

LYMPHOID NEOPLASIA

Epigenetic regulator genes direct lineage switching in *MLL/AF4* leukemia

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KEY POINTS

- Myeloid relapse can originate from various differentiation stages of *MLL/AF4*⁺ ALL.
- Dysregulation of epigenetic regulators underpins fundamental lineage reprogramming.

The fusion gene *MLL/AF4* defines a high-risk subtype of pro-B acute lymphoblastic leukemia. Relapse can be associated with a lineage switch from acute lymphoblastic to acute myeloid leukemia, resulting in poor clinical outcomes caused by resistance to chemotherapies and immunotherapies. In this study, the myeloid relapses shared oncogene fusion breakpoints with their matched lymphoid presentations and originated from various differentiation stages from immature progenitors through to committed B-cell precursors. Lineage switching is linked to substantial changes in chromatin accessibility and rewiring of transcriptional programs, including alternative splicing. These findings indicate that the execution and maintenance of lymphoid lineage differentiation is impaired. The relapsed myeloid phenotype is recurrently associated with the altered

expression, splicing, or mutation of chromatin modifiers, including *CHD4* coding for the ATPase/helicase of the nucleosome remodelling and deacetylation complex. Perturbation of *CHD4* alone or in combination with other mutated epigenetic modifiers induces myeloid gene expression in *MLL/AF4*⁺ cell models, indicating that lineage switching in *MLL/AF4* leukemia is driven and maintained by disrupted epigenetic regulation.

Introduction

Translocation of mixed lineage leukemia (*MLL*) with 1 of >130 alternative partner genes is a recurrent cytogenetic finding in both acute myeloid leukemia (AML) and acute lymphoblastic leukemia (ALL) and is generally associated with a poor prognosis.^{1,2} Among the most common translocations is t(4;11)(q21;q23), forming the *MLL/AF4* (also known as *KMT2A/AFF1*) fusion gene. Unique among *MLL* rearrangements (*MLLr*), *MLL/AF4* is almost

exclusively associated with pro-B cell ALL and is prototypical of infant ALL, where it carries a very poor prognosis.¹ However, despite this general lymphoid presentation, *MLL/AF4* leukemias have an intriguing characteristic, that of lineage-switched relapses. Lineage-switched acute leukemias lose their lymphoid-specific features and gain a myeloid phenotype upon relapse.³⁻⁵ Alternatively, *MLL/AF4* leukemias may harbor distinct lymphoid and myeloid populations at the same time and thus are classified as

mixed-phenotype acute leukemias (MPALs) of the bilineage subtype.⁶

Lineage plasticity has been associated with the loss of original therapeutic targets.^{7,8} To understand the molecular basis of lineage promiscuity and switching, we examined a unique cohort of *MLL/AF4*⁺ lineage-switched acute leukemia presentation/relapse pairs and MPALs. We demonstrate that disruption of the epigenetic machinery, including the nucleosome remodelling and deacetylation (NuRD) complex, is associated with the loss of lymphoid restriction. Lineage switching is then enacted through redistribution of transcription factor binding and chromatin reorganization. These findings provide novel insight into factors that may prove critical to the effective implementation of lineage-specific, epitope-directed therapies, such as chimeric antigen receptor T-cell (CAR T-cell) or bispecific T-cell-engaging antibody approaches.

Methods

Patient samples and data

Patients were diagnosed by local hematology specialists according to contemporary clinical diagnostic criteria based on morphology and immunophenotypic analysis. All patient samples were collected at the point of diagnosis, remission after treatment, or relapse and were stored with written informed consent for research in 1 of 6 centers (Newcastle Haematology Biobank, Newcastle, United Kingdom; University Hospital Schleswig-Holstein, Kiel, Germany; Dmitry Rogachev National Medical Research Center of Pediatric Hematology, Oncology and Immunology, Moscow, Russia; Haematological Malignancy Diagnostic Service, Leeds, United Kingdom; Princess Maxima Center for Pediatric Oncology, Utrecht, The Netherlands; and Cincinnati Children's Hospital Medical Center, Cincinnati, OH). Mononuclear cells were isolated from bone marrow or peripheral blood by density centrifugation followed by immediate extraction of DNA or RNA, or cryopreservation in the presence of 10% v/v dimethyl sulfoxide.

Samples were requested and used in accordance with the ethics approvals granted to each of the local/institutional ethics review boards (NRES [National Research Ethics Service] Committee North East, Newcastle & North Tyneside 1, United Kingdom, reference 07/H0906/109 + 5; Medical Faculty Christian-Albrechts University, Kiel, reference A 103/08; Dmitry Rogachev National Medical Research Center, Moscow, Russia, references MB2008: 22.01.2008, MB2015: 22.01.2015, and ALL-REZ-2014: 28.01.2014; Haematological Malignancy Research Network, Yorkshire, United Kingdom, reference 04/Q1205/69; Haematological Malignancy Diagnostic Service, Leeds, United Kingdom, reference 14/WS/0098; Erasmus MC METC (Medisch Ethische Toetsings Commissie), The Netherlands, reference MEC-2016-739; and Institutional Review Board (IRB) of Cincinnati Children's Hospital, Cincinnati, OH, reference 2010-0658), and in accordance with the Declaration of Helsinki. Each patient and sample was allocated an anonymized reference, and no identifiable information was shared.

Additional methods are described in the supplemental Methods, available on the *Blood* Web site.

Results

Characterization of *MLL/AF4* acute leukemias with lineage switching

We focused on lineage switches that originally presented as ALL and relapsed as AML and MPALs presenting with distinct lymphoid and myeloid populations. Lymphoid and myeloid phenotypes were defined by morphology and by expression of either B lymphoid (CD19, CD22, and CD79A) or myeloid (CD33, CD117/KIT, and CD64/FCGR1A) antigens (Figure 1A; supplemental Table 1). To exclude de novo and therapy-associated AMLs, which are unrelated to the original ALL and do not share the initiating event, the lymphoid and myeloid presentations and relapses had to display identical *MLL/AF4* breakpoints as proof of genetic relationship (Figure 1B; supplemental Figure 1, supplemental Table 1). Using these definitions, we collected a cohort of 12 cases of *MLL/AF4* ALL comprising 6 infant, 2 pediatric, and 2 adult patients who relapsed with AML, including 1 infant (LS10) who relapsed after B-lineage-directed blinatumomab treatment and 2 infant *MLL/AF4* MPALs (supplemental Table 1).

Lineage-switched leukemia is associated with transcriptional reprogramming

We hypothesized that lineage switching would be linked with changes in gene expression. Because the changes in transcriptome composition may include altered regulation of both transcription and mRNA maturation,⁹ we compared gene expression and splicing between lymphoid and myeloid populations from lineage switching and patients with MPAL. Cluster analysis of differential gene expression robustly separated both population types (Figure 2A). We identified 1374 upregulated (adjusted $P < .01$; log fold change >2) and 1323 downregulated genes in the AML lineage switches and the myeloid populations of patients with MPAL linked to reduced lymphoid and increased myeloid gene expression (Figure 2B, supplemental Table 2). Changed gene expression included the loss of lymphoid genes, such as *PAX5*, *EBF1*, *CD19*, *CD20* (*MS4A1*) and *CD22*, diminished gene expression of immunoglobulin genes, genes involved in the VDJ recombination (*RAG1*, *RAG2*, and *DNTT*), and gain-of-myeloid genes, including *CLEC12A*, *PRAM1*, and *CSF3R* and members of the *CEBP* transcription factor family (Figure 2C-D; supplemental Figure 2A-B).¹⁰⁻¹² Moreover, almost 30% of direct bona fide target genes of *MLL/AF4* including *PROM1*, encoding the stem cell markers *CD133*, *IKZF2*, and *HOXA7* showed lower expression in myeloid cells despite sharing the same *MLL/AF4* isotype (supplemental Figure 3A-D, supplemental Table 2).¹³⁻¹⁵ These data show that lineage switching also involves differential *MLL/AF4*-driven gene expression.

