Insight into genetic predisposition to chronic lymphocytic leukemia from integrative epigenomics

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Genome-wide association studies have provided evidence for inherited genetic predisposition to chronic lymphocytic leukemia (CLL). To gain insight into the mechanisms underlying CLL risk we analyze chromatin accessibility, active regulatory elements marked by H3K27ac, and DNA methylation at 42 risk loci in up to 486 primary CLLs. We identify that risk loci are significantly enriched for active chromatin in CLL with evidence of being CLL-specific or differentially regulated in normal B-cell development. We then use in situ promoter capture Hi-C, in conjunction with gene expression data to reveal likely target genes of the risk loci. Candidate target genes are enriched for pathways related to B-cell development such as MYC and BCL2 signalling. At 14 loci the analysis highlights 63 variants as the probable functional basis of CLL risk. By integrating genetic and epigenetic information our analysis reveals novel insights into the relationship between inherited predisposition and the regulatory chromatin landscape of CLL.
Chronic lymphocytic leukemia (CLL) is an indolent B-cell malignancy that has a strong genetic component, as evidenced by the eight-fold increased risk in relatives of CLL patients. Our understanding of CLL predisposition has been transformed by genome-wide association studies (GWAS), which have identified alleles at 43 loci influencing risk. Elucidating the function of these risk loci is an important step toward the development of testable hypotheses regarding the biological processes involved in CLL predisposition and pathogenesis. Most GWAS signals, however, map to the non-coding genome and elucidating the mechanisms through which these non-coding variants exert their effect has proven challenging. Integration of chromatin immunoprecipitation–sequencing (ChIP-seq), chromatin accessibility, DNA methylation, and gene expression profiles with GWAS data has revealed novel biological insights for other diseases. Thus far, understanding the molecular basis of CLL susceptibility loci has been limited because genome-wide information on the CLL regulome has not been available for a large CLL series.

In this study, we sought to address this deficiency by analyzing a large, genetically and epigenetically well-characterized CLL series, together with epigenome data of the normal B-cell lineage. First, by characterizing the epigenomes of CLL and different stages of B-cell differentiation we gained insight into the developmental basis of CLL risk. Second, we explored the CLL risk loci using histone modification profiles, chromatin accessibility assessed by assay for transposase-accessible chromatin using sequencing (ATAC-seq), DNA methylation, and gene expression data in conjunction with DNA genotypes in the same CLL cases. Furthermore, we used three-dimensional chromatin data to explore physical interactions between the risk loci and their candidate target genes. These analyses allowed for a detailed interpretation of GWAS signals for CLL and to infer the molecular mechanisms through which loci operate.

**Results**

The active chromatin landscape at CLL risk loci. The majority of GWAS risk loci map to non-coding regions of the genome and influence gene regulation. Hence, to gain insight into the biological basis underlying genetic CLL predisposition, we evaluated profiles of three histone marks related to active regulatory elements at each locus in CLL cells (determined by H3K27ac, H3K4me3, and H3K4me1 ChIP-seq marks). We first considered each of the 42 non-HLA risk loci solely on the basis of the strongest single-nucleotide polymorphism (SNP) association with CLL in our recent meta-analysis. One third (14/42) of the risk SNPs localize to either active promoters or enhancers in CLL cells. Moreover, on the basis of linkage disequilibrium (LD; $r^2 \geq 0.2$) with the sentinel SNP, 93% (39/42) of the risk loci featured SNPs mapping to an active regulatory element. Specifically, on average 81% (34/42) of the LD-defined CLL risk loci showed enrichment of active promoters and/or enhancers per CLL sample. This enrichment of CLL-related active regulatory elements at CLL risk loci contrasts with the significantly lower enrichment observed at both colorectal (CRC) and breast cancer (BC) GWAS loci (Fig. 1b).

