Acute megakaryoblastic leukemia (AMKL) is a form of acute myeloid leukemia (AML) associated with a poor prognosis. The genetics and pathophysiology of AMKL are not well understood. We generated a knockin mouse model of the one twenty-two–megakaryocytic acute leukemia (OTT-MAL) fusion oncogene that results from the t(1;22)(p13;q13) translocation specifically associated with a subtype of pediatric AMKL. We report here that OTT-MAL expression deregulated transcriptional activity of the canonical Notch signaling pathway transcription factor recombination signal binding protein for immunoglobulin κ J region (RBPJ) and caused abnormal fetal megakaryopoiesis. Furthermore, cooperation between OTT-MAL and an activating mutation of the thrombopoietin receptor myeloproliferative leukemia virus oncogene (MPL) efficiently induced a short-latency AMKL that recapitulated all the features of human AMKL, including megakaryoblast hyperproliferation and maturation block, thrombocytopenia, organomegaly, and extensive fibrosis. Our results establish that concomitant activation of RBPJ (Notch signaling) and MPL (cytokine signaling) transforms cells of the megakaryocytic lineage and suggest that specific targeting of these pathways could be of therapeutic value for human AMKL.

### Introduction

Acute megakaryoblastic leukemia (AMKL) is a heterogeneous subtype of acute myeloid leukemia (AML) and is more frequent in children than in adults (1–3). The molecular basis of AMKL is poorly understood in adults, whereas 2 major molecular subtypes are recognized in pediatric AMKL. The first group is represented by Down syndrome (DS) patients with both transient myeloproliferative disease (transient MPD) and AMKL who present with acquired GATA-binding protein 1 (GATA1) mutations, resulting in an N-terminal truncated GATA1 short (GATA1s) protein (4). The second group occurs in infants and is associated with the t(1;22)(p13;q13) chromosomal translocation, resulting in expression of the one twenty-two megakaryocytic acute leukemia (OTT-MAL) (also known as RBM15-MKL1) fusion protein (5–7). MAL is a cofactor of the serum response factor (SRF) endowed with strong transcriptional activation properties (8), and its localization and function are deregulated by fusion with OTT (9). OTT is related to the SHARP transcription factor that has been shown to interact with recombination signal binding protein for immunoglobulin κ J region (RBPJ), the transcription factor involved in response to the canonical Notch pathway, and repress transcription of its target genes in absence of Notch signaling (10–13).

The thrombopoietin (TPO) receptor myeloproliferative leukemia virus oncogene (MPL) and its ligand, TPO, are essential for the proliferation of megakaryocyte (Mk) progenitors (MkPs) and their differentiation into mature platelet-producing Mks (14). Ligand stimulation of MPL results in activation of the Janus kinases JAK2 and TYK2, followed by docking and phosphorylation of a spectrum of signaling proteins (15–17). The relevance of this pathway to pathogenesis of AMKL is highlighted by demonstration of activating point mutations in JAK2 or JAK3 (18–20) and in MPL (21). In addition, secondary AMKL may also occur with leukemic transformation of chronic myeloproliferative syndromes, including polycythemia vera, essential thrombocytosis, and myeloid metaplasia, that are frequently associated with activating mutations in JAK2 or MPL (22, 23). These observations indicate that activating mutations in the TPO/MPL/JAK signaling pathway play an important role in Mk malignancies in humans.
To gain insight into the requirements for AMKL leukemogenesis, we generated a conditional knockin allele of Ott-MAL in mice. We observe that OTT-MAL expression from its endogenous promoter results in aberrant megakaryopoiesis associated with enhanced self renewal and activation of RBPJ-mediated transcription. In addition, analysis of gene expression data from infants with AMKL supports a role for deregulation of Notch signaling in transformation of the megakaryocytic lineage. We further observe that expression of OTT-MAL with concomitant Notch signaling deregulation cooperates with an activating allele of MPL to induce a highly penetrant AMKL that recapitulates the human phenotype.

Results

Conditional expression of Ott-MAL in a knockin mouse model. We engineered a conditional Ott-MAL knockin mouse model that allowed for expression of the fusion protein from the endogenous murine Ott promoter after Cre recombinase-mediated excision of a STOP cassette flanked by loxP recombination sites (floxed-STOP; Figure 2A). Southern blot analysis of XmnI-digested ES cell DNA identified clones with homologous recombination (Figure 1B and C). Functional integrity of the targeted construct was confirmed by transducing Ottlox/WT ES clones with adenovirus encoding the Cre recombinase, followed by selection of OttCre-MAL excised clones based on Southern blot analysis (Figure 1D). Conditional expression of the fusion mRNA and protein was confirmed by RT-PCR and Western blot analysis, respectively (Figure 1E and F). Animals with germline transmission of the Ottlox allele were crossed to an EIIA-Cre transgenic mouse line, resulting in in vivo excision of the floxed-STOP cassette, with germline transmission of the OttCre-MAL allele. OttOtt-MAL/WT animals (term OM) were observed at predicted Mendelian ratios and were backcrossed into the C57BL/6 background for further analysis.