The analysis of RNA isoform compositions showed that lineage switching is associated with altered splicing, comprising changes in intron retention and differential usage of exons and exon-exon linkages (Figure 3A; supplemental Tables 3 and 4). Interestingly, 85% of all differentially used exon-exon linkages were noncanonical and mainly consisted of exon skipping and complex splicing events (Figure 3A-B; supplemental Table 4). Pathway analysis revealed an enrichment of alternatively spliced genes in immune pathways, including antigen processing and membrane trafficking, suggesting that alternative splicing is

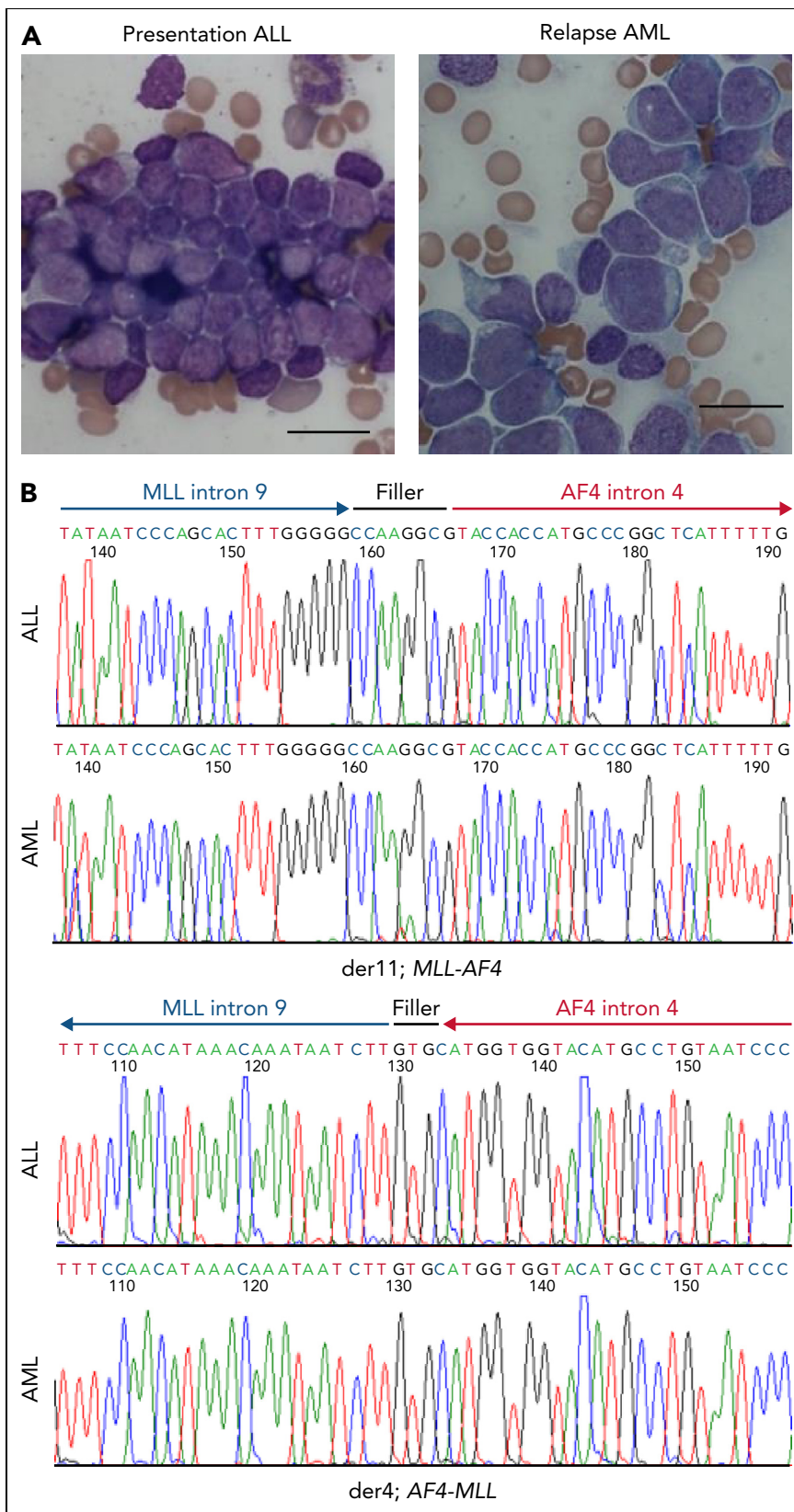


Figure 1. Characterization of MLL/AF4 lineage-switched cases. (A) Morphological change from lymphoblastic leukemia (left) to acute monoblastic/monocytic leukemia (right) in patient LS01. Bar represents 20 μ m. (B) Sanger sequencing of MLL/AF4 and reciprocal AF4/MLL fusions in LS01 presentation ALL (top) and relapsed AML (bottom) identifies a common breakpoint with identical filler sequences in the ALL and AML samples.

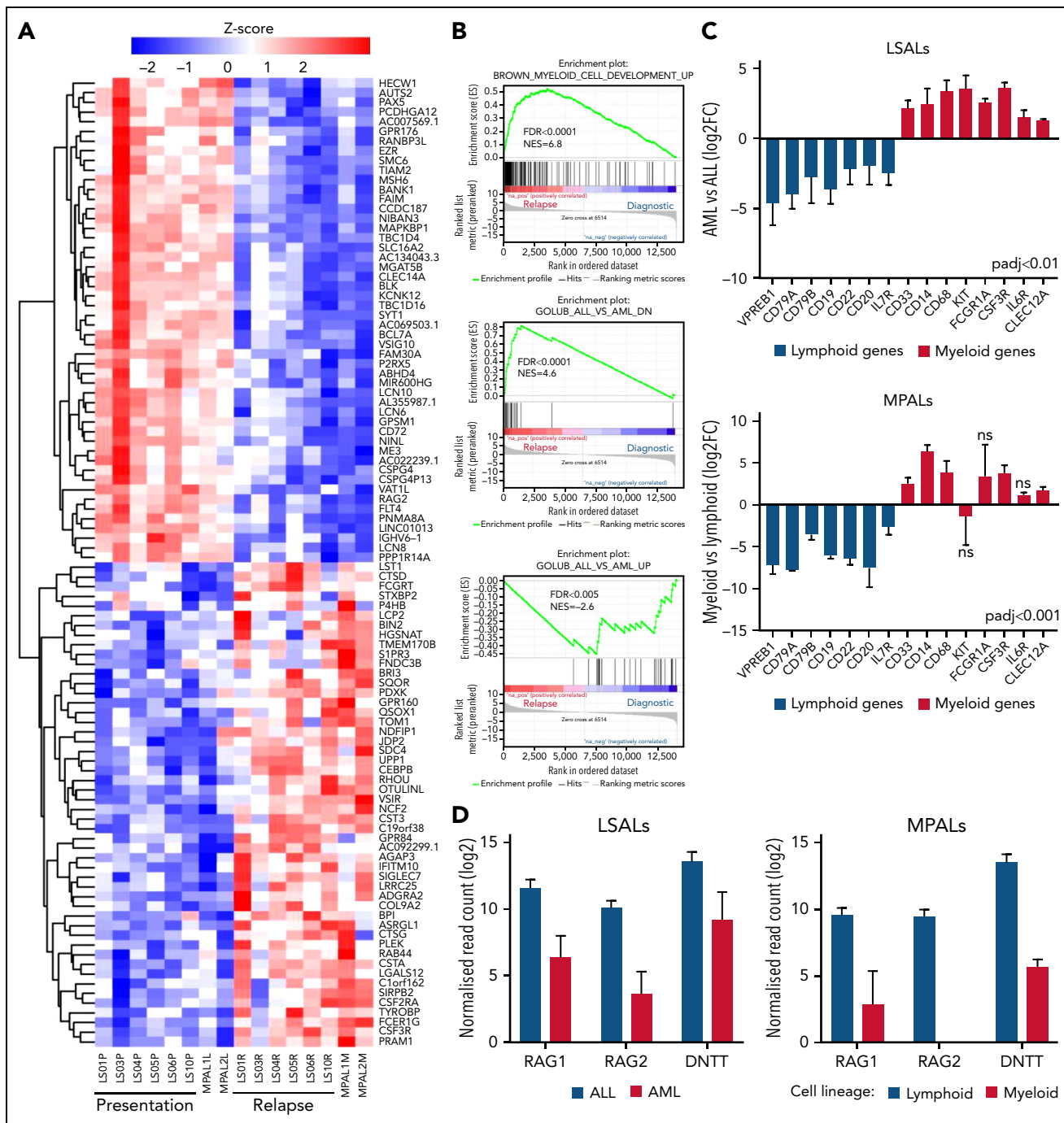


Figure 2. Transcriptional reprogramming in lineage-switched and MPAL cases. (A) Heat map showing the top 100 differentially expressed genes between ALL and AML from 6 lineage-switched (LS01, LS03, LS04, LS05, LS06, and LS10) and 2 MPAL cases, ranked by Wald statistics. (B) Enrichment of myeloid growth and differentiation signature in relapse samples (left) identified by gene set enrichment analyses (GSEA), also pointing to downregulation of genes highly correlating with acute lymphoblastic leukemia (middle and right). GSEAs were performed based on data derived from 6 lineage-switched samples. FDR, false-discovery rate; NES, normalized enrichment score. Differential expression of lineage-specific (C) and immunoglobulin recombination machinery genes (D) in lineage-switched and MPAL cases. Error bars show standard error of the mean for lineage-switched cases and ranges for 2 MPAL cases.

linked to the change from a lymphoid to a myeloid differentiation state (Figure 3C).

