Hoxa11-mediated reduction of cell migration contributes to myeloid sarcoma formation induced by cooperation of MLL/AF10 with activating KRAS mutation in a mouse transplantation model

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Abstract

The molecular mechanism of myeloid sarcoma (MS) formation remains unclear. Our clinical and mouse model findings indicate that cooperation of KMT2A (MLL) translocation (MLL-t) with activating N-K-RAS mutations promotes MS formation in a shorter latency. To improve the understanding of MS formation, in this study, we performed scanning cell trafficking analysis and demonstrated that cells harboring cooperating mutations migrated more slowly to omental adipose tissues and more cells were retained in adipose tissues in vivo. Comparison of transcriptome profiling among three pairs of mouse MLL/AF10(OM-LZ) leukemia cell lines harboring activating and wild-type KRAS identified 77 differentially expressed genes (DEGs) with >1.5-fold change. Functional annotation of these 77 DEGs using Gene Ontology (GO) enrichment analysis followed by cluster analysis revealed that GO terms related to development/differentiation have the highest enrichment score. The roles of Hoxa10 and Hoxa11, two genes which mapped to this cluster, were further characterized. Silencing Hoxa10 and Hoxa11 in cells harboring cooperating mutations prolonged the survival and reduced MS formation, respectively, in the recipient mice. Data of imaging cell trafficking as well as competitive engraftment and clonal expansion analyses indicated that silencing or overexpressing Hoxa11 in mouse leukemia cells affected cell migration and retention in omental adipose tissue. Although silencing Hoxa11 in leukemia cells did not affect Cxcr4 expression, it resulted in increased transwell migration, motility in confined spaces 3 μm in size, and cell protrusion. Our results revealed that Hoxa10 plays an important role in survival and Hoxa11 contributes to MS formation in MLL-t acute myeloid leukemia with activating KRAS mutation.

Keywords: MLL translocation, KRAS mutation, Hoxa11, Hoxa10, Myeloid sarcoma, Migration

Introduction

Myeloid sarcoma (MS), previously called chloroma, myeloblastoma, or granulocytic sarcoma, is a rare manifestation of hematologic malignancies presenting as extramedullary soft tissue masses [1,2]. MS can be found in patients either as the only clinical presentation or concomitantly with, before, or after a diagnosis of acute myeloid leukemia (AML), myelodysplastic syndrome, or myeloproliferative neoplasm [1,2]. The incidence of MS in
patients with AML is approximately 0.8% to 9% [3,4], and it can occur at many sites, the most frequently reported of which are skin, lymph node, gastrointestinal tract, bone, testis, peritoneum, and central nervous system [5-7]. Several chromosomal abnormalities are recurrently detected in patients with MS, including KMT2A (MLL) translocations (MLL-t, t(8;21), inv(16), t(8;16)(p11;p13), monosomy 7, and trisomy 8 [2,5-7]. Patients with AML harboring t(8;21) or MLL-t have significantly lower overall survival in the MS group than the non-MS group [8,9]. The most frequently detected genes with mutations in patients with MS are N-K-RAS, NPM1, and DNMT3A [10-12]. In a previous study, we found that 35 (60%) of patients with AML and MLL/AF10 and N-K-RAS mutations had MS formation [13]. Using a retroviral transduction/transplantation mouse model, we demonstrated that cooperation of MLL/AF10 with activating KRAS (KRASG12D) induced MS formation in a shorter latency [13]. Our results supported that MLL-t and KRAS mutations are associated with MS formation, and additionally, MS formation is associated with shorter survival.

The molecular mechanism of MS formation remains unclear. Among the chromosomal abnormalities frequently detected in patients with MS, MLL-t and t(8;16)(p11;p13) involve genes with epigenetic regulating activities. MLL encodes a histone H3 lysine 4 methyltransferase that is essential for regulating gene expression during early development and hematopoiesis. Translocations of MLL lead to misregulation of MLL downstream genes, such as HOX11–HOX10 and subsequently, impair hematopoietic lineage commitment and induce leukemia development [14,15]. Cells from the human leukemia cell line THP-1, possessing MLL/AF9 and activating NRAS, have high expression levels of HOX11, HOX10, HOXA7, and HOXA4. These genes were downregulated during phorbol 12-myristate 13-acetate-induced monocyte-macrophage differentiation [16]. The t(8;16)(p11;p13) translocation resulted in the fusion of MYST3 and CREBBP, both of which exhibit histone acetyltransferase activity. Gene expression profile analysis of AML cases with t(8;16)(p11;p13) revealed that they were clustered near those with MLL-t and highly expressed HOXA11, HOX10, RET, PERP, and GGA2 [17]. Increased expression of HOX10 and HOXA11 was reported in both types of AML. HOX10 expression in AML is associated with poor prognosis [18,19]. Additionally, Hoxt10 overexpression-induced AML was confirmed using a mouse model [20]. In contrast, the t(7;11)(p15;p15) translocation, which results in the fusion of NUP98 and HOXA11, was identified in patients with AML, chronic myeloid leukemia, and juvenile myelomonocytic leukemia [21,22], suggesting that Hoxa11 is closely related to leukemogenesis. Whether these two genes are involved in MS formation has not been investigated.

