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Accessibility
Depletion of RUNX1/ETO in t(8;21) AML cells leads to genome-wide changes in chromatin structure and transcription factor binding

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INTRODUCTION

Chromosomal rearrangements are a hallmark of hematopoietic malignancies. Many of these translocations generate aberrant transcriptional regulators that reproducibly lead to defined blocks in differentiation.1 A subclass of acute myeloid leukemia (AML) is associated with a group of chromosomal translocations that each result in disruption of the function of the core factor-binding (CBF) complex, which consists of a heterodimer of RUNX transcription factor family members and their binding partner CBFβ.2 The best-characterized CBF-type translocation is t(8;21), which fuses RUNX1 (AML1), a gene that is essential for normal hematopoiesis,3 with the gene encoding transcriptional co-repressor ETO (RUNX1T1 or MTG8).4,5 The ectopic expression of RUNX1/ETO redirects the specific gene expression program of normal precursor cells.6 The mechanism of such deregulation is based on RUNX1/ETO interfering with normal RUNX1 function.7,11 However, the molecular details of this interference are poorly understood on a genome-wide level. Several studies have demonstrated that RUNX1/ETO acts as a constitutive repressor by recruiting histone deacetylase complexes, and that it interferes directly with other transcriptional regulators of hematopoiesis such as C/EBPα.12-16 The presence of RUNX1/ETO can alternatively lead to gene activation.15-20

The t(8;21) translocation is a leukemia-initiating event, and fusion gene sequences can be found long before the onset of leukemia in the blood from newborn children.21 However, the induction of fully developed AML in t(8;21) patients requires secondary genetic alterations,22-25 which complicates the establishment of in vitro model systems for gain-of-function studies. In order to evaluate the suitability of RUNX1/ETO as a therapeutic target, it is therefore necessary to identify its target sites at the genome-wide level in actual leukemic cells, to define its individual role in reprogramming gene expression networks and to determine whether its continued presence is required for maintaining deregulation. As RUNX1/ETO and RUNX1 occupy the same binding sites, it is also important to elucidate how chromatin and RUNX1 occupancy respond to loss of RUNX1/ETO binding at its target genes. To this end, we determined genome-wide patterns of RUNX1/ETO occupancy in primary cells from patients and t(8;21) cell lines, and compared histone acetylation profiles, RNA-Polymerase II occupancy and RUNX1 binding and gene expression before and after siRNA-mediated depletion of RUNX1/ETO. Global loss of RUNX1/ETO binding results in widespread and complex changes in chromatin structure patterns, RUNX1 occupancy and

Keywords: acute myeloid leukemia; RUNX1/ETO; epigenetic regulation; chromatin; integrated analysis of high-throughput data

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The (t(8;21)) translocation fuses the DNA-binding domain of the hematopoietic master regulator RUNX1 to the ETO protein. The resultant RUNX1/ETO fusion protein is a leukemia-initiating transcription factor that interferes with RUNX1 function. The result of this interference is a block in differentiation and, finally, the development of acute myeloid leukemia (AML). To obtain insights into RUNX1/ETO-dependant alterations of the epigenetic landscape, we measured genome-wide RUNX1- and RUNX1/ETO-bound regions in t(8;21) cells and assessed to what extent the effects of RUNX1/ETO on the epigenome depend on its continued expression in established leukemic cells. To this end, we determined dynamic alterations of histone acetylation, RNA-Polymerase II binding and RUNX1 occupancy in the presence or absence of RUNX1/ETO using a knockdown approach. Combined global assessments of chromatin accessibility and kinetic gene expression data show that RUNX1/ETO controls the expression of important regulators of hematopoietic differentiation and self-renewal. We show that selective removal of RUNX1/ETO leads to a widespread reversal of epigenetic reprogramming and a genome-wide redistribution of RUNX1 binding, resulting in the inhibition of leukemic proliferation and self-renewal, and the induction of differentiation. This demonstrates that RUNX1/ETO represents a pivotal therapeutic target in AML.

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Identification of DNaseI- and ChiP-sequencing peaks

The raw sequence data were aligned to the hg18 assembly (NCBI Build 36.1) using BWA, and data were displayed using the UCSC Genome Browser. Regions of enrichment of DNaseI and ChiP data were identified using MACS software. De novo motif analysis was performed using HOMER.

Microarray gene expression data analysis

Microarray gene expression data were analyzed in GenomeStudio software (Illumina) with background subtraction. The raw data output was analyzed using the Lumi R package and quantile normalization. The 10% threshold (P-value < 0.1) was applied to all data. Genes were selected with at least twofold change in expression.

