

Functional relevance of circRNA aberrant expression in pediatric acute leukemia with *KMT2A::AFF1* fusion

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Key Points

- New data on circRNAome dysregulation in *KMT2A::AFF1* ALL indicate aberrant transcripts from *KMT2A* signature genes and many new disease loci.
- Functional study disclosed the oncogenic role in *KMT2A::AFF1* ALL of circFKBP5, circKLHL2, circNR3C1, and circPAN3.

Circular RNAs (circRNAs) are emerging molecular players in leukemogenesis and promising therapeutic targets. In *KMT2A::AFF1* (*MLL::AF4*)-rearranged leukemia, an aggressive disease compared with other pediatric B-cell precursor (BCP) acute lymphoblastic leukemia (ALL), data about circRNAs are limited. Here, we disclose the circRNA landscape of infant patients with *KMT2A::AFF1* translocated BCP-ALL showing dysregulated, mostly ectopically expressed, circRNAs in leukemia cells. Most of these circRNAs, apart from circHIPK3 and circZNF609, previously associated with oncogenic behavior in ALL, are still uncharacterized. An in vitro loss-of-function screening identified an oncogenic role of circFKBP5, circKLHL2, circNR3C1, and circPAN3 in *KMT2A::AFF1* ALL, whose silencing affected cell proliferation and apoptosis. Further study in an extended cohort disclosed a significantly correlated expression of these oncogenic circRNAs and their putative involvement in common regulatory networks. Moreover, it showed that circAFF1 upregulation occurs in a subset of cases with *HOXA KMT2A::AFF1* ALL. Collectively, functional analyses and patient data reveal oncogenic circRNA upregulation as a relevant mechanism that sustains the malignant cell phenotype in *KMT2A::AFF1* ALL.

Introduction

Circular RNAs (circRNAs), transcripts in which by backsplicing the splice donor site is covalently bound to an upstream acceptor site, are key molecular players whose dysregulation and functions affect

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All the RNA-seq data of the study, including samples of leukemia cells of patients with B-cell precursor acute lymphoblastic leukemia harboring *MLL::AF4* rearrangements, patient-derived xenografts and the RS4;11 cell line, and CD34⁺ hematopoietic stem

cells (GSE213172) and CD19⁺ B cells of healthy donors (GSE110159), are deposited in the Gene Expression Omnibus database (accession number GSE213990).

The full-text version of this article contains a data supplement.

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almost all cancer hallmarks^{1,2} and attract high interest in studies aiming to discover new disease mechanisms and therapeutic targets.³⁻⁶

The prominent function of circRNAs is their ability to act as efficient microRNA (miRNA) sponges, regulating axes involving miRNAs.⁷⁻⁹ CircRNAs can interact with a variety of RNA-binding proteins¹⁰⁻¹² and even generate functional peptides¹³⁻¹⁶ not coded by linear transcripts.

CircRNAs are pervasively expressed in the hematopoietic compartment,^{1,17,18} and we and others have shown the dysregulation and oncogenic potential of circRNAs in acute leukemias.¹⁸⁻²²

In addition, recent discoveries have opened new perspectives for the circRNA role in leukemias with *KMT2A/MLL* rearrangements (*KMT2Are*).^{1,2,23,24} In leukemia with *KMT2A::MLL3* (*MLL::AF9*) translocation, oncogenic fusion circRNAs exhibited transforming activity linked to an enhancement of the oncogenic potential of the fusion protein.²⁵ Furthermore, preliminary targeted investigation of specific circRNAs in B-cell precursor acute lymphoblastic leukemia (BCP-ALL) with *KMT2Are*¹⁸ and of the expression dysregulation of circRNAs expressed by the translocation partner genes in *KMT2A::MLL1* ALL²⁶ provided evidence that *MLL* and several translocation partner genes express circRNAs in hematopoietic cells and that *KMT2Are*, beyond generating fusion circRNAs, can affect circRNA expression in general.²⁶

Infant *KMT2A::AFF1* leukemia is an aggressive disease with short event-free survival compared with other pediatric BCP-ALLs,²⁷ and its unique features, including extensive spread beyond the hematopoietic compartment, resistance to therapies, tendency to relapse, and myeloid switch,²⁸ motivated a thorough investigation of the biology underlying this malignancy ever since the first clinical report of a large set of patients carrying this translocation published in the early 80s.²⁹ The low frequency of somatic mutations in *KMT2A::AFF1*³⁰ indicated the fusion protein as the main disease driver, even if *KMT2A::AFF1*-induced leukemogenesis was difficult to model.³¹⁻³³ Several studies disclosed the impact of *KMT2A::AFF1* on gene expression, defining 2 different types of *KMT2A::AFF1* signatures^{34,35} and identifying the chimera direct targets.³⁶⁻³⁸ Less is known about the transcripts expressed by *KMT2A* fusion genes. In *KMT2A::AFF1* ALL, all whole transcriptome studies focused on linear transcripts only, and data about circRNA expression and roles in *KMT2A::AFF1* ALL are very limited. We recently detected fusion circRNAs in the RS4;11 cell line and in patients harboring *KMT2A::AFF1* translocation.³⁹ A circAFF1 isoform contributing to release miR-128-3p inhibition on *KMT2A::AFF1* protein expression has been implicated in leukemogenesis.⁴⁰

This study defining the *KMT2A::AFF1* ALL genomic landscape of circRNA expression was instrumental in identifying, through extensive functional investigation, multiple oncogenic circRNAs whose upregulation in malignant cells play an active role in the disease.

Materials and methods

Patients and samples

Leukemia cells from bone marrow samples at the diagnosis of 4 infant patients with BCP-ALL carrying *KMT2A::AFF1*re and 2 additional patient-derived xenograft samples were obtained (supplemental Methods).

CD34⁺ hematopoietic stem cells from cord blood samples of 3 healthy donors (HDs) and CD19⁺ healthy B cells/B lineage cells sorted using a fluorescence-activated cell sorter from peripheral blood mononuclear cells of 4 HDs¹⁸ were used as normal counterparts (GSE213990).

An enlarged cohort of pediatric patients with BCP-ALL at diagnosis, including cases with *KMT2A::AFF1* translocation (N = 45) and wild-type *KMT2A* (N = 13), was collected to assess the expression of circRNAs under investigation by real-time quantitative polymerase chain reaction (qRT-PCR), in comparison with bone marrow samples of HDs (N = 5). All samples were collected with informed consent, according to the Declaration of Helsinki.

RNA isolation and sequencing

RNA was isolated using TRIzol (Thermo Fisher Scientific, Waltham, MA) following the manufacturer's instructions. RNA quality was assessed with Agilent 2100 Bioanalyzer (Santa Clara, CA) keeping only samples with RNA integrity number > 7. RNA was quantified using the Qubit Assay (Thermo Fisher Scientific).

RNA libraries were prepared with the TruSeq Stranded Total RNA Ribo-Zero Gold kit and sequenced with an Illumina HiSeq2000 (San Diego, CA) at an average sequencing depth of 90 million reads per sample (paired-end reads 100-150 bp).