To improve the understanding of MS formation, in this study, we compared the transcriptome profiling between mouse MLL/AF10(OM-LZ) leukemia cells harboring wild-type and oncogenic KRAS and discovered that Hoxa10 and Hoxa11 were differentially expressed. Additionally, the functions of Hoxa10 and Hoxa11 in survival and MS formation were investigated using a mouse model, and the roles of Hoxa11 in cell migration, retention, motility, and protrusion formation were examined.

Materials and methods

Cell lines and cell culture conditions

Mouse MLL/AF10 leukemia cell lines have been generated by retroviral transduction of MLL/AF10(OM-LZ) and wild-type (KRASWT) or activating KRAS (KRASG12D) to 5-Fluorouracil-enriched C57BL/6 (B6; National Laboratory Animal Center, Taiwan) mouse bone marrow (BM) cells in our previous studies [13,23,24]. Of these mouse leukemia cell lines, 12G cells harbor MLL/AF10(OM-LZ) solely, AK2G and AK3G cells harbor MLL/AF10(OM-LZ) and KRASG12D, and Akw1G cells harbor MLL/AF10(OM-LZ) and KRASWT. Mouse leukemia cell lines were maintained in RPMI 1640 medium (Gibco, Grand Island, NY, USA) containing 20% fetal bovine serum (FBS; HyClone, Logan, UT, USA), 2 mM L-glutamine, 100 μM 2-mercaptoethanol, and 10 ng/ml interleukin-3 (R&D Systems, Minneapolis, MN, USA).

Imaging cell trafficking analysis

CellVue NIR815-labeled cells were prepared according to the manufacturer’s instructions (LI-CORE Biosciences, Lincoln, NE, USA). Briefly, leukemia cells were washed using medium without FBS and resuspended in diluted C reagent containing CellVue NIR815 dye (LI-CORE) at a final concentration of 10⁻⁶ M for 5 min. The dye uptake reaction was stopped by adding an equal volume of FBS to the cell suspension. Labeled cells were washed to remove the unincorporated CellVue NIR815 dye and then imaging using Pearl Image (LI-CORE) or by fluorescence microscopy (TissueGnostics GmbH, Vienna, Austria) to ensure successful labeling. The labeled cells were intraperitoneally (ip) injected into sublethally irradiated B6 mice (1 × 10⁵ cells/mouse). Mouse injected with unlabeled cells served as blank controls. In vivo fluorescence images of the transplanted mice were collected using Pearl Image at 5 min, 1.5 h, 24 h, 7 days, 25 days, and 32 days. During the imaging period, mice were anesthetized with 2.5% isoflurane gas in the oxygen flow (1.5 l/min). In all imaging experiments, reproducibility was confirmed with 3–5 independent mice. The fluorescent signals of organ biopsies from transplanted mice were detected using Xenogen IVIS Spectrum (Caliper Life Science, Hopkinton, MA, USA) or Pearl Image. Organ and tissue biopsies were also formaldehyde-fixed, paraffin-embedded, sectioned, and stained by hematoxylin and eosin staining reagents according to standard protocols.

Microarray analysis and Gene Ontology (GO) enrichment analysis

Trizol reagent (Invitrogen, Carlsbad, CA, USA) was used to extract total RNA of leukemia cells. The RNA was amplified, labeled, and then hybridized with a mouse genome 430A Array chip (12G vs. AK3G), 430 2.0 Array chip (12G vs. AK2G), or Clariom D Array chip (AKw1G vs. AK3G) (Affymetrix, Santa Clara, CA, USA) according to the manufacturer’s instructions. This procedure was performed by the staff of the Genomic Medicine Research Core Laboratory at Chang Gung Memorial Hospital in Linkou, Taiwan. The microarray data are available at the NCBI GEO website (accession numbers: GSE82156 and GSE134586) [24,25]. Transcriptome Analysis Console (TAC) software 4.0 (Affymetrix) was used for differential expression analysis. Heatmaps were generated using TreeView Cluster 3.0 (http://bonsai.hgc.jp/~mdehoon/software/cluster/) and JavaView v.1.6.4 (http://www.javaview.de). Differentially expressed genes (DEGs) with more than 1.5-fold change between paired leukemia cell lines were functionally annotated with Gene Ontology (GO) enrichment analysis using online Database for Annotation, Visualization, and Integrated Discovery (DAVID) v6.8 annotation tools (https://david.ncifcrf.gov/). Statistical significance was evaluated using Fisher’s exact test and adjusted by Bonferroni correction for multiple testing. The GO biological process (GO_BP) terms with P < 0.05 were considered statistically significant.