Gene set enrichment analysis (GSEA) was performed using the stand-alone application (Broad Institute) by ranking genes according to their correlation (Pearson) with the F1 metagene, the metagene highly expressed in the cells treated with siRNA for RUNX1/ETO. David/EASE analysis was performed using the online tool at http://david.abcc.ncifcrf.gov.

A more detailed description of Methods is given in Supplementary Material.

RESULTS

Identification of high-confidence binding sites for RUNX1/ETO and RUNX1 in t(8;21) cells

In order to perform a comprehensive and stringent determination of the chromatin and expression state of all genomic regions that respond to the expression of RUNX1/ETO, we employed Kasumi-1 cells, a well-studied model of t(8;21) AML. This cell line was obtained from a patient in second relapse after chemotherapy and bone marrow transplantation, and thus represents the most aggressive form of t(8;21) AML. Kasumi-1 does not express wild-type ETO and still carries one intact RUNX1 allele. This permits the discrimination between wild-type and translocated RUNX1 using antibodies against either the C terminus of RUNX1 recognizing only wild-type RUNX1, or the ETO moiety recognizing only the RUNX1/ETO fusion. We identified a genome-wide set of target sites for RUNX1 and RUNX1/ETO by ChiP followed by high-throughput sequencing (ChiP sequencing; Figure 1, Supplementary Figure 1). To determine how RUNX1/ETO target genes respond to loss of RUNX1/ETO binding and to evaluate the specificity of peak calling, we performed ChiP sequencing after specific RUNX1/ETO depletion using a siRNA that targets the RUNX1/ETO junction within the transcript as well as both a mismatch siRNA and mock transfection as controls. This knockdown model system has been extensively validated and characterized, and routinely leads to >70% reduction of all forms of the RUNX1/ETO protein (Figure 1a). A single siRNA electroporation yielded a maximal RUNX1/ETO transcript knockdown after 48 h. We therefore harvested siRNA-treated cells at this time point for ChiP experiments.

To identify high-confidence RUNX1/ETO targets regions, we first selected a set of 4598 ETO ChiP peaks that were shared between mock-transfected and mock-treated cells (Figure 1b). In all, 97% of these ETO ChiP peaks disappeared after the specific knockdown of RUNX1/ETO (Figure 1b, Supplementary Table 1). We have previously shown at specific genes that RUNX1/ETO targets regions, we first selected a set of 4598 ETO ChiP peaks that were shared between mock-transfected and mock-treated cells (Figure 1b). In all, 97% of these ETO ChiP peaks disappeared after the specific knockdown of RUNX1/ETO (Figure 1b, Supplementary Table 1). We have previously shown at specific genes that RUNX1/ETO targets regions, we first selected a set of 4598 ETO ChiP peaks that were shared between mock-transfected and mock-treated cells (Figure 1b). In all, 97% of these ETO ChiP peaks disappeared after the specific knockdown of RUNX1/ETO (Figure 1b, Supplementary Table 1). We have previously shown at specific genes that RUNX1/ETO targets regions, we first selected a set of 4598 ETO ChiP peaks that were shared between mock-transfected and mock-treated cells (Figure 1b). In all, 97% of these ETO ChiP peaks disappeared after the specific knockdown of RUNX1/ETO (Figure 1b, Supplementary Table 1).
High-confidence peaks were also found between the telomerase protein gene \textit{TERT} and \textit{CLPTM1L}, a locus associated with several types of cancer\textsuperscript{42} (Supplementary Figure 1A, Supplementary Table 4). Selected peaks were validated manually in Kasumi-1 cells (Supplementary Figures 1B and C) and in primary cells from a \textit{t(8;21)} AML patient (Supplementary Figures 1D–F), with and without \textit{RUNX1}/ETO knockdown.

To validate our genome-wide cell line ChIP-sequencing data, we compared the \textit{RUNX1}/ETO-binding pattern of Kasumi-1 cells to that of primary, patient-derived \textit{t(8;21)} AML cells. As with \textit{t(8;21)} cell lines, the wild-type ETO protein is neither expressed in \textit{t(8;21)}-negative nor in \textit{t(8;21)}-positive AML cells,\textsuperscript{36,43,44} which agrees with a lack of DHSs at the \textit{ETO} (\textit{RUNX1T1}) locus in all hematopoietic cell types studied here (data not shown). With primary cells from two patients (patients A and B), we obtained a large number of small peaks, many of which disappeared when analyzed at higher stringency (data not shown), but 2629 peaks occurred in both patients. In all, 76\% of joint peaks intersected with DHS from the two other \textit{t(8;21)} patients (no. 1 and no. 2, Supplementary Figure 2A).

