The t(4;11)(q21;q23) translocation is associated with high-risk infant pro-B-cell acute lymphoblastic leukemia and arises prenatally during embryonic/fetal hematopoiesis. The developmental/pathogenic contribution of the t(4;11)-resulting MLL-AF4 (MA4) and AF4-MLL (A4M) fusions remains unclear; MA4 is always expressed in patients with t(4;11)-positive B-cell acute lymphoblastic leukemia, but the reciprocal fusion A4M is expressed in only half of the patients. Because prenatal leukemogenesis manifests as impaired early hematopoietic differentiation, we took advantage of well-established human embryonic stem cell-based hematopoietic differentiation models to study whether the A4M fusion cooperates with MA4 during early human hematopoietic development. Co-expression of A4M and MA4 strongly promoted the emergence of hemato-endothelial precursors, both endothelial- and hemogenic-primed. Double fusion-expressing hemato-endothelial precursors specified into significantly higher numbers of both hematopoietic and endothelial-committed cells, irrespective of the differentiation protocol used and without hijacking survival/proliferation. Functional analysis of differentially expressed genes and differentially enriched H3K79me3 genomic regions by RNA-sequencing and H3K79me3 chromatin immunoprecipitation-sequencing, respectively, confirmed a hematopoietic/endothelial cell differentiation signature in double fusion-expressing differentiating hematopoietic cells. Importantly, chromatin immunoprecipitation-sequencing analysis revealed a significant enrichment of H3K79 methylated regions specifically associated with HOX-A cluster genes in double fusion-expressing differentiating hematopoietic cells. Overall, these results establish a functional and molecular cooperation between MA4 and A4M fusions during human hematopoietic development.
**Introduction**

The mixed-lineage leukemia (*MLL*) gene encodes for an H3K4 histone methyltransferase important in hematopoietic development. The human *MLL* gene is frequently rearranged in acute leukemia and typically confers a dismal outcome. Of particular interest is the t(4;11)(q21;q22) translocation, which encodes the fusion proteins MLL-AF4 (MA4) and AF4-MLL (A4M), and is associated with infant B-cell acute lymphoblastic leukemia (B-ALL). This t(4;11)− infant leukemia is characterized by a very brief latency, raising the question of how it evolves so quickly. Moreover, the exceptionally high concordance rate of t(4;11)− B-ALL in monozygotic twins infants suggests that all the necessary (epi)genetic events required for leukemogenesis are accomplished prenatally, during embryonic/fetal hematopoietic development. However, our understanding of t(4;11)-mediated developmental effects is limited due, at least in part, to the variety of phenotypes and long latency observed in currently available t(4;11) mouse models. These different phenotypes likely result from targeting a cell in the wrong developmental stage or not addressing the impact of secondary hits, leaving open questions about the developmental impact of the t(4;11) translocation during early human development.

The functional and molecular contribution of the reciprocal fusion genes resulting from the derivative translocated chromosomes remains obscure in cancer. The MA4 fusion is always expressed in t(4;11)− B-ALL patients, whereas the reciprocal fusion A4M is expressed in only half of the patients. Importantly, t(4;11)− cell lines display addiction to MA4 but not to A4M, and although A4M was not sufficient to initiate leukemia in cord blood-derived CD34+ cells, it was nevertheless capable of initiating B-ALL in mice without the requirement of MA4, indicating that it contributes to the t(4;11)-driven leukemogenesis. Strikingly, a very recent clinical study has unraveled an independent prognostic value for MA4 expression in t(4;11)− infant B-ALL, thus adding a new piece to the puzzle. Thus, the developmental/pathogenic contribution of the t(4;11) resulting reciprocal fusion A4M remains enigmatic.

Human embryonic stem cells (hESC) represent a powerful tool for modeling different developmental aspects of human disease that cannot otherwise be addressed by analyses of patients’ samples or mouse models. Given that prenatal leukemogenesis manifests as impaired early hematopoietic differentiation, modeling hematopoietic differentiation in hESC may represent a promising in vitro approach to study the onset of hematopoiesis and the mechanisms underlying early human hematopoietic development. During hESC differentiation, a primitive population of CD45− hematopoietic-primed precursors (HEP) arises and further differentiates into CD45+ hematopoietic and mature endothelial cells. Beyond its pathogenic role in acute leukemias, the *MLL* gene has also been implicated in endothelial cell maturation, and endothelial dysfunction was recently linked to disease outcomes in childhood leukemias. We previously reported that MA4 favors the emergence of endothelial-primed HEP but not hemopoietic HEP from hESC. Here, we took advantage of well-established hESC-based differentiation systems to study whether the A4M fusion cooperates with MA4 during early human hematopoietic and endothelial development. We report a functional and molecular cooperation between MA4 and A4M fusions, which results in enhanced hemato-endothelial output during human embryonic development.