Interestingly, lineage switching also affected total expression and the composition of alternatively spliced fusion transcript isoforms for both *MLL/AF4* and *AF4/MLL*. For instance, we detected in relapse tissue from patient LS01 a fusion variant

skipping *MLL* exon 9 (supplemental Figure 3E). In addition, we observed changes in transcription and splicing for genes regulating the chromatin landscape. Several epigenetic regulators, including the polycomb PRC1-like complex component *AUTS2* and the SWI/SNF complex component *BCL7A* were down-regulated in myeloid compared with lymphoid cells (Figure 2A). Several other spliceosome and SWI/SNF members were either

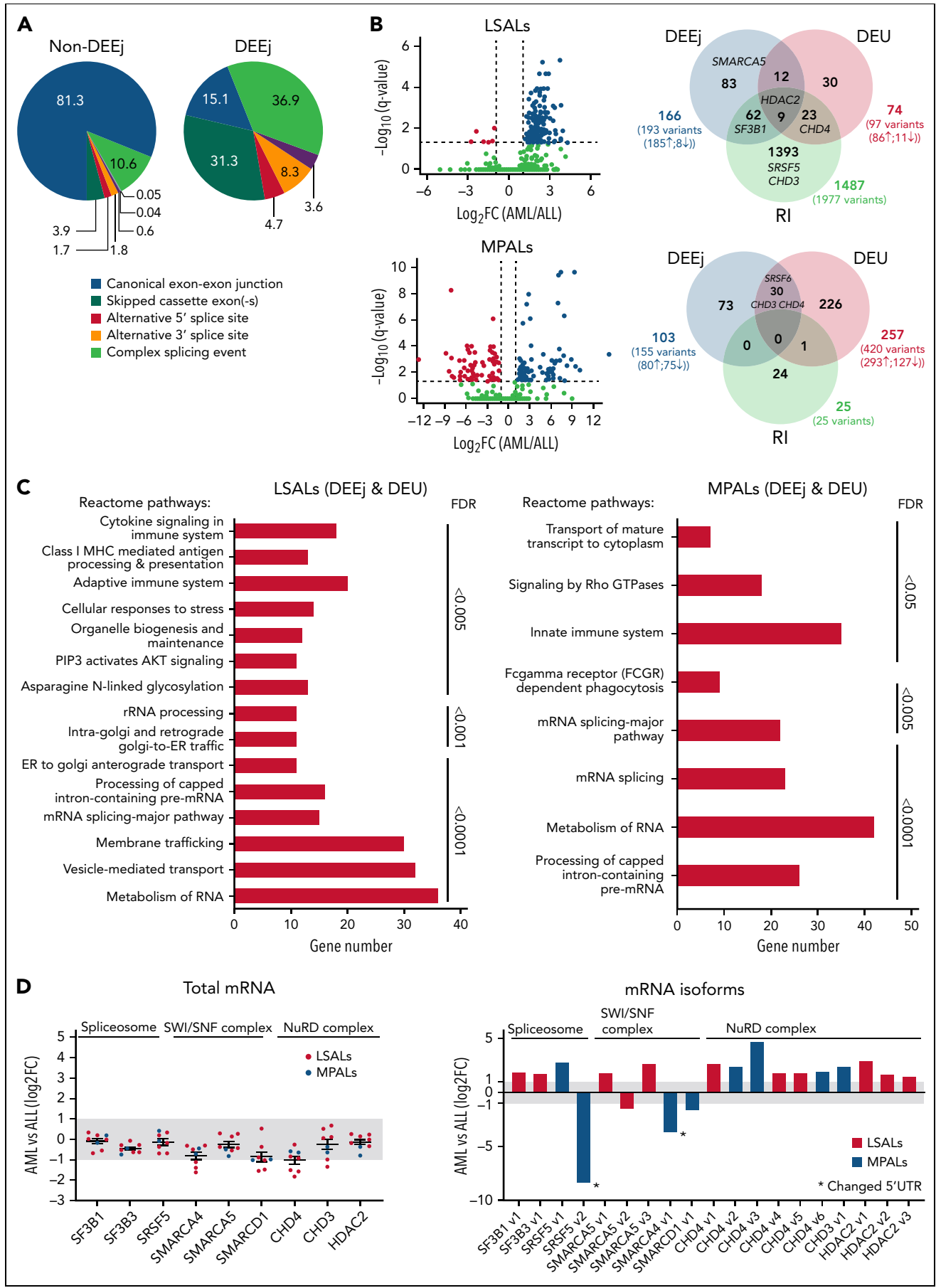


Figure 3.

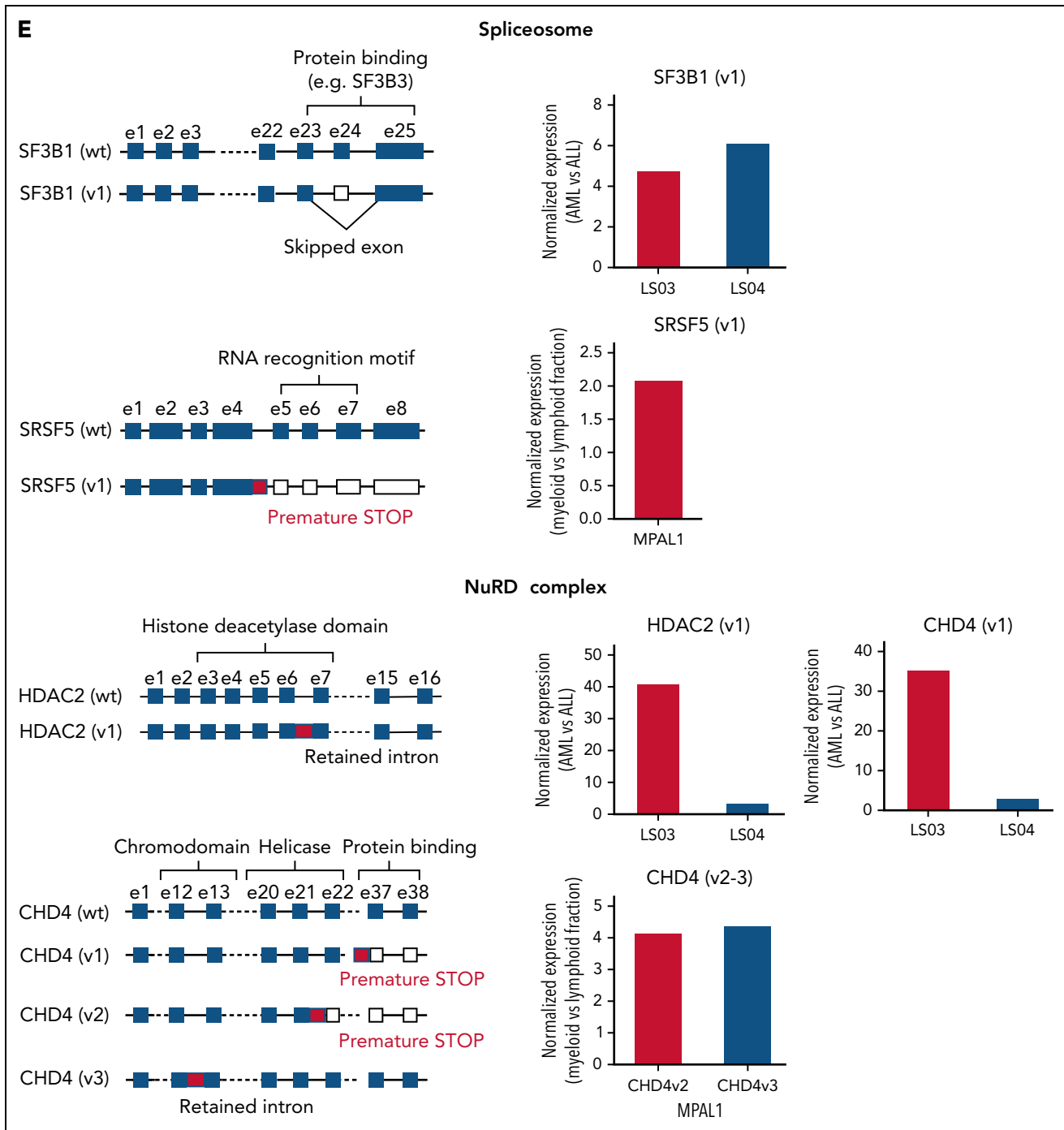


Figure 3 (continued) Alternative splicing in lineage-switched and MPAL cases. (A) Pie charts showing the classification of nondifferential (non-DEE) and differential (DEE) exon-exon junctions. Shown are the percentages of splicing events assigned to a particular mode of splicing. A complex splicing event corresponds to several (2 or more) alternative splicing incidents occurring simultaneously in the same sample. (B) Volcano plots demonstrating differential usage of exon-exon junctions in the transcriptome of AML/myeloid vs ALL/lymphoid cells of cases of lineage switching (LS01, LS03, and LS04) or MPAL. The vertical dashed lines represent twofold differences between the AML and ALL cells, and the horizontal dashed line shows the FDR-adjusted q -value threshold of .05 (left). Venn diagrams (right) showing distribution of splice variants identified as significantly changed in AML (or myeloid fraction of patients with MPAL), including DEEs, differential exon usage (DEU), and retained introns (RI). (C) Enrichment analysis of affected signaling pathways by the DEEs and DEU in the lineage-switched acute leukemia (LSAL) AML relapse and myeloid compartment of patients with MPAL. Pathway enrichment analysis was performed with g:Profiler (<https://biit.cs.ut.ee/gprofiler/gost>) at the highest significance threshold, with multiple-testing correction (g:SCS algorithm). (D) Fold \log_2 change of total transcript levels among genes affected by alternative splicing (left) and of differentially spliced variants in lineage-switched and myeloid compartments of patients with MPAL (right). (E) Representation of impact of alternative splicing on mRNA composition and open reading frames (ORFs) of select genes. Column graphs on the right indicate corresponding fold changes of variant expression between AML (or myeloid) and ALL (or lymphoid) populations in 2 tested lineage-switched cases (LS03 and LS04) and 1 case of MPAL.