\begin{figure}[ht]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Identification of high-confidence binding sites for \textit{RUNX1}/ETO and \textit{RUNX1}. (a) Time course of \textit{RUNX1}/ETO knockdown in Kasumi-1 cells. Top panel: real-time PCR analysis of mRNA expression; bottom panels: immunoblot detection of \textit{RUNX1}/ETO protein. \textit{\textbeta}-\textit{ACTIN} served as loading control. Time points are indicated at the bottom. \textit{RUNX1}/ETO transcript levels are recovering 48 h after siRNA electroporation, whereas protein levels remain low for another 24 h. siRE, \textit{RUNX1}/ETO siRNA; siMM, mismatch siRNA; control, mock-transfected cells. (b) Intersection analysis of \textit{RUNX1}/ETO peaks. The Venn diagram shows the overlap between \textit{RUNX1}/ETO peaks in mock, mismatch siRNA or \textit{RUNX1}/ETO siRNA-treated Kasumi-1 cells. The vast majority of \textit{RUNX1}/ETO peaks were common to mock and siMM-treated Kasumi-1 cells, and >97\% of the common peaks disappeared after \textit{RUNX1}/ETO depletion. (c, d) Identification of high-confidence peaks for \textit{RUNX1}/ETO and \textit{RUNX1} in Kasumi-1 cells. More than 75\% of both the \textit{RUNX1}/ETO and \textit{RUNX1} peaks colocalize with DHS in Kasumi-1 cells, thus constituting high-confidence peaks. (e) UCSC Genome Browser image depicting the human \textit{NFE2} and \textit{SPI1 (PU.1)} loci demonstrating a colocalization of \textit{RUNX1}/ETO peaks in patient cells, specific \textit{RUNX1}/ETO and \textit{RUNX1} peaks in Kasumi-1 cells as well as DHS.}
\end{figure}
1G, Supplementary Table 1) and thus represented high-confidence peaks. More than 50% of genes bound by RUNX1/ETO in Kasumi-1 cells were also specifically bound in patient cells (Supplementary Table 5). This high degree of concordance was also confirmed by comparing RUNX1/ETO binding to specific genes across the genome (Figure 1e, Supplementary Figure 1A), including known targets of RUNX1/ETO such as LAT2.45

RUNX1/ETO- and RUNX1-binding sites overlap only partially in t(8;21) cells. Both RUNX1 and RUNX1/ETO showed a similar pattern of distribution of binding sites for the subsets of peaks located within 1.5 kb of transcription start sites, and 60% of the RUNX1/ETO peaks overlapped with RUNX1 peaks (Figures 2a and b). The unbiased de novo identification of enriched consensus binding motifs showed that, in contrast to RUNX1-unique peaks, both RUNX1/ETO-associated and RUNX1/ETO-unique peaks were enriched in motifs for E-box-binding proteins (Figure 2c), in agreement with observations that these factors form stable interactions with the NHR1 domain of RUNX1/ETO.12 The same motifs were found when examining the entire set of RUNX1- or RUNX1/ETO-bound sequences in Kasumi-1 cells (Supplementary Figures 2A and B). Both RUNX1/ETO- and RUNX1-binding regions were enriched for RUNX1 consensus sequences, but only 30% of RUNX1-unique peaks contained such motifs, indicating that at

```
| motif   | match | score (log p-value) |
|---------|-------|---------------------|
| RUNX    | -1318 |                     |
| ETS     | -1158 |                     |
| ETS     | -1130 |                     |
| RUNX    | -567  |                     |
| ETS     | -437  |                     |
```

Figure 2. Identification and characterization of RUNX1/ETO and RUNX1 target regions in t(8;21) cells. (a) Positional distribution of RUNX1- (left) or RUNX1/ETO- (right) binding sites relative to the transcription start site (TSS) of their nearest gene. (b) Intersection of RUNX1 and RUNX1/ETO peaks in Kasumi-1 cells. The Venn diagram shows the numbers of high-confidence peaks bound by RUNX1 and RUNX1/ETO. (c) De novo motif discovery performed on the set of regions bound by the RUNX1 and/or RUNX1/ETO in Kasumi-1 cells identified enriched RUNX consensus and different types of ETS factor-binding sites. E-box motifs were significantly enriched in peaks either unique for RUNX1/ETO or common to RUNX1/ETO and RUNX1. (d) RUNX, ETS, E-box and C/EBP consensus sequence were mapped back to all regions bound by RUNX1/ETO and RUNX1. A large proportion of regions bound by RUNX1 did not contain RUNX (TGYGGT) and/or E-box (CANNTG) consensus binding motifs, whereas most regions contain ETS (GGAW) sites.
sites not occupied by RUNX1/ETO, RUNX1 may bind via interaction with other transcription factors. These are likely to be ETS family members, which present the top-scoring motif in this peak population (Figure 2c). In contrast, up to 70% of shared and unique RUNX1/ETO-bound sequences contained a RUNX1 motif (Figure 2d), suggesting that the fusion protein needs to