**Methods**

**Vector construction and lentiviral transduction**

The cDNA for MA4 and A4M were subcloned into the pRL-EF1α-PGK-NEO vector. Both fusions have been described previously ([Online Supplementary Figure S1A](#)). We used the following lentivectors containing either neomycin or dTo for cell selection: pRL-EF1α-PGK-NEO (empty vector; EV), pRL-EF1α-MA4-PGK-NEO (MA4) and pRL-EF1α-A4M-PGK-dTo (A4M). VSVG-pseudotyped lentiviral particles were generated in 293T cells using standard transfection protocols and concentrated by ultracentrifugation. hESC were infected overnight with concentrated EV or MA4 lentivirus plus 8 μg/mL polybrene. Viral supernatants were washed away the next day, and EV- and MA4-transduced hESC were then selected with G418 (50-100 μg/mL) for 3 weeks. For dual transduction of MA4 and A4M fusions, G418-resistant MA4-expressing hESC were infected with A4M-expressing viruses. EV/G418-selected hESC were also transduced with A4M alone. Transgene expression was confirmed for all the genotypes (Figure 1).

**Human embryonic stem cell culture and characterization of transgenic human embryonic stem cell lines**

hESC (AND1 cell line) were maintained undifferentiated on a layer of irradiated human mesenchymal stem cells in complete knockout Dulbecco modified Eagle medium containing 20% knockout serum replacement and 8 ng/mL basic fibroblast growth factor. The medium was changed daily, and cells were passaged weekly by dissociation with 1:1 collagenase IV/dispase. Cultures were visualized daily by phase contrast microscopy. Approval for hESC work was obtained from the Spanish National Embryo Ethical Committee. The pluripotency of transgenic hESC was characterized by flow cytometry using antibodies against SSEA-3, SSEA-4, TRA-1-60, TRA-1-81, OCT4, NANOG, SOX2, CRIPTO, and DNMT3B as well as transgene expression (MA4 and A4M) were analyzed by quantitative real-time polymerase chain reaction ([2CR](#)) ([Online Supplementary Table S1](#) shows the primers and PCR conditions used).

**Hematopoietic differentiation from human embryonic stem cells by embryoid body formation**

Undifferentiated hESC were treated with collagenase IV/dispase for 1 h at 37°C. To examine embryoid body (EB) formation, cells were transferred to low-attachment plates and incubated overnight in differentiation medium (knockout Dulbecco modified Eagle medium supplemented with 20% fetal bovine serum, 1% non-essential amino acids, 1 mmol/L L-glutamine, and 0.1 mmol/L β-mercaptoethanol). The medium was changed the next day to the same differentiation medium supplemented with the following hematopoietic cytokines: 300 ng/mL stem cell factor, 300 ng/mL Flt3L, 10 ng/mL interleukin-3, 10 ng/mL interleukin-6, 50 ng/mL granulocyte - colony-stimulating factor and 25 ng/mL bone morphogenetic protein-4 (all from R&D). EB were dissociated at different time points during development using collagenase B and enzyme-free Cell Dissociation Buffer (Invitrogen). Dissociated cells were stained with anti-CD34-FITC, anti-CD31-APC or anti-CD45-PE-Cy7, CD31-BV510, anti-
glycophorin A, anti-CD43-FITC, anti-CD45-APC antibodies (all from BD Biosciences) and 7-actinomycin D, and analyzed using a FACS Canto flow cytometer. Colony-forming unit assays were performed on days 10 and 15 of EB differentiation by plating 60x10^4 EB cells onto serum-free methylcellulose H4435 (Stem Cell Technologies). Colonies were scored after 12 days. Colony-forming unit assays were performed on days 10 and 15 of EB differentiation by plating 60x10^4 EB cells onto serum-free methylcellulose H4435 (Stem Cell Technologies). Colonies were scored after 12 days.