OTT-MAL expression results in abnormal fetal and adult hematopoiesis but induces AMKL with low penetrance. As the OTT-MAL fusion is restricted to infant leukemias and is likely to occur in utero, we first analyzed fetal liver hematopoiesis using flow cytometry. Plating of purified Lineage Sca1+ c-Kit+ (LSK) cells from E12.5 fetal livers revealed a significant increase in colony-forming efficiency due to expansion of multipotent myeloid progenitors (CFU–granulocyte-macrophage-erythroid-Mk [CFU-GEMM]) in OM versus WT cells (Figure 2A), although no differences were observed in the absolute number of hematopoietic progenitors or CD41+ cells between OM and WT fetal liver at day E12.5 (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI35901DS1; and data not shown). Furthermore, replating of LSK cell–derived primary colonies from OM embryos showed an increase in both Mk-containing (CFU-GEMM) and pure Mk (CFU-Mk) secondary colonies (Figure 2B), whereas no CFU-Mk were obtained with plating of WT cells. Identification of Mk-containing colonies was confirmed by staining cytopsins of single colonies for acetylcholinesterase (AchE) activity (data not shown). These data indicate that OTT-MAL expression results in aberrant differentiation of HSC toward the megakaryocytic lineage in vitro.

We next investigated the effect of OTT-MAL expression on adult hematopoiesis. As OM animals did not show hematologic abnormalities during the first 6 months of life (data not shown), we studied an older cohort of 15 OM, 10 Ottlox/WT, and 10 WT littermate animals aged 18–24 months. Since extramedullary hematopoiesis leading to splenomegaly is a hallmark of AMKL in humans, we first evaluated the number of progenitors in splenocytes and observed an increase in OM animals (Figure 2C). In addition, these cells also showed enhanced replating activity in the majority of OM animals (8/15), whereas no replating activity was observed in cells derived from WT littermate (0/10) or Ottlox animals (0/10) (Figure 2C). Colonies derived from OM animals at the fifth round of replating showed AchE+ cells, indicating their megakaryocytic nature (data not shown). In addition, histopathologic analysis revealed that animals with increased replating activity of splenocytes had abnormalities in the hematopoietic system including extramedullary

Figure 1

Ott-MAL knockin approach. (A) Mouse Ott locus (showing splicing between exon 1 and 2). OTT-MAL targeting construct/Ottlox allele, and OM allele are represented. OMF and OMR show locations of primers used for detection of the fusion transcript by RT-PCR. X, XmnI site. (B and C) Southern blot analysis of ES clone DNA using XmnI digestion show correct homologous recombination on the 5’ and 3’ sides of the targeting construct, respectively. Rec, recombinated. (D–F) Southern blot, RT-PCR, and Western blot analysis, respectively, on ES clones obtained after adenov-Cre–mediated excision of the Hygro-STOP cassette. Arrow indicates the OTT-MAL (OM) fusion protein. Asterisk indicates nonspecific band; IVT, in vitro transcription control.
Figure 2
OTT-MAL induces abnormal fetal and adult hematopoiesis and AMKL with low penetrance. (A) Plating efficiency of LSK cells. Mean ± SD of the number and type of colony counted after 7 days in triplicate is shown. (B) Colonies from A were replated into fresh methylcellulose plates and were scored 7 days later. Mean ± SD of 3 independent experiments is shown. Mk, CFU-Mk; E, blast-forming unit–erythroid; GM, CFU–granulocyte-macrophage. (C) Serial replating assays of splenocytes from 18- to 24-month-old OM, Ottlox/WT, and WT littermate animals. Mean value of the number of colonies is indicated. (D) Kaplan-Meier survival curves of secondary (II prv) and tertiary (III prv) recipients injected with $1 \times 10^6$ splenocytes from 6127 (WT) and 6133 (OM). (E) Immunohistochemistry on BM sections from secondary recipients using vWF antibody. Cells expressing vWF show a dark brown coloration. Original magnification, ×100; ×1000 (insets). (F) Proliferation of 6133 cells in absence or presence of SCF (10 ng/ml). Of note, proliferation of 6133 cells was not dependent on TPO. (G) Wright-Giemsa staining of the 6133 cell line. Original magnification, ×1000. (H) Flow cytometrical immunophenotype of 6133 cells. Percentages of live cells are indicated.
hematopoiesis in the spleen and liver (6/15) and frank leukemia (2/15: nos. 6133 and 8904). These leukemias had phenotypic attributes of human AMKL, including infiltration of the BM, spleen, liver, and kidney with an admixture of immature megakaryocytic and erythroid elements as well as megakaryoblasts circulating in the peripheral blood (Supplemental Figure 2). The disease was transplantable into secondary and tertiary recipients, with a median survival of 45 days and 24 days, respectively (Figure 2D). Secondary recipients showed a phenotype similar to that of primary animals, with infiltrating leukemic blasts that were vWF+, confirming the megakaryoblastic nature of the disease (Figure 2E and Supplemental Figure 3).