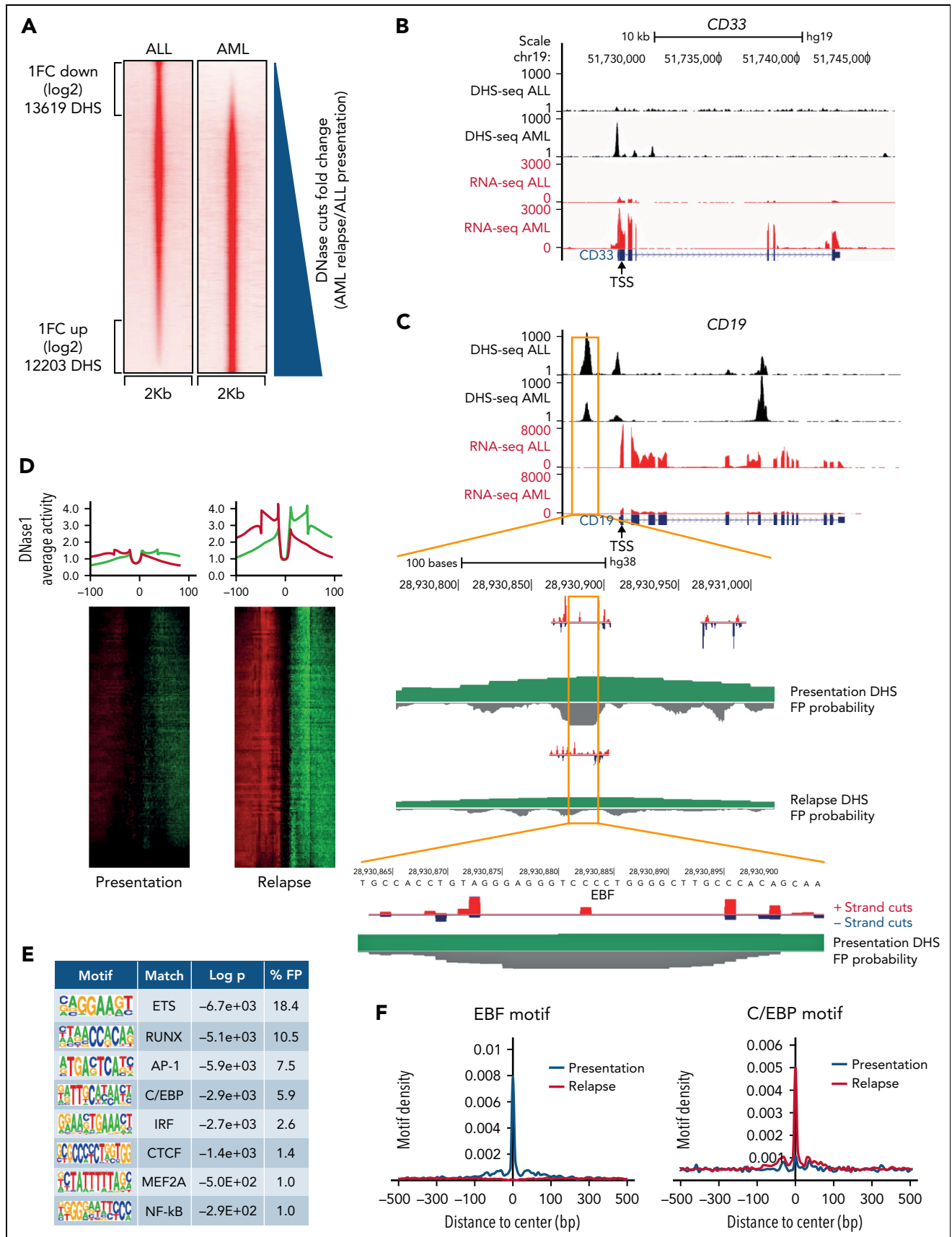


Figure 4.

differentially expressed or spliced. Among all NuRD complex members, only *CHD4* demonstrated differential expression, whereas *CHD4*, *CHD3*, and *HDAC2* were differentially spliced in AML relapse cells or the myeloid subpopulations of MPALs (Figure 3D-E; supplemental Table 4). For instance, *CHD4* encoding the ATPase/helicase subunit of the histone-modifying NuRD complex showed a significantly lower expression in AML relapses of patients with lineage switching, but was differentially spliced in patients with MPAL, resulting in premature stops or intron retention, most likely leading to loss of function isoforms.

Reorganization of chromatin accessibility and transcription factor binding site occupancy upon lineage switching

The substantial gene expression changes, including those affecting epigenetic regulators and lineage-determining transcription factors, prompted us to link transcriptional changes to altered genome-wide chromatin accessibility. High-resolution DNase I-hypersensitive site mapping, combined with digital footprinting analysis using the Wellington algorithm,¹⁶ uncovered multiple differentially accessible genes, including the hematopoietic surface marker genes *CD33* and *CD19* and transcription factors (Figure 4A-C; supplemental Figure 4A-B). These alterations occurred both at locations distal and proximal to transcription start sites indicating the involvement of enhancers and promoters (Figure 4D; supplemental Figure 4C). Digital footprinting is now generally accepted to highlight factors that are essential for regulating specific cell fates.¹⁷⁻¹⁹ These analyses showed that changes in chromatin accessibility after lineage switching were linked to an altered pattern of occupancy of transcription factor binding sites (Figure 4E; supplemental Figure 4D), with a loss of occupancy of consensus binding sites for lymphoid transcription factors, including EBF and PAX5, and a corresponding increased occupancy of binding motifs for myeloid factors including C/EBP family members (Figure 4E-F). We also observed a redistribution of footprinted sites for transcription factors controlling both lymphoid and myeloid maturation, such as RUNX, AP-1, and ETS family members, to alternative cognate motifs (Figure 4E; supplemental Figure 4D).^{20,21} This finding is exemplified by decreased accessibility of a region located 1 kb upstream of the *CD19* transcription start site with concomitant loss of EBF binding site occupancy at this element (Figure 4C). In summary, the transition from lymphoid to myeloid immunophenotype is associated with genome-wide alterations in chromatin accessibility and occupancy of transcription factor binding sites.

The mutational landscape of lineage switching

Next, we examined the mutational landscape of lineage-switched *MLL/AF4* leukemias by performing exome sequencing on the entire cohort. In agreement with previously reported mutation rates in *MLL*-rearranged leukemias, presentation ALLs displayed a relatively quiet mutational landscape, with a median

of 25 nonsynonymous somatic single nucleotide variants (SNVs) or insertions/deletions (indels) (supplemental Figure 5A-B; supplemental Table 5).^{10,22} Most of them were subclonal with <30% of the reads. The group of AML relapses showed a mean of 92 SNVs and indels. However, this increase was related to the more heterogeneous composition of the relapse group: 2 cases (LS07AML and LS08AML) carried mutated DNA polymerase genes resulting in increased mutational burden. We observed this phenotype in only 2 of 10 relapses, arguing against this phenomenon being a general requirement for the lineage switch.

In general, we found only a limited overlap between mutations in ALL presentation and AML relapses (Figure 5A-B; supplemental Table 5). Although ALL mutations were not associated with genes belonging to specific functional pathways, AML-specific mutations were associated with the regulation of transcription and chromatin binding and modification, further emphasizing the notion of transcriptional reprogramming during lineage switching. Most of the subclonal mutations identified in presentation samples were subsequently lost at relapse, indicating alternative subclones as the origin of relapse. This included *KRAS* and *NRAS* mutations, which have been shown to confer a worse clinical outcome to infants with *MLL*-rearranged ALL (Figure 5C).²³ Also the MPALs harbored many mutations that were exclusively found in either the lymphoid or myeloid subpopulation indicating the presence of subclones with a lymphoid and myeloid bias (Figure 5A-B). These combined data show that lymphoid and myeloid leukemia phenotypes are associated with distinctive mutation signatures both in lineage switches and in MPALs.

