

Delta-like 4 is the essential, nonredundant ligand for Notch1 during thymic T cell lineage commitment

Ute Koch,¹ Emma Fiorini,² Rui Benedito,³ Valerie Besseyrias,⁴ Karin Schuster-Gossler,⁵ Michel Pierres,⁶ Nancy R. Manley,⁷ Antonio Duarte,³ H. Robson MacDonald,² and Freddy Radtke¹

¹Ecole Polytechnique Fédérale de Lausanne, Swiss Institute for Experimental Cancer Research, 1066 Epalinges, Switzerland

²Ludwig Institute for Cancer Research, Lausanne Branch, University of Lausanne, 1066 Epalinges, Switzerland

³Faculdade de Medicina Veterinária, 1300-477 Lisboa, Portugal

⁴Department of Biomedicine, Institute of Physiology, University of Basel, 4056 Basel, Switzerland

⁵Institute for Molecular Biology OE5250, Medizinische Hochschule Hannover, 30625 Hannover, Germany

⁶Centre d'Immunologie de Marseille-Luminy, Parc Scientifique de Luminy, Case 906, 13288 Marseille, Cedex 9, France

⁷Department of Genetics, University of Georgia, Athens, GA 30603

Thymic T cell lineage commitment is dependent on Notch1 (N1) receptor-mediated signaling. Although the physiological ligands that interact with N1 expressed on thymic precursors are currently unknown, in vitro culture systems point to Delta-like 1 (DL1) and DL4 as prime candidates. Using *DL1-* and *DL4-lacZ* reporter knock-in mice and novel monoclonal antibodies to DL1 and DL4, we show that DL4 is expressed on thymic epithelial cells (TECs), whereas DL1 is not detected. The function of DL4 was further explored in vivo by generating mice in which *DL4* could be specifically inactivated in TECs or in hematopoietic progenitors. Although loss of DL4 in hematopoietic progenitors did not perturb thymus development, inactivation of *DL4* in TECs led to a complete block in T cell development coupled with the ectopic appearance of immature B cells in the thymus. These immature B cells were phenotypically indistinguishable from those developing in the thymus of conditional *N1* mutant mice. Collectively, our results demonstrate that DL4 is the essential and nonredundant N1 ligand responsible for T cell lineage commitment. Moreover, they strongly suggest that N1-expressing thymic progenitors interact with DL4-expressing TECs to suppress B lineage potential and to induce the first steps of intrathymic T cell development.

CORRESPONDENCE

Freddy Radtke:
Freddy.Radtke@epfl.ch

The thymus is continuously seeded by progenitors derived from hematopoietic stem cells, which reside in the BM. These progenitors migrate via the blood stream into the thymus, where they adopt a T cell fate, proliferate, and differentiate into mature functional T cells. This differentiation process is characterized by multiple developmental stages. The earliest thymic progenitors lack surface expression of CD4 and CD8 and are therefore referred to as double-negative (DN) thymocytes. They subsequently up-regulate both CD4 and CD8 coreceptors (double positive [DP]) before undergoing positive and negative selection, and maturing to CD4 and CD8 single-positive (SP) thymocytes that emigrate to the periphery. Immature DN thymocytes can be subdivided into four subpopulations according to the surface expression

of CD117, CD44, and CD25. The most immature thymocyte progenitors (DN1) express CD117 and CD44 and are negative for CD25, followed by the DN2 population, which up-regulates CD25, and the DN3 cells, which down-regulate CD117 and CD44 before generating DN4 thymocytes lacking expression of all three markers (1, 2).

Over the last decade, many reports highlighted the importance of the evolutionarily conserved Notch cascade for the lymphoid system (3). Mammals possess 4 Notch receptors (N1–4), which are activated by two classes of

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transmembrane bound ligands named Jagged 1 and 2, and Delta-like (DL) 1, 3, and 4. Notch signaling is initiated upon ligand receptor interaction, which results in the proteolytic release of the Notch intracellular cytoplasmic domain (NICD) of Notch receptors. The liberated NICD translocates to the nucleus and binds to the transcription factor RBP-J (also known as CSL), thereby converting it from a transcriptional repressor into an activator. Mastermind-like proteins are required for Notch signaling, as they bind to the NICD–RBP-J complex and recruit additional coactivators.

