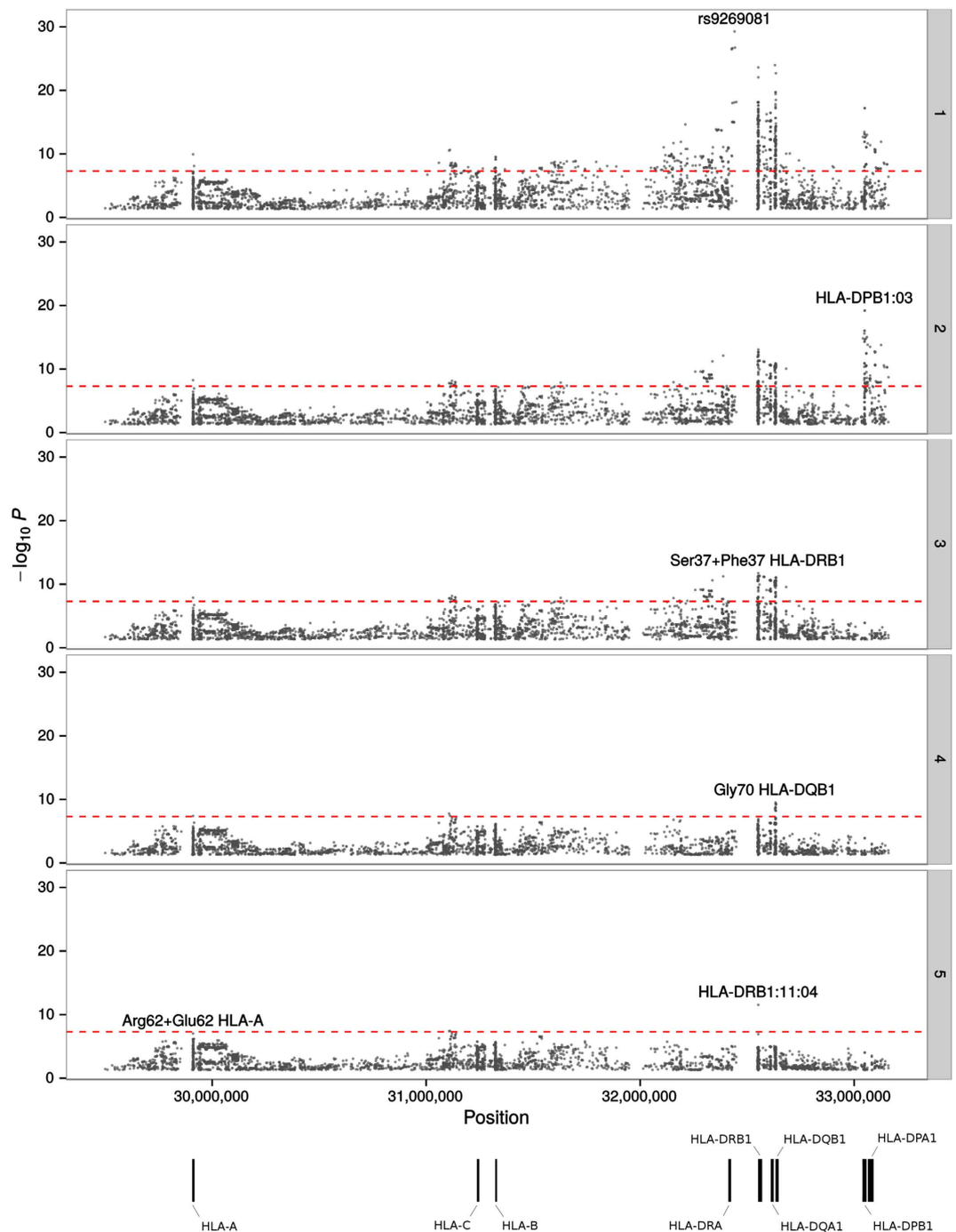


Figure 3. Manhattan plot representation of the step-wise conditional logistic regression of risk of BCM in the HLA region. (1) Unconditioned test of the HLA region. (2) Results of the HLA region after conditioning on rs9269081. (3) Results of the HLA region after conditioning on rs9269081 and HLA-DPB1:03. (4) Results of the HLA region after conditioning on rs9269081, HLA-DPB1:03 and Ser37 + Phe37 HLA-DRB1. (5) Results of the HLA region after conditioning on rs9269081, HLA-DPB1:03, Ser37 + Phe37 HLA-DRB1 and Gly70 HLA-DQB1. The $-\log_{10}(P)$ of the combined logistic regression test P -values are plotted against their physical chromosomal position. The broken red line represents the genome-wide level of significance ($P < 5 \times 10^{-8}$).