CircRNA detection and quantification from RNA-seq data

CircRNA identification and quantification and calculation of the circular-to-linear proportion (CLP) from RNA sequencing (RNA-seq) data were obtained with CirComPara version 0.6.3^{41,42} as described in supplemental Methods. Ensembl GRCh38 human genome and annotation version 93 was used for all analyses.

CircRNA expression was normalized with the regularized logarithm method and corrected using "sva."⁴³ Differential expression (DE) was assessed by DESeq2 (version 1.22.2)⁴⁴ with the parametric model, including Wald significance tests and no independent filtering, using an adjusted (Benjamini Hochberg) *P* value ≤ .05 as the significance threshold.

Cell lines

RS4;11 and SEM cell lines harboring *KMT2A::AFF1* (Deutsche Sammlung von Mikroorganismen und Zellkulturen, Braunschweig, Germany) were cultured in RPMI 1640 (Gibco, Life Technologies, CA) medium supplemented with 10% fetal bovine serum (Gibco), 1% L-glutamine and 1% penicillin/streptomycin and maintained at 500 000/mL concentration at 37°C in a 5% CO₂ incubator.

CircRNA silencing

Silencer select small interfering RNAs (siRNAs) were designed and synthesized by Thermo Fisher Scientific, specifically targeting the backsplice site of each circRNA, and tested to exclude off-targets among known human transcripts. Silencer Select Negative Control (Thermo Fisher Scientific) no.1 siRNA was used as the negative control (sirNEG).

RS4;11 and SEM cell lines were transfected using Amaxa 4D-Nucleofector instrument (Lonza, Davis, NC) and Mirus electroporation solution (Ingenio Electroporation Solution) with an optimized siRNA concentration of 30pMol (siRNA sequence are reported in supplemental Table 1).

SEM (EN-138 electroporation program) and RS4;11 (DS-130 program) cell lines were transfected using 16-well strips, with a cell (400 000 cells per well)-siRNA-Mirus final volume of 20 μ L. After electroporation, we seeded 1 000 000 cells per mL. At least 3 independent experiments were performed for all tests.

Cell viability, proliferation, and apoptosis assays

Cell lines were collected 24 hours and 48 hours after transfection and subjected to different *in vitro* functional assays to measure cell proliferation (5-ethynyl-2'-deoxyuridine incorporation) and apoptosis rate (annexin V-propidium iodide [PI]) as described in supplemental Methods.

RNA extraction, RNase R treatment, RT-PCR, and qRT-PCR

RNA was extracted from transfected cell lines using TRIzol (Thermo Fisher Scientific) and quantified using Qubit Assay (Thermo Fisher Scientific). For the validation of circRNAs presence in cell lines, RNA was treated with RNase R (Epicenter Biotechnologies, Madison, WI) to enhance circRNA abundance at a ratio of 2 U/ μ g. A total of 1 μ g of RNA was retrotranscribed using Random hexamers (Invitrogen and Thermo Fisher Scientific) and SuperScript II Reverse Transcriptase (Invitrogen).

To confirm the circRNA silencing efficiency, we performed the qRT-PCR using Platinum Sybr Green qPCR SuperMix-UDG (Invitrogen) carried out for 40 cycles and divergent primers to detect the backsplice site of each circRNA molecule (supplemental Table 2).

qRT-PCR with convergent primers (supplemental Table 3) was also performed to monitor the linear transcript expression. The $2^{(-\Delta\Delta Ct)}$ method was applied to obtain circRNA or linear transcript expression using sirNEG results as calibrators. *GAPDH* was used as a reference gene. Statistical analysis of experimental results was performed using Prism 8 (GraphPad Software Inc, La Jolla, CA).

CircRNA expression in patients was quantified using quantitative reverse transcription polymerase chain reaction (qRT-PCR) with divergent primers (supplemental Table 2), as previously described, without RNase R treatment. Bone marrow samples from HDs were used as a calibrator to calculate $2^{(-\Delta\Delta Ct)}$.

CircRNA functional predictions

The CRAFT tool⁴⁵ was applied to predict miRNA binding sites of circRNAs and to retrieve validated miRNA target genes (supplemental Methods). Functional enrichments have been calculated with the ClusterProfiler 4.0 package.

Gene expression profiling of patients harboring *KMT2A::AFF1*

Gene expression profiling data already available in the Laboratory of Onco-Hematology at University of Padova (GSE77416; pediatric *KMT2A::AFF1* ALL samples from GSE13204) were processed as described in supplemental Methods.

Results

CircRNA expression in patients harboring *KMT2A::AFF1*

The first aim of this study was to disclose the circRNAome of *KMT2A::AFF1* BCP-ALL. Analysis with CirComPara⁴² of

high-depth ribodepleted RNA-seq data in a discovery cohort of infant patients with *KMT2A::AFF1* (N = 4) identified 7596 circRNAs expressed from 3297 loci.

The 100 most expressed circRNAs account for about 18% of the expression in *KMT2A::AFF1* samples, whereas 1050 isoforms cover 50% of circRNA expression (supplemental Figure 1). Among the most expressed circRNAs in *KMT2A::AFF1* leukemia (Figure 1A), there are both circRNAs with well-characterized functions in other diseases, such as the oncogenic circHIPK3⁴⁶ and circZNF609,⁴⁷ and less-characterized molecules, such as circRNF220 and circNR3C1.^{48,49}

In line with previous reports,¹⁸⁻²⁰ about half of circRNA host genes expressed multiple circular isoforms. Eleven genes can generate at least 15 circRNAs expressed in the discovery sample set, with the highest number of isoforms detected for *UBAP2* (27 circRNAs), *ZCCHC7* (21), and *ANKRD36* (20) (supplemental Figure 2A). A single predominant circRNA more highly expressed than the other isoforms was observed for 5 of these genes, including *ANKRD36*, *PAN3*, and *AKT3*, whereas 3, such as *BIRC6* and *ANKRD36C*, presented 2 circRNAs with similarly high expression and the other circRNAs with medium to low expression (supplemental Figure 2B). The remaining genes presented multiple isoforms with comparably high expression.