A cytokine-dependent cell line was derived from leukemic cells of animal no. 6133 (Figure 2F). Similar to primary leukemic cells from this animal, the 6133 cell line displayed an immature phenotype (Figure 2G). These cells were positive for c-Kit as well as several megakaryocytic markers, including CD41 and CD42b, and were negative for myeloid markers, including Mac1 and FcεR1a.
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OTT-MAL activates RBPJ-mediated transcriptional activity. OTT belongs to a family of proteins that interact with the transcription factor RBPJ (11–13, 24), and constitutive activation of RBPJ-mediated transcription due to Notch signaling has been reported to result in increased self renewal and cancer (25, 26). To better understand the mechanism of megakaryocytic transformation by OTT-MAL, we determined whether OTT-MAL could interfere with RBPJ-mediated transcriptional regulation. Coimmunoprecipitation experiments in 293T cells showed that OTT-MAL formed a complex with RBPJ that required the RNA recognition motif (RRM) domains of OTT-MAL (Figure 3, A and B). Furthermore, transactivation assays in 293T cells using an RBPJ luciferase reporter showed that OTT-MAL activated RBPJ-mediated transcription in a dose-dependent manner, whereas expression of OTT resulted in a subtle inhibition of RBPJ transcription (Figure 3C) (13). Transactivation of the RBPJ reporter by OTT-MAL was dependent on the transactivation domain (TAD) of MAL and the RRM domains of OTT (Figure 3, B and C).

To investigate whether the RBPJ target genes could be transcriptionally activated by OTT-MAL in vivo, we first crossed OM animals with an established transgenic Notch reporter (TNR) mouse model, in which the expression of a GFP reporter is under the control of RBPJ response elements (27, 28). GFP fluorescence was higher in lineage-negative hematopoietic cells from double OM+TNR animals than in cells from control TNR animals (Figure 3D). In addition, primary lineage negative hematopoietic cells from OM animals showed increased expression of Hes1, Hes5, and Dtx1 transcripts compared with WT animals (Figure 3E), confirming activation of RBPJ transcription by OTT-MAL in vivo. Of note, Gata1 and Gata2 transcripts were also upregulated in OM versus WT cells (Supplemental Figure 4A). These results demonstrate increased RBPJ transcriptional activity in the hematopoietic progenitors of OM animals with respect to WT controls.

We next assessed whether OTT-MAL activation of RBPJ transcription is involved in the proliferation of 6133 AMKL cells. We performed communoprecipitation to confirm that endogenous OTT-MAL but not MAL interacted with RBPJ in the 6133 AMKL cell line (Figure 3F). In addition, chromatin immunoprecipitation analyses indicated that MAL and RBPJ were present at the HES1 promoter in 6133 cells but were not present at the HES1 promoter in control Ba/F3 cells that do not express OTT-MAL (Figure 3G). As confirmation, we also transduced 6133 cells with a dominant negative (dn) form of RBPJ (dnRBPJ) that is unable to bind DNA (29) or with a dn Mastermind-like 1 (dnMAML1) mutant that entraps intracellular Notch (ICN) and RBPJ in transcriptionally inactive complexes when Notch signaling is activated (30, 31). Compared with empty vector control, dnRBPJ expression specifically inhibited the growth of 6133 AMKL cells but not control...
Ba/F3 cells (Figure 3H and Supplemental Figure 4). In contrast, dnMAML1 or γ-secretase inhibitors had no significant effect on growth of 6133 or Ba/F3 cells (Supplemental Figure 4 and data not shown), suggesting that the expression of RBPJ target genes is not dependent on the activation of Notch receptors but is due to direct interaction between OTT-MAL and RBPJ. Taken together, these results indicate that RBPJ-mediated transcription is aberrantly induced by OTT-MAL in vitro and in vivo and is required for the growth of OTT-MAL–transformed AMKL cells.