Methods

Subjects and GWAS datasets. We used data generated from GWAS of CLL, HL, and MM performed in European populations which have been the subject of previous publications^{10–12,14,16}. Briefly, the MM-UK GWAS comprised 2,282 cases (1,060 male; mean age at diagnosis: 64 years) recruited through the UK Medical Research Council (MRC) Myeloma-IX and Myeloma-XI trials. The MM-GER GWAS comprised 1,508 cases (867 male; mean age at diagnosis: 59 years) recruited by the German Multiple Myeloma Study Group (GMMMG) coordinated by the University Clinic, Heidelberg. The HL-UK GWAS comprised 622 cases ascertained through: (i)

the Royal Marsden Hospital National Health Service Trust Family History study during 2004–2008 ($n = 104$, 63 male; mean age at diagnosis: 38 years); and (ii) an ongoing national study of HL in females ($n = 518$, mean age at diagnosis: 23 years) conducted by the Institute of Cancer Research (ICR). The HL-GER GWAS comprised 1,001 HL cases (597 male; mean age at diagnosis: 35 years) ascertained by the German Hodgkin Study Group during 1998–2007. The CLL-UK1 GWAS comprised 517 cases: (i) 155 cases (95 male; mean age at diagnosis: 59 years) from ICLLC; and (ii) 362 cases (269 male; mean age at diagnosis: 63 years) from the Leukaemia Research CLL-4 trial. CLL-UK2 comprised 1,403 cases collected from two ongoing initiatives: (i) 1,111 cases collected through a UK national study of CLL genetics coordinated by the ICR (712 male; mean age at diagnosis: 63 years); and (ii) 292 cases collected through the Newcastle CLL Consortium (181 male; mean age at diagnosis: 66 years) from patients attending six haematology units in the UK.

Collection of blood samples and clinical information from subjects was undertaken with informed written consent and relevant ethical review board approval at respective institutions, in accordance with the tenets of the Declaration of Helsinki. Specifically, approval for the CLL data was approved by the UK Multi-Research Ethics Committee (MREC 99/1/082). For the MM data, the Myeloma-IX trial was approved by the Medical Research Council Leukaemia Data Monitoring and Ethics committee (MREC 02/8/95, ISRCTN68454111), the Myeloma-XI trial by the Oxfordshire Research Ethics Committee (MREC 17/09/09, ISRCTN49407852), and the GMMG study was approved by the University of Heidelberg Ethical Commission (229/2003, S-337/2009, AFmu-119/2010). For the HL data, approval was obtained from the Multi-Research Ethics Committee (MREC 03/1/096) for the UK data, and the Ethics Committee of the University of Cologne for the German data. All methods and experimental protocols were performed in accordance with relevant guidelines and regulations.

Genotyping of cases was performed using Illumina arrays: CLL-UK1 on 317K array, HL-UK on 660w-Quad BeadChip, and CLL-UK2, HL-GER and all MM samples using Omni-express arrays (Illumina, San Diego, CA, US). For the UK controls, we used publicly accessible data generated by the Wellcome Trust Case Control Consortium (WTCCC), the 1958 Birth Cohort (also known as the National Child Development Study) and UK Blood Service⁷⁶. Genotyping of both sets of controls was conducted using Illumina Human 1.2M-Duo Custom_v1 Array BeadChips. For the German studies we utilised controls from the Heinz Nixdorf Recall study genotyped using Illumina OmniExpress array⁷⁷.

Full details of the genotyping of cases and quality control can be found in previously published work^{10,14,16}. Briefly, general genotyping quality control assessment was as previously described⁷⁸ and all SNPs presented in this study passed the required thresholds. Duplicate samples were used to check genotyping quality. SNPs and samples with <95% SNPs genotyped were eliminated from the analyses. Genotype frequencies at each SNP were tested for deviation from the Hardy–Weinberg equilibrium and rejected at $P < 10^{-5}$. The number of samples and variants that passed quality control is provided in Supplementary Table 1.

We have previously confirmed an absence of systematic genetic differences between cases and controls^{10,14,16}. Prediction of the untyped SNPs was carried out using IMPUTEv2 based on a merged reference panel from UK10K (April 2014 release) and from the 1000 Genomes Project (phase 1 v3)^{79,80}. Association meta-analyses only included markers with info scores > 0.4, imputed call rates/SNP > 0.9 and MAFs > 0.005.