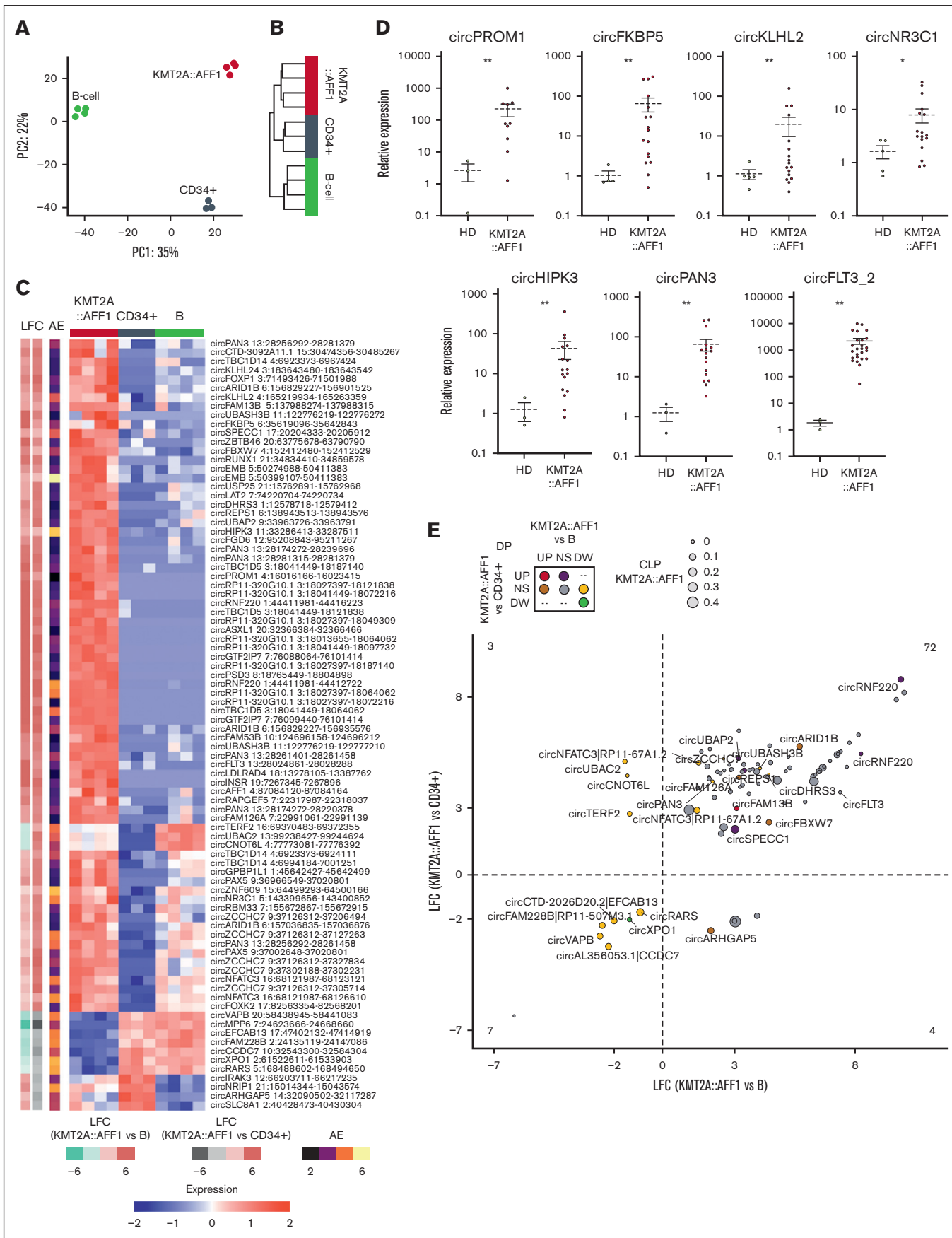
Beyond the absolute expression (AE) of circRNAs, we also considered the CLP, indicating the relative abundance of each circRNA with respect to the overlapping host gene transcripts. Figure 1B shows the group of 67 circRNAs with the highest CLP in *KMT2A::AFF1* leukemia and AE. CircGUSBP2 accounted for almost all the expression of its host gene, circC90orf84 (alias SHOC1) and circNBPF19 were more abundant than their linear counterparts (CLP > 0.5). CircZNF609, circSMARCA5, circMAN1A2, and circHIPK3 were among the circRNAs with the highest CLP and particularly high AE.

Dysregulation of circRNA absolute and relative expression level in *KMT2A::AFF1* leukemia

Next, to identify dysregulated circRNAs in *KMT2A::AFF1* possibly specific to leukemia, the circRNAome of *KMT2A::AFF1* leukemia was compared with CD34⁺ hematopoietic stem cells and B cells from HDs. For this comparison, 8607 expressed circRNAs, derived from 3602 loci, were used.

Unsupervised analyses of circRNA expression by principal component analysis (Figure 2A) and hierarchical clustering (Figure 2B) clearly separated *KMT2A::AFF1* ALL samples from the 2 healthy cell populations as well as CD34⁺ cells from B cells. In terms of circRNAome, *KMT2A::AFF1* ALL is closer to CD34⁺ cells than to B cells. In line, DE analysis identified 206 circRNAs that significantly varied (109 upregulated and 97 downregulated) in *KMT2A::AFF1* ALL when compared with CD34⁺ samples and 1116 (714 upregulated and 402 downregulated) when compared with B cells.

Of particular interest, a group of 86 circRNAs was DE in *KMT2A::AFF1* ALL compared with both healthy cell populations (Figure 2C; supplemental Table 4). Seven of these circRNAs were expressed in leukemia cells at an intermediate level compared with the 2 control populations: circTERF2, circUBAC2, and circCNOT6L being more expressed in cells from patients with



circRNA upregulation was prevalently concordant with the increase in linear counterpart expression.

Are dysregulated circRNAs expressed by known *KMT2A* signature genes?

Functional enrichment showed that host genes of the circRNAs dysregulated in *KMT2A::AFF1* ALL samples in comparison with both normal populations are significantly enriched in leukemia-associated genes (DISGenet signatures of acute myeloid leukemia with ryeodysplasia-related changes: *ASXL1*, *FLT3*, and *RUNX1*; primary myelofibrosis: *ASXL1*, *FLT3*, *RARS*, *NR3C1*, *RUNX1*, and *FKBP5*; and precursor B-cell lymphoblastic leukemia-lymphoma: *FLT3*, *INSR*, *PAX5*, *NR3C1*, and *RUNX1*).

Next, we assessed if these leukemia-associated genes are part of the known *KMT2A::AFF1* signature defined by the linear transcriptome studies based on microarrays and RNA-seq or DNA-binding by CHIP-seq data.

We collected 15 human or murine⁵⁰⁻⁵⁵ signatures from public databases of genes upregulated in association with *KMT2A* rearrangements in general or specifically with the *KMT2A::AFF1* fusion, and assembled a *KMT2A*/*KMT2A::AFF1* meta-signature of 1881 genes (supplemental Table 6). Custom gene set enrichment analysis (GSEA) using this meta-signature demonstrated a significant enrichment of genes with circRNAs upregulated in *KMT2A::AFF1* ALL (Figure 3A). In fact, 121 (6.4%) signature genes expressed 1 or more circRNAs DE in *KMT2A::AFF1* ALL, most of them being upregulated in 1 or both comparisons (supplemental Figure 4).

Up to 6 (PAN3) upregulated circRNAs were displayed by 16 host genes previously included in at least 1 signature (Figure 3B). Most DE circRNAs we detected were not expressed by genes previously associated with *KMT2A* rearrangements and represent novel findings of loci potentially associated with this disease.

More than 80% of the signature gene circRNAs had a poorly altered CLP in *KMT2A::AFF1* ALL, including circHIPK3, circNR3C1, circRNF220, and circTBC1D14, with AE markedly increased in leukemia cells, whose previously reported gene upregulation could be linked to the circRNA overexpression.