An RBPJ pathway signature is upregulated in t(1;22)(p13;q13) AMKL. This mouse model expressing OTT-MAL indicates that aberrant activation of the RBPJ transcription factor, a mediator of the canonical Notch signaling pathway, is important for leukemic transformation of the Mk lineage. To confirm the role of the pathway in human AMKL, we analyzed the expression of RBPJ pathway genes in human AMKL associated with t(1;22)(p13;q13) (OM-AMKL) compared with DS-AMKL, the other molecularly defined subgroup of childhood AMKL. Using published global expression data from AMKL samples (32), we performed Gene Set Enrichment Analysis (GSEA) using a list of genes implicated in the RBPJ pathway (Supplemental Table 1). GSEA results showed that a RBPJ pathway signature was significantly enriched in OM-AMKL compared with DS-AMKL (Figure 4A). The most upregulated genes in OM-AMKL included Notch1, Hes1, and Gata1.
inhibitors (Figure 5G). Taken together, these results indicate that megakaryoblastic transformation and increases led to a rapid increase in Hes1 transcript but not Gata1 transcript that proliferation was mediated by activation of the TPO/MPL signaling pathway rather than specifically by the MPL W515L mutant. To express the OTT-MAL fusion.

In both the WT and OM contexts, the animals showed comparable degrees of leukocytosis, polycythemia, thrombocytopenia, and splenomegaly, whereas the degree of hepatomegaly was significantly increased when OM cells were used (Figure 6, A and B, and Supplemental Figure 6). In contrast, when MPL W515L and OTT-MAL were coexpressed, flow cytometric analysis of splenocytes showed a significant shift toward immature progenitors as assessed by a marked increase in c-Ki67+ staining in OM+MPL W515L animals (Figure 6C and Supplemental Figure 7A), and the disease phenotype was more severe in this context, as indicated by a substantial increase in the number of CD41+, CD71+Ter119+, and Mac1+Gr1+ cells when compared with expression of MPL W515L alone. We next performed multiparameter flow analysis on BM or spleen cells, respectively, to phenotypically characterize hematopoietic progenitor populations. In contrast with WT+MPL W515L animals, which showed an increase in the common myeloid progenitor (CMP) population, OM+MPL W515L animals showed a preferential increase in the Mk-erythrocyte progenitor (MEP) population in both BM and spleen (Figure 6D and Supplemental Figure 8). In addition, MkP that were specifically engaged in the Ml differentiation downstream of the MEP (37) were markedly expanded in OM+MPL W515L animals compared with WT+MPL W515L animals (Figure 6E and Supplemental Figure 8).

Histopathologic analysis of BM, spleen, and liver from WT+MPL W515L or OM+MPL W515L animals showed infiltration with an admixture of megakaryocytic, erythroid, and myeloid elements (data not shown). Immunohistochemistry for vWF on spleen sections showed an increase in the number of mature polyploid Mk s in WT+MPL W515L animals, whereas OM+MPL W515L animals showed, in addition, the presence of numerous hypolobated immature megakaryoblasts (Figure 6F and Supplemental Figure 9). Together, these data demonstrate that in the WT C57BL/6 background, MPL W515L induces an MPD characterized by thrombocytosis and associated with expansion of the CMP population. In contrast, coexpression of MPL W515L with OTT-MAL resulted in expansion of the MEP and MkP progenitor populations and evidence of transformation of the MPD phenotype into AMKL.

To further annotate malignant transformation, secondary transplantation was performed into sublethally irradiated recipients using 1 × 10^6 splenocytes from primary animals. Secondary OM+MPL W515L animals developed a disease with leukocytosis and disease latency similar to that observed in the primary OM+MPL W515L animals (Figure 7, A and C). However, in striking contrast with primary recipients, secondary recipients developed thrombocytopenia (Figure 7B). In control experiments, secondary WT+MPL W515L or OM+MPL W515L animals showed no sign of disease for over 200 days, indicating that the MPL W515L-induced MPD was not transplantable. Compared with control animals, flow cytometry analysis of secondary OM+MPL W515L splenocytes showed abnormal CD41+, CD42b+, CD41/CD61+, or CD9+ populations, a fraction of which were also c-Ki67+ (Figure 7D, Supplemental Figure 7B, and Supplemental Figure 10). These cells also expressed the CD71 marker but not Ter119, Mac1, or Gr1 markers. In agreement with these observations, histopathologic analysis demonstrated that BM, spleen, and liver were heavily infiltrated with immature hypolobated vWF+ megakaryoblasts (Figure 7E and Supplemental Figure 10), with a striking reduction in median polyploidy (Figure 7F) and fibrosis (Figure 7G and Supplemental Figure 11). Southern blot analysis of DNA from BM cells showed over 10 viral integrations in primary and secondary animals and suggests that the disease is oligo/polyclonal (Supplemental Figure 12). Transplantation into tertiary recipients resulted in a similar phenotype as in the secondary recipients of OM+MPL W515L cells, albeit with a shorter latency (median = 38 days; Figure 7C). Collectively, these data show that coexpression of OTT-MAL and an activated MPL mutant induces AMKL with characteristic features of the human disease.
Figure 6
Coexpression of MPL\textsuperscript{W515L} with OTT-MAL results in AMKL in vivo. (A) Representative pictures of spleens from animals transplanted with OM or WT BM cells transduced with MPL\textsuperscript{W515L} at 60 days after transplant. (B) Mean ± SD of liver weights (n = 5). (C) Flow cytometry analysis of splenocytes at 60 days after transplantation. Experiments were performed in triplicate, and a representative analysis is shown. Percentages of live cells are indicated. (D) Multiparameter flow analysis of BM CMP, granulocyte-macrophage progenitor (GMP), and MEP populations. Experiment was performed in duplicate and a representative analysis gated on PI\textsuperscript{–}Lin\textsuperscript{–}Sca\textsuperscript{–}c-Kit\textsuperscript{+} cells is shown. Percentages of PI-Lin-Sca-Kit\textsuperscript{+} cells are indicated. (E) Multiparameter flow analysis of the BM MkP population. MkP are defined as PI-Lin-Kit\textsuperscript{+}CD9\textsuperscript{+}Fc\textgamma{RII/III}CD41\textsuperscript{+}. Percentages of Lin-Kit\textsuperscript{+}CD9\textsuperscript{+} cells are indicated. (F) vWF immunohistochemistry on spleen sections. Central panels show maturing polyploid Mks. Far right panel shows abnormal immature hypoploid Mks. Original magnification, ×1000.
Discussion