Reverse transcription-quantitative polymerase chain reaction (RT-qPCR)

Total RNA of mouse leukemia cells was reverse transcribed into complementary DNA (cDNA) using random hexamers and Superscript II reverse transcriptase (Life Technologies, Rockville, MD, USA) according to the manufacturer’s protocol. Quantitative PCR was performed using SYBR Green PCR master mix and analyzed using an ABI Prism 7900 system (Applied Biosystems, Foster City, CA, USA). The primer sets were 5′-GAA AAC CTC GCT TCC TCC GA-3′ and 5′-ATA AGG GCA GCC CTT TTG GC-3′ for mouse Hoxa11, 5′-CCA AGG GCC AAA ATG CAG CC-3′ and 5′-CTG CTC GTG CTT GTG GTA AG-3′ for Hoxa10, and 5′-TTC ACC
ACC ATG GAG AGG GC-3' and 5'-GGC ATG GAC TGT GGT CAT GA-3' for Gapdh. The thermal cycling conditions were as follows: 95°C for 2 min, followed by 40 cycles of 95°C for 15 s and 65°C for 30 s. The gene expression level of the target gene was normalized by the housekeeping gene Gapdh. Fold change was calculated as $2^{-\Delta\Delta Cm}$ [26].

**Gene knockdown by short hairpin RNA (shRNA)**

To generate stable Hoxa10 and Hoxa11 knockdown cell lines, AK3G cells were infected with a lentivirus-expressing shRNA against Hoxa10 (The RNAi Consortium shRNA clone numbers TRCN000012515 and TRCN000012516) or against Hoxa11 (TRCN0000070753 and TRCN0000070755) at a multiplicity of infection of one. The lentiviruses were obtained from the National RNAi Core Facility at the Institute of Molecular Biology/Genomic Research Center, Academia Sinica, Taiwan. Infected cells were selected in RPMI medium containing puromycin (2.5 μg/ml) for 2 weeks. AK3G cells stably transfected with blank lentiviral vector pLKO.1-empty or pLKO_TRC025 were used as negative controls.

**Western blotting**

Western blotting was performed as described previously [13]. Twenty micrograms of total protein lysate was loaded in each lane. The primary antibody against mouse Hoxa10 (1:5000, GXB-CEOCD3) was obtained from GenWay (San Diego, CA, USA), whereas those for Hoxa11 (1:5000, NBP1-80228) and β-actin (1:10000, sc-47778) were obtained from Novus Biologicals (Littleton, CO, USA) and Santa Cruz Biotechnology (Santa Cruz, CA, USA), respectively. The horse radish peroxidase (HRP)-conjugated secondary antibody (1:10000, C04001) was obtained from Croyez Bioscience (Taipei, Taiwan). Chemiluminescence was generated using a Western Lightning Plus ECL Kit (Perkin Elmer, Skovlund, Denmark) and captured using X-ray film or Analytik Jena™ UVP ChemStudio PLUS and analyzed by VisionWorks Touch Software 9.0 (Analytik Jena US LLC, Upland, CA, USA).

**BM transplantation assay**

The BM transplantation assay has been reported previously [25,27]. Briefly, leukemia cells were ip injected into male B6 mice (1 × 10⁶ cells/mouse) that received either a sublethal dose of γ-irradiation (5.25 Gy) (AKw1G and AK5G, n = 5 each; AK3G-shV and AK3G-shH10-2, n = 10 each) or no irradiation (AK3G-shV and AK3G-shH11-2, n = 10 each). Mice ip injected with normal saline (n = 5) served as a blank control. Mouse complete blood count of peripheral blood was assessed weekly by hemocytometry using a Hemavet 950 (Drew Scientific, Oxford, CT, USA) or BC-5000 (Mindray, Shenzhen, China). Physical examination was performed daily. Mice were sacrificed at the moribund state, which was characterized by leukocytosis, with hunched posture, weakness, shortness of breath, and 20% body weight loss. BM, peripheral blood, ascites, organs, and tumor masses were collected and weighed.

**Flow cytometry analysis**

To detect CXCR chemokine receptor-4 (Cxcr4) expression, leukemia cells were incubated with allophycocyanin (APC)-conjugated anti-mouse CD184 (Cxcr4) antibody (0.5 μg per 10⁶ cells in 100 μl volume, #146507; BioLegend, San Diego, CA, USA). Cells incubated with an isotype antibody (0.5 μg per 10⁶ cells in 100 μl volume, #400612; BioLegend) served as a negative control. Data were acquired using a FACS Canto II flow cytometer (Becton-Dickinson, San Jose, CA, USA) and analyzed using FlowJo v10.4 software (Tree Star Software, San Carlos, CA, USA).