![Diagram](image1)

**Figure 3.** RUNX1 in t(8;21)-positive and -negative leukemic cells associates with different binding sites motifs. (a) Identification of high-confidence RUNX1 peaks from blasts from a AML patient with a normal karyotype (KN-AML). Venn diagram showing the intersection of RUNX1 peaks with DHS. (b) Positional distribution of RUNX1-binding sites in KN-AML blasts relative to the transcription start site (TSS) of their nearest gene. (c) Intersection of RUNX1 peaks in KN-AML cells and in Kasumi-1 cells. The Venn diagram shows the numbers of high-confidence peaks bound by RUNX1. (d) RUNX1 consensus sequences were mapped back to all regions bound by RUNX1 in t(8;21)-negative AML blasts. A large proportion of regions exclusively bound by RUNX1 contain RUNX, ETS and E-box consensus binding motifs as well as consensus binding motifs for C/EBP and AP1. (e) De novo motif discovery performed on the set of regions bound by RUNX1 in t(8;21)-negative KN-AML blasts identifies enriched RUNX and ETS consensus sequences as well as E-box, AP1- and C/EBP-binding sites.
Figure 4. Knockdown of RUNX1/ETO leads to a redistribution of RUNX1 binding. Kasumi-1 cells were electroporated with either mismatch control siRNA (siMM) or RUNX1/ETO siRNA (siRE). Two days after siRNA electroporation, RUNX1 binding was measured by ChIP sequencing in both populations. (a) Top panel: Venn diagram showing the appearance of new RUNX1-binding sites after RUNX1/ETO depletion. Bottom left: Venn diagram showing the overlap between RUNX1/ETO-bound regions and de novo RUNX1 sites, demonstrating that the latter are distinct from sites previously bound by RUNX1/ETO. Bottom right: Venn diagram showing the overlap between RUNX1/ETO-bound sequences and sites bound by RUNX1 before and after RUNX1/ETO depletion. (b) Example of alterations in RUNX1 binding. Left panel shows UCSC browser images depicting one gene (NFIL3) at a site not previously bound by RUNX1/ETO and another gene (FGR) showing a de novo RUNX1 with a RUNX1/ETO-bound site showing an increase in RUNX1 binding at this site after knockdown.

Figure 5. Analysis of RUNX1/ETO-dependent gene expression patterns. (a) Hierarchical clustering of genes responding by an at least twofold change in expression levels to RUNX1/ETO knockdown in Kasumi-1 cells. The heat map shows early upregulated (Group I), downregulated (Group II) and late upregulated genes (Group III) over a time course of 10 days. At the bottom of the heat map note non-clustered genes that are either upregulated more than threefold or show a more complex response pattern (Group IV). Expression levels were compared between RUNX1/ETO siRNA and mismatch siRNA-treated cells. Time points are indicated on the top of the heat map. Dark red indicates highly upregulated genes and black indicates highly downregulated genes. (b) Effect of RUNX1/ETO depletion on gene expression in primary t(8;21) AML blasts. The graph shows real-time PCR-based validation of early responding genes. The columns represent the means for three t(8;21) AML patients and the error bars the s.e.m. Inset: immunoblot showing siRNA-mediated depletion of RUNX1/ETO in blasts from a t(8;21) AML patient. (c) GSEA ranked according to the correlation of genes with a metagene (F1) summarizing the gene expression profile of Kasumi-1 cells after RUNX1/ETO knockdown. From left to right: significant enrichment of gene sets downregulated in human hematopoietic stem cells upon RUNX1/ETO overexpression and enrichment of genes determined in this study to have a high-confidence RUNX1/ETO-binding site with a corresponding DHS in the region 2 kb upstream of the start of transcription in Kasumi-1 cells. P, nominal P-value; q, false discovery rate. (d) The numbers of upregulated and downregulated genes upon RUNX1/ETO depletion in Kasumi-1 cells. The bottom row indicates genes with RUNX1/ETO peaks. (e) Classification of RUNX1/ETO target genes. The columns show the percentage of genes with high-confidence RUNX1/ETO peaks of all genes with changed gene expression upon RUNX1/ETO knockdown. siMM, mismatch siRNA; siRE, RUNX1/ETO siRNA.
be recruited by directly anchoring it to DNA. These results contrast to recently published findings by Maiques-Diaz et al.,\textsuperscript{46} who reported a substantially lower fraction of RUNX1 motifs in RUNX1/ETO peaks. This discrepancy may be due to the fact that in these experiments a ChIP-chip proximal promoter array was used.

