Brief report

FLT3 expression initiates in fully multipotent mouse hematopoietic progenitor cells

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Lymphoid-primed multipotent progenitors with down-regulated megakaryocyte-erythroid (MkE) potential are restricted to cells with high levels of cell-surface FLT3 expression, whereas HSCs and MkE progenitors lack detectable cell-surface FLT3. These findings are compatible with FLT3 cell-surface expression not being detectable in the fully multipotent stem/progenitor cell compartment in mice. If so, this process could be distinct from human hematopoiesis, in which FLT3 already is expressed in multipotent stem/progenitor cells. The expression pattern of Flt3 (mRNA) and FLT3 (protein) in multipotent progenitors is of considerable relevance for mouse models in which prognostically important Flt3 mutations are expressed under control of the endogenous mouse Flt3 promoter. Herein, we demonstrate that mouse Flt3 expression initiates in fully multipotent progenitors because in addition to lymphoid and granulocyte-monocyte progenitors, Flt3-Mk- and E-restricted downstream progenitors are also highly labeled when Flt3-Cre fate mapping is applied. (Blood. 2011;118(6):1544-1548)

Introduction

Several recent observations suggested that the first lineage restriction step of mouse HSCs does not result in strictly separated common myeloid and common lymphoid commitment pathways. Rather, the earliest step in lymphopoiesis appears to result in the establishment of primitive lymphoid-primed multipotent progenitors (LMPPs) with down-regulated megakaryocyte-erythroid (MkE) transcriptional priming and lineage potentials, but sustained granulocyte-monocyte (GM) and lymphoid potentials.1,2 In further support of the lymphoid commitment pathway sustaining GM (but not MkE) potential, the earliest thymic progenitors have combined T and GM potential.3,4 The existence of adult and fetal GM-lymphoid–restricted mouse MPPs has been confirmed through alternative approaches,2,5-8 and recently a similar progenitor was also identified in human hematopoiesis.9,10 The restriction of mouse LMPPs to the fraction of LIN–SCA1+/KIT+ (LSK) cells with high cell–surface FMS-like tyrosine kinase receptor 3 (FLT3) expression and of long-term self-renewing HSCs to LSK cells lacking detectable cell-surface FLT3 expression11-13 raises the question as to what stage of lineage commitment FLT3 (protein) and Flt3 (mRNA) expression is initiated. Notably, the expression of Flt3 in human hematopoiesis appears to initiate already in multipotent stem or progenitor cells with sustained MkE potential,14,15 differing from the apparent expression pattern in multipotent progenitors in mice. If so, it could have important implications for mouse models in which activating Flt3 mutations, which are among the most common mutations in human acute myeloid leukemia,16 are expressed under control of the mouse Flt3 promoter.17,18

To more conclusively establish at which level in the mouse HSC and MPP hierarchy Flt3 mRNA expression is initiated, we investigated Flt3 expression at the single-cell level and also applied a Flt3-Cre fate mapping approach19 to establish to what degree progenitors of different cell lineages are derived through Flt3-expressing multipotent stem and progenitor cells.

Methods

Mice

Wild-type (W7) mice were on a pure C57BL/6 (CD45.2 or CD45.1) background. Flt3-Cre and R26R-EYFP mice have been described.19,20 Animal experiments were approved by ethics committees at Lund University and University of Oxford and by the United Kingdom Home Office.

Fluorescence-activated cell staining and sorting

BM cells and thymocytes were stained with fluorescence-conjugated antibodies as previously described21,22 and analyzed on a FACS LSRII (BD Biosciences) or sorted on a FACSAriaIIu Special Order Research Web site; see the Supplemental Materials link at the top of the online article).

Transplantation assays

Lethally irradiated (9 Gy) C57BL/6 CD45.1 mice were transplanted with 2 million BM cells from 7- to 11-week-old Flt3-Cre+/R26R-EYFP+/ mice. CD45.2+LSKEYFP+ or CD45.2+LSKEYFP– cells were transplanted into secondary lethally irradiated CD45.1 recipients.

MkE potential

MkE potential was evaluated as previously described.5 For details, see supplemental Methods.

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Gene expression analysis

Multiplex single-cell PCR was performed as previously described. Quantitative RT-PCR analysis was performed with the use of a dynamic array platform (BioMark; Fluidigm Corporation). See supplemental Methods for details, including lists of primer sets.