Chromatin state enrichment in CLL essentially defines five groups of risk loci characterized by: (i) active regulatory elements only ($n = 15$ loci), (ii) active regulatory elements and transcribed regions ($n = 10$ loci), (iii) weak regulatory elements only ($n = 10$ loci), (iv) weak regulatory elements and transcribed regions ($n = 3$ loci), and (v) poised and inactive regions ($n = 4$ loci). Since the epigenome of B cells varies according to their maturation state, we examined H3K27ac profiles at the risk loci in CLL and a range of normal B-cell subpopulations (Fig. 1c). H3K27ac signals at 10 risk loci (represented by 30 H3K27ac peaks) were significantly increased in CLL cells as compared to normal B-cell subpopulations, a significantly higher number than expected ($P = 3.5 \times 10^{-5}$, Fisher’s exact test, 11 peaks based on 1000 permutations). These included five regions of de novo activation previously documented in CLL (Supplementary Data 5). Overall, 93% of the LD-defined risk loci (39/42, represented by 282 H3K27ac peaks) displayed differential H3K27ac profiles across CLL and the different normal B-cell subpopulations, also significantly more than that expected ($P = 3.0 \times 10^{-4}$, Fisher’s exact test, 239 peaks based on 1000 permutations). Collectively, our data show that CLL risk loci are enriched for CLL-related regulatory elements being CLL specific or having differential regulation across CLL and B-cell development, as exemplified by the loci marked by the SNPs rs11637565 and rs35923643 (Fig. 1d).

To gain further insight into the regulatory mechanisms underpinning the CLL risk loci, we performed a quantitative trait locus (QTL) analysis using 487 genotyped CLL cases, with corresponding H3K27ac ChIP-seq profiles ($n = 97$), chromatin accessibility data assessed by ATAC-seq ($n = 99$), and 450k Illumina DNA methylation profiles ($n = 486$) (Fig. 2a and Supplementary Data 6). Nine of the risk loci were typed by H3K27ac, 14 risk loci by chromatin accessibility, and 28 risk loci by DNA methylation QTLs (13 accessibility QTL peaks overlap with 10 of the H3K27ac QTL peaks, covering 5 risk loci), the direction of effect of associated epigenetic changes was concordant. Furthermore, overlapping DNA methylation QTLs within these regions showed opposite effects, i.e., higher DNA methylation with lower activity/accessibility (Supplementary Data 7). These data are consistent with risk alleles mediating their effects by influencing chromatin activity.

Biological mechanisms and significance of CLL risk loci. We next focused on the underlying biological mechanisms through which genetic variants at CLL risk loci shape the regulatory genome. Both our study (Fig. 1a) and published data indicate that cancer GWAS risk variants map to regulatory regions, suggesting that germline genotypes may mediate their effect through altered transcription factor (TF) binding. In view of this, we assessed whether TF binding may be affected by genetic variation at chromatin-accessible regions of CLL risk loci. We used motifBreaker to predict disruption of TF-binding motifs and examined lymphoblastoid cell line (LCL) ChIP-seq data from the ENCODE project (Supplementary Data 8). This analysis revealed disruption of SP1- and NFKB-binding sites at the risk alleles of SNPs rs13149699 ($r^2 = 0.81$ with sentinel SNP rs57214277) and rs539846 ($r^2 = 0.95$ with sentinel SNP rs8024033), respectively, as previously reported. We also detected disruption of SP1-binding sites at risk alleles of five other risk loci, associated with sentinel SNPs rs1002015, rs11637565, rs142215530, rs210143, and rs489882, as well as disruption of other TFs relevant for B-cell development, including PAX5 and MEF2 TF family members. Furthermore, we observed increased binding affinities of the TFs previously associated with de novo active regions in CLL, that is, FOX, NFAT, and TCF/LEF family members, at risk alleles of 16 risk loci, associated with sentinel SNPs rs11637565, rs4368253, and rs4869818, among others.
These findings underline the role of these TFs in CLL pathogenesis. In addition, as risk variants may exert their pathogenic effect by altering the accessibility of affected alleles, we studied the existence of allelic imbalance at ATAC-seq peaks. The power to detect allelic imbalance in chromatin accessibility was restricted to sites with common variants and relatively large effects. Nevertheless, 10% of the risk loci (4/42) showed evidence of allelic imbalance for chromatin accessibility at a total of 9 SNPs (Supplementary Data 9). Eight of these SNPs (89%) overlapped with accessibility QTLs, a significantly larger fraction than for SNPs that did not show allelic imbalance (35/180, 19%, $P = 3.1 \times 10^{-5}$, Fisher’s exact test).