\begin{table}
\centering
\begin{tabular}{|c|c|c|c|c|c|}
\hline
\textbf{Time (h)} & 48 & 96 & 168 & 240 \\
\hline
\textbf{Regulation} & up & down & up & down & up & down \\
\hline
\textbf{Total} & 90 & 68 & 174 & 106 & 293 & 256 & 656 & 741 \\
\hline
\textbf{Direct RUNX1/ETO targets} & 46 & 13 & 63 & 27 & 106 & 57 & 152 & 112 \\
\hline
\end{tabular}
\caption{Gene expression changes after knock-down.}
\end{table}
Differential binding of RUNX1 in AML cells without a CBF complex mutation

The incomplete overlap between RUNX1 and RUNX1/ETO peaks prompted us to investigate whether RUNX1 bound to different sequences in the human genome in RUNX1/ETO-expressing cells compared to AML cells without CBF mutations. To this end we identified genome-wide high-confidence RUNX1-binding sites in AML cells with a normal karyotype (KN-AML) from a patient that expressed a full-length RUNX1 protein (data not shown) but displayed a block at a similar differentiation stage as the t(8;21) patients and had a surface marker expression profile highly similar to the two t(8;21) AML samples (Supplementary Table 2). Similar to Kasumi-1 cells, 90% of all RUNX1 peaks in the KN-AML blast cells colocalized with DHSs (Figure 3a). Direct RUNX1 targets included genes controlling hematopoietic differentiation such as CSF1R, LAT2, and LMO2 (Supplementary Figure 3). The KN-AML RUNX1 peaks showed a similar pattern of distribution around the transcription start sites as the RUNX1 peaks from Kasumi-1 cells (Figure 3b), but their genome-wide distribution and enriched motif composition was strikingly different. When directly compared (Figure 3c), only 26% of the KN-AML RUNX1-binding sites were found in Kasumi-1 cells. GATA3, GCHFR and KLF13 are examples for this notion (Supplementary Figure 3). Most importantly, in contrast to t(8;21), cells, the RUNX1 peaks unique for KN-AML showed a highly significant enrichment for binding sites for inducible factors such as AP1, C/EBP and NF-κB (Figures 3d and e).

RUNX1/ETO knockdown leads to a shift in the pattern of RUNX1 occupancy

Our RUNX1 ChIP data indicate that the genome of hematopoietic precursor cells contains a large number of functional RUNX1-binding sites which are differentially occupied in t(8;21) AML and KN-AML cells and are associated with different binding site motifs. We therefore tested by ChIP sequencing whether depletion of RUNX1/ETO led to an alteration of RUNX1 occupancy in RUNX1/ETO expressing cells (Figure 4). The comparison between RUNX1 binding before and after knockdown showed that the depletion of RUNX1/ETO led not only to an increase in the number of RUNX1 peaks, but also to the formation of a large number of de novo RUNX1-binding sites (Figure 4a), as exemplified by the transcription factor gene NFI/IL3, which is upregulated by RUNX1/ETO depletion (Figure 4b, right panel). Only 10% of de novo peaks overlapped with RUNX1/ETO-bound sites (Figure 4a, bottom left). In contrast, a much larger proportion (34%) of peaks shared between knockdown and control samples were originally bound by RUNX1/ETO (Figure 4a, bottom right). Interestingly, although at most tested target regions depletion of RUNX1/ETO led to an upregulation of RUNX1 binding and gene expression (an example of this is shown with another upregulated gene, FGR, a member of the SRC kinase family, in Figure 4b, left panel), this was not always the case (Supplementary Figure 4). For instance, PU.1 (SPI1) represents a class of genes, where RUNX1/ETO binding did not or only marginally affect RUNX1 occupancy. The PU.1 upstream regulatory element contains an upstream enhancer element (PU.1E) that is bound by RUNX1 at multiple sites, and this binding is vital for enhancer activity,47 but it is also bound by RUNX1/ETO (this study). At this element RUNX1 binding did not respond to RUNX1/ETO knockdown, and gene expression was barely altered, which may indicate a complex binding pattern with multiple occupancies of both RUNX1/ETO and RUNX1.