CircAFF1 upregulation is linked to HOXA *KMT2A::AFF1* ALL

Using RNA-seq analysis in the discovery cohort, we detected 21 circular isoforms of the *AFF1* gene (Figure 4A). The most expressed circRNA formed by backsplicing of exons 3 and 4 (4:87046166-87047594, named circAFF1_A), previously shown

to be upregulated in *KMT2A::AFF1* ALL,⁴⁰ was not significantly increased in leukemia cells according to our RNA-seq data. Instead, we detected a significant upregulation of a circular isoform formed by backsplicing of exon 5 with 7 (4:87084120-87091829, circAFF1_B) and a very short one (circularized exon 5; 4:87084120-87084164, circAFF1_C). Further quantification by qRT-PCR in an extended pediatric patient cohort (32 *KMT2A::AFF1* ALL and 12 BCP-LL non-*KMT2A*-rearranged and bone marrow samples from 5 HDs) showed a higher expression of circAFF1_B in *KMT2A::AFF1* ALL than in non-*KMT2A* BCP-ALL, however, without a dramatic increase compared with the normal counterpart (Figure 4B). The average expression of circAFF1_A was significantly higher in *KMT2A::AFF1* ALL than in controls and non-*KMT2A* cases (Figure 4B). Notably, in the *KMT2A::AFF1* ALL cohort, circAFF1_A expression was heterogeneous (Figure 4C), and stratifying the cases based on circAFF1_A expression, we detected a different expression profile in the 2 groups. The genes more upregulated in cases with high circAFF1_A expression (greater than the median; Figure 4D) were significantly enriched, according to GSEA based on C2CGP MSigDB, in genes characterizing the *KMT2A* signatures, including *HOX* cluster genes (Figure 4E). On the contrary, patients with less circAFF1_A expression showed an upregulation of *IRX1* and *IRX2* genes (Figure 4D).

Collectively, these data indicate that circAFF1_A expression is associated with the previously identified gene expression *KMT2A* subgroups (ie, HOXA and IRX subtype).

Functional screening identified the oncogenic role of circFKBP5, circKLHL2, circNR3C1, and circPAN3 in *KMT2A::AFF1* ALL in vitro

Next, we conducted an extensive functional screening using a loss-of-function approach in vitro to identify oncogenic circRNAs in *KMT2A::AFF1* cells (Figure 5A). We prioritized 27 circRNAs to be functionally studied, for which high expression in *KMT2A::AFF1* ALL was confirmed by additional RNA-seq data of patient-derived xenograft samples and the RS4;11 cell line (supplemental Figure 5), including the 7 (circPROM1, circFLT3, circNR3C1, circPAN3, circKLHL2, circFKBP5, and circHIPK3) previously validated in the extended cohort (Figure 2).

We designed a custom siRNA panel, including 2 siRNAs targeting each of the selected circRNA molecules, specific for the backsplicing region and without any predicted human off-target transcripts, plus sirNEG siRNAs (supplemental Table 1). Thirteen circRNAs were silenced with an efficiency higher than 75% in both RS4;11 and SEM cell lines 24 hours after transfection by at least 1 of the siRNAs tested (Figure 5B).

Figure 2. The *KMT2A::AFF1* circRNAome is highly dysregulated. (A) Principal component (PC) analysis and (B) hierarchical clustering using Euclidean distance of samples from patients with *KMT2A::AFF1* ALL and CD34⁺ cells and B-cell populations from HDs, based on circRNA expression profiles. (C) Heat map of standardized expression of the 86 circRNAs differentially expressed when comparing *KMT2A::AFF1* ALL vs B cells and CD34⁺ cells from HDs; (left) the log₂ fold change (LFC) in the 2 comparisons and the average AE in all samples are shown. (D) CircRNA expression quantification by RQ-PCR in an extended patient cohort (relative expression calculated as 2^{-ΔΔCT} values using glyceraldehyde-3-phosphate dehydrogenase as reference and bone marrow samples from at least 3 HDs as calibrator; Mann-Whitney *U* test *P* values: **P* < .05 and ** *P* < .01); N = 10 MLL:AF4 BCP-ALL for circPROM1; N = 17 *KMT2A::AFF1* BCP-ALL for circFKBP5, circKLHL2, circNR3C1, circHIPK3, and circPAN3; and N = 27 MLL:AF4 BCP-ALL for circFLT3_2. (E) Relation between dysregulation of circRNA expression and proportion in *KMT2A::AFF1* ALL for the same 86 DE circRNAs in panel C. The dot size represents the CLP value in patients with *KMT2A::AFF1*, the dot color indicates DP variation categories in the 2 comparisons; names of circRNAs DP in at least 1 comparison (*KMT2A::AFF1* vs B cells and/or *KMT2A::AFF1* vs CD34⁺ cells) are shown, and the number of circRNAs in each quadrant is indicated. UP, upregulated; NS, unvaried; DW, decreased.