AMKL is generally associated with a poor prognosis, and its frequency may be underrepresented because of its complex clinical presentation and the difficulty in interpreting diagnostic material (38). Challenges in understanding the molecular basis of AMKL derive in part from the difficulty in obtaining adequate tissue samples derived from the fibrotic BM of patients. The identification of the OTT-MAL fusion in a subset of AMKL (5–7) provided a basis for development of a murine model of disease that could be used to increase understanding of the molecular pathogenesis of AMKL and potentially to provide insights into alternative therapeutic approaches.

We have observed that expression of OTT-MAL from its endogenous promoter results in abnormal megakaryopoiesis during embryonic and adult development but rarely causes AMKL. Several observations could explain the low efficiency of transformation of the Mk lineage in the context of the OTT-MAL fusion alone. It is possible that mutations associated with infant mega-
karyoblastic leukemia preferentially target the fetal Mk compartment but have minimal phenotypic effects and transforming efficiency in the adult hematopoietic compartment. Biological differences between adult and fetal Mk precursors (39) as well as different microenvironmental influences may thus be crucial for transformation of the megakaryocytic lineage. In support of this hypothesis, other human Mk lineage disorders, including thrombocytopenia associated with the thrombocytopenia and aplasia of the radius (TAR) syndrome or the transient MPD in DS patients, are observed in infancy but often resolve spontaneously during childhood (40, 41). In addition, similar findings were reported for a Gata1 knockout mouse model, in which developmentally restricted abnormalities in megakaryopoiesis were observed but not leukemic transformation (42).

The observation that OTT-MAL animals develop malignancy at a low frequency underscores the requirement for cooperating mutations to achieve transformation to AMKL. In support of this hypothesis, several mutations in tyrosine kinases and signaling molecules have been identified in patients with AMKL and Mk malignancies (18, 19, 22, 23). We used a candidate gene approach to address the potential for cooperativity based on previously reported activating alleles in other megakaryocytic malignancies. Our data indicate that OTT-MAL cooperates only with a limited subset of activated signaling molecules. In particular, we observed that coexpression of OTT-MAL with an MPL mutant generates a fully penetrant and rapidly fatal disease closely resembling human AMKL. MPL is essential for normal Mk development, and activating MPL mutations are found in a spectrum of Mk disorders, including AMKL (21, 23, 43). Our data indicate that a basis for complementation of OTT-MAL with activated MPL might lie in the strong activation of the MAPK pathway. In keeping with this hypothesis, MAPK pathway activation has been shown to favor Mk differentiation (44, 45) and to control the activity of serum response factor and its cofactors (8, 9, 46), suggesting that it could synergize with OTT-MAL. Together, these observations suggest that AMKL pathogenesis by OTT-MAL may be associated with pathogenetic events leading to the activation of the MAPK pathway.