RUNX1/ETO depletion can lead to both activation and repression of its direct target genes

In order to examine the effects of RUNX1/ETO depletion on gene expression at the global level and to correlate these changes with identified RUNX1/ETO target genes, we generated expression profiles during a time course of sustained RUNX1/ETO knockdown for 10 days in Kasumi-1 cells (Figure 5, Supplementary Table 6, Supplementary Figure 5A). Hierarchical clustering identified four groups of genes differentially responding to knockdown, consisting of early (Group I) and late (Group III) upregulated genes, downregulated genes (Group II) as well as genes showing a complex pattern of response (Group IV; Figure 5a). These results were validated by manual analysis with a selection of early responding genes both in Kasumi-1 cells and in primary t(8;21) cells obtained from three different AML patients (Figure 5b, Supplementary Figure 5B).

To compare global gene expression profiles, we produced two coordinate regulated metagenes F1 and F2 in an unsupervised manner that summarized the expression of RUNX1/ETO-dependent genes in a single score using non-negative matrix factorization.48,49 The F1 and F2 metagenes were highly expressed in Kasumi-1 cells with and without RUNX1/ETO knockdown, respectively. We also tested the relevance of the RUNX1/ETO-dependent global gene expression changes for another t(8;21)-positive cell line, SKNO-1, and for primary t(8;21)-positive AML. Kasumi-1-derived RUNX1/ETO-dependent metagene expression was mirrored and validated in the SKNO-1 and primary AML data sets, demonstrating an excellent concordance between the RUNX1/ETO-associated global gene expression changes in these three different t(8;21)-positive cell types (Supplementary Figure 5C). Furthermore, RUNX1/ETO-associated gene expression changes inversely correlated with gene expression changes observed in previous knockdown studies and therefore represent the primary human CD34+ cells (Figure 5c, Supplementary Figures 5D and E).50-53 These combined analyses confirmed that RUNX1/ETO-associated shifts in gene expression in Kasumi-1 faithfully reflect gene expression features in the human t(8;21) AML.

The metagene was also used to rank genes for the purpose of GSEA according to their correlation with the metagene score (Supplementary Table 7). Genes with high-confidence RUNX1/ETO peaks were highly enriched in the RUNX1/ETO knockdown signature (Figure 5c). However, not all genes responding to RUNX1/ETO knockdown were associated with RUNX1/ETO peaks. Genes responding early at day 2, such as CST7, IGFBP7 and MT5S1, consisted of 50% direct RUNX1/ETO target genes, whereas this was true for only 20% of the late responding genes at day 10 (Figures 5d and e). Moreover, almost 80% of all early responding target genes were upregulated upon RUNX1/ETO knockdown, in contrast to 60% of all late responding target genes (Figures 5d and e). This indicates that downstream effects of the removal of RUNX1/ETO contribute rapidly and progressively to the reorganization of the transcriptional network.

The depletion of RUNX1/ETO leads to complex changes in the histone acetylation pattern and RNA Polymerase II occupancy

As RUNX1/ETO recruits histone deacetylases to its target promoters13,14 and alters gene transcription, we examined the immediate consequences of RUNX1/ETO depletion on histone H3 lysine 9 acetylation (H3K9Ac) and occupancy of the elongating form of RNA Polymerase II (RNA Pol II). We therefore performed ChIP experiments with RUNX1/ETO-depleted and control cells at day 2 after knockdown, and integrated RNA Pol II and H3K9Ac data in Kasumi-1 cells on RUNX1/ETO target loci with the gene expression data (Figures 6a and b, Supplementary Table 6). After 2 days of knockdown, half of the early upregulated genes, including IGFBP7, NFE2 and PRG2, exhibited more than twofold increased RNA Pol II occupancy, whereas most downregulated genes such as CD34 and 75% of the late upregulated genes generally showed no or little change in RNA Pol II binding at this stage (Supplementary Figure 6A).