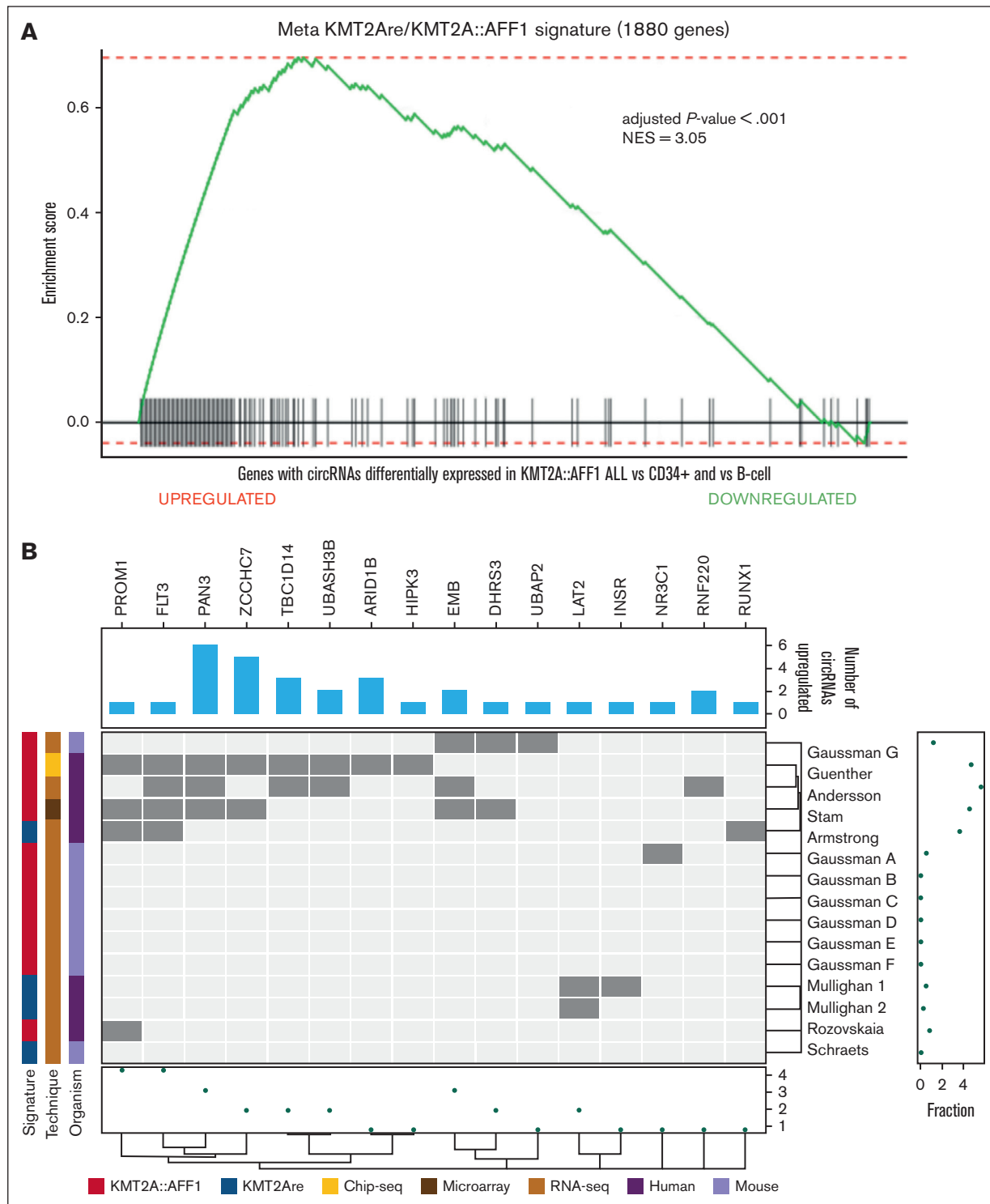


Figure 3. Relation between circRNAs dysregulated in *KMT2A::AFF1* ALL and known *KMT2Are* signature genes. (A) Custom GSEA of the host genes expressing the 86 circRNAs most dysregulated in *KMT2A::AFF1* using custom assembled *KMT2Are/KMT2A::AFF1* meta-signature of 1881 genes previously associated with the disease. (B) Bar plot of the number of circRNAs dysregulated in *KMT2A::AFF1* expressed from genes belonging to the *KMT2Are/KMT2A::AFF1* meta-signature. The heat map shows the genes belonging to each original signature (dark gray); the columns are clustered according to the Dice distance, and the rows according to the Sokal and Michener distance; the marginal dot plots display the number signatures that contain each gene (bottom) and the fraction of genes of each signature that host at least 1 circRNA (right); information about the original signatures considered is given by the left bars. NES, normalized enrichment score.

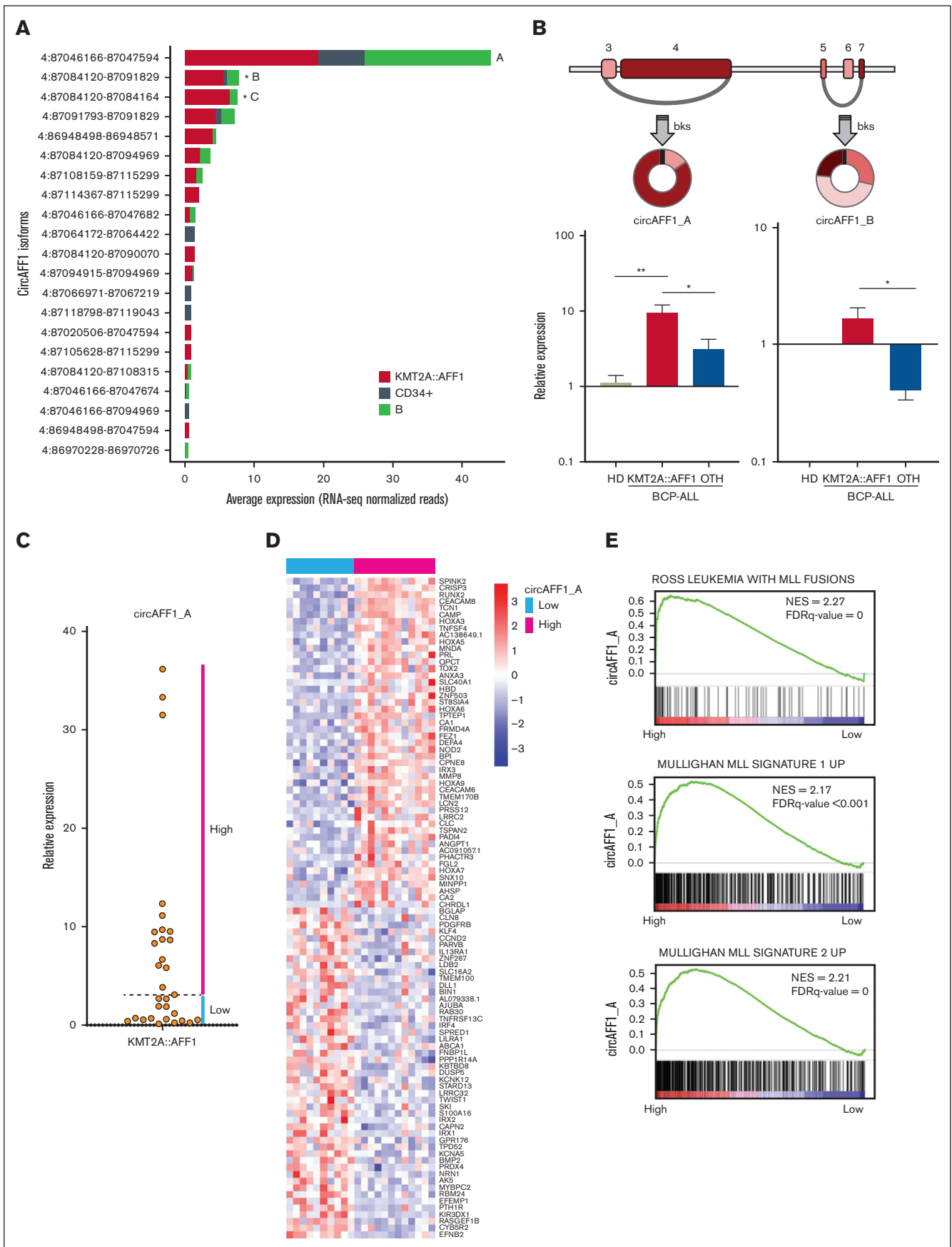


Figure 4.

A first functional test revealed an increased percentage of apoptotic cells after silencing of circFKBP5, circKLHL2, circNR3C1, circPAN3, and circPROM1 (Figure 5B). Silencing of circFKBP5, circKLHL2, circNR3C1, and circPAN3 (Figure 5C, top) did not alter the expression of the linear counterpart (Figure 5C, bottom). Further in vitro experiments were performed considering these 4 circRNAs, using, for each of them, the siRNA with the best silencing performance.

