

Interplay between Flt3L and IL-7 in early haematopoietic development

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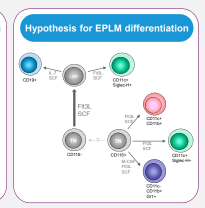


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Fit3Ltg
IL-7K0xFit3Lt

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Abstract

The long standing model of haematopoiesis describes single routes of differentiation towards individual fates and decisions were seen to be irrevocable. However, latest research points towards a more flexible decision making process. We are especially interested in early events in haematopoietic progenitor cell specification with focus on the EPLM cell population early progenitors with lymphoid and myeloid potential - which are defined as cKitlowB220+CD19 NK1.1. Using the cell surface markers Ly6D, Siglec-H, CD11c, and CD115 we were able to distinguish five subpopulations with certain differentiation biases. Now we are studying their individual developmental potentials and possible precursor-product relationships using different in vitro as well as in vivo approaches. While the Ly6D single-positive population seems to be biased towards the B-lymphoid lineage and retains pDC potential, the CD115⁺ portion of Ly6D SiglecH-CD11c⁻ cells shows efficient myeloid, cDC, and pDC differentiation. The CD115 counterpart gives rise to B cells and pDCs. Additionally, we assess the potential instructive role of the cytokines Flt3L and IL-7 in the decision-making process. For this purpose we generated a complete set of knock-out and transgenic mice (and the respective combinations). We found that the B-cell deficiency of IL-7 knock-out mice is rescued by increased levels of Flt3L and Bcl-2, indicating that IL-7 acts mainly as proliferation factor.



Excess Flt3L rescued B-cell specific

transcription factor expression

Excess Flt3L or Bcl-2 rescued

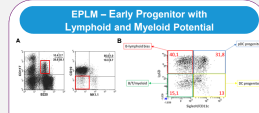
committed B-cell stages

rexpression of Flt3L or Bcl-2 in IL-7 knock-out mice rescued early committee es in the BM. (A) cKit*CD19* preBI and (B) cKit*CD19*1gM- preBII cells were ctable again. (C) Peripheral B-cell receptor expressing cells were restored as

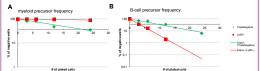
Conclusions

1. EPLM consist of five subpopulations showing differentiation biases TripleNegative CD115⁻ EPLM show B-cell as well as pDC developmental potential with Flt3L and IL-7

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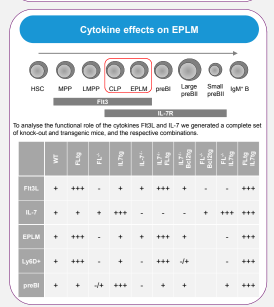


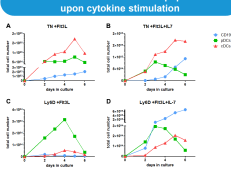
(A) The EPLM are characterised as CD117^{low}B220^oCD19NK1.1. They constitute about 0,1% of nucleated BM cells. (B) The cell surfers markers Ly6D, CD11c, as well as Siglec-H further subdivide them into four different subopoliations. More recently we found within the triple negative cells equal fractions of CD115⁺ and CD115⁺ cells.



In vitro as well as in vivo assays revealed diverging differentiation biases of the two CD11c/Siglec-H negative subpopulations. In limiting dilution on ST2 (A) or OP9 (B) stroma cells + IL-7, respectively, the triple negative compartment had some lymphoid and strong myeloid potential, whereas the Ly6D single-positive EPLM lost the latter and almost ly gave rise to lymphoid cells

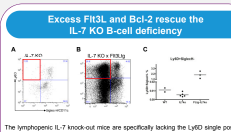
possible precusor-product relationship with the triple negative EPLM giving rise to the Ly6D⁺ ones



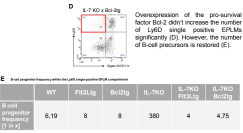


Differentiation of the EPLM subpopulations

5x10⁴ triple negative or Ly6D* single positive EPLM were stimulated with 0,1µg/ml SCF, 0,05µg/ml Flt3L + 10% IL-7. The cultures were analysed daily for the generation SCF. 0.05µg/ml Ht3L + 10% IL-7. The cultures were analysed daily for the generation of CD19' B cells, CD11*CD1b* cDCs, and CD11*C5jlec-H* pDCs. B cells arise late and in low numbers from triple negative EPLM (A), and IL-7 seems to be needed for their expansion (B). Ly6D* EPLM were most potent in generating CD19' cells upon addition of FI18L and IL-7 (D). cDCs were obtained almost exclusively from TN EPLM (A) and B) and FI18L was sufficient. pDCs derived from TN and Ly6D+ EPLM even upon addition of FI19L exclv (A) to CDS addition of FIt3L only (A to D).



ymphopenic IL-7 knock-out mice are specifically lacking the Ly6D single positive I compartment (A), which we think is the direct precursor of CD19° committed B Excess FIt3L rescues this deficiency in numbers as well as in terms of B-cell rsor frequency (B, C, and E). EPI M co cells. Excess Elt31 re:



Acknowledgements

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TripleNegative CD115⁺ EPLM give rise to myeloid cells, cDCs, h as well as some pDCs Ly6D EPLM are lymphoid biased – B cells and pDCs 2. Excess Flt3L or Bcl-2 rescue the B-cell deficient phenotype of IL-7 knock-out mice Ly6D single positive EPLM

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- B-cell specific transcription factor expression
- Early committed B cells
- Peripheral B-cell receptor positive B cells

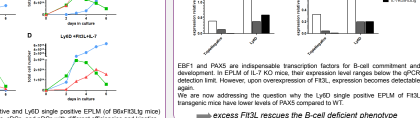
of IL-7 KO mice

3. IL-7 is not necessarily required for B-cell commitment, but it acts as proliferation factor making B cell development more efficient

References

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Upon sorting the triple negative and Ly6D single positive EPLM (of B6xFlt3Ltg mice) gave rise to committed B cells, cDCs, and pDCs with different efficiencies and kinetics.