A more complex pattern of changes at RUNX1/ETO peaks was seen when measuring H3K9Ac. Although after 2 days of knockdown most upregulated genes displayed a rapid increase in H3K9Ac, we also found an increase at 50% of the downregulated
**Figure 6.** RUNX1/ETO silencing leads to changes in RNA-Polymerase II occupancy and the histone H3K9 acetylation pattern at RUNX1/ETO target genes. (a) Comparison of RNA Pol II occupation with transcriptional profiling reveals a substantial correlation between changes in gene expression and changes in RNA Pol II association upon RUNX1/ETO knockdown. RNA Pol II occupation was analyzed 48h after siRNA treatment, ranked according to fold change in occupancy and compared with changes in gene expression during a time course of 10 days with siRNA treatment. Dark red indicates high occupancy or upregulation, black low occupancy or downregulation, respectively. (b) The H3K9Ac pattern correlates partially with changes in gene expression associated with RUNX1/ETO depletion. H3K9Ac occupation was analyzed and compared analogously to (a). (c, d) UCSC Genome Browser image of the human ZBTB16 and WT1 loci depicting ChIP-Seq tags for RUNX1/ETO, RNA-Pol II and H3K9Ac in mock-treated (control), mismatch siRNA (siMM) and RUNX1/ETO siRNA (siRE)-treated Kasumi-1 cells. (e) Heat map resulting from an unsupervised clustering of H3K9Ac- and RUNX1/ETO-binding sites with and without RUNX1/ETO knockdown (siMM and siRE, respectively) as well as without transfection (control) showing two groups of sequences and their genomic location. In each lane, the ChIP enrichment score is shown 10 kb upstream and downstream from the peak center. (f) Integration of the sequence enrichment of each group into a density plot showing the location of RUNX1/ETO-binding sites with and without knockdown in relation to H3K9Ac. Dark blue: H3K9Ac (siMM); pink: H3K9Ac (siRE); green: RUNX1/ETO (control); yellow RUNX1/ETO (siMM); light blue: RUNX1/ETO (siRE).
genes (Figure 6b). This group of down-regulated genes partially overlapped with genes displaying an increase in RNA Pol II occupancy (Figure 6a, Supplementary Figures 6B and C). Such peaks were often localized in introns (for example, ZBTB16 or WT1, Figures 6c and d) indicating that not all RUNX1/ETO peaks that are associated with alterations of gene expression colocalize with classical promoter or enhancer elements. This diversity was also reflected in reporter gene assays of the promoters of target genes (Supplementary Figure 6D): Many, but not all, of these promoter elements directly responded to transactivation by RUNX1 and repression by RUNX1/ETO.

We next used our histone acetylation data to examine the chromatin architecture and the genomic distribution of RUNX1/ETO-binding sites (Figures 6e and f). Unsupervised clustering showed that RUNX1/ETO-binding sites could be clustered into two groups (Figure 6e). The largest group contained intergenic and intragenic sites that are most likely to be enhancers, as these elements are also DNase I hypersensitive. The analysis of the histone H3K9 acetylation profile before and after knockdown (Figure 6f) showed that in both groups the RUNX1/ETO complex was flanked by acetylated histones, demonstrating that histone acetylation was not absent in such elements. This is consistent with findings that described the presence of histone acetyl transferases at these sites. However, the depletion of RUNX1/ETO led to a strong increase in H3K9Ac, while maintaining a protein complex between nucleosomes, demonstrating that RUNX1/ETO does not lead to the formation of inactive chromatin, but shifts the balance between active and inactive chromatin.

RUNX1/ETO depletion induces myeloid differentiation and silences leukemic self-renewal programs

Finally, we asked which pathways and transcriptional programs responded to regulation by RUNX1/ETO. Functional annotation clustering separated RUNX1/ETO-bound genes from those exclusively bound by RUNX1. In both primary cells and cell lines, genes bound by the fusion protein are involved in processes, such as cytoskeletal organization, cell adhesion and cell cycle control, whereas genes solely bound by RUNX1 cluster with genes for nuclear structures, translation-associated processes and RNA processing (Supplementary Table 8). Genes responding to RUNX1/ETO knockdown showed a link with cytoskeleton, cell adhesion and migration, similar to those containing high-confidence RUNX1/ETO peaks (Supplementary Table 8). Moreover, separate analysis for upregulated and downregulated genes suggested an association of upregulated genes with differentiation, whereas downregulated genes are involved in proliferation and cell cycle regulation (Supplementary Table 9). These combined analyses suggest that the downregulated RUNX1/ETO genes are preferentially involved in proliferation and cell cycle progression, whereas upregulated genes are involved in myeloid differentiation. Indeed, suppression of RUNX1/ETO in t(8;21) AML
cell lines impairs clonogenicity and proliferation, and leads to cell cycle arrest in G1 phase.26

This notion is further supported by gene expression profile comparison and GSEA. Metagene-based comparison with global gene expression profiles from cells of different normal human hematopoietic differentiation stages26 demonstrate that Kasumi-1 cells depleted for RUNX1/ETO change their combined gene expression pattern from a unique specific subset remotely related to lymphoid and early myeloid precursors towards that of the normal granulocyte/monocyte precursor, monocyte and neutrophil subsets (Figure 7a).