(Fig. 3, Supplementary Data 8). These findings underline the role of these TFs in CLL pathogenesis.
We finally sought to infer the biological relevance of risk loci using two complementary approaches. First, we performed a gene expression QTL (eQTL) analysis to identify candidate target genes using GeneChip Human Genome U219 gene expression data from 452 CLL cases. Since genomic spatial proximity and long-range chromatin looping interactions are central to the regulation of gene expression, eQTL analysis was performed considering the expression levels of genes located within respective topologically associating domains (TADs) spanning the LD regions (Supplementary Data 1). We detected 36 eQTLs (defined by 62 probe sets assigned to 37 target genes) associated with 23 risk loci (Fig. 2a, b and Supplementary Data 7). Included were many genes with established roles in CLL oncogenesis (e.g. LEF1), B-cell development (e.g. IRF8), or apoptosis (e.g. BCL2L11, CASP8, FAS, BMF). Of note, three genes highlighted by the eQTL analysis, LEF1 (rs7690934), IPCEF1 (rs4869818), and DMRTA1 (rs1679013), have been previously identified as target genes of de novo active regions in CLL (Supplementary Data 5) and show higher expression in CLL compared with normal B cells. Such observations suggest a role for these genes in both CLL predisposition and pathogenesis. Overall, 36 of the 42 risk loci (86%) showed a significant QTL in at least one of the four layers analyzed (Fig. 2b).

Second, we made use of in situ promoter capture Hi-C (CHi-C) data to examine chromatin looping interactions linking the LD-defined risk loci to promoters of candidate target genes. We analyzed CHi-C information from CLL cells as well as from naive and total B cells (Supplementary Data 10). As well as confirming the interaction between the rs2466029-associated LD region and MYC at 8q24.21, the looping interactions implicate a number of other genes with established roles in B-cell development. Notably, these include BCL2 as the target gene of rs77551289/rs4987852 variation at 18q21.33 and BCL6 as the target gene of rs73192661 variation at 3q28. Overall, there was supportive evidence from the CHi-C data for chromatin interactions between the LD-defined CLL risk loci with promoters of 15 of the 37 candidate genes (41%) identified in the eQTL analysis, including TLE3 as a target of the rs11637565 LD region as well as UBR5 as a target of the rs2511713LD region (Fig. 4).

Discussion

Characterization of the regulatory elements that are perturbed by CLL GWAS risk variants is central to efforts to define the mechanisms through which these variants operate and to reconstruct the biological networks that underlie CLL tumorigenesis. However, genome-wide epigenomic characterization of CLL in combination with the availability of genetic data in large CLL series has so far been lacking. Here we have addressed this deficit by using a large, genetically well-characterized CLL series for which genome-wide data of multiple epigenetic layers were recently generated. We combined the genetic and epigenetic data to determine in unprecedented detail genotype-dependent patterns of chromatin accessibility, activity, and DNA methylation in CLL. Overall, our approach has allowed us to infer potential functional variants at 14 of the CLL risk loci (Supplementary Data 11). Moreover, our analyses underline that risk loci in CLL affect genes that participate in interconnecting cellular pathways that are central to B-cell function, including immune response (SP140, BCL6, OAS1, and IRF8), apoptosis (BCL2L11, CASP8, CFLAR, FAS, BMF, and BCL2), and Wnt signaling (UBR5, TLE3, and LEF1). This information serves to illustrate the value of delineating the functional mechanisms and target genes underlying risk loci as such data can have an impact on the successful development of new therapeutic agents. In this respect, it is notable that the recently introduced CLL treatment Ibrutinib inhibits BTK, a kinase with a key role in B-cell function, while Venetoclax targets the antiapoptotic activity of BCL2. In conclusion, our approach has allowed us to generate a refined regulatory map of CLL risk loci offering a more granular annotation of their functional impact, which will facilitate future experimental validation of potential causal variants.