**Cell cycle and apoptosis analysis**

For cell cycle analysis of hESC-derived HEP and CD45+ cells, day 15 EB were dissociated and harvested cells were fixed overnight in 70% ice-cold ethanol. Cells were then washed in phosphate-buffered saline and incubated with anti-CD31-FITC, anti-CD34-PE-Cy7 and anti-CD45-APC antibodies for 15 min. Cells were then suspended in propidium iodide-containing buffer and acquired and analyzed on a FACS Canto-II using Modfit LT4.0 software, discriminating between quiescent cells (G0/G1), cycling cells (S-phase) and G2/M cells. Apoptosis was assessed with an Annexin-V Apoptosis Detection kit (BD Biosciences). Human embryonic stem cell-OP9 co-cultures

hESC-OP9 co-cultures were performed as described elsewhere. OP9 stroma was prepared by plating OP9 cells in gelatin-coated dishes, and allowing them to overgrow as a monolayer. hESC were prepared as a suspension of small aggregates using collagenase IV:dispase. One-tenth of this suspension was plated on top of the 8-day overgrown OP9 stroma. Media were replaced on

**Figure 1. Characterization of transgenic human embryonic stem cells expressing the reciprocal fusion A4M together with MA4.** (A) RNA-sequencing and quantitative real-time polymerase chain reaction (qRT-PCR) validation revealed that ~45% (11/25) of the patients with t(4;11)+ B-cell precursor acute lymphoblastic leukemia do not express the reciprocal fusion A4M. (B) Left, Phase-contrast morphology of representative colonies from each transgenic human embryonic stem cell (hESC) line. Right, Reverse transcriptase polymerase chain reaction analysis (RT-PCR) confirming expression of both fusions in undifferentiated hESC. (C) qRT-PCR expression of the pluripotency genes OCT4, SOX2, NANOG, CRIPSCO, and DNMT3B. (D) Representative FACS data confirming expression of the pluripotency surface markers SSEA-3, SSEA-4, TRA-1-60, and TRA-1-81. BCP-ALL: B-cell precursor acute lymphoblastic leukemia; RNA-seq: RNA-sequencing; pos: positive; neg: negative; EV: empty vector; C+: positive control.
the next day and one-half volume media changes were performed every other day thereafter. Hematopoietic differentiation was assessed by flow cytometry at day 9 of co-culture. Accordingly, hESC-OP9 co-cultures were treated with collagenase IV/TrypLE and cells were dissociated and filtered through a 70-um strainer. Cell suspensions were stained with anti-mouse CD29-FITC and anti-human CD34-PE and CD45-APC antibodies. The proportion of HEP (CD34+CD31+CD45) and total blood cells (CD45+) were analyzed within the CD29 hESC-derived cell population. Hemogenic and endothelial HEP were distinguished based on CD34 and CD43 expression.40

Culture of FACS-isolated hemato-endothelial precursors in MS5 stroma or liquid culture

Day 9 human hESC-OP9 co-cultures were dissociated as above and both CD45+ cells and HEP were analyzed. FACS-purified HEP (CD29+CD34+CD31+CD45) were plated onto MS5 stroma or in liquid culture for 30 or 16 days, respectively, in differentiation medium with hematopoietic cytokines (50 ng/ml stem cell factor, 50 ng/ml Flt3L, 10 ng/ml interleukin-8, 20 ng/ml interleukin-7). The medium was changed every 7 days, and the emergence of CD45+ hematopoietic cells was analyzed by FACS.

Endothelial differentiation of hemato-endothelial precursors

HEP (2×10⁶) from day 9 human hESC-OP9 co-cultures were seeded onto 0.1% gelatin-coated plates in complete EGM-2 medium with microvasculature supplements (Lonza) for 7 days. Cells were then fixed, permeabilized and stained with rabbit anti-human VE-cadherin (Cayman), mouse anti-human endothelial nitric oxide synthase (BD Biosciences), and mouse anti-human endothelial nitric oxide synthase (BD Biosciences), and mouse anti-human von Willebrand factor (DAKO) followed by Alexa 488-conjugated anti-rabbit or Cy3-conjugated anti-mouse antibodies (Jackson ImmunoResearch) antibodies. Nuclei were counterstained with 4',6-diamidino-2-phenylindole. Images were obtained using an inverted fluorescence microscope. Day 7 differentiating cells were trypsinized and cell suspensions were stained with anti-human CD31-FITC and CD144-PerCP-Cy5.5 antibodies.

Mouse transplantation and analysis of hematopoietic-endothelial engraftment

NOD/Scid-IL2 receptor-γ− (NSG) mice were housed under sterile conditions. The Animal Care Committee approved all mouse protocols. Briefly, cord blood-derived CD34+ hematopoietic stem and progenitor cells (5×10⁵ cells) or cells from day 15 EB (5×10⁵ cells) were transplanted into the bone marrow as described previously.40

Animal health was monitored throughout the entire experiment. Mice were killed 10 weeks after transplantation and cell suspensions were analyzed by FACS for human chimerism using anti-HLA-ABC-FITC, anti-CD31-PE, CD144-PerCP-Cy5.5, and anti-CD45-APC antibodies.