Perturbation of CHD4 and PHF3 disrupts lymphoid development in MLL/AF4-expressing cells

To identify factors contributing to the lineage plasticity in *MLL/AF4*⁺ leukemic cells, we compared all genes demonstrating differential expression, alternative splicing or mutation in the AML relapse (Figure 6A). This comparison highlighted 8 genes common to all patients with lineage-switched relapse. One common gene was *CHD4*, which codes for the ATPase/helicase subunit of the NuRD, a multiprotein transcriptional corepressor complex with both histone deacetylase and adenosine triphosphate (ATP)-dependent chromatin remodelling activity. NuRD is critical for determination of lymphoid lineage by interacting with the transcription factor IKZF1.²⁴⁻²⁶ *CHD4* shows significantly lower expression in lineage-switched AML when compared with ALL presentation and is differentially spliced in the MPAL cases (Figures 3E and 6B). Finally, although *CHD4* mutations have been reported in <1.5% *MLL*-germline childhood cases of ALL,²⁷ as with the R1068H mutation found in the relapse of patient LS01, these variants commonly affect highly conserved

Figure 4. Chromatin reorganization and differential transcription factor binding underpins lineage switching. (A) DNase I hypersensitive site sequencing identified 13 619 sites with a log₂-fold reduction and 12 203 sites with a log₂-fold increase after lineage switching to AML. Relative peak heights in the AML sample were plotted against those of the ALL sample. (B) A University of California, Santa Cruz (UCSC) Genome Browser screenshot displaying differential expression at lineage-specific loci (red tracks) accompanied by altered DNase I hypersensitivity (black tracks) proximal to the transcriptional start site (TSS) of *CD33*. (C) UCSC Genome Browser screenshot for *CD19* zoomed in on an ALL-associated DHS with EBF occupation as indicated by high-resolution DHS-seq and Wellington analysis. FP, footprint. (D) Heat maps showing distal DHS regions specific for AML relapse on a genomic scale. Red and green indicate excess of positive and negative strand cuts, respectively, per nucleotide position. Sites are sorted from top to bottom in order of decreasing footprint occupancy score. (E) De novo motif discovery in distal DHSs unique to AML relapse as compared with ALL relapse, as shown in panel D. (F) EBF1 and C/EBP binding motifs demonstrate differential motif density in presentation ALL and relapsed AML. DHS, DNase-hypersensitive site.

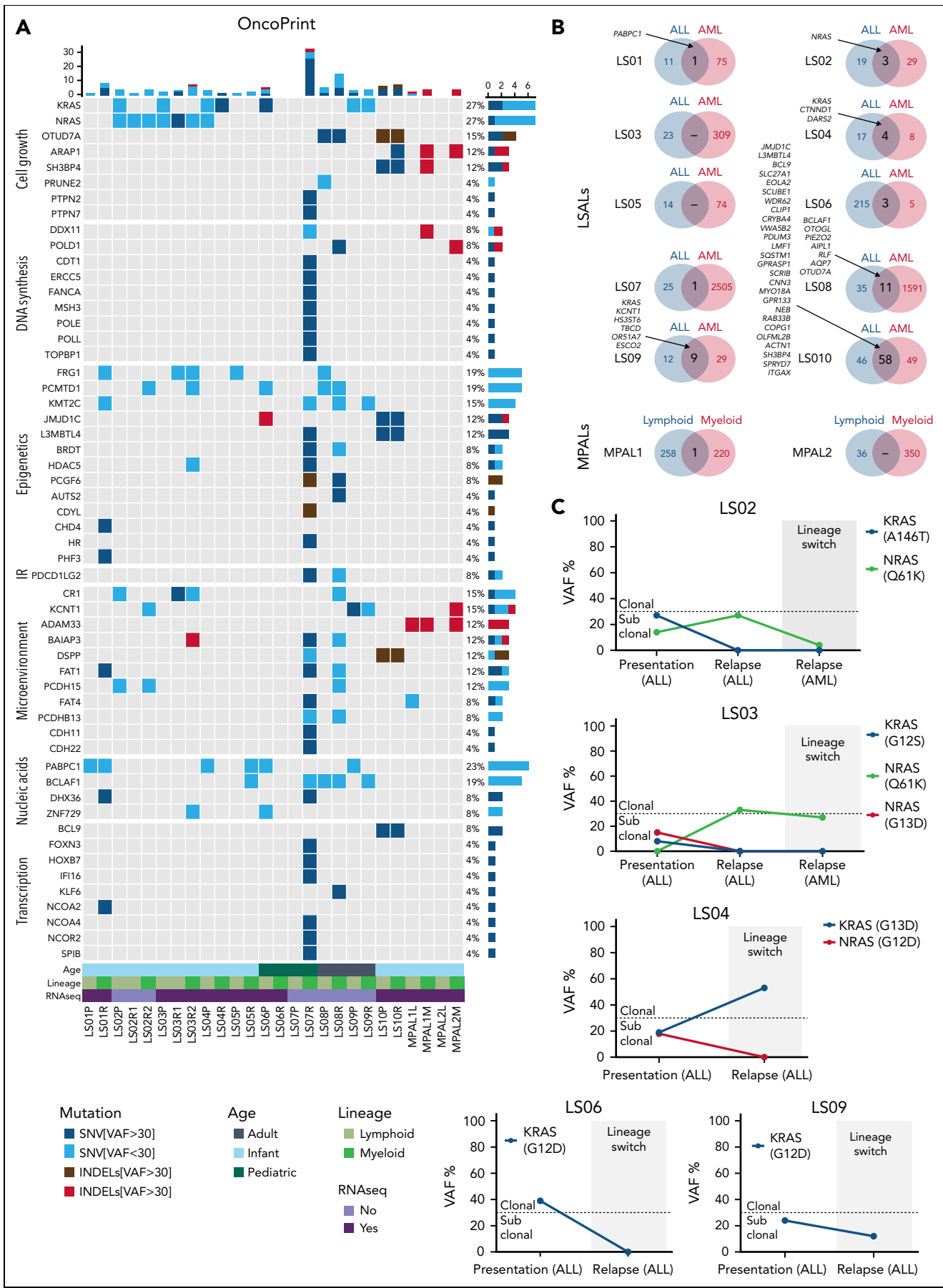


Figure 5.

residues in the helicase/ATPase domains and are thought to disrupt its activity (Figure 6C; supplemental Figure 5C).²⁸⁻³⁰ In contrast, recurrent mutations in other NuRD complex members have not been described in ALL, and no other NuRD complex member was clonally mutated in our cohort (supplemental Table 5).

We therefore hypothesized that CHD4 was vital in maintaining lineage fidelity in *MLL/AF4*-positive ALL. To test this idea, we performed knockdown experiments in the *MLL/AF4*-expressing and CD33⁻ ALL cell line SEM, where we included *ACAP1*, *DHX36*, *NCOA2*, *PHF3*, and *PPP1R7* as 5 additional genes with potentially deleterious mutations in patient LS01 (supplemental Figure 6A). Reverse engineering of a mutual gene network from 216 ALL and AML gene expression data sets identified *CHD4* and *PHF3*, a cofactor in RNA Pol II-mediated transcription,³¹ as the most relevant network components of the mutated genes investigated (*PHF3*: 21 edges, $P = .010$; *CHD4*: 12 edges, $P = .0005$; supplemental Figure 6B, supplemental Table 6).^{32,33}

Only knockdown of *CHD4* and of *PHF3* induced robust expression of the myeloid surface marker CD33 with a combined knockdown resulting in an even stronger CD33 expression (supplemental Figures 6A,C). Moreover, knockdown of either *CHD4* or *PHF3* also increased CD33 levels in RS4;11, another *MLL/AF4* ALL cell line, but not in the 2 *MLL*-germline ALL cell lines 697 and REH (supplemental Figure 6D), indicating that loss of *CHD4* or *PHF3* may only affect CD33 in the context of *MLL/AF4*. Finally, the combined knockdown of *CHD4* and *PHF3* in PDX from diagnostic ALL cells significantly increased the fraction of CD33⁺ cells from 8% to more than 25% (supplemental Figure 6E). These combined data suggest that *CHD4* and *PHF3* restrict *MLL/AF4*⁺ leukemic cells to a lymphoid phenotype.

To examine the role of additional mutations of chromatin modifiers found in our cohort, we investigated the impact of the PRC1 members *PCGF6* and *AUTS2*, genes with known roles in B-lymphoid malignancy³⁴ and mutated in LS07RAML and LS08RAML (Figure 5A). Although knockdown of *AUTS2* did not change the CD33 levels, depletion of *PCGF6* increased CD33 surface expression in SEM cells, further supporting the notion of epigenetic factors in regulating lineage determination in ALL (supplemental Figure 6F).