Statistical analysis. The association between variants with cancer risk in each of the six GWAS was evaluated by logistic regression under a log additive model using SNPTEST v2⁷⁹. In the MM-GER study, genomic inflation due to population stratification was detected ($\lambda > 1.1$), so the per-allele odds ratios (ORs) were adjusted using principal components obtained from smartPCA⁸¹.

To investigate pleiotropy globally, we generated stratified Q-Q plots of association signals in one cancer stratified by the P -values in a second cancer³⁷, for every combination of BCM. Leftward inflation in the null line is indicative of a higher degree of pleiotropy between the two tumours than expected by chance.

Subset meta-analysis was conducted using the R statistical package ASSET (association analysis based on subsets) which explores all possible subsets of “non-null” studies to identify the strongest association signal and then evaluates the significance of the signal while accounting for multiple tests required by the subset search^{35,36}. One-tailed tests are subsequently combined to produce a 2-sided test statistic. Although ASSET has the advantage of accounting for subsets of studies with no effects and/or effects in opposing directions, where a large majority of effects are in one direction it will have lower power compared to the conventional fixed-effect analysis. The number of overlapping subjects in the GWAS (*i.e.* controls from WTCCC⁷⁶ and Heinz-Nixdorf⁷⁷ controls) were used as a covariate when estimating standard errors³⁵. Imputed SNPs that showed significant associations were genotyped using standardised Sanger sequencing methods to confirm the imputation fidelity.

HLA imputation and analysis. To determine whether specific coding variants within HLA genes contributed to the diverse association signals, we imputed the classical HLA alleles (A, B, C, DQA1, DQB1, DRB1) and coding variants across the HLA region using SNP2HLA⁴⁵. The imputation was based on a reference panel from the Type 1 Diabetes Genetics Consortium (T1DGC) consisting of genotype data from 5,225 individuals of European descent with genotyping data of 8,961 common SNPs and indel polymorphisms across the HLA region, and four digit genotyping data of the HLA class I and II molecules. This reference panel has been used previously and showed high imputation quality for the HLA region in other studies^{45,82,83}.

To identify independent effects, dependency analyses by step-wise logistic regression were carried out by conditioning on the strongest association signal in the specific BCM. The index SNP at each region was included as a covariate, and the association statistics were recalculated for the remaining test SNPs. This process was repeated until no SNPs reached the minimum level of significance. The criteria for declaring an independent effect were defined as $P < 5 \times 10^{-8}$.

Functional prediction. LD between SNPs were calculated with VCFtools⁸⁴ using data from the UK10K (April 2014 release) and the 1000 Genomes Project (phase 1 v3)^{79,80}. These data were plotted using visPIG⁸⁵.

To explore the epigenetic profile of genomic location associated with BCM, we used ENCODE histone modification data and HaploReg and RegulomeDB^{86,87} to examine whether any of the SNPs or their proxies (*i.e.* $r^2 > 0.8$ in the 1000 Genomes EUR reference panel) annotate transcription factor binding or enhancer elements.

To examine enrichment in specific TF binding across risk loci we adapted the variant set enrichment method of Cowper-Salari *et al.*⁸⁸ Briefly, for each risk locus, a region of strong LD (defined as $r^2 > 0.8$ and $D' > 0.8$) was determined, and these SNPs were termed the associated variant set (AVS). TF ChIP-seq uniform peak data was obtained from ENCODE for the GM12878 cell line, and included data for 82 TF. For each of these marks the overlap of the SNP in the AVS and the binding sites was determined to produce a mapping tally. SNPs with the same LD structure as the risk associated SNP were randomly selected to calculate a null mapping tally. A null distribution was produced by repeating this process 10,000 times, and approximate *P*-values were calculated as the proportion of permutations where the null mapping tally was greater or equal to the AVS mapping tally. An enrichment score was calculated by normalising the tallies to the median of the null distribution. Thus the enrichment score is the number of standard deviations of the AVS mapping tally from the mean of the null distribution tallies.

eQTL analysis. The presence of potential eQTL was investigated through the use of several public data sets, namely the Blood eQTL browser⁴⁶ in whole blood, and MuTHER⁴⁷ and Geuvadis/1000 Genomes⁴⁸ in lymphoblastoid cell lines (LCL).