RNA- and chromatin immunoprecipitation-sequencing

Details of the RNA- and chromatin immunoprecipitation-sequencing and analysis are provided in the Online Supplementary Methods.

Statistical analysis

All data are expressed as mean ± standard error of mean. Statistical comparisons were performed using GraphPad Prism software with the nonparametric Mann-Whitney test, two-tailed P-value (with 95% confidence interval). Statistical significance was defined as a P-value <0.05.

Results

Co-expression of A4M and MA4 does not hijack pluripotency

We showed very recently that only 45% of t(4;11)+ B-ALL patients express the reciprocal fusion A4M, whereas MA4 is consistently expressed in all t(4;11)+ B-ALL patients (Figure 1A).10 Here, we generated transgenic hESC lines expressing “MA4 alone”, “A4M alone”, or MA4+A4M (double fusion), (Figure 1B and Online Supplementary Figure S1B). EV (control)- and MA4-hESC were established by G418 selection.41 G418-resistant EV- or MA4-expressing hESC were then transduced with A4M/dTo-expressing lentiviruses and greater than 90% transduction efficiency was achieved. Transgenic hESC lines were maintained for more than 50 passages and retained hESC-like morphology (Figure 1B, left), transgene expression (Figure 1B, right), and expression of pluripotency-associated transcription factors (Figure 1C) and surface markers (Figure 1D). All hESC genotypes formed teratomas in NSG mice (data not shown).53,54 Thus, (co-)expression of A4M and/or MA4 is compatible with hESC pluripotency.

A4M and MA4 co-operate to promote emergence of hematopoietic-endothelial precursors and enhance blood production

Hematopoietic differentiation was assessed using two distinct and well-established differentiation systems: EB formation45,46 (Figure 2) and OP9 co-culture (Figure 3). During differentiation, a population of primitive HEP arises, which is responsible for further hematopoietic and endothelial commitment.30,31 (Figures 2A and 3A). We investigated whether co-expression of A4M and MA4 affects hESC-derived hematopoiesis by analyzing the emergence of HEP during EB development in hESC individually expressing the single fusions or the double fusion. We observed a pronounced (~5- to 10-fold; P<0.05) increase in HEP at days 7 and 10 of development in EB expressing the double A4M and MA4 fusion over those expressing single fusions (Figure 2B, upper-left panel). We next assessed whether co-expression of A4M and MA4 influences subsequent hematopoietic commitment of HEP. The kinetics of emergence and output of both total CD45+ hematopoietic cells and CD45+CD31+ hematopoietic progenitors was faster (EB day 10) from double fusion-expressing hESC than from equivalent single fusion-expressing cells, achieving a 2- to 3-fold higher hematopoietic output by day 15 of EB development (Figure 2B). Furthermore, double fusion-expressing HEP massively accelerated (EB day 10) the emergence of clonogenic hematopoietic progenitors as compared to single fusion-expressing HEP (Figure 2B, bottom-right panel). According to our previous work, if the kinetics of human EB differentiation is extended, allowing for a continuum of HEP-to-blood transition, MA4-expressing human EB display enhanced HEP production coupled to impaired blood output (EB day 20) (Online Supplementary Figure S2A) and clonogenic potential (EB day 15) (Figure 2B). We confirmed stable expression of ectopic MA4 and A4M upon EB differentiation, supporting the link between genotype and phenotype (Figure 2C).

We also investigated hematopoietic differentiation using the OP9 differentiation system (Figure 3A,B), and by plating FACS-sorted HEP in either hematopoietic liquid
culture (Figure 3C) or onto MS5 feeders (Figure 3D). After 10 days on OP9 stroma, double fusion-expressing hESC yielded a 10-fold higher number of CD45+ hematopoietic cells than did single fusion-expressing hESC (Figure 3B). Moreover, when HEP were FACS-sorted from day 9 OP9 co-cultures and allowed to differentiate into CD45+ blood cells, the yield of CD45+ cells was up to 60-fold higher in double fusion-expressing HEP than in single fusion-expressing HEP (Figure 3C,D). Encouraged by these results, we next investigated whether ectopic expression...
of both A4M and MA4 confers in vivo engraftment capacity to hESC-derived hematopoietic derivatives. To do this, we transplanted $5 \times 10^5$ hESC hematopoietic derivatives from each genotype into myeloablated NSG mice, finding that, despite regulating hematopoietic development in vitro, double fusion-expression did not confer in vivo engraftment to hESC hematopoietic derivatives (Figure 2D).