To establish a direct link between *CHD4/PHF3* binding to the upregulation of myeloid genes, we investigated the impact of *CHD4* or *PHF3* perturbation on gene expression and chromatin organization by performing RNA-sequencing (RNA-seq), ATAC-seq, and chromatin immunoprecipitation-seq for *CHD4* in SEM cells and the *MLL* germline cell line 697 (Figure 6D; supplemental Figure 7A-B). In this analysis, we ranked the ATAC-seq and chromatin immunoprecipitation-seq signals according to their fold changes alongside the control patterns, which

demonstrated that ATAC-seq analysis of control-treated SEM cells show a very similar pattern to *CHD4* binding (Figure 6D), confirming that this factor is a global regulator of chromatin accessibility. Knockdown of both factors caused a shift in the overall chromatin accessibility pattern, as shown by clustering analysis (supplemental Figure 7A-B, bottom) suggesting that the postknockdown cells shifted their cisome and thus their identity, whereby *CHD4* knockdown resulted in a gain of open chromatin sites (Figure 6D, top). The knockdown of *PHF3* caused both a loss and a gain of open chromatin sites (Figure 6D, bottom). Gene set enrichment analysis demonstrated a strong correlation of these gene expression changes in SEM cells after knockdown of *CHD4* and *PHF3* and lineage-switching cases (supplemental Figure 7C-D). However, these changes were particular to *MLL/AF4* cells, because in *MLL* germline 697 cells, *CHD4* knockdown-induced changes in chromatin accessibility were not linked to altered gene expression, and knockdown of *PHF3* did not affect chromatin accessibility (Figure 6D, right).

Knockdown of *CHD4* or *PHF3* in SEM cells changed chromatin structure and reduced expression of *CD79B*, *RAG2*, *VPREB1*, and *CD22*, whereas concomitantly increasing transcription of *CEBPA*, *LYZ*, *SIRPA*, and *CD33* (Figure 6E; supplemental Figure 8A-B). However, 697 cells showed neither a change in immunophenotype nor altered expression of these genes, suggesting that *CHD4*- and *PHF3*-mediated changes in gene expression correlate with the presence of an *MLL* fusion gene.

Given that the relapse-initiating cell may arise within an uncommitted, *MLL/AF4*-translocated HSPC population, we assessed the impact of loss of *CHD4* and *PHF3* in a human cord blood model that harbors a chimeric *MLL/Af4* fusion.³⁵ Knockdown of either *CHD4* or *PHF3* under lymphoid culture conditions significantly impaired lymphoid differentiation potential, whereas knockdown of both *CHD4* and *PHF3* disrupted differentiation entirely (Figure 6F-G; supplemental Table 7). Transcriptomic analysis of the sorted populations revealed that CD33⁺ cells exhibited a metagene expression pattern similar to *MLLr* AML, whereas the pattern describing CD19⁺ cells was most similar to *MLLr* ALL, confirming that changes in surface marker expression were associated with the corresponding changes in the transcriptomic profiles (supplemental Figure 6G).

Taken together, our data show the important role of *CHD4* and *PHF3* in the epigenetic control of lymphoid lineage maintenance in *MLL/AF4*⁺ leukemia. In particular, dysregulation of *CHD4*/NuRD was mediated by mutation, downregulation of expression, and differential splicing across the entire cohort. These data support a role for these factors in the lineage-determining capacity of *MLL/AF4*, whereas their loss undermines execution and maintenance of the lymphoid lineage fate.

Figure 5. Molecular characterization of lineage-switched *MLL/AF4* leukemias. (A) Whole-exome-sequencing data showing genes recurrently mutated within the analyzed cohort and genes clonally mutated in relapsed cases belonging to the same function protein complexes (eg, DNA polymerases, epigenetic complexes, and transcriptional regulators). Data are presented according to the disease time point/cell lineage and age of the patient. Depicted are major SNVs/indels that were found in >30% of reads and minor SNVs/indels present in <30% reads. (B) Comparison of total mutation load (SNVs and indels) identified in patients at presentation (ALL) and relapse (AML) disease stage or lymphoid and myeloid fraction in MPALs. Listed are common SNVs predicted (by Condel scoring) to have deleterious effects. (C) Evolution of *KRAS/NRAS* mutation-carrying cells during the lineage-switching process. Clonal vs subclonal mutations were defined based on variant allele frequencies (VAFs) of identified hits at setup cutoff of 30%.

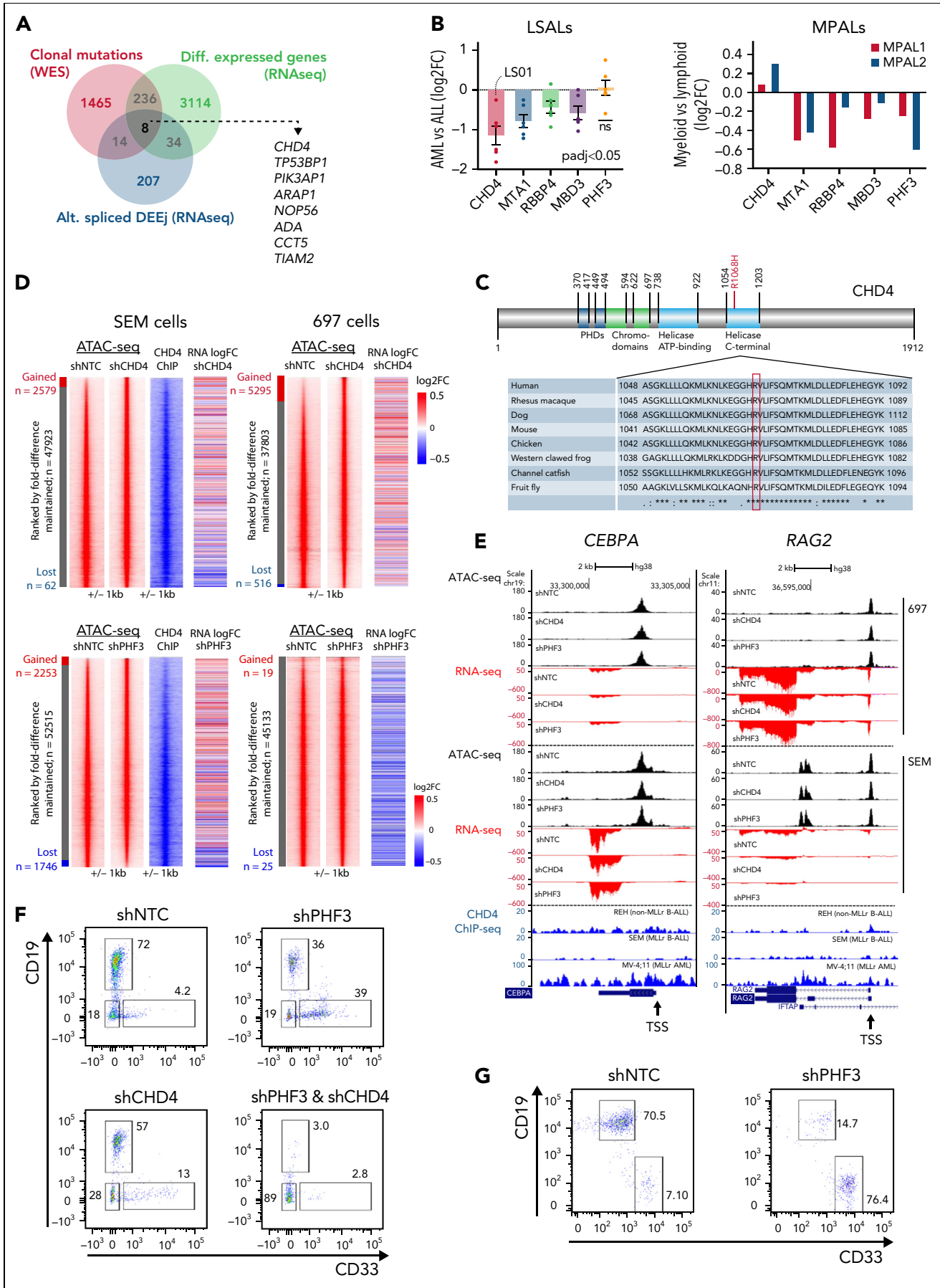


Figure 6.

Clonal evolution of AML relapse

The observed cooperation of *CHD4* and *PHF3* in the control of lineage determination predicted that both mutations co-occur in the same cell. Furthermore, because both mutations may be necessary for the lineage switch in patient LS01, we hypothesized that they should be detectable in the most immature populations of this AML sample for which we had viable cellular material. We therefore investigated the order of acquisition of these secondary mutations within the structure of the normal hematopoietic hierarchy. Dissecting the relapsed AML sample using cell sorting, we isolated HSC-, MPP-, LMPP- and GMP-like, as well as more differentiated populations, followed by targeted deep sequencing that examined *MLL/AF4* and 12 SNVs, including mutated *CHD4* and *PHF3*, that were unique to the relapse sample. The fusion oncogene was found in the multipotential progenitor population (MPP, CD34⁺CD38⁻CD45RA⁻CD90⁺) and in the lymphoid-primed multipotent progenitorlike population (LMPP; CD34⁺CD38⁻CD45RA⁺), with lymphoid, myeloid, but not megakaryocyte-erythroid potential. (supplemental Figure 9A-B; supplemental Table 8). When examining the presence of the 12 SNVs across the different populations, only *PHF3* and *CHD4* mutations were present within the purified MPP-like fraction with variant allelic frequency ≥ 0.3 (Figure 7A, supplemental Table 8). In contrast, LMPP- and GMP-like populations contained all 12 SNVs at high variant allelic frequency. These findings place the *CHD4* and *PHF3* mutations among the earliest genetic events in this patient during the evolution of lineage-switched relapse. Moreover, they suggest, at least for this patient, an MPP-like or even more immature cell population as the origin of relapse.