GSEA showed that knockdown of RUNX1/ETO resulted in significant downregulation of cell cycle progression, telomere maintenance genes and TERT target genes (Figure 7b, Supplementary Figure 7A, Supplementary Table 7). In particular, the inverse correlation between RUNX1/ETO knockdown and TERT target gene expression supports a model of RUNX1/ETO-controlled telomerase activity, a key factor of normal and malignant self-renewal.37 In both t(8;21)-positive Kasumi-1 and patient-derived primary AML cells, RUNX1/ETO binds to a region between TERT and CLPTM1L implying TERT as a direct target gene of RUNX1/ETO (Supplementary Figure 1A). Indeed, sustained suppression of RUNX1/ETO caused an increasingly pronounced inhibition of TERT expression over time in Kasumi-1 and in SKNO-1 cells (Figure 7c, Supplementary Figure 7B). Interestingly, in spite of being highly activated particularly in MLL-rearranged leukemias, HOXB and late HOXA genes may not be part of the RUNX1/ETO-driven self-renewal program.51 In both t(8;21)-positive cell lines and AML blasts, several HOXA members including HOXA9 as well as the complete HOXB locus showed low or absent expression levels and a lack of DHSs (Supplementary Figure 7C).

**DISCUSSION**

Gain-of-function studies using ectopic expression of RUNX1/ETO in t(8;21)-negative leukemic and non-leukemic cells have yielded
insights into the molecular mechanisms of initiating leukemogenesis, whereas loss of function studies in actual leukemic t(8;21) cells (including the Kasumi-1 cells described here) inform about the role of RUNX1/ETO in maintaining leukemia. Our experiments and novel integrative data analyses demonstrate a substantial concordance between these two approaches, indicating that integral parts of the t(8;21)-specific leukemiation-initiating program are also required for maintaining the leukemic phenotype. In both primary cells and cell lines, RUNX1/ETO binds genes associated with the control of the cell cycle and cell structure. Concordantly, siRNA-mediated depletion of RUNX1/ETO affects transcriptional programs associated with myeloid differentiation, proliferation and self-renewal, in addition to those promoting cell cycle progression and DNA synthesis.

Our siRNA knockdown shows that RUNX1/ETO binding leads to large-scale, but reversible, alterations throughout the epigenome. H3K9Ac at RUNX1/ETO-binding sites was mostly increased after knockdown, which is consistent with a repressive role of the fusion protein at these sites. However, RUNX1/ETO depletion was also associated with upregulation of gene expression. In a recent study, RUNX1/ETO was shown to also regulate gene activation in a p300-dependent manner. Concurrent with this observation, when combined with gene expression data, RUNX1/ETO depletion concurred with strikingly complex regulatory patterns, with an increase of H3K9Ac and RNA Pol II binding alterations being associated with both upregulated and downregulated genes. An illustrative example for this notion is the CD34 locus (Supplementary Figure 6C, lower panel), which contains an intragenic RUNX1/ETO site and is downregulated by RUNX1/ETO depletion, but shows increased histone acetylation at the main promoter. At the WT1 locus, RUNX1/ETO binds to an element downstream of the main start site that contains a bidirectional promoter driving an alternative WT1 transcript and an inhibitory antisense intronic transcript. RUNX1/ETO may therefore maintain WT1 expression by repressing the expression of non-coding RNAs. This is of significant clinical interest because the level of WT1 transcription is a prognostic factor in leukemia diagnosis. Moreover, we also found widespread siRNA-mediated changes in H3K9Ac at RUNX1/ETO peaks that were not associated with alterations of steady-state mRNA levels. These elements may represent sites that respond to the upregulation of myeloid regulators such as C/EBPα and prime genes for the onset of myeloid differentiation. Follow-up of these observations is outside of the scope of this study, but our datasets provide a wealth of resources for experiments unravelling the mechanistic details of such changes.

Another important result from this study is our finding that the depletion of RUNX1/ETO and subsequent cell differentiation is associated with a redistribution of RUNX1-binding activity throughout the genome. We observed a large number of new binding sites distinct from those previously bound by RUNX1/ETO. The induction of myeloid differentiation after RUNX1/ETO depletion therefore involves not only loss of repression, but also increased recruitment of transcriptional activators to additional sites. Currently, we do not know how RUNX1 is directed to new sites, but it is likely that this involves the interaction of RUNX1 with other transcription factors whose activity is altered by RUNX1/ETO, such as C/EBPα and PU.1. Our observation of a differential binding of RUNX1 is also important in the context of leukemogenesis in general because it implies that mutations of RUNX1 that are widespread in leukemogenesis and cause specific disease phenotypes may differentially affect alternate subsets of genes depending on how the interaction with cooperating transcription factors is altered at each gene. Therefore, one of the future challenges in leukemia research will be to unravel the differential activity of transcription factors in a system-wide manner and to model the regulatory consequences of such differences for each specific type of leukemia.

In conclusion, our study demonstrates that epigenetic alterations mediated by RUNX1/ETO are reversible at a global scale by solely interfering with its function, emphasizing the feasibility of targeted therapeutic approaches either using siRNA or small molecules.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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