Results and discussion

SLAM markers CD150 and CD48 allow separation of HSCs (LSKCD150⁺CD48⁻) and MPPs (LSKCD150⁻CD48⁻) in the BM LSK compartment. By using a PE-conjugated antibody excited by a high-powered green laser to enhance the detection of cell-surface FLT3, we found that only a small fraction (6%) of LSKCD150⁺CD48⁻ cells expressed detectable FLT3 at low levels, whereas most LSKCD150⁻CD48⁻ MPPs expressed FLT3 at greater but variable levels, and a fraction of LSKCD150⁻CD48⁻ cells expressed FLT3 at intermediate levels (Figure 1A). By using a highly sensitive single-cell PCR, we found that a small fraction (12.6%) of HSCs defined through another cell-surface phenotype (LSKCD34⁺FLT3⁻) and a larger fraction (41.0%) of LSKCD34⁺FLT3⁻ MPPs expressed Flt3 transcripts (Figure 1B).
Figure 2. Flt3 fate mapping reveals that most erythroid and megakaryocyte progenitors are derived from Flt3-expressing progenitors. (A) EYFP expression in early thymic progenitors (LIN- CD25 -Kit+; bottom left) and BM B220+ B-cell progenitors (bottom right) of Flt3-Cre<sup>Cre<sup>YFP<sup></sup></sup>-/R26R<sup>EYFP<sup></sup></sup>- mice. (B-C) Representative FACS profiles of EYFP expression in myeloid progenitor subsets (B) or LSKCD150<sup>-</sup>CD48<sup>-</sup> HSCs (also gated as FLT3<sup>-</sup> C) in the BM of Flt3-Cre<sup>Cre<sup>YFP<sup></sup></sup></sup>-/R26R<sup>EYFP<sup></sup></sup>- mice. Numbers in histograms are mean percentages from analysis of 3-6 mice. GMP indicates GM progenitor. (D) Quantitative RT-PCR analysis of Flt3-Cre<sup>Cre<sup>YFP<sup></sup></sup></sup>-/R26R<sup>EYFP<sup></sup></sup>- recipients and purity analysis after sorting. (H) Five thousand purified LSKEYFP<sup></sup> cells were transplanted into lethally irradiated CD45.1 recipients and 24 single EYFP<sup></sup> cells toward total BM cells. To the left FACS profiles from representative mice. Percentages are mean values relative to total BM cells. To the right mean (SD) contribution in a total of 4-6 recipients of each of transplanted LSKEYFP<sup></sup> and LSKEYFP<sup></sup> cells toward total CD45.2<sup></sup> BM cells.
Despite undetectable cell surface FLT3, suggesting that Flt3 transcriptional activation might initiate already in the HSC and MPP compartments. Moreover, the results of multiplex single-cell PCR demonstrated that Flt3+ HSCs and MPPs were not only transcriptionally activated for GM but also MK/E lineage programs, although less than Flt3- cells (Figure 1C).

Because these findings were compatible with the transcriptional initiation of Flt3 already in the MPP or even HSC compartment and cell-surface FLT3 expression in MPPs, we next investigated the Mk and E potentials of LSK cells, separated on the basis of different levels of FLT3 expression (Figure 1D). In agreement with previous studies,1,2 we found that LSKFLT3hi LMPPs (25% greatest FLT3-expressing cells within the LSK compartment) lacked detectable Mk and E potential, whereas a large fraction of LSKFLT3lo cells produced colonies with mixed GM, MK, and/or E potential (Figure 1E-F). LSKFLT3lo and in part also LSKFLT3int cells produced MK- and E-containing colonies (Figure 1E-F), in agreement with previous studies in the fetal liver,2 suggesting that the MKE potential is gradually down-regulated with increasing cell surface FLT3 expression.

We next adapted cre-loxP fate mapping, in which mice expressing CRE recombinase under control of the mouse Flt3 promoter18 were crossed with mice with a loxP-flanked transcriptional termination sequence preceding the Eryf gene, under control of the ubiquitous Rosa26 promoter.20 When intercrossed, cells expressing Flt3 and all their progeny (irrespective of Flt3 expression) will express Eryf. As expected, on the basis of the importance of FLT3 in early lymphoid development,25 virtually all early thymic progenitors and BM B220+ B-cell progenitors were EYFP+ (Figure 2A), and as recently shown,21 also most preGM and GM progenitors (Figure 2B). Most notably, all stages of Mkp and E progenitors also expressed EYFP at high frequencies (Figure 2B). Because Mk and E progenitors lack detectable FLT3 expression,2 these findings are compatible with MPPs expressing Flt3. In further support of this, a fraction (23%) of LSKCD150+CD48- cells were EYFP+, although in contrast to other progenitor subsets, most cells in the LSKCD150+CD48- HSC compartment were EYFP- (Figure 2C).