Cellular origin of lineage-switched relapse

To examine whether lineage-switched relapse regularly arises from lymphoid primed or even earlier leukemic populations, we examined whether relapsed AML cells contained and even shared B-cell receptor (BCR) rearrangements with the preceding ALL. To interrogate the developmental stage at which the myeloid relapse arose, we analyzed BCR rearrangements with RNA-seq and whole-exome-seq-derived data.³⁶ All cases of ALL showed classic oligoclonal rearrangements of BCR loci, supporting the lymphoid lineage decision (supplemental Figure 9C, supplemental Table 9). We observed 3 distinct patterns for AML relapses (Figure 7B). Pattern 1 comprises AML cells with no BCR rearrangements, implying the presence of a relapse-initiating cell residing in a primitive precursor population before early

DJ recombination. This pattern was seen with patient LS01 and, together with the presence of *CHD4* and *PHF3* mutations, strongly supports an MPP-like population as a putative origin of relapse (Figure 7A). As a second pattern, we found unrelated BCR rearrangements, which may indicate either aberrant rearrangement in a myeloid cell or relapse initiating from B-lymphoid cells committed to undergo rearrangement, or a transdifferentiated minor ALL clone with an alternative rearrangement (Figure 7C; cases LS03, LS06, LS07, LS08, MPAL1, and MPAL2). Interestingly, this pattern is found in a relapse after blinatumomab treatment (LS10) suggesting that immune escape may occur by direct transdifferentiation (Figure 7C). Pattern 3 shows shared BCR rearrangements between diagnostic and relapse material, which suggests a transdifferentiated myeloid relapse from the major ALL clone (cases LS05 and LS09). These data demonstrate that AML relapses can originate from different stages of lymphoid leukemogenesis.

Discussion

This study describes impaired epigenetic control as being central to the phenomenon of lymphoid-myeloid lineage switching in *MLL/AF4* leukemia and demonstrates a heterogeneous cellular origin of relapse. The comparison of BCR rearrangements between matched ALL presentation and AML relapse cases demonstrates that whereas relapse can evolve directly from pro-B-like ALL blast populations, in keeping with the general self-renewal capacity of ALL cells,³⁷ it can alternatively originate within the hematopoietic stem and progenitor cell (HSPC) compartment. Indeed, the identification of *MLL/AF4*-expressing MPP-like cells shows that lineage-switched relapse can originate from very immature hematopoietic progenitor populations. This finding agrees with recently published data pointing at MPP cells as the origin of *MLL/AF4* leukemia³⁸ and is in line with transcriptomic similarities between t(4;11) ALL and Lin-CD34⁺CD38⁻CD19⁻ fetal liver cells, again suggesting HSPCs as the cells of origin.²³ Furthermore, the identification of *MLL/AF4* within HSPC populations is consistent with the recent identification of an early lymphoid progenitor, ELP-like signature specifically in *MLL*-rearranged ALL.³⁹ Nevertheless, and in agreement with previously published findings for MPALs,⁶ the data derived from the present cohort strongly support a non-lineage committed progenitor compartment as a source for lineage-switched relapse. However, we cannot exclude additional cells of origin of *MLL/AF4* ALL.

Figure 6. Epigenetic modulatory genes influence lineage-specific expression profiles. (A) Intersection between identified hits of clonal mutations (variant allelic frequency [VAF] >30%), differentially expressed genes and alternatively spliced, differentially used exon-exon junctions (adj. $P < .01$) in lineage-switched myeloid relapse/myeloid fraction of MPALs, present in the analyzed cohort. (B) Fold change in expression of NuRD complex members (*CHD4*, *MTA1*, *RBBP4*, and *MBD3*) and *PHF3*, after lineage-switched relapse (left) and in MPAL cases (right). (C) *CHD4* structure; the R1068H mutation (red) is located in the critical helicase domain of *CHD4* at a highly conserved residue. *Number of positions that have a single, fully conserved residue; colon, conservation between groups of strongly similar properties, scoring >.5 in the Gonnet PAM 250 matrix; period, conservation between groups of weakly similar properties, scoring ≤ 0.5 in the Gonnet PAM 250 matrix. (D) Identification of regions of differential chromatin accessibility before and after knockdown of *CHD4* and *PHF3* depicted in red in *MLLr* SEM cells (left) and non-*MLLr* 697 cells (right). For all reads, the fold change in ATAC-peak height was calculated relative to the control (shNTC) and ATAC peaks from knockdown cells were plotted according to their fold change vs the control signals. *CHD4* ChIP density plots from SEM cells (depicted in blue) were plotted along with the corresponding DNA regions of the shNTC control. Differentially expressed genes associated with changing ATAC peaks (\log_2FC analyzed vs shNTC) identified in each cellular variant are represented by heat maps included at the right side of each gene (for SEM and 697 cells). (E) UCSC genome browser screenshots representing differential chromatin accessibility (ATAC-seq) and gene expression level (RNA-seq) in the myeloid *CEBPA* and the lymphoid *RAG2* loci after *CHD4* and *PHF3* knockdown in *MLLr* SEM cells and non-*MLLr* 697 cells. ChIP-seq traces representing normal *CHD4* occupancy in non-*MLLr* B-ALL (REH cells), *MLLr* B-ALL (SEM cells) and *MLLr* AML cells (MV-4;11) are shown as a reference at the bottom of each screenshot. TSS, transcriptional start site is depicted for each gene. (F) Expression of the lineage-specific cell surface markers CD19 (lymphoid) and CD33 (myeloid) after culture of *MLL/AF4*-transformed hCD34⁺ cord blood progenitor cells in lymphoid-permissive conditions. Knockdown of *PHF3*, *CHD4*, or a combination disrupts the dominant lymphoid differentiation pattern in nontargeting control (shNTC). (G) *PHF3* knockdown is capable of influencing the surface marker expression after a longer incubation period (33 days). *CHD4* knockdown impaired cellular survival upon longer in vitro culture (data not shown). ChIP, chromatin immunoprecipitation.