For all the 4 tested circRNAs, the annexin-PI assay from at least 3 independent replicates confirmed an increased ratio of apoptotic cells upon in vitro silencing in *KMT2A::AFF1* specimens with respect to sirNEGs, both 24 hours and 48 hours (Figure 5D). The effect was particularly marked for circFKBP5 and circKLHL2. Moreover, the EdU-incorporation assay highlighted a tendency toward reduced cell proliferation after knockdown of these 4 circRNAs, with a reduction of $\leq 50\%$ compared with that in the control (Figure 5E). A high effect was observed upon circKLHL2 silencing, with a significant reduction at both time points (24 hours $P = .0032$ and 48 hours $P = .0003$). Results supported an oncogenic role of circFKBP5, circKLHL2, circNR3C1, and circPAN3 ectopically expressed in *KMT2A::AFF1* ALL.

Next, the high correlation between the expression profiles of these 4 oncogenic circRNAs in the enlarged cohort of patients with *KMT2A::AFF1* at diagnosis (Figure 6A; supplemental Figure 6) motivated a further investigation of their possible overlapping networks of action, potentially involving common miRNAs and target genes. For each of the 4 circRNAs, binding sites were predicted for 1 up to 14 miRNAs, for a total of 21 miRNAs (Figure 6B). CircNR3C1 shared binding sites for miR-370-3p with circKLHL2 and for miR-3198 with circPAN3. Despite the limited number of predicted binding miRNAs shared between the 4 circRNAs, we found 205 validated miRNA target genes potentially commonly regulated by at least 3 of these 4 circRNAs, mainly through different miRNAs (Figure 6B-C). Upon GSEA, these genes were significantly (false discovery rate < 0.05) associated with histone modification and methylation, key biological processes affected by the *KMT2A::AFF1* chimera complex (Figure 6D) and the KMT2A wild-type protein (Figure 6E; supplemental Table 7). According to *KMT2A::AFF1* CHIP-seq data available for SEM and RS4;11 cell lines,⁵⁶ only 14 of the genes targeted by miRNAs potentially sequestered by the circRNAs under consideration are direct *KMT2A::AFF1* chimera targets (Figure 6E), including *ARID5B* and *RUNX1*, which are strongly associated with this leukemia. Overall, these data suggest that the identified ectopically expressed oncogenic circRNAs could operate as a reinforcement of the *KMT2A::AFF1* chimera activity, although prevalently by regulatory action on genes not under the direct control of the fusion protein.

Discussion

In this study, we described for the first time, to our knowledge, the circRNAome of pediatric *KMT2A::AFF1* translocated leukemia, bringing to light new players that sustain the molecular mechanisms contributing to leukemogenesis.

Through deep analysis of RNA-seq data with cutting-edge bioinformatic methods, we observed that the circRNA landscape is dysregulated in patients with *KMT2A::AFF1* ALL, with circRNAs aberrantly expressed in malignant cells compared with the normal counterpart. Most of the dysregulated circRNAs were ectopically expressed in patients, thus being candidate oncogenic molecules.

Interestingly, several circRNAs are aberrantly overexpressed and derived from host genes associated with *KMT2A::AFF1* gene signatures, including *PROM1*, *FLT3*, *PAN3*, *ZCCHC7*, *TBC1D14*, *HIPK3*, and *RNF220*.^{57,58} Nevertheless, most of the altered circRNAs we detected did not derive from genes previously associated with *KMT2A*, in general, or with the *KMT2A::AFF1* fusions, in particular, and indicated several new loci, including *FKBP5* and *KLHL2*, with a potential role in this disease. An orthogonal analysis of the CLP in *KMT2A::AFF1* ALL showed that most (80%) of the circRNAs overexpressed in leukemia cells followed the expression profiles of their linear counterparts, whereas we also found genes in which the circRNAs accounted for a significantly increased expression proportion in *KMT2A::AFF1* than in healthy cells. Collectively, these observations support the added value of the study of circRNA expression and potential associations to the molecular mechanism of this disease. The observed excess of upregulated circRNAs in leukemia cells can be at least partly explained by the activation of transcriptional programs in *KMT2A::AFF1* ALL because several of them are expressed from known chimera target genes. Nevertheless, the CLP increase in malignant cells indicates that disturbances of posttranscriptional processes, including RNA splicing dysregulation, can play a role.

Several circular isoforms of the *AFF1* gene were expressed in the samples, including circAFF1_A previously shown to be upregulated in *KMT2A::AFF1* ALL.⁴⁰ Here, we disclosed that patients can be classified according to the expression level of circAFF1_A and that the cases with the highest expression of this circRNA present a specific gene expression profile, with a marked upregulation of *KMT2A* signature genes, particularly *HOX* cluster genes. A low circAFF1_A expression was instead associated with upregulation of the *IRX1* and *IRX2* genes. Of novelty, our examination of a sizable cohort disclosed that the features associated with high expression of circAFF1_A are not a general characteristic of *KMT2A::AFF1* ALL but specific to a patient subgroup. Several studies have reported that patients with t(4;11) can be grouped based on *HOXA* gene expression and that the missing expression

Figure 4. CircAFF1_A and circAFF1_B isoform structure and expression in the extended patient cohort. (A) Expression levels of the circAFF1 isoforms in samples from patients with *KMT2A::AFF1* ALL and CD34⁺ cells and B cells populations from HDs, according to RNA-seq analysis (significant DE is indicated by *). (B) Schematic representation of circAFF1_A and circAFF1_B structure (exons number according to ENST00000395146 transcript; colored boxes represent exons; gray segments, introns; and arches, circRNAs;) and bar graphs of expression levels quantified by qRT-PCR in bone marrow samples from HDs (N = 3), in pediatric patients with BCP-ALL, 31 with (*KMT2A::AFF1*, N = 31 for circAFF1_A and N = 17 for circAFF1_B) and 13 without (OTH) *KMT2A::AFF1* rearrangement (mean \pm standard error of the mean is shown; Mann-Whitney U test P value * $P < .05$ and ** $P < .01$). (C) circAFF1_A level stratify patients into high- and low-expressors groups. (D) Patients with high and low circAFF1_A expression level are characterized by different genes expression profiles according to microarrays analysis (the top 50 ranked genes differentially expressed are shown in the heat map); (E) *MLL* signatures resulted positively enriched among genes differentially expressed between patients with high and low circAFF1_A expression. bks, backsplice site; FDR, false discovery rate.