The increased hematopoietic output of double fusion-expressing hESC might be the consequence of transgene-mediated proliferation/survival of the emerging HEP or CD45$^+$ cells. To address this, we analyzed cell cycle distribution (Online Supplementary Figure S2B) and apoptosis (Online Supplementary Figure S2C) within both HEP and the CD45$^+$ cell population. No differences in the proportions of either cycling HEP or CD45$^+$ cells were detected between genotypes (25–36% for HEP and 35–41% for CD45$^+$ cells) (Online Supplementary Figure S2B). Apoptotic levels were similarly low in the different genotypes of HEP (6–8%) and CD45$^+$ cells (5–7%) (Online Supplementary Figure S2C). Collectively, these results show that A4M cooperates with MA4 to induce HEP specification and blood commitment, without hijacking proliferation or survival of HEP.

**A4M and MA4 cooperate to enhance endothelial cell fate from hemato-endothelial precursors**

We next addressed the developmental impact of A4M in endothelial maturation from HEP. We hypothesized that co-expression of A4M and MA4 in HEP may concomitantly promote subsequent endothelial and hematopoietic commitment or skew the hemato-endothelial commitment in favor of hematopoiesis. To test this, we analyzed the ability of HEP to differentiate into mature endothelial cells. OP9-hESC co-cultures were dissociated on day 9 of development and HEP were FACS-sorted and cultured for 1 week in endothelial-promoting conditions (Figure 4A). The expression of the mature endothelial markers VE-cadherin (CD144), von Willebrand factor, endothelial nitric oxide synthase and CD31 was then analyzed. Irrespective of the genotype, HEP cultured in endothelial conditions attached, became spindle-shaped, and formed VE-cadherin$^+$ endothelial-like structures co-expressing endothelial nitric oxide synthase, von Willebrand factor and CD31 (Figure 4B,C, top panel). However, double fusion-expressing HEP were more prone to differentiate into mature endothelial cells than were single fusion-expressing HEP. Accordingly, they yielded a 20-fold higher number of VE-cadherin$^+$ endothelial-like structures (Figure 4C, top panel) and CD144$^+$CD31$^+$ endothelial cells (Figure 4C, bottom panel). Interestingly, endothelial cells (HLA.ABC$^+$CD31$^+$CD45$^+$CD45$^+$CD45$^+$) were found in the bone marrow of mice transplanted with double fusion-expressing hESC blood derivatives at levels ~4-fold higher than those in mice transplanted with single fusion-expressing cells (Figure 4D).

Within CD34$^+$CD31$^+$CD45$^+$ HEP, two subpopulations of phenotypically and functionally distinct HEP can be distinguished based on the expression of CD34 and CD43: hemogenic HEP (CD34$^+$CD43$^+$CD45$^+$) and endothelial HEP (CD34$^+$CD43$^+$CD45$^+$) (Figure 5A). We thus analyzed the contribution of both HEP populations to the superior hematopoietic and endothelial differentiation observed in double fusion-expressing HEP. Co-expression of A4M and MA4, but not single fusions, robustly enhanced the emergence of both endothelial and hemogenic HEP (Figure 5B,C). The identity of hemogenic and endothelial HEP was confirmed by the specific expression of markers.
expression of early hematopoietic and endothelial master genes (Figure 5D). Thus, A4M cooperates with MA4 to promote hematopoietic and endothelial cell fate.

**Genome-wide transcriptomic and H3K79 methylation profiles support the developmental cooperation between A4M and MA4**

To identify patterns of gene expression that might provide a molecular explanation of the functional cooperation between A4M and MA4 in hematopoietic specification, we performed RNA-sequencing analysis on FACS-purified EV-, MA4-, A4M- and double fusion-expressing HEP from day 15 EB. Figure 6A shows a heatmap representation of the hierarchical clustering of the 335 genes differentially expressed between the four genotypes (Online Supplementary Table S2). There is a clear transcriptomic transition towards a hematopoietic/endothelial gene signature from EV-HEP to double fusion-expressing HEP. Single fusion-expressing HEP clustered interspersed between EV and double-fusion HEP. The biological func-