**Transwell migration assay**

Cells (1 × 10⁵) were added to the insert of a 24-well Transwell plate (5-μm pore size; Corning Costar, Cambridge, MA, USA), and RPMI 1640 medium with stromal derived factor-1 (SDF-1) (150 ng/ml; PeproTech, Rocky Hill, NJ, USA) was added to the bottom chamber. After 6-h incubation, the cells in the insert and bottom chamber were counted using a hemacytometer. The values shown are expressed as the mean ± standard deviation (SD) of three independent experiments.

**Cell motility and protrusion in confined spaces**

To observe leukemia cell motility and protrusion formation in confined spaces (3 μm in size), the polydimethylsiloxane (PDMS) slab-based approach [28] was performed with some modifications. Briefly, a PDMS layer (SYLGARD 184 Kit; Dow Corning, Midland, MI, USA) was polymerized in a 30-mm glass-bottom culture dish (GeneDireX, Taiwan). An 8-mm hole was made by putting a polystyrene cloning cylinder (Sigma-Aldrich, St Louis, MO, USA) into the PDMS during polymerization. Cells were stained using a CellTrace Far Red Cell Proliferation Kit (Thermo Fisher, Waltham, MA, USA) according to the manufacturer’s instructions. A 50-μl cell suspension (5 × 10⁵) and 0.5 μl of latex beads (10%) (3-μm in diameter; SI-LB30, Sigma-Aldrich) were mixed thoroughly and pipetted into the hole. The cells and beads were moved into the space underneath the PDMS slab by lifting it using a pipette tip. The dish was then placed on the stage of a ZEISS Axiovert 200M inverted fluorescence microscope (Carl Zeiss AG, Oberkochen, Germany) equipped with a digital camera. After 30 min to ensure that the beads were holding tightly to create a confined space between the PDMS slab and the dish bottom, images were acquired every 6 s for 20 min. To observe the directional migration and protrusion of cells in response to SDF-1, 100 μl of SDF-1 (150 ng/ml in RPMI 1640 medium) was added to the hole, and after a 10-min incubation, the images were acquired every 6 s for 20 min.
Table 1
Survival and extramedullary myeloid sarcoma formation in the recipient mice.

| Mouse number | Survival (range) (days)* | AKw1G | AK5G | P_value | AK3G-shV | AK3G-shH10-2 | P_value | AK3G-shV | AK3G-shH11-2 | P_value |
|--------------|--------------------------|-------|------|---------|----------|--------------|---------|----------|--------------|---------|
| Mouse number | Survival (range) (days)* | 5     | 5    | 0.0332  | 10       | 10           | <0.001  | 10       | 10           | 0.4738  |
| MS occurrence (%)# | 0% | 100% | 0.0079 | 90% | 89% | 1.0000 | 100% | 100% | 1.0000 |
| Gonad (g)* | 0.94 | 1.06 | 0.1202 | 1.24 | 0.88 | 0.1480 | 1.41 | 1.15 | 0.3404 |
| Gonad/TBW (%)* | 3.24% | 3.57% | 0.3479 | 4.7% | 4% | 0.1237 | 4.3% | 4% | 0.5119 |
| OM (g)* | 0.78 | 1.86 | 0.0227 | 2.01 | 1.01 | 0.0057 |
| OM/TBW (%)* | 3.02% | 5.82% | 0.213 | 6.5% | 3.5% | 0.0045 |
| GI-OM (g)* | 3.82 | 3.32 | 0.9927 |
| GI-OM/TBW (%)* | 13.9% | 13.6% | 0.9741 |
| Ascites (%)# | 0% | 100% | <0.001 | 50% | 30% | 0.6500 | 80% | 10% | 0.0050 |

* T test or log-rank test
# Fisher’s exact test

Fig. 1. Migration and retention of AK3G cells at extramedullary sites. (A) Imaging analysis of mice ip injected with normal saline (blank) or 1:1 mixed 12G and AK3G cells by Peral Imager at indicated time points. CellVue NIR815-labeled mice were under anesthesia during image capture. (B) Imaging analysis of excised omentum (i) and gonadal adipose tissue from blank, 12G, and AK3G mice by Xenogen IVIS Spectrum 7 days post-transplantation. Arrows indicate fluorescent signals. (C) Histologic features of omental (i) and gonadal (ii) adipose tissues obtained from blank, 12G, and AK3G mice 7 days post-transplantation (× 40, H&E stain). (D) Imaging analysis of mice ip injected with CellVue NIR815-labeled AKw1G or AK3G cells by Peral Imager at indicated time points. (E) Imaging analysis of excised organs from AKw1G and AK3G mice by Pearl Imager 32 days post-transplantation (upper panel). The arrangement of organs is shown in the lower panel. L, liver; S, spleen; G, stomach; I, intestine; B, bladder; Go, gonad. Color bars (A, B, D, E) indicate the total fluorescence radiant efficiency.

Ethical requirements

All animal experiments were performed according to the protocol approved by the Animal Research Committee of Chang Gung Memorial Hospital (IACUC No. 2015033001).