Because the Cre transgene was inserted into the first exon of the Flt3 locus in Flt3-Cre mice19 we could not easily distinguish between transcriptional and genomic expression of Cre in purified EYFP+ HSCs. However, in agreement with EYFP expression in a fraction of the LSKCD150+CD48- HSC compartment, quantitative PCR analysis demonstrated expression of Flt3 mRNA in LSKCD150+CD48-EYFP+ cells that was greater than in LSKCD150+CD48-EYFP- cells and, as expected, lower than in LMPPs that express high levels of cell-surface FLT3 (Figure 2D). Furthermore, single-cell PCR analysis detected Flt3 mRNA expression in almost one-half of EYFP+ LSKFLT3-CD150+CD48- HSCs, in contrast to EYFP+ erythroid progenitors, which lacked Flt3 mRNA expression as reported previously (Figure 2E).21 These findings demonstrate that unlike detectable FLT3 cell-surface protein expression, Flt3 transcriptional activity already initiates in the phenotypic pluripotent HSC compartment, and in agreement with being progeny of HSCs/MPPs, although not expressing Flt3 themselves, MKE progenitors sustain EYFP expression when Flt3-Cre fate mapping is applied.

Notably, the expression of the MKE-related genes WFF and Gfi1b was down-regulated in EYFP+ compared with EYFP- LSKCD150+CD48- cells, whereas expression of Runx3, implicated in lymphoid development,26 was increased (Figure 2F). These findings suggest that Flt3-Cre/EYFP expression marks cells that have initiated a lineage priming (down-regulation of MKE and up-regulation of lymphoid-related transcripts), preparing cells for transition into a LMPPP-like state.

Finally, to investigate whether the expression of Flt3 mRNA in a fraction of LSKCD150+CD48- cells reflects that Flt3 transcription already initiates in functionally defined HSCs, irradiated mice underwent transplantation with Flt3-Cre/o/EYFP+/R26R-EYFP/+ unfractionated BM cells. After 8 weeks, a mean of 58% of donor-derived LSK cells were EYFP+, and LSK-EYFP- and LSK-EYFP+ cells were purified from reconstituted mice and transplanted competitively into secondary recipients to assess for functional HSC activity (Figure 2G). After 6 months, only LSK-EYFP- cells contributed robustly to the short-lived myeloid lineage, whereas LSK-EYFP+ cells gave much lower overall and in particular myeloid reconstitution (Figure 2H), suggesting that few if any self-renewing HSCs express Flt3 in steady state and on cell cycling after transplantation.

Herein, single-cell mapping of Flt3 mRNA expression and Flt3-Cre fate mapping demonstrated that Flt3 expression initiates in fully multipotent mouse progenitors, because not only lymphoid and GM progenitors but also most Mk and E restricted progenitors were EYFP+ in Flt3-Cre/o/EYFP+/R26R-EYFP+ mice, despite lacking cell-surface FLT3 and Flt3 mRNA expression. Thus, these data demonstrate that Flt3-driven Cre protein expression and function precedes FLT3 protein expression in multipotent progenitors.

Unlike in LSKFLT3hi LMPPPs, Flt3 is coexpressed at the single-cell level with MKE transcriptional programs in MPPs, and a fraction of LSK cells with low and intermediate (but not high) levels of cell surface FLT3 sustain MKE potential. In agreement with this, stem cell reconstitution patterns were most compatible, with Flt3 expression being activated in the transition from self-renewing HSCs to MPPs, perhaps as a result of cell-cycle activation.

The present studies also reinforce the existence of GM-lymphoid restricted LSKFLT3hi LMPPPs, establishing that MKE transcriptional priming and potential is gradually down-regulated in MPPs with increasing FLT3 expression, underscoring the importance of defining LMPPs on the basis of high FLT3 expression2,3 and/or alternative markers.6-8

Our findings suggest that in mouse knock-in models,2,7 Flt3-itd will be activated already in fully multipotent progenitors. This finding is of considerable importance because Flt3 expression in humans has been suggested to initiate at the multipotent stem/progenitor stage,14,15 although it is difficult to extrapolate from mouse to human because human HSCs have yet to be as stringently defined as in mice.

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Authorship

Contribution: N.B.-V. and S.E.W.J. designed and conceptualized the overall research, analyzed the data, and wrote the manuscript; N.B.-V. performed the transplantations, phenotypic FACS analysis, sorting, and cell-culture experiments; P.W. S.D., M.L., and T.B.-J. performed FACS analysis of stem/progenitors and peripheral blood; P.W., and M.L., and H.F. performed cell sorting; P.W. and A.H. performed the gene expression analysis; and S.L. performed MkE assays.

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