For myeloma plasma cell eQTL analysis, we included a German ($n = 658$) and a UK ($n = 183$) case series which had been the subject of a previous eQTL analysis⁴⁹ and 608 cases of a recently published US GWAS¹³. Gene expression profiling of CD138-purified plasma cells using Affymetrix U133 2.0 plus arrays was performed as described^{89–91}. Pre-processing of expression data was done as previously published⁴⁹. Briefly, we used the Affymetrix U133 2.0 plus array custom (CDF) (v17) mapping to Entrez genes⁹² as chip definition file and excluded microarray probes binding to polymorphic sites. Expression data were normalized using GC-RMA. We only included genes with \log_2 expression > 3.5 in at least 95% of samples of each set. After quality control and excluding autosomal genes, expression data for 8,505 genes was available. The filtered set was analysed using probabilistic estimation of expression residuals (PEER)⁹³ to infer known and hidden intervening variables, such as cytogenetic subgroups.

For the Geuvadis and MM plasma cell data, the relationship between SNPs and expression of genes located within 1 Mb was analysed using the Matrix eQTL⁹⁴ package under a linear model. In all the datasets, SNPs in LD ($r^2 > 0.8$) with the potential pleiotropic associations were explored, and were included where FDR adjusted $P < 0.05$.

Network analysis. Pathway enrichment analysis was performed using the Improved Gene Set Enrichment Analysis for Genome-wide Association Study (i-GSEA4GWAS v2)⁹⁵. This tool also performed a functional annotation analysis on these pathways by identifying the top SNPs that map to the pathway genes, and determining if any of these SNPs fall within ENCODE peak data, namely DNase-seq peaks of open chromatin, FAIRE peaks of open chromatin, TFBS SPP-based peaks, TFBS PeakSeq-based peaks and Histone peaks. In addition, eQTLs were determined using several eQTL databases, namely eQTL Browser, GTEX and seeQTL. Common networks were identified using GeneMANIA⁵⁸. This database collated data on protein and genetic interactions from a number of sources, including BioGRID, InterPro, Reactome, and Ensembl.

Promoter capture Hi-C data. To map risk SNP to interaction involving promoter contacts and identify genes involved in HL susceptibility, we analysed previously published promoter capture Hi-C data on the GM12878 cell line as a model B-cell⁹⁶. The promoter capture Hi-C interactions were used to functionally annotate GWAS SNPs to seek for evidences of looping between the SNPs and the promoters of nearby genes. Reads from technical replicates were combined before processing and valid pairs were identified using HICUP⁹⁷. Two biological replicates were analysed to assure reproducibility and significant interactions were determined using CHiCAGO⁹⁸.

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Author Contributions

P.J.L., A.S., J.S.M., M.H. and R.S.H. designed the study. R.S.H. and G.M. obtained financial support in the UK, K.H. and H.G. obtained support in Germany. P.J.L., A.S., J.S.M. and R.S.H. drafted the manuscript. P.J.L., A.S., J.S.M. and M.H. performed the principal statistical and bioinformatic analysis. G.O. and O.L. performed further bioinformatics analyses. P.B. coordinated UK laboratory analysis and H.E.S. performed sample processing. A.S. performed Sanger sequencing. G.J.M. performed ascertainment and collection of case study samples for MM-UK GWAS. D.C.J. managed and prepared Myeloma IX and XI case study DNA samples. M.K. oversaw MM-UK GWAS data processing. H.G., N.W., B.C., M.I.d.S.F., H.T., K.H. and A.F. coordinated and managed the German DNA samples and data. P.H. and M.M.N. performed and coordinated the GWAS of German cases and controls. K.-H.J. and L.E. ascertained and managed the HNR sample. N.W. performed eQTL analysis on plasma cells. A.J.S. provided samples for HL-UK GWAS, R.C. provided data on samples for HL-UK GWAS. P.H. and M.M.N. were responsible for GER-HL GWAS analysis; E.P.v.S. and A.E. were responsible for German HL patient samples. D.C. performed recruitment of CLL samples. For Newcastle, J.M.A. and D.J.A. conceived of the Newcastle CLL Consortium; J.M.A. obtained financial support, supervised laboratory management, genotyping of cases and data handling; N.J.S. and T.R. performed sample management of cases and data handling; and G.H.J., G.S., R.J.H., A.R.P., D.J.A., J.C., J.R.B., G.P., C.P. and C.F. developed protocols for recruitment of individuals with CLL and sample acquisition and performed sample collection of cases. All authors contributed to the final paper.

Additional Information

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