Statistical analyses

The statistical significance of differences in gene expression levels of the two groups based on cDNA microarray data was compared by the Mann–Whitney test. Survival analysis was conducted according to the...
Kaplan–Meier method. The statistical significance of differences in organ weight and transwell migration was assessed using the Student’s t-test. All statistical analyses were performed using SPSS software version 20.0 for Windows (IBM Corp., Armonk, NY, USA). A P value less than 0.05 indicated statistical significance.

Results

Survival and MS formation in mice transplanted with MLL/AF10 leukemia cells harboring wild-type or mutant KRAS

The mouse MLL/AF10(OM-LZ) leukemia cell line (12G) and MLL/AF10(OM-LZ) cells harboring KRAS<sup>G12C</sup> (AK2G, AK3G, and AK5G) or KRAS<sup>G12V</sup> (AKw1G) were generated by retroviral transduction in our previous studies [13,23,24] and in this study (Supplementary Figure 1A). Our previous studies have demonstrated that mice transplanted with AK2G or AK3G cells had shorter survival compared with those transplanted with 12G. Additionally, 100% of the AK2G and AK3G mice, but not 12G mice, formed MS in omental-mesentery, gonadal, and subcutaneous adipose tissues [13]. In this study, we further demonstrated that mice transplanted with AK5G cells had shorter survival than those injected with AKw1G cells (Supplementary Figure 1B). Like AK2G and AK3G mice, AK5G mice, but not AKw1G mice, had significant MS formation in omental-mesentery adipose tissues (Supplementary Figure 1C; Table 1).

In vivo imaging to monitor cell trafficking

To investigate the process of MS formation in adipose tissues, in vivo imaging analysis was performed to monitor leukemia cell trafficking. The results showed that in mice ip injected with CellVue NIR815-labeled AK3G cells, dispersed fluorescent signals in the peritoneal cavities gradually condensed after 1.5 h, with the signals gaining strength 24 h post-transplantation (Fig. 1A, right column). Excised organs from AK3G mice 7 days post-transplantation showed that the fluorescent signals were mainly localized in the adipose tissue around the gastric and splenic veins (omentumal adipose tissue) (Fig. 1B-i, right column) or the epididymis and testis.

Fig. 2. Upregulation of Hoxa10 and Hoxa11 in MLL/AF10 cells harboring KRAS<sup>G12C</sup>. (A) Heat map representing the relative gene expression levels of 77 shared DEGs between paired cell lines based on cDNA microarray data. Two Affymetrix chips were used in paired cell lines: the 430A (12G vs. AK3G and 12G vs. AK2G) and the Clariom D (AKw1G vs. AK3G). Raw values were log2-transformed and centered relative to the median. A heat map was obtained using Cluster 3.0 and Java TreeView v1.1.6r4. The color bar depicts the color contrast level of the heat map. Red and green indicate high and low expression levels, respectively. (B) Top 10 enriched GO_BP terms for 77 DEGs. All GO terms listed in the table show significant enrichment (all P < 0.05). Red and blue indicate genes that are upregulated or downregulated, respectively, in MLL/AF10 cells harboring KRAS<sup>G12C</sup>. (C) Two clusters of redundant annotation GO_BP terms. GO enrichment and clustering analyses were performed using online DAVID v6.8 annotation tools.
Fig. 3. Silencing Hoxa10 of AK3G cells prolonged survival of leukemic mice. (A, B) The mRNA level of Hoxa10 in 12G, AKw1G, AK2G, and AK3G cells (A), and AK3G-shV, AK3G-shH10-1, and AK3G-shH10-2 cells (B) were determined by RT-qPCR analysis. Data shown are representative of three independent experiments. Error bars indicate SD of the mean. Fold changes are indicated above the bars. (C) The protein level of Hoxa10 in AK3G-shV and AK3G-shH10-2 cells was determined by Western blotting. β-actin was used as the loading control. (D) Kaplan–Meier survival curves of sublethally γ-irradiated mice intraperitoneally injected with AK3G-shV or AK3G-shH10-2 cells. A P value less than 0.05 indicates statistical significance. (E) Gross view of excised gastrointestinal tract and omental-mesentery (GI-OM) adipose tissue from representative AK3G-shV and AK3G-shH10-2 mice. (F) Aligned dot plot showing the percentage weight of GI-OM adipose tissue over the total body weight of AK3G-shV and AK3G-shH10-2 mice. Horizontal lines represent the median, which was not statistically different between AK3G-shV and AK3G-shH10-2 mice (13.9% vs. 13.6%, respectively; P = 0.974).