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**Figure 4. Enhanced endothelial cell fate from hemato-endothelial precursors co-expressing MA4 and A4M.** (A) Scheme of hemato-endothelial precursor (HEP) endothelial differentiation and phenotypic characterization. (B) FACS-sorted HEP from day 9 human ESC-OP9 co-cultures were cultured in EGM2 medium for 5 days and analyzed by immunofluorescence for VE-cadherin, endothelial nitric oxide synthase and von Willebrand factor. (C) Top, Endothelial-like structures were identified and quantified based on VE-cadherin staining (white dotted-lined areas in B, top panel). Bottom, Frequency of CD45-CD31+CD144+ endothelial cells quantified by flow cytometry. (D) In vivo endothelial engraftment potential (HLA.ABC+CD31+CD144+CD45-) analyzed in bone marrow of NSG mice 8 weeks after transplantation of HEP. Data are presented as mean ± standard error of mean from five independent experiments. *P<0.05. bFGF: basic fibroblast growth factor; EV: empty vector; CB: cord blood.
tions affected by genes differentially expressed in MA4-, A4M- and double fusion-expressing HEP relative to EV were classified by Ingenuity Pathway Analysis software. The top significantly enriched functional categories included "hematological system development and function", "cancer" and "hematological disease" (Figure 6B). Statistical (-log_{p-value}) power shows distant effects of MA4 and A4M; however, co-expression of both fusions seems to establish a molecular balance/developmental cooperation in promoting blood-endothelial specification from hPSC. Strikingly, biofunctions specifically associated with hematologic lymphoid malignancies (not with non-hematologic cancer) were predicted to be activated (positive z-score) exclusively in double fusion-expressing HEP, further suggesting a molecular cooperation between MA4 and A4M in development and infant leukemia. (Figure 6C).

The C-terminal partners of MLL fusions normally interact with the histone methyltransferase DOT1L, which is the sole histone methyltransferase catalyzing histone 3 lysine 79 (H3K79) methylation, a chromatin modification widely associated with the dysregulated expression of HOX-A cluster genes in MLL leukemias. We thus performed genome-wide chromatin immunoprecipitation-sequencing analysis of the H3K79 trimethylation (H3K79me3) profiles in control, MA4-, A4M- and double fusion-expressing hESC-derived blood derivatives (Figure 7, Online Supplementary Figure S3A, Online Supplementary Table S3). In line with the RNA-sequencing data, functional analysis of the differential H3K79me3 peaks specific for double fusion-expressing cells revealed significant gene ontology functional categories associated with "definitive hematopoiesis", "myeloid and erythroid differentiation/homeostasis" and "endothelial cell development" (Figure 7A, Online Supplementary Figure S3B). This further supports the developmental cooperation between A4M and MA4 in promoting hematopoietic specification.

Finally, we analyzed the H3K79me3 profiles at genomic loci of the classical MLL target genes reported by Guenther et al. Non-HOX-A classical MLL targets such as RUNX1, LMO2, ADMA10, and KDM6A showed a slight although non-significant, enrichment of H3K79me3 in MA4-expressing hESC, validating our chromatin immunoprecipitation-sequencing approach (Figure 7B). However, enrichment of H3K79me3 in HOX-A cluster genes was observed exclusively in A4M-expressing cells although it was statistically significant only in double
fusion-expressing differentiating hESC (false discovery rate <0.1) (Figure 7C). As such, HOX-A genes were upregulated in double fusion-expressing hematopoietic clonogenic progenitors (Online Supplementary Figure S3C). No differential enrichment of the repressive H3K4me2/3 mark was observed in either HOX-A or non-HOX genes in double fusion-expressing cells (Online Supplementary Figure S4). Collectively, these data suggest that the deregulated expression of HOX-A genes in MLL leukemias may be imposed by the reciprocal A4M fusion through H3K79 methyltransferase activity. In support of this, a recent RNA-sequencing study performed in 42 infants with t(4;11)- B-ALL enrolled in the Interfant treatment protocol, revealed that 45% of t(4;11)- patients express the A4M...
fusion, and that HOX-A cluster genes are exclusively expressed in this AF4-MLL-expressing subgroup of t(4;11)+ patients, who do in fact have a significantly more favorable clinical outcome.19

Discussion

From etiological and pathogenic standpoints, infant cancer is distinct from adult cancer and should be studied as a developmental disease.6,7,16 A biologically and clinically intriguing infant cancer is the t(4;11)+ B-ALL, which is associated with a dismal outcome.20,21 Evidence in support of its prenatal origin comes from studies in monozygotic twins and archived blood spots, providing compelling evidence of a single prenatal cell as the origin for t(4;11), and also from recent genome-wide studies demonstrating that this infant leukemia has one of the lowest frequencies of somatic mutations of any sequenced cancer.22 The stable genome of these patients suggests that in infant developmental cancer, one “big-hit” might be sufficient to cause overt disease, supporting a key contribution of the prenatal cell-of-origin during a critical developmental window of stem cell vulnerability in leukemogenesis. However, despite its aggressiveness and short latency, our current understanding about its etiology, pathogenesis and cellular origin is still limited.2,4,14,16,52 Importantly, a recently developed xenograft model which represents the most bona fide model for t(4;11)+ B-ALL so far, has revealed the instructive role of MLL-AF4 in cord blood-derived CD34+ cells.14