Methods

Patients. The clinical and biological characteristics of the 502 CLL patients studied are detailed in Supplementary Data 6. Cases were defined as IGHV-mutated when the identity of immunglobulin genes was >98%. Tumor samples were obtained pre-therapy. All patients gave informed consent for their participation in the study following the International Cancer Genome Consortium (ICGC) guidelines and the ICGC Ethics and Policy committee, and this study was approved by the clinical research ethics committee of the Hospital Clinic of Barcelona.
were obtained from Law et al.2. Sentinel SNPs at 169 BC risk loci were obtained by imputation: 5 out of 38 SNPs with info score < 0.85 excluded. The imputation excluded samples with ancestry, excessive heterozygosity, or sample relatedness. Five of the 38 imputed SNPs were subject to poor imputation with an information measure <0.85. The imputation fidelity check using the WGS data resulted into the exclusion of one CLL case with >10% discordance between the imputed and WGS data. In the lower part of the panel, summary results for each type of analysis are shown. Complete missing genotypes for remaining 145 out of 487 cases with WGS data: Including genotypes of 5 SNPs with imputation score < 0.85

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Fig. 2 Quantitative trait locus (QTL) analysis and results overview. a Workflow describing the generation of genotypes for 42 non-HLA risk single-nucleotide polymorphisms (SNPs) in 487 chronic lymphocytic leukemia (CLL) patients for use in QTL analyses. The genotypes were assigned using a combination of directly genotyped (n = 4 independent SNPs) or imputed (n = 38 independent SNPs) SNP6.0 array data (502 independent CLL patients) and whole-genome sequencing (WGS) data (n = 146 independent CLL patients). Fourteen of the 502 CLL cases were excluded owing to non-European ancestry, excessive heterozygosity, or sample relatedness. Five of the 38 imputed SNPs were subject to poor imputation with an information measure <0.85. The imputation fidelity check using the WGS data resulted into the exclusion of one CLL case with >10% discordance between the imputed and WGS data. In the lower part of the panel, summary results for each type of analysis are shown. b Per locus overview of QTL analysis results. Loci are denoted by their respective sentinel SNPs (with their location in GRCh38) and shaded boxes indicate the presence of ≥1 significant QTL at false discovery rate <0.05 for each of the four layers analyzed: assay for transposase-accessible chromatin using sequencing (QTLs marked by dark red shading); H3K27ac (orange); DNA methylation (blue); gene expression (green). In addition, on the right, the candidate target genes identified by expression QTL analysis are indicated

Datasets. Whole-genome sequencing (WGS) and SNP6.0 array data (Thermo Fisher Scientific) of, respectively, 146 and 502 patients were obtained from Puente et al.14. Chromatin state segmentations (200 bp bins) of 7 CLLs and 15 normal B-cell samples as well as variance-stabilized (vat) data signals of H3K27ac and ATAC-seq determined by DESeq2 (corrected for signal proportion of tags (SPOT) score) were obtained from Michailidou et al.16. Sentinel SNPs at 78 CRC risk loci were obtained from Schmit et al.15. LD regions were determined as the merged LD regions (r² ≥ 0.2) of all SNPs in the 1000 Genomes Project and UK10K, excluding regions containing CLL, BC, or CRC risk loci (Supplementary Data 12). The difference in chromatin state enrichment for CLL risk regions covered by chromatin state segmentation as previously defined14,15,16 could not be mapped to the GRCh38 genome build and was therefore excluded. Three CRC risk loci were excluded (rs7758229, rs6061231, rs11064437) as they have only been mapped to the GRCh38 genome build and was therefore excluded. Three CRC risk loci (rs554219, 4:84370124, rs7695891) were excluded (rs7758229, rs6061231, rs11064437) as they have only been reported in Asian populations.