Identification of DEGs and GO enrichment analysis

To identify the genes responsible for MS formation by leukemia cells in adipose tissues, we compared transcriptome profiling between three pairs of MLL/AF10(OM-LZ) leukemia cells harboring KRASwt and KRAS<sup>G12C</sup> (12G vs. AK3G, 12G vs. AK2G, and AKw1G vs. AK3G) using cDNA microarray data. Seventy-seven shared DEGs with more than 1.5-fold change were identified, consisting of 25 upregulated and 52 downregulated genes in AK2G/AK3G cells (Fig. 2A). Functional annotation of the 77 DEGs was performed using DAVID Bioinformatics Resources v6.8. The top 10 GO_BP terms overrepresented in these 77 DEGs included three related to immune/inflammatory responses, two related to receptor signaling, two related to development/differentiation, one related to endocytosis, one related to intermediate filament bundle assembly, and one related to myocyte adhesion-cell communication (all P < 0.05) (Fig. 2B). Clustering analysis of all GO_BP terms showed that the terms related to development were the most enriched (Fig. 2C). Among the genes mapped to development, Hoxa10 and Hoxa11 were upregulated in AK2G and AK3G cells (Fig. 2A). Since high expression of Hoxa10 and Hoxa11 was observed in AML with MLL-t or t(8;16)(p11;p13), and these two types of AML are frequently associated with MS formation (13, 17), we selected Hoxa10 and Hoxa11 for further investigation of their roles in MS formation and survival.

Involvement of Hoxa10 in survival

To confirm the differential expression of Hoxa10 in MLL/AF10(OM-LZ) leukemia cells based on cDNA microarray data, RT-qPCR analysis was performed and confirmed that AK2G and AK3G cells have higher expression
levels of Hoxa10 compared to 12G and AKw1G cells (Fig. 3A). To investigate the role of Hoxa10, we first generated stable Hoxa10 knockdown AK3G cells by lentiviral transduction of shRNAs targeting Hoxa10. Measurement of Hoxa10 mRNA levels of lentiviral-transduced AK3G-shH10-1 and AK3G-shH10-2 cells revealed 35% and 45% reduction, respectively, compared to control AK3G-shV cells (Fig. 3B). The successful generation of Hoxa10 knockdown clones was further confirmed by Western blotting (Fig. 3C). Data of the BM transplantation assay revealed significantly longer survival of the mice transplanted with AK3G-shH10-2 cells than those with AK3G-shV cells (median 65.5 days vs. 55 days, respectively; P < 0.001) (Fig. 3D and Table 1). However, no differences in MS formation in omental-mesenteric or gonadal adipose tissues were observed between the mice transplanted with AK3G-shH10-2 cells and control AK3G-shV cells (Fig. 3, E–F and Table 1). These results indicated that the upregulation of Hoxa10 by cooperation of MLL/AF10(OM-LZ) with KRASG12C contributes to MS formation.

**Involvement of Hoxa11 in MS formation**

RT-qPCR analysis confirmed the induction of Hoxa11 expression in AK2G, AK3G, and AK5G cells compared to 12G and AKw1G cells (Fig. 4A). Stable Hoxa11 knockdown AK3G clones (AK3G-shH11-1 and AK3G-shH11-2) were generated by lentiviral transduction of shRNAs targeting Hoxa11 to AK3G cells. Compared to control AK3G-shV cells, the reductions of Hoxa11 mRNA and protein levels in AK3G-shH11-1 and AK3G-shH11-2 cells were confirmed by RT-qPCR and Western blotting, respectively (Fig. 4, B–C). It is of note that, although the Hoxa11 knockdown efficiencies of the cells are significant (35% in AK3G-shH11-1 and 25% in AK3G-shH11-2 cells compared to AK3G-shV cells), the expression level of Hoxa11 in AK3G-shH11-2 cells did not drop to the level of no expression as that of 12G or AKw1G cells (Fig. 4, A–B). Data of the BM transplantation assay revealed no significant differences in survival between mice transplanted with AK3G-shV and AK3G-shH11-2 cells (Fig. 4D and Table 1). Nevertheless, compared to AK3G-shV mice, AK3G-shH11-2 mice had significantly reduced MS size in omental adipose tissue (Fig. 4, E–F and Table 1). Our results indicated that the induction of Hoxa11 expression by cooperation of ML1/AF10(OM-LZ) with KRASG12C contributes to MS formation.