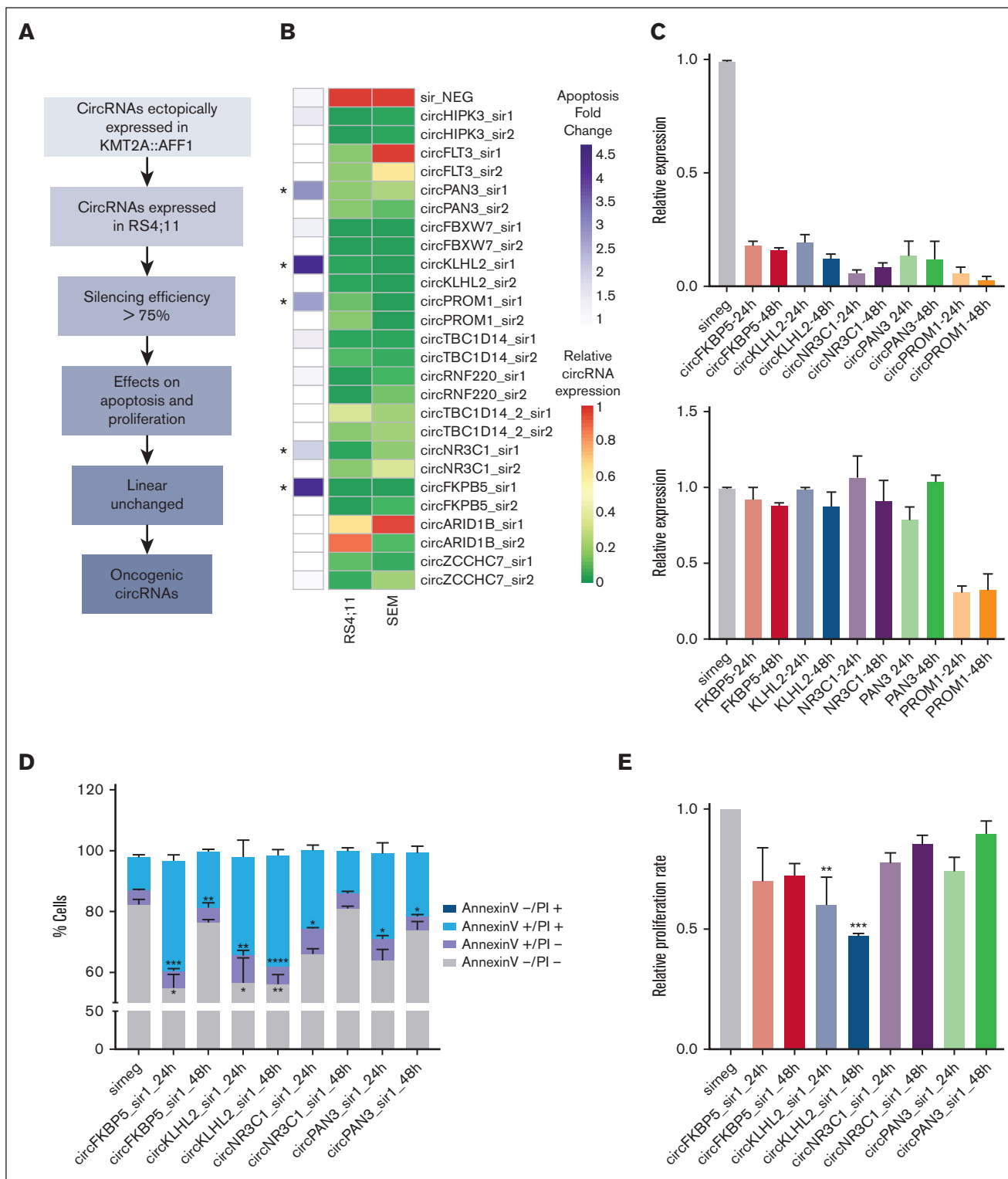


Figure 5. Functional screening of circRNAs using a loss-of-function approach in vitro. (A) Workflow of the strategy used for circRNA prioritization and functional screening. (B) Results of the semimassive siRNA screening: the heat map on the left shows the average relative apoptotic rate with respect to sirNEG (measured as relative percentage of annexin and propidium iodide (PI) marked, ie, the sum of annexin-positive/PI-positive and annexin-positive/PI-negative; *indicates LFC > 1.5) upon silencing of 13 circRNAs with high siRNA efficiency (right columns; at least 75% of circRNA expression reduction in SEM and RS4;11 cell lines; relative circRNA expression measured by qRT-PCR and expressed as $2^{-\Delta\Delta Ct}$ using sir_NEG as calibrator and *GAPDH* as reference gene). (C) Relative circular RNA (top) and linear transcript (bottom) expression in the silenced samples compared to the sirNEG (gray bar). (D) Percentage of annexin-PI marked cells and (E) relative cell proliferation rate according to EdU assay upon circRNA silencing at 24 hours and 48 hours. For each bar, data are reported as the mean \pm standard error of the mean of at least 3 independent replicates in RS4;11.