Definition of risk loci. Sentinel SNPs at the 42 non-HLA risk CLL GWAS loci were obtained from Law et al.14. Sentinel SNPs at 169 BC risk loci were obtained from Michailidou et al.16. Sentinel SNPs at 78 CRC risk loci were obtained from Schmit et al.15. LD regions were defined by r² ≥ 0.2 with each sentinel SNP. LD metrics were calculated using LDLink32 for the European populations of the 1000 Genomes project (Phase 3 data). Three BC risk loci (rs554219, 4:84370124, rs7758229) were absent in the 1000 Genomes data and therefore excluded (LD regions could not be calculated). One BC risk locus (rs1876636) could not be mapped to the GRCh38 genome build and was therefore excluded. Three CRC risk loci were excluded (rs7758229, rs6061231, rs11064437) as they have only been reported in Asian populations.

Linking risk loci to chromatin states. Chromatin states at risk loci were assigned using the chromatin state segmentations as previously defined12,14,15. For each CLL sample, the chromatin state enrichment at a CLL risk region, E, was calculated as

\[
E_i = \log_2 \frac{C_i}{C_{B}} \tag{1}
\]

where \(i\) is a CLL risk allele, \(j\) is a chromatin state, \(C_i\) is the fraction of the LD region (r² ≥ 0.2) covered by chromatin state j, and \(C_B\) is the fraction of the background regions covered by chromatin state j. These background regions were determined as the merged LD regions (r² ≥ 0.2) of all SNPs in the 1000 Genomes Project and UK10K, excluding regions containing CLL, BC, or CRC risk loci (Supplementary Data 12). The difference in chromatin state enrichment for CLL risk loci with BC and CRC risk loci was calculated using Wilcoxon rank-sum test. No covariates were tested. To account for multiple testing, a Bonferroni correction was applied.

Comparison of H3K27ac signal among CLL and normal B cells. Differential signal intensity of non-individual H3K27ac peaks was calculated using DESeq235,16. H3K27ac counts at these loci were mined from Beekman et al.12,16. The analysis was performed by contrasting CLL with each of the normal subpopulations samples using nbinomWaldTest, correcting for the SPOT score; per sample, its condition (CLL or the corresponding normal B-cell subpopulation) and the SPOT score were introduced into the model as per Beekman et al.12,16. Regions with a false discovery rate (FDR) < 0.01 were considered significantly enriched.