One of the most intriguing pathogenetic insights gleaned from the OTT-MAL mouse model is the involvement of the RBPJ transcription factor in transformation of the megakaryocytic lineage through interaction with OTT-MAL and aberrant increases of RBPJ transcriptional activity. Activation of RBPJ-mediated transcription by OTT-MAL is also required for the growth of AMKL cells derived from leukemic animals. Thus, the transcriptional program engaged by RBPJ is important for sustaining the proliferation of OTT-MAL–transformed cells. Of note, OTT was recently reported to modulate Notch-induced transcriptional activation and thereby affect myeloid differentiation (13, 47). The N-terminal portion of OTT, containing the RRM domains, is required for interaction of OTT-MAL with RBPJ, and the TAD of MAL (8) is necessary for activation of RBPJ transcription, suggesting that OTT-MAL aberrantly recruits transcriptional coactivators to regulatory elements of genes normally controlled by RBPJ in the absence of Notch signaling (Figure 7H). Although RBPJ activation by OTT-MAL may not strictly mimic activation by the canonical Notch pathway and the essential RBPJ targets for leukemogenesis by OTT-MAL remain to be identified, this mechanism is reminiscent of constitutive activation of RBPJ transcription by several viral proteins including EBNA2 (Epstein-Barr virus) and RTA (Kaposi virus) (48).

RBPJ is an essential mediator of the canonical Notch pathway that has been implicated in normal developmental processes (33, 49) and in cancer (26, 50). We have recently demonstrated that canonical Notch signaling specifies Mk fate of HSCs through RBPJ-dependent mechanisms (51). We now show that several RBPJ pathway genes are specifically upregulated in human AMKL patient cells associated with the t(1;22)(p13;q13) as compared with DS-AMKL, the other subgroup of childhood AMKL. In our knockin model, we show that OTT-MAL–expressing hematopoietic progenitors have a differentiation bias toward the megakaryocytic lineage. Together, these results support the notion that aberrant activation of RBPJ transcription by OTT-MAL plays a role specifically in Mk transformation by skewing commitment of hematopoietic progenitors preferentially to the megakaryocytic lineage.

In toto, this knockin model of OTT-MAL expression provides important biological insights into what we believe is a novel mechanism of leukemogenesis of the Mk lineage and should be a useful tool for testing therapeutic approaches for the treatment of human AMKL that may include inhibition of RBPJ and the MAPK pathway, among others.

Methods

Targeting construct. A 15-kb genomic DNA fragment containing the murine Ott locus was obtained by screening of a phage λ mouse genomic DNA library. A 6.9-kb EcoRI-BamHI fragment representing the 5′ homology arm was subcloned into a pBluescript KS vector along with a 5.4-kb BamHI-AvrII fragment representing the 3′ homology arm. The human MAL cDNA (NCBI nucleotide sequence number AJ297257) was obtained by screening of a phage λ leukocytes cDNA library (Stratagene; gift from S. Gisselbrecht and G. Courtois, INSERM, Paris, France). An Xhol-BamHI fragment containing exons 4 to 15 of MAL was introduced into the pBS302 vector (Life Technologies) in 3′ of the transcriptional termination STOP cassette. A Salt-Sall fragment containing a loxP site was then introduced in between the STOP cassette and the MAL cDNA. A 2.1-kb EcoRI-HindIII fragment containing part of the Ott first intron with a BgIII restriction enzyme site and Ott second exon was then introduced in the pBS302 vector upstream of the STOP cassette. A BamHI-BamHI fragment containing a floxed Hygromycin resistance cassette was then introduced into the BgIII site, creating a loxP-Hygromycin-loxP-Ott Exon2-loxP-MAL cDNA. A BamHI fragment containing this loxP-Hygromycin-loxP-Ott Exon2-loxP-MAL cDNA was finally introduced into the BamHI site of the vector containing the 5′ and 3′ homology arms. All intermediates and final constructs were entirely resequenced.

ES cell culture and screening. A purified 21-kb Sall-NotI targeting fragment was electroporated into R1 ES cells by the Dartmouth College Transgenic Core (Dartmouth, New Hampshire, USA). For Southern blot analyses, genomic DNA from individual clones was digested with the restriction enzyme XmnI. Ottlac/WT ES cell clones were infected with an adenovirus encoding the Cre recombinase (MOI of 100). Fresh medium was added 2 hours later. 24 hours after infection, cells were trypsinized and plated at 1,000 cells/100-mm Petri dish. Individual colonies were replated after 5 days and expanded, and DNA was extracted for screening by Southern blot as described above. C57BL/6 blastocyst injection of ES cells presenting a normal karyotype was performed by the Brigham and Women’s Hospital Transgenic Core. Chimeras were bred with C57BL/6 mice (Charles River) for germline transmission. Ottlac/WT animals were bred with Eila-Cre transgenic mice (The Jackson Laboratory) to achieve transmission of the OTT-OTT-MAL allele through germline excision. Subsequently, OM mice were backcrossed at least 5 generations with WT C57BL/6 animals.
Robust multiarray analysis (RMA) algorithm (52) using BioConductor.