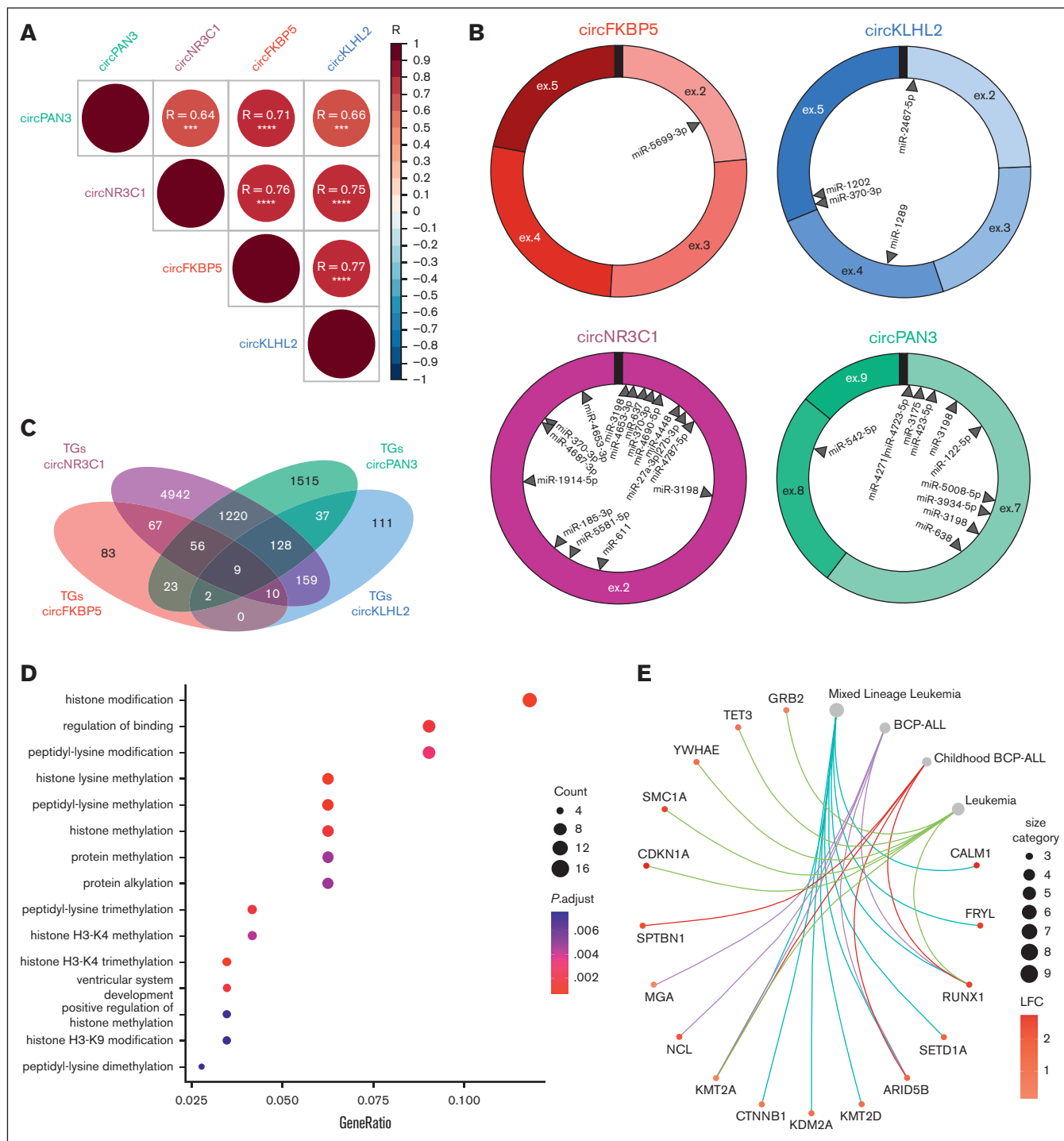


Figure 6. Correlated expression and putative functions of oncogenic circFKBP5, circKLHL2, circNR3C1, and circPAN3 ectopically expressed in *KMT2A::AFF1* ALL. CircFKBP5, circKLHL2, circNR3C1, and circPAN3 (A) pairwise expression correlation in a cohort of 17 pediatric patients with *KMT2A::AFF1* ALL; (B) miRNA-binding sites predicted by CRAFT; (C) venn diagram of the overlap of validated target genes (TGs) of miRNAs potentially bound by circFKBP5, circKLHL2, circNR3C1 and circPAN3; overrepresentation of (D) biological processes; and (E) diseases among genes commonly linked, through miRNAs, to at least 3 of the 4 oncogenic circRNAs identified and with positive LFC comparisons of *KMT2A::AFF1* ALL samples with CD34⁺ cells from HDs.

of these genes is correlated with high *IRX1* or *IRX2*.^{34,59} Indeed, we demonstrated that high expression of circAFF1 is associated with a high *HOXA* expression, *AFF1* overexpression, and a

KMT2A-leukemia related signature, emphasizing the explanation of its role to regulate *KMT2A::AFF1* expression.⁴⁰ Differently, in patients with activation of the IRX homeobox protein, either due to

the absence of *AFF1::KMT2A* expression or to its repression, circ*AFF1*-mediated regulation can be lacking.

We did not see upregulation in *KMT2A::AFF1* ALL of the circRNAs expressed from *KMT2A*, in line with our previous data in *KMT2A::MLLT1* (*MLL::ENL*) cases²⁶ and the recent interesting finding that a circ*KMT2A* isoform overexpressed before disease onset plays a role in the acquisition of *KMT2A*.⁶⁰

Among the circRNAs most aberrantly expressed in *KMT2A::AFF1* ALL, circHIPK3⁶¹⁻⁶⁴ and circZNF609^{47,65} had been previously described as being deregulated in other tumors and developmental processes. Regarding pediatric leukemias, we showed the upregulation of circHIPK3 in different subtypes of BCP-ALL¹⁸ and a possible oncogenic role of circZNF609 in T-cell ALL.²⁰ Other circRNAs upregulated in *KMT2A::AFF1* ALL are lacking functional information to our knowledge, particularly circFKBP5. CircKLHL2 was shown to be upregulated in multiple myeloma.⁶⁶ CircNR3C1 involvement in tumorigenic processes was described in different contexts but not in leukemias. An oncogenic role of circPAN3 in acute myeloid leukemia has been reported, linking its overexpression to doxorubicin resistance.⁹ Beyond circPAN3, it is worth noting that other circRNAs previously associated with acute myeloid leukemia,⁶⁷ such as circFBXW7²¹ and circRNF220,⁶⁸ are overexpressed in *KMT2A::AFF1* ALL.

Because most of the identified upregulated circRNAs were not previously characterized, we set up a functional screening using specific circRNA silencing, through which we identified new circRNAs with oncogenic potential in *KMT2A::AFF1* ALL in vitro. Starting with 27 circRNAs, for about half of them, we obtained an efficient silencing in 2 cell lines. The specific silencing of circFKBP5, circKLHL2, circNR3C1, and circPAN3 had an effect on the cell apoptosis rate. More in-depth functional investigation provided robust data about the oncogenic roles of these circRNAs in *KMT2A::AFF1* ALL in vitro. The silencing of each of these 4 circRNA, particularly circFKBP5, circKLHL2, significantly increased the ratio of apoptotic cells, indicating that their overexpression sustains the survival of *KMT2A::AFF1* ALL cells. In addition, reduced cell proliferation upon the silencing of these circRNAs was observed, with the highest effect linked to circKLHL2 silencing. Overall, these results support an oncogenic role of each of these 4 circRNAs in *KMT2A::AFF1* ALL, which is the most important highlight of our study.

In our data, several circRNAs were altered in malignant cells, and at least 4 of them were proven to have oncogenic properties by functional investigation. In our view, the attribution of leukemogenic potential to single circRNA molecules can be overly simplistic and somewhat fail to account for the pleiotropic roles of these molecules and for the possible interplay of circRNAs concurrently

dysregulated in malignant cells. Along this line and because of the observation of a significant correlation between oncogenic circRNA expression in the extended cohort of *KMT2A::AFF1* ALL cases, we explored the possible regulatory networks controlled by the oncogenic circFKBP5, circKLHL2, circNR3C1, and circPAN3 in this malignancy. Integrative analysis of the miRNA binding potential of these circRNAs and of the expression of validated target miRNA genes disclosed a set of genes for which upregulation could be sustained by circRNA overexpression. These genes showed a notable enrichment of leukemia-associated genes, most of which are not under direct control of the *KMT2A::AFF1* chimera complex. These observations, together with the results of our functional studies in vitro, suggest oncogenic circRNA upregulation as an additional mechanism that sustains the malignant cell phenotype in *KMT2A::AFF1* ALL. In the future, more in-depth functional studies, including in vivo experiments, will be needed to provide a clearer vision of the leukemogenic potential of circRNAs in this neoplasm. Nevertheless, our data are laying new bases for the development of innovative RNA-based therapies.

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Authorship

Contribution: S. Bortoluzzi, S. Bresolin, and G.t.K. conceived the study; S. Bortoluzzi and S. Bresolin supervised the study development; E.B., P.G., A.M.V., G.C., A.B., F.L., B.B., and L.H.M. provided patient and control samples; C.T.P., G.L., A.C. and V.S. performed experiments; A.D.M., S. Bresolin, and S. Bortoluzzi did bioinformatic analysis, E.G. and A.B. provided bioinformatics methods. C.T.P., A.D.M., S. Bresolin and S. Bortoluzzi prepared tables and figures; A.D.M., S. Bresolin and S. Bortoluzzi wrote the manuscript; and all authors revised and approved the manuscript.

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