**Hoxa11 in leukemia cell migration and retention in vivo and in vitro**

In vivo imaging cell trafficking analysis of mice injected with a 1:1 premix of AK3G-shH11-2 and AK3G-shV cells, using CellVue-NIR815 to label cells of one of the two cell lines at a time, revealed that the fluorescent signals of AK3G-shH11-2 cells rapidly condensed within 5 min, were maintained for 1.5 h, and then decreased 24 h post-transplantation (Fig. 5A, right column). In contrast, the fluorescent signals of AK3G-shV cells condensed after 1.5 h and were maintained for 24 h post-transplantation (Fig. 5A, left column). Excised organs derived from mice 7 days post-transplantation showed that
Fig. 5. Silencing Hoxa11 of AK3G cells enhanced cell migration and reduced cell retention in omental adipose tissue. (A) Imaging cell trafficking analysis of mice ip injected with CellVue NIR815-labeled AK3G-shV or AK3G-shH11-2 cells by Perla Imager at indicated time points. Mice were under anesthesia during image capture. (B) Imaging analysis of excised organs from AK3G-shV and AK3G-shH11-2 mice 7 days post-transplantation by Perla Imager. Color bars (A, B) indicate the total fluorescence radiant efficiency. (C) Competitive engraftment and clonal expansion analysis of AK3G-shV and AK3G-shH11-2 cells in the omental (OM) adipose tissue of recipient mice on day 1 and week 4. AK3G-shV and AK3G-shH11-2 cells were initially mixed at a 1:1 ratio (1 x 10^6 cells/cell line). The arrowhead indicates the 79th nucleotide (C for AK3G-shH11-2 and G for AK3G-shV). (D) Flow cytometric analysis of Cxcr4 receptor protein levels of AK3G-shV and AK3G-shH11-2 cells. Data were acquired using a FACs Canto II flow cytometer and analyzed using FlowJo software. (E) Transwell migration of AK3G-shV and AK3G-shH11-2 cells stimulated by SDF-1 (150 ng/ml). Cells were incubated in a Transwell plate (5-μm pore size) for 6 h. Transmigrated and nonmigrated cells were counted using a hemacytometer. The values shown are expressed as the mean ± SD of three independent experiments. A P value less than 0.05 indicates statistical significance. (F) Cell motility (i) and protrusion (ii) of AK3G-shV and AK3G-shH11-2 cells stimulated by SDF-1 (150 ng/ml) for 0 to 20 min in a confined space (3 μm in size).

The fluorescent signals of AK3G-shH11-2 cells were weaker than that of AK3G-shV cells in omental adipose tissue (Fig. 5B). The reduction of AK3G-shH11-2 cells in omental adipose tissue was further confirmed by competitive engraftment and clonal expansion assay. The amount of PCR product derived from AK3G-shH11-2 cells was less than AK3G-shV cells in omental adipose tissue 1 day and 4 weeks post-transplantation (Fig. 5C). Our results revealed that silencing Hoxa11 of AK3G cells enhances cell migration through the adipocyte septe of omental adipose tissue or the endothelium of blood vessels.

In a previous study, we generated Hoxa11-overexpressing 12G cells (12G-H11-1) by retroviral transduction of full-length Hoxa11 into 12G cells [24], BM transplantation assay revealed that mice transplanted with 12G-H11-1 cells, similar to control 12G-V1 cells, did not induce MS formation [24]. However, in vivo imaging cell trafficking analysis revealed that strong fluorescent signals of 12G-H11-1 cells were detected in omental adipose tissue 5 min and 1.5 h post-transplantation with the strength of the signals reduced after 24 h, whereas low fluorescent signals of 12G-V1 cells were detected at all time points (Supplementary Fig. 2A). Data of competitive engraftment and clonal expansion assay showed that 12G-H11-1 cells were the major cells located in omental and gonadal adipose tissue after 24 h (Supplementary Fig. 2B), further supporting that Hoxa11 reduces leukemia cell migration through adipocyte septa or endothelium and prolongs cell retention in adipose tissues.

Cxcr4/SDF-1 axis-induced cell polarization plays a critical role in hematopoietic stem cell and leukemia cell migration and homing to BM [29,30]. However, flow cytometry analysis showed no significant differences in Cxcr4 expression levels between AK3G-shV and AK3G-shH11-2 cells (Fig. 5D), suggesting Hoxa11-induced reduction in cell migration was not via reducing of Cxcr4/SDF-1 interaction. Next, we compared cell migration between AK3G-shV and AK3G-shH11-2 cells using SDF-1-stimulated transwell migration assay. Our data showed a 29.7% increase by AK3G-shH11-2 cells in transwell migration through a membrane with 5 μm pores (Fig. 5E). Under confined spaces (3 μm in size) with SDF-1-stimulated directional migration, the moving distance of AK3G-shH11-2 cells was significantly further than AK3G-shV cells (Fig. 5F). Additionally, protrusion of AK3G-shH11-2 cells was more active than that of AK3G-shV cells (Fig. 5, F-i). These results indicated that Hoxa11 expression in AK3G cells reduces cell migration and prolongs cell retention in adipose tissue, at least partly, by reducing cell motility and protrusion.
Discussion

The molecular mechanism of MS formation has not been thoroughly investigated. In a previous study, we demonstrated that the upregulation of Adgra3 (previously named Gpr125) in leukemia cells harboring MLLAF10 and KRASG12C-enhanced homotypic cell-cell adhesion in vitro and MS formation in vivo [13]. In this study, we demonstrated that the upregulation of Hoxa11 in leukemia cells enhanced cell retention in omental adipose tissue and MS formation in vivo. Data of SDF-1-stimulated directional transwell migration and cell motility in confined spaces revealed that Hoxa11 expression reduced leukemia cell migration and protrusion. Furthermore, the observed reduction in cell motility in leukemia cells was not via reducing of SDF-1/Cxcr4 interaction. These results indicate that MS formation is associated with increased cell-cell adhesion and reduced migration of leukemia cells.