Assignments of genotypes. Genotypes of (i) 42 sentinel SNPs, (ii) 225 bi-allelic SNPs located in ATAC-seq peaks analyzed in the QTL analysis being in LD with the sentinel SNP, and (iii) 94,233 bi-allelic background SNPs were assigned from
SNP6.0 array data (using normal DNA) and WGS data (using normal and/or tumor DNA). The first set of peaks was used for QTL analyses, the latter two for the allelic imbalance analysis (see also Supplementary Data 9 and the paragraph on allelic imbalance in this "Methods" section). Genotyping data from the 502 CLL cases were subject to standard quality control36, resulting in the exclusion of 14 cases prior to imputation because of non-European ancestry (using the HapMap version 2 CEU, JPT/CHB, and YRI populations as reference), excessive heterozygosity, or relatedness. SNPs with a call rate <95%, a minor allele frequency <1%, or displaying significant deviation from Hardy–Weinberg equilibrium (i.e., P < 10^-6) were also excluded. Whole-genome imputation was performed using the IMPUTE2 v2.3 software37 with a merged reference panel consisting of data from the 1000 Genomes Project (phase 1 integrated release 3, March 2012) and UK10K34. No genotypes were assigned to poorly imputed SNPs with an information measure <0.85 (sentinel SNPs, n = 5; SNPs for allelic imbalance, n = 31; SNPs with low information measures were not included in the background set of 94233 SNPs). No genotypes were assigned to SNPs residing in regions with copy number alterations in the corresponding CLL samples (as determined from the SNP6.0 array data of the matched tumor DNA). Imputation probabilities were converted to SNP genotypes using PLINK v1.938, using an uncertainty threshold of >0.1 to preclude genotype assignment. In order to check the fidelity of imputation and to look for potential mismatches in the CLL samples, we examined WGS data from 146 CLL cases for the sentinel SNPs (the cases overlapping with the 488 patients prior to imputation) and 38 CLL cases for the SNPs for the allelic imbalance analysis and the background SNPs (the cases overlapping with the 99 samples with ATAC-seq data). For the WGS data, SNPs were called with the MAQ consensus model implemented in SAMTools in those coordinates with at least 10 reads with a minimum mapping quality of 30 and base quality of 3014. No genotypes were assigned following imputation (SNP information measure <0.85 or genotype probability <0.90), generating matrices of 42 sentinel SNPs genotyped in up to 487 cases for QTL analyses (Fig. 2a) and of 225 SNPs as well as 94233 background SNPs genotyped in up to 99 cases for allelic imbalance analysis.

**Potential functional SNP (sentinel SNP)**

| SNP       | Transcription factor | Risk allele | Binding affinity risk allele | Motif |
|-----------|----------------------|-------------|-------------------------------|-------|
| rs210142  | SPI1                 | C           | Decreased                     |       |
| rs12591150| PAX5                 | T           | Decreased                     |       |
| rs4787033 | MEF2A                | T           | Decreased                     |       |
| rs7761411 | NFATC1               | T           | Increased                     |       |
| rs8083367 | FOXI1                | G           | Increased                     |       |

**Fig. 3** Transcription factor (TF)-binding motif analysis. Examples of altered TF-binding motifs at potential functional risk loci. The affected nucleotides in the different motifs are marked by an asterisk. The sequence surrounding the single-nucleotide polymorphism (SNP) is located under the motif graphs, with the different possible alleles of the potential functional SNP indicated below each other.

**QTL analysis.** QTL analyses were performed using genotype data for the sentinel SNPs in conjunction with ATAC-seq (n = 99)12, H3K27ac (n = 97)12, methylation (n = 486)14, and gene expression data (n = 452)14 from CLL samples. Input data were vst-normalized data signals (corrected for SPOT score) for ATAC-seq and H3K27ac, M-values for DNA methylation, and GC-RMA-normalized for gene expression. ATAC-seq and H3K27ac QTL analyses were performed for peaks present in at least 10% (with a minimum of two) of the patients in one or more of the following subgroups: homozygous non-risk, heterozygous risk, or homozygous risk based on the sentinel SNP genotype. eQTL analysis was restricted to expressed probes (GC-RMA levels >4.5) again in at least 10% (with a minimum of two) of the patients in one or more of the following subgroups: homozygous non-risk, heterozygous risk, or homozygous risk based on the sentinel SNP genotype. To account for confounding factors, we implemented the probabilistic estimation of expression residuals (PEER) method39 using 10, 10, 45, and 65 factors for ATAC-seq, H3K27ac, methylation, and gene expression data, respectively. Inverse normal-transformed PEER-computed residuals were used as input for QTL analyses, which were conducted using Matrix eQTL under a linear model40. TADs for the eQTL analysis were defined using in situ Hi-C data from the LCL GM1287841 and were mined from Beekman et al.51 (http://resources.idibaps.org/paper/the-reference-epigenome-and-regulatory-chromatin-landscape-of-chronic-lymphocytic-leukemia).