Studies using primary cells from t(4;11)+ B-ALL patients are incapable of addressing the developmental genesis of the hematopoietic system. Recent data suggest that fetal liver lymphoid-primed multipotent progenitors may provide the developmental prerequisites for the initiation of t(4;11)+/MLL-AF4 infant leukemia.23 Because leukemogenesis manifests as a blockage or altered cell differentiation, the hematopoietic differentiation of hESC may represent a promising in vitro model for studying the onset of hematopoiesis and the earliest events leading to the specification of the hematopoietic cells.24 Previous studies have addressed the oncogenic role of leukemic fusion genes in hESC-derived hematopoiesis.25-27 We previously explored the developmental impact of the prenatal fusion MA4 in hESC hematopoietic development,28 and found that MA4 expression promotes the emergence of endothelial-primed HEP and further endothelial commitment, but hijacks the specification of hemogenic-primed HEP, impairing hematopoietic output.10

Figure 7. H3K79 methylation profiles at genomic loci of MLL targets in MA4-, A4M- and double fusion-expressing human embryonic stem cell-derived blood derivatives. (A) Gene ontology enrichment of differential H3K79me3 peaks specific for double fusion-expressing cells. (B,C) Representative profiles for chromatin immunoprecipitation-sequencing using anti-H3K79me3 antibody at genomic regions of typical non-HOXA (B) and HOXA MLL targets (C).
The functional and molecular contribution of the reciprocal fusion genes resulting from the derivative translocated chromosomes remains obscure in cancer. The MA4 fusion is always expressed in t(4;11) B-ALL patients, whereas the reciprocal fusion A4M is expressed in only 45-50% of the patients. Here, we took advantage of well-established hESC-based differentiation systems to study whether the A4M fusion cooperates with MA4 during early human hematopoietic and endothelial development. Co-expression of A4M and MA4 strongly promoted the emergence of both endothelial-primed and hemogenic-primed HEP. Moreover, the double fusion-expressing HEP specified into significantly higher numbers of both hematopoietic and endothelial cells, irrespective of the in vitro differentiation protocol used and without affecting survival or proliferation, indicating a functional (developmental) cooperation between MA4 and A4M fusions during human hematopoietic development. This notion was confirmed by genome-wide transcriptomic analysis of differentiating HEP. These developmental biology studies support previous evidence suggesting that A4M contributes to the pathobiology of t(4;11) B-ALL. Accordingly, Birden et al. reported that A4M-transduced murine hematopoietic stem cells developed pro-B-ALL, whereas co-transduction with MA4 and A4M resulted in mixed lineage leukemia. Moreover, studies from Milne’s laboratory demonstrated that RUNX1 is directly activated by MA4 and the RUNX1 protein interacts with the A4M protein, suggesting a mechanism of cooperation between the two fusion genes at the molecular level.

In the embryo, definitive hematopoiesis cannot occur in the absence of endothelial cell development. Definitive hematopoietic stem cells in both mice and humans emerge from the hemogenic endothelium by a process known as endothelial-to-hematopoietic transition. Hematopoietic differentiation of hESC occurs through the generation of HEP, from which both endothelial and hematopoietic cells then originate. Here, co-expression of A4M and MA4 in HEP concomitantly promoted endothelial and hematopoietic commitment rather than skewing the hematopoieto-endothelial commitment in favor of one lineage over the other. This finding is important because beyond its pathogenic role in acute leukemias, the MLL gene is implicated in endothelial cell maturation, and endothelial dysfunction was recently linked to disease outcome in childhood leukemias. Furthermore, other leukemia fusion oncogenes as well as lymphoma-specific genetic aberrations have been found in endothelial cells from patients with chronic myeloid leukemia and B-cell lymphoma, suggesting that endothelial cells may be part of the neoplastic clone. In addition, bone marrow-derived mesenchymal stem cells from infant t(4;11) B-cell ALL were recently found to harbor and express the t(4;11) translocation. The existence of a common embryonic precursor for mesenchymal stem cells and endothelial cells has been recently demonstrated by hESC-directed differentiation. The finding of such a common embryonic precursor, and the presence of t(4;11) in both leukemic blasts and bone marrow mesenchymal stem cells of infant patients, suggests that the t(4;11) translocation arises and has a developmental impact on early pre-hematopoietic precursors. As a technical caveat, it is important to emphasize that MA4 and A4M were sequentially transduced to allow for antibiotic selection and homogeneously-transduced hESC cultures. However, in double fusion-expressing differentiating blood cells, MA4 was never individually expressed in the absence of A4M. When hematopoietic differentiation of hESC was induced by EB formation both fusions were readily co-expressed, ruling out a biased functional phenotype driven by the sequential expression of each transgene in distinct developmental windows.