To our knowledge, there is no report linking HOXA11 with cell migration in hematopoietic malignancies. However, the effect of HOXA11 on cell migration was recurrently reported in solid tumor cells, but with controversial results. HOXA11 overexpression reduces tumor cell migration in endometrial cancer and renal cell carcinoma [31,32], but enhances tumor cell migration in gastric and breast cancer cells [33,34]. These data indicated that HOXA11 exhibits positive or negative control on migration is depending on the context of tumor cell type. Further studies are needed to reveal Hoxa11-mediated diverse mechanisms underlying cell migration. During cell migration, the first step of the process is the formation of cell protrusions, including actin polymerization-driven lamellipodia and contractility-driven blebs [35]. Among the 77 DEGs identified in this study, 6 genes (Gas7, Hck, Mshb1, Myo1f, Plk1, and Csf1r) were mapped to GO_BP terms related to actin filament/bundle polymerization or positive regulation of actin cytoskeleton reorganization (Supplementary Table 1). All of these genes were downregulated in MS-inducing AK2G and AK3G cells (Fig. 2A). Whether the downregulation of these genes affects actin polymerization-driven lamellipodia and contractility-driven bleb formation and whether these genes are regulated by Hoxa11 need further investigation.

In this study, we demonstrated that silencing Hoxa10, but not Hoxa11, in AK3G cells prolonged the survival of recipient mice (Figs. 3D and 4D). Our results support the clinical observations that Hoxa10 overexpression is associated with poor prognosis [18,19]. However, in a previous study, we showed that silencing Hoxa11 expression in MLLAF10(OM-LZ) leukemia cells harboring PTEN1G503A (APm-1) prolonged the survival of recipient mice [24]. These contradictory results may be due to differences in the knockdown efficiency of Hoxa11 in APm-1 and AK3G leukemia cells. In Hoxa11 knockdown APm-1 cells (APm-1-shH11-2), Hoxa11 expression was reduced to the level of cells harboring PTEN1G503A (APw-1 cells). In contrast, gene expression was only partially reduced in Hoxa11 knockdown AK3G cells (Fig. 4, A–B). Therefore, we cannot exclude the possibility that Hoxa11 expression is also critical for the survival of AK3G leukemia mice. MS is considered a poor prognostic factor in AML. Although our results showed that mice transplanted with Hoxa11 knockdown AK3G cells had reduced MS formation, it did not influence survival compared to those transplanted with control cells (Table 1). Based on the same reason, we cannot exclude the possibility that MS formation is correlated with survival in our mouse model. Our results showed that ectopic expression of Hoxa11 in 12G-H11 cells did not induce MS formation in omental adipose tissue, suggesting Hoxa11 alone is insufficient to induce MS formation. More studies are needed to clarify the biological pathway and the molecular mechanism of Hoxa11, as well as Adgra3, in MS formation, which may contribute to the development of new targeted therapies to treat patients with AML and MS.

Supplementary Figure 1. Cooperation of MLLAF10(OM-LZ) with KRASG12C-induced myeloid leukemia with shorter latency and induced MS formation at extramedullary sites. (A) Flowchart for establishing the MLLAF10(OM-LZ) cell line (12G) and the MLLAF10(OM-LZ) cell lines harboring KRASG12C (AKw1G) or oncogenic KRASG12C (AK2G, AK3G, and AK5G). (B) Kaplan–Meier survival curves of mice ip injected with AKw1G (n = 5) and AK5G (n = 5) cells. Mice ip injected with normal saline served as a blank control. (C) Gross view of excited GI-OM adipose tissue from blank, AKw1G, and AK5G mice. Arrows indicate tumor masses.

Supplementary Figure 2. Ectopic expression of Hoxa11 in 12G cells prolonged cell retention in adipose tissue. (A) Imaging analysis of mice ip injected with CellVue NIR815-labeled AK3G-shV or AK3G-shH11-2 cells by Peral Imager at indicated time points. Mice were under anesthesia during image capture. (B) In vivo competitive engrafment and clonal expansion analysis of 12G-V and 12G-H11 cells in OM and Go adipose tissues of recipient mice on the first day: 12G-V and 12G-H11 cells were initially mixed in a 1:1 ratio by cell number. The arrowhead indicates the 62nd nucleotide (C for 12G-H11 and G for 12G-V).

Declaration of Competing Interest

The authors declare no competing financial interests.

CRediT authorship contribution statement

Jen-Fen Fu: Conceptualization, Data curation, Funding acquisition, Investigation, Methodology, Project administration, Validation, Writing – original draft. Chih-Jen Wen: Methodology, Data curation, Formal analysis, Visualization. Tsung-Hai Yen: Methodology, Software, Visualization. Lee-Yung Shih: Conceptualization, Data curation, Resources, Writing – review & editing.

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