**Promoter Chi-C.** Raw data of previously generated in situ promoter capture Hi-C (Chi-C) on CLL, naive, and total B cells was used12,21. Reads were aligned to the GRCh37 build using bowtie2 v2.2.642 and identification of valid di-tags was performed using HiCUP v0.5.943. To declare significant contacts, HiCUP output was processed using CHICAGO v1.1.841 and were mined from Beckman et al.12 (http://resources.idibaps.org/paper/the-reference-epigenome-and-regulatory-chromatin-landscape-of-chronic-lymphocytic-leukemia).

**Allelic imbalance.** A beta-binomial test was used to assess the allelic imbalance at all bi-allelic SNPs that (i) were in LD (r^2 ≥0.2) with the sentinel SNP and located in ATAC-seq peaks used for the QTL analysis (n = 225), and (ii) had at least 3 heterozygous samples and at least 10 reads, leading to a final number of 189 SNPs
Fig. 4 Characterization of quantitative trait loci (QTLs) and looping interactions at the 15q23 and 8q22.3 chronic lymphocytic leukemia (CLL) risk loci. (Top panels) Boxplots illustrating associations between a rs11637565 and b rs2511713 genotypes and chromatin accessibility (assay for transposase-accessible chromatin (ATAC) QTL), histone H3 lysine 27 acetylation (H3K27ac QTL), DNA methylation (methylation QTL), and candidate target gene transcript levels (expression QTL) in CLL patients (number of independent CLL samples for ATAC: n = 99, H3K27ac: n = 97, DNA methylation: n = 486, and transcript levels: n = 452). (Lower panels) Chromosomal positions of sentinel single-nucleotide polymorphisms (SNPs; rs11637565/rs2511713, gray dots), QTL peaks/probes, and coding genes (black rectangles). The linkage disequilibrium (LD) regions, encompassing SNPs in LD = 0.3, were indicated by light blue rectangles. Also shown are looping interactions (pink arcs) from the LD regions to the promoters of TLE3 and UBR5. Pink rectangles represent interacting HindIII domains. Topologically associating domains (dark blue rectangles, TAD) containing all features are truncated for clarity; their extension beyond the figure boundaries is indicated by arrowheads.

to be assessed. Reads with mapping bias were removed using WASP and bwa 0.7.15. In brief, the nucleotide sequence of the reads overlapping the investigated SNPs was modified to investigate potential mapping bias. More specifically, the nucleotides at known SNP positions within these reads were substituted by nucleotides of the other possible allele(s). Reads that after modification failed to remap in the same position in the genome were discarded. PCR duplicates were removed using the WASP script rmdup_pe.py. Thereafter, for each analyzed SNP the number of reads containing the two different possible alleles was determined, pooling all heterozygote samples. To control for possible errors in genotyping, the number of reads containing the two alternative allele together. The beta-binomial parameters were estimated using bi-count ratio of 0.1

values were corrected for multiple testing (FDR < 0.1). The central tendency of allelic imbalance was defined per SNP by calculating the allele ratio of the reference allele, while its 95% confidence intervals was calculated using the normal approximation.

TF motif and binding analysis. To investigate TF binding site disruption, the motifbreaks package was used. Scores were calculated using the relative entropy algorithm. P values were calculated using the score, given the score distribution using the position weight matrices. SNPs were also annotated for bound TFs using ENCODE ChIP-seq data for LCLs.

GWAS datasets. Meta-analysis of three GWAS datasets (UK-CLL1, UK-CLL2 and the NHL-GWAS Consortium) was conducted as previously described. NHL-GWAS Consortium data were a subset of those previously published and were downloaded from dbGAP phs000801.v2.p1 (sub-study phs000802.v2.p1). After standard QC procedures, final GWAS datasets comprised of UK-CLL1:
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