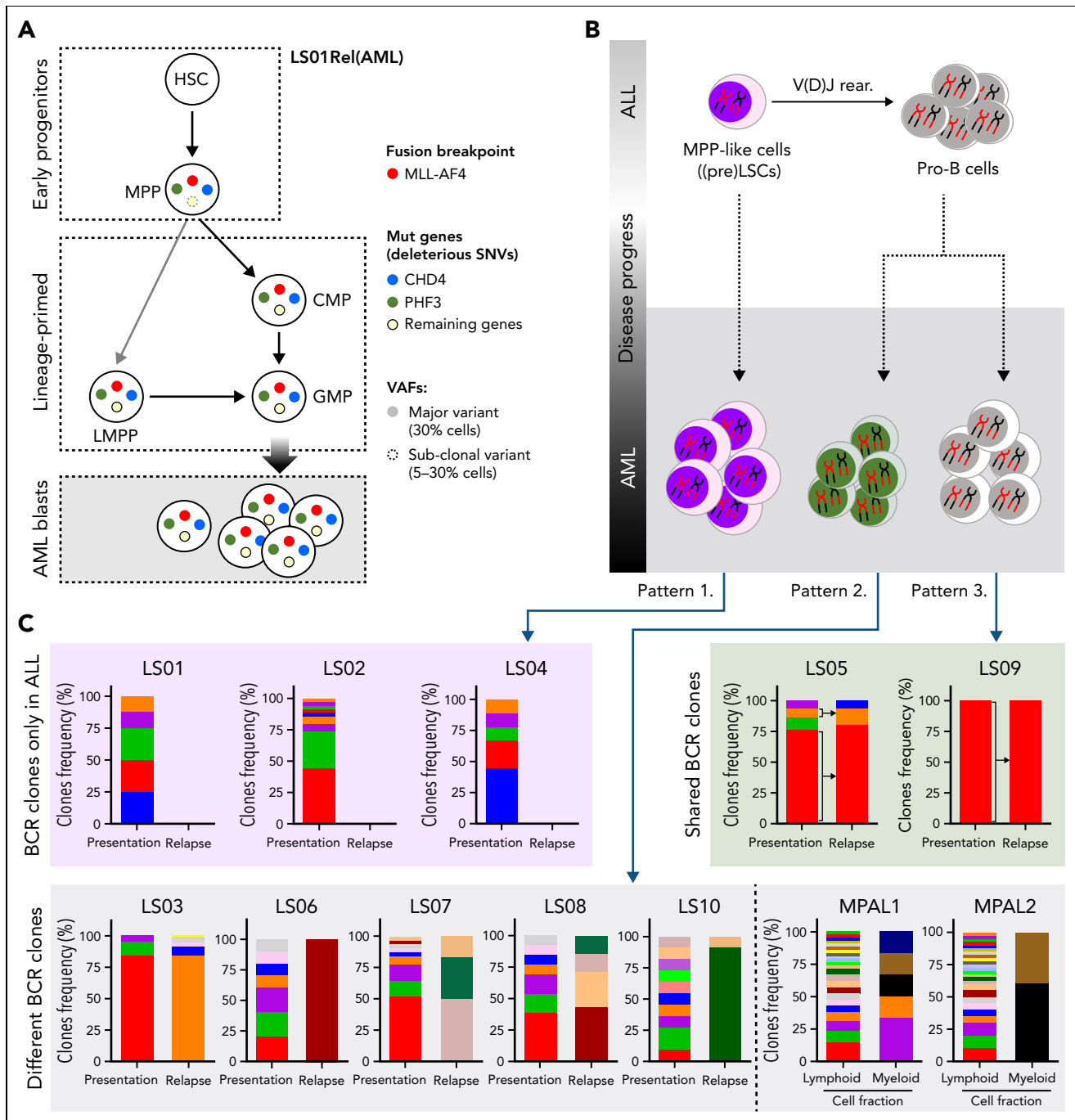


Figure 7. Hematopoietic stem/progenitor populations carry *MLL/AF4*. (A) Summary of *MLL/AF4* positivity and 12 SNVs exclusive for the AML relapse, within different populations analyzed in patient LS01RAML. Circles with solid colors indicate VAF >30%, light dashed circle indicates VAF 5% to 30%. Remaining genes (open circle) represent the 10 other SNVs (of 12 SNVs) which showed the same pattern in the frequency of mutation acquisition (described in supplemental Table 8). (B) Summary of the proposed model of the origin of lineage switched relapse. Evaluation of B-cell receptor repertoires on ALL (presentation) and AML (relapse) lineage switched, and MPAL cases identified 3 different patterns. Pattern 1, with clonotypes on the ALL only. Pattern 2, B-cell receptor-containing clones on ALL and AML, but distinct to each other. Pattern 3, B-cell receptor-containing clones shared between ALL and AML. (C) BCR clone frequencies identified in whole-exome-seq data with application of MixCR software in all lineage-switched acute leukemia (LSAL) and MPAL cases. VAF, variant allele frequency.

Irrespective of the cellular origin of the relapse, lineage switching is associated with a major rewiring of gene regulatory networks. At the level of transcriptional control, the decision for lymphoid development relies, not only on the activation of a lymphoid transcriptional program, but also on the silencing of a default myeloid program.⁴⁰ That decision is enacted by lymphoid master regulators including *EBF1*, *PAX5*, and *IKAROS*,

which represent genes commonly mutated in precursor B-ALL and do not just upregulate B-cell specific genes, but also repress the myeloid program.⁴⁰⁻⁴⁴ *Pax5*^{-/-} pro-B cells, which lack lymphoid potential, although capable of erythromyeloid differentiation in vitro, maintain expression of early B-cell transcription factors *EBF1* and *E2A (TCF3)*.⁴⁰ In contrast, we showed that lineage switched *MLL/AF4* pro-B leukemic relapse is associated

with a significant reduction in expression of these earliest B lymphoid transcription factors, which links to changes in the *MLL/AF4* transcriptional program, ultimately establishing a myeloid differentiation fate. Unfortunately, we were not able to directly prove changes in transcription factor binding and associated changes in histone modifications because of the lack of available primary patient samples. However, high-resolution DNase-hypersensitive site-seq clearly demonstrated changes in chromatin accessibility and loss of occupation of the corresponding transcription factor binding sites.

The opposite scenario is observed when myeloid transcription factors are expressed in B-lymphoid cells.⁴⁵ Overexpression of *C/EBP α* efficiently reprograms such cells into macrophages by suppressing lymphoid genes. *CEBPA* is strongly upregulated after *CHD4* knockdown (Figure 6E) and is likely to be a driving force behind lineage switching. Taken together, these published and newly presented data confirm that (1) the balance between lymphoid and myeloid transcription factors is instructive for lineage choice, and (2) the downregulation of the myeloid program is essential for the maintenance of lymphoid fate.

How can the mutation of global chromatin regulators cause a switch in cell fate? Similar to the *Pax5* knockout, loss of IKAROS DNA-binding activity prevents lymphoid differentiation.²⁶ NuRD cooperates directly with IKAROS to repress hematopoietic stem cell self-renewal and myeloid differentiation, permitting early lymphoid development.^{26,46,47} Lineage switching was associated with heterozygous mutation; reduced expression; or, in the case of 2 MPALs, alternative splicing of *CHD4* and other NuRD components. These gene dosage effects are consistent with reports showing that complete loss of *CHD4* impairs normal and leukemic proliferation^{48,49} and myeloid and lymphoid differentiation of HSPCs and causes exhaustion of HSC pools,⁴⁶ indicating that basal *CHD4* expression is essential for maintaining AML. Moreover, a partial inhibition of *CHD4* supported induction of pluripotency in iPSCs, whereas a complete deletion eliminated cell proliferation, demonstrating that lowering *CHD4* expression may facilitate lineage promiscuity.⁵⁰

Recent studies have identified core NuRD and PRC1 complex members as being direct targets of *MLL/AF4* binding.^{51,52} Moreover, NuRD components including *CHD4* were shown to be part of an MLL supercomplex.⁵³ We therefore hypothesize that epigenetic regulator genes are recruited by lineage-specific factors during *MLL/AF4* leukemogenesis and mediate fundamental lineage-specific decision-making processes, in this case, the repression of the myeloid lineage program. Multiple routes to their dysregulation may result in escape from this lineage restriction and may be enacted at different stages of hematopoiesis. However, importantly and in keeping with a previous murine study of lineage conversion after CAR T-cell therapy, we did not identify evidence of relapse from a pre-existent myeloid clone.⁵⁴

Of substantial clinical importance, lineage switching results in the loss of B-cell surface markers (eg, CD19), providing an alternative mechanism for relapse after CAR T-cell or blinatumomab therapy,^{55,56} in addition to mutations, alternative splicing, and T-cell trogocytosis.⁵⁷⁻⁵⁹ Whereas these therapies target lineage-specific surface markers, lineage-switched (pre) leukemic progenitor populations escape epitope recognition and provide a potential clonal source for the relapse.⁶⁰ As recognition of

lineage switching after CD19 CAR T-cell therapy grows, 2 recent studies have highlighted the particular vulnerability of patients with *MLLr* ALL.^{54,61,62} Given the increasing use of advanced immunological therapies, a detailed understanding of the molecular processes underlying lineage determination and switching will be critical for developing new strategies to avoid this route to clinical relapse. In this report, we highlight an important role of epigenetic regulatory complexes in the context of *MLL/AF4* leukemia.

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Authorship

Contribution: O.H., S. Bomken, and C.B. conceived the study; O.H., C.B., R.T., K.S., P.M., S. Bomken, A.P., C.M., A.K., Z.K., J.B., V.B., R.M., J.V., J.M.A., and S.L. developed the methodology; S.N., J.Y.H.-K., V.V.G., A.K., D.W., and P.C. were responsible for the software programming; S.N., M.A., J.H.K., V.V.G., A.K., D.W., P.C., P.K., C.B., and O.H. performed the formal analysis; R.T., K.S., P.M., A.P., C.M., P.S.C., H.J.B., S.G.K., A.K., S.A.A., M.R.I., E.K.S., P.E., H.M., A.E., N.M.-S., S.E.F., Y.S., D.P., and P.C. performed the investigations; J.S., F.V., E.Z., A.S., J.C.M., L.J.R., C.E., O.A.H., S. Bailey, R.S., N.M., M.C., V.B., R.M., M.W., C.J.H., C.A.C., D.S., Y.O., M.J.T., P.N.C., J.C.M., M.H., C.B., and O.H. obtained the resources; S.N., D.W., and P.C. were responsible for data curation; S. Bomken, O.H., C.B., R.T., and K.S. wrote the manuscript; O.H., S. Bomken, J.M.A., J.V., and C.B. supervised the study; and O.H., J.V., S. Bomken, C.B., P.N.C., J.M.A., and E.Z. acquired the funding.

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Footnotes

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O.H. would like to dedicate this paper to Fritz Eckstein on the occasion of his 90th birthday.

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Requests for additional specific data/materials should be made to Olaf Heidenreich (o.t.heidenreich@prinsesmaximacentrum.nl).

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