Mechanistically, a putative function of A4M is to activate chromatin, rendering a chromatin landscape similar to that present during stem cell development. It is currently unknown how A4M is able to reprogram chromatin, but it does contain the SET domain disrupted from its "specification domain", the N-terminal portion of MLL, which binds to MEN1 and LEDGF, shaping the gene targeting module of the MLL gene. When A4M is expressed, the N-terminal portion is substituted by the AF4 N-terminal, which is the crucial domain (AF4N) that binds to and strongly activates RNA polymerase II (RNAPII) for transcriptional elongation. Overexpression of either AF4, AF4N or the fusion protein A4M induces robust RNAPII-dependent gene transcription by overwriting the elongation control process in a dominant fashion. Since gene transcription per se and in particular "sterile" transcription is a powerful mechanism for chromatin activation, A4M could potentiate MA4 to skew normal and leukemic hematopoietic cell fate decisions. This also explains why MA4 has a more prominent role in the disease than the reciprocal A4M. If A4M functioned to initiate this process only by itself, then it would become obsolete after fulfilling the "chromatin opening job". However, transcription factors such as MA4, RUNX1 or others then establish the transcriptional program leading to leukemogenesis. This is reflected in the enhanced hematopoietic potential of double fusion-expressing hESC and the enriched H3K79me3 activation mark in HOX-A cluster genes exclusively when MA4 and A4M are co-expressed. Thus, A4M prepares other transcription factors to become oncoproteins.

Molecularly, C-terminal-partners of MLL fusions (AF4, AF9, ENL) interact with DOT1L, which is the sole histone methyltransferase catalyzing H3K79 methylation, a chromatin modification widely associated with the dysregulated expression of the HOX-A gene cluster in MLL-rearranged leukemias. Here, chromatin immunoprecipitation-sequencing analysis of differentially enriched H3K79me3 genomic regions confirmed a hematopoietic/endothelial cell differentiation signature in double fusion-expressing HEP, and revealed a significant enrichment of H3K79 methylated regions specifically associated with HOX-A cluster MLL target genes (but not with non-HOX-A MLL targets) in double fusion-expressing differentiating hematopoietic cells. This is in line with the recently found significant positive correlation between the upregulation of the HOX-A gene cluster and the expression of A4M in primary t(4;11) infant B-cell ALL samples, and with previous studies identifying that approximately one-half of t(4;11) patients do not have an activated HOX-A signature. This may explain why MA4 failed recently to bind to HOX-A genes to regulate HOXA gene expression. Collectively, MA4 and A4M might cooperate through a complex molecular interaction to control HOX-A gene regulation. In conclusion, we describe a functional and molecular cooperation between MA4 and A4M fusions during human hematopoietic development, and demonstrate how hESC-based hematopoietic differentiation represents a promising system to explore the developmental impact of the chimeric
proteins resulting from chromosomal translocations, which remains obscure in human leukemia.

Acknowledgments

Financial support for this work was obtained from the European Research Council (CoG-2014-646903 and PoC-2018-811220) and the Generalitat de Catalunya (SGR330 and PERIS 2017-2019) to PM, the Spanish Ministry of Economy and Competitiveness (SAF2016-80481-R and SAF2016-76758-R) to PM and IV, the Spanish Association against Cancer (AECC-Cl-2015) and Fero Foundatio to CB, the Health Institute Carlos III (ISCIII/FEDE, PI17/01028 and PI17/01028) to CB and PIJR, the NIHR GOSH BRC and Great Ormond Street Hospital Children’s Charity to JdB, and Bloodwise and Cancer Research UK to BG. RM and PM were also supported by the Deutsche José Carreras Leukämie Stiftung. PM also acknowledges financial support from the Obra Social La Caixa-Fundació Josep Carreras. R-T-R is supported by a fellowship from the Spanish Association of Cancer Research (AECC). RV-M is supported by a Torres Quevedo fellowship from the Spanish Ministry of Science and Innovation (PTQ-16-08623). P.M is an investigator of the Spanish Cell Therapy cooperative network (TERCEL).

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