For immunoprecipitation experiments, full-length or mutant HA-tagged OTT-MAL constructs were transfected into HEK293 cells. Protein lysates (1,000 µg) were collected 48 hours later by Dual-Luciferase Reporter Assay Kit (Promega), and histograms showed normalized results to pRL-TK luciferase. TNR mice (27, 28) expressing the eGFP under the control of 4 tandem copies of the RBPJ binding sites were obtained from The Jackson Laboratory (no. 005854) and crossed to OM mice.

Microarray data analysis. For microarray analysis, we used a published data set of AMKLs (32) (Gene Expression Omnibus accession number GSE4119). We collected raw global gene expression values (from Affymetrix GeneChipsU133A) for t(1;22)(p13;q13) AMKL (OM-AMKL) samples and DS-AMKL samples. We processed these samples with the robust multiarray analysis (RMA) algorithm (52) using BioConductor software, version 2.3 (53). Using the processed data, we then performed GSEA (http://www.broad.mit.edu/gsea/) (54) to look for enrichment of a list of Notch pathway genes (Supplemental Table 1). The GSEA algorithm identified a leading edge set of the genes that best distinguished OM-AMKL from DS-AMKL.

Immunoprecipitation and Western blot analysis. Cells were lysed in 20 mM Tris, pH 7.4, 1% Triton X-100, 150 mM NaCl, 1 mM EDTA, 10% glycerol (Complete; Roche), 1 mM PMSF, 1 mM sodium orthovanadate, and 0.1 mM phenylarsine oxide. Lysates were normalized for protein content (BCA protein assay; Pierce, Thermo Scientific). For direct Western blot, 500 µg (Figure 1) of total proteins was used following standard protocols. Anti-MAL antibodies (home-made no. 2346 directed against human MAL Ct portion or sc-21558; Santa Cruz Biotechnology Inc.) were used. For immunoprecipitation experiments, full-length or mutant HA-tagged OTT-MAL constructs were transfected into HEK293 cells. Protein lysates (1,000 µg) collected 48 hours later were immunoprecipitated with either RBPJ (sc-28713; Santa Cruz Biotechnology Inc.) or STAT5 (sc-835; Santa Cruz Biotechnology Inc.). Eluate was analyzed by Western blot for HA (2362; Cell Signaling Technology) and RBPJ. 50 µg of protein lysate was analyzed by direct Western blot to check for input. PD98059 (Calbiochem) was resuspended in DMSO, and an identical concentration of OM-AMKL was used as control.

Chromatin immunoprecipitation analysis. 2 × 10^6 6133 or Ba/F3 cells were used for each chromatin immunoprecipitation, with 2 µg of each antibody and following standard procedures.

Hes1 promoter primers used for quantitative PCR were as follows: Hes1-F: GCCAAGCCGAGAGAGAGGATTG; and Hes1-R: AGGAGAGAGAGTAGACAGGGGATTC. Flow cytometric analysis. Antibodies were purchased from BD except for FcRRII/III-PECy7 (Abcam) and goat anti-rabbit Alexa Fluor 647 (Molecular Probes; Invitrogen). Staining for flow cytometry was performed in 1x PBS with 2% PBS, except when otherwise mentioned. HSCs were stained for multiparameter flow analysis as previously described (20, 55), then analyzed on a highly modified double-laser (488-nm/350-nm, Enterprise II; and 647-nm, Spectrum; DakoCyton) FACS (Moflo-MLS; DakoCyto- mation or FACSARia; BD Biosciences). Cells populations were defined as follows: HSC, Lin IL7R Sca-1 `1-C Kin`; CMP, Lin Sca-1 `1-C Kin` CD34 `FcRRII/III`; granulocyte-macrophage progenitor (GMP), Lin Sca-1 `1-C Kin` CD34 `FcRRII/ III`; MEP, Lin Sca-1 `1-C Kin` CD34 `FcRRII/III`. 100 HSC (Lin Sca-1 `1-C Kin`) were double sorted directly onto plates containing MethoCult M3434 (STEMCELL Technologies) supplemented with 10 ng/ml rhLH-11, 10 ng/ml rmGM-CSF, and 10 ng/ml rmTPO (STEMCELL Technologies). Colonies on triplicate plates were scored 7 days later.

MkP analysis was performed as previously described (37) with different fluorochrome-coupled antibodies to accommodate the presence of GFP in the cells: PECy5.5-conjugated lineage antibodies (Gr1, Ter119, CD19, B220, CD4, CD8), and CD3; biotin-CD9/APC-Cy7-avidin; CD41-PE; e-Kir-APC; and FcRRII/III-PECy7. MkPs were defined as Lin `1-C Kin` CD9 `FcRRII/III` CD41.

For ploidy analysis, cells were first stained with CD41 antibody, then stained with APC-conjugated secondary antibody. Cells were then incubated 30 minutes in a 0.1% sodium citrate solution containing 50 µg/ml of RNAse and 50 µg/ml of propidium iodide. Analysis of ploidy was performed using 1 × 10^6 CD41+ cells. For phospho-flow analysis, 6133 cells were transduced with the different mutant viruses and analysis was performed 48 hours later. Cells were starved for 2 hours in RPMI 1640 plus 1% BSA for 2 hours, fixed with 1% methanol-formaldehyde (Polysciences Inc.), permeabilized with 100% methanol, and stained with phospho-specific antibodies (Cell Signaling Technologies); p-ERK1/2 (no. 9101), p-STAT3 (no. 9131), p-STAT5 (no. 9351), p-S6 (no. 2211) followed by secondary goat anti-rabbit Alexa Fluor 647 antibody (Molecular Probes; Invitrogen). Analysis was gated on GFP+ cells. Acquisition of the data was performed on a FACS Calibur (BD Biosciences); data were analyzed with FlowJo software, version 8.5.3.

Colony assay. Single-cell suspensions were prepared from E12.5 fetal livers by passing cells sequentially through 18-gauge, 21-gauge, and 27-gauge needles. Cells were incubated with Red Blood Cells Lysis Buffer (Puregene; QIAGEN) for 5’ on ice and then resuspended in culture medium.

For myeloid colony assays, 2 × 10^4 cells were plated in MethoCult 3434 (STEMCELL Technologies) supplemented with TPO, and IL-11 and colonies were scored after 7 days. For Mk colony assays, 2.2 × 10^4 cells were mixed with Megacult-C (STEMCELL Technologies) containing TPO, IL-11, IL-3, and IL-6, according to manufacturer’s recommendation, and plated onto 2 double-chamber culture slides; colonies were enumerated after 8 days.

Staining and immunohistochemistry. AChE staining of Megacult cultures or cytospun cells was performed following manufacturer’s recommendations (STEMCELL Technologies). Blood smears were stained with Wright-Giemsa Stain (Sigma-Aldrich). H&E staining, reticulin staining, and 1000 images were obtained with a Nikon Eclipse E400 microscope equipped with SPOT RT color digital camera (model 2.1.1; Diagnostic Instruments Inc.). The microscope was equipped with a 10x/22 NA ocular lens. x100 images were obtained with a 10x/0.25 objective lens, x1000 images were obtained with a 100x/1.4 objective lens with oil (Trak-300; Richard-Allan Scientific). Images were analyzed using Adobe Photoshop 6.0.

Transplantation and viral infection. 1 × 10^6 spleenocytes from 6133 (OTT<sup>WT-MAL</sup>) and 6127 (WT littermate) animals were injected into sublethally irradiated SCID recipients (C.B. 17 Fox Chase SCID, CD1<sup>7SC.M</sup> Taconic). For cooperation studies, viral supernatants were obtained as described (19, 20). 6133 cells were mixed with viral supernatants, spininfected for 90 minutes at 1800 g, and returned to the incubator for 3 to 5 hours. 48 hours after infection, cells were transferred to SCF-free medium, and viable cells were counted every day using trypan blue staining. For BM transplant, 8- to 10-week-old OTT<sup>WT-MAL</sup> or WT littermate donor mice

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were injected with 5-FU 5 days prior to BM collection. On day 0, primary BM cells were obtained from femurs and tibiae and cultured overnight in RPMI 1640 supplemented with 10% FBS+IL-3, IL-6, and SCF after lysis of red blood cells. Cells were mixed with identical titer viral supernatants 3 times on day 1 and day 2 and spininfected for 90 minutes at 1800 g each time. After the second spinfection, 1 × 10³ cells were injected in the tail veins of lethally irradiated C57BL/6 recipients. Nonlethal eye bleeds were performed using EDTA or heparin-coated capillary tubes, and blood counts were performed within 30 minutes on a HemaVet HV950FS (Drew Scientific Group). Approval for the use of animals in this study was granted by the Children’s Hospital Boston Animal Care and Use Committee (Boston, Massachusetts, USA).

Statistics. 2-tailed Student’s t tests were performed using the program GraphPad Prism, version 4.0c (GraphPad Software); P < 0.05 was considered significant.

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Address correspondence to: Olivier A. Bernard, INSERM EM10210, Tour Pasteur—Hôpital Necker, 149 rue de Sèvres, 75743 Paris Cedex 15, France. Phone: 33-1-44-49-58-63; Fax: 33-1-42-19-27-40; E-mail: olivier.bernard@insERM.fr. Or to: D. Gary Gilliland, Brigham and Women’s Hospital, 1 Blackfan Circle, Boston, Massachusetts 02115, USA. Phone: (617) 355-9092; Fax: (617) 355-9093; E-mail: ggilliland@rics.bwh.harvard.edu.


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