Multiple genetic loss- and gain-of-function experiments show that signaling mediated through the N1 receptor plays an important role for T cell lineage commitment and maturation within the thymus. Inducible inactivation of *N1* in BM progenitors results in a block in T cell development and ectopic B cell development in the thymus, suggesting that N1 instructs an early thymic progenitor to adopt a T cell fate (4, 5). An identical phenotype is observed in mice in which the *RBP-J* gene was inactivated in BM progenitors (6), strongly indicating that T cell specification is mediated by N1–RBP-J-dependent signaling. Interference with Notch signaling by transgenic expression of modulators (such as *Fringe*, *Deltex*, or *Nrarp*) or dominant-negative forms of transcriptional coactivators (*mastermind-like 1*) also blocks T cell development concomitant with B lymphopoiesis in the thymus (for review see reference 3). Reciprocal gain-of-function studies overexpressing NICD in BM progenitors result in ectopic T cell development at the expense of B cell development (3). Thus, both loss- and gain-of-function studies demonstrate that N1 is essential for T lineage commitment. In addition, N1–RBP-J signaling promotes the differentiation of pro-T cells into pre-T cells within the thymus by controlling rearrangement of the TCR β locus (7) through regulating chromatin accessibility (8), thereby assuring the successful generation of a pre-TCR complex, which is essential for thymocyte development.

Although N1 is unequivocally the key Notch receptor involved in T lineage commitment and thymic T cell maturation in vivo, the ligands that could be the physiological partners of N1 in these processes are still a matter of debate. The thymic epithelial microenvironment expresses all *Notch* ligands (mainly assessed by RT-PCR), except *DL3*, which is hardly found in this organ (9–12). However, both Jagged ligands can be excluded from playing an essential role, as *Jagged2*^{-/-} and, more recently, *Jagged1*^{-/-} mice have been shown to have normal T cell development (13, 14), thus implicating DL1 and/or DL4 ligands. Historically, DL1 has been favored as the potential N1 ligand in T cell fate specification, because DL1-expressing stromal cells can support the complete development of mature functional T cells from BM precursors in vitro (10). Moreover, when BM progenitors are co-cultured on stromal cells overexpressing DL1, B cell development is blocked (15). Surprisingly, in contrast with these promising data obtained in vitro, conditional inactivation of *DL1* in thymocytes and/or thymic epithelial cells (TECs) does not inhibit T cell development (16). This discrepancy may be caused by the in vivo presence of DL4, which shares a high degree of homology

with DL1. Data obtained in vitro support this hypothesis, because overexpression of DL4 in stromal cells can also direct T lymphopoiesis (16, 17). In addition, overexpression of DL4 in BM precursors transferred into γ -irradiated mice leads to ectopic T cell development in the BM as well as a severe defect in B cell development (17–19). Moreover, we recently described a hierarchy of Notch–Delta interactions showing that the avidity of DL4 to bind to immature thymocytes is much higher than that of DL1. DL4 binding is high from the DN1 to DN3 stages and declines in DN4 to become undetectable in subsequent DP and SP thymocytes. This binding pattern parallels the functional requirements of Notch signaling during thymocyte development (20, 21), suggesting that DL4 might play an important role during T cell development within the thymus.

In this report, we explore the function of DL4 within the thymus using a conditional loss-of-function approach. Our data show that *Foxn1-Cre*-mediated inactivation of *DL4* within the thymic epithelium leads to a complete block in T cell development accompanied by ectopic B cell development within the thymus, which phenocopies mice with inducible inactivation of *N1* in BM progenitors. Thus, DL4 is essential for T cell fate specification and thereby represents the physiological partner of N1 during this process.

RESULTS AND DISCUSSION

Because DL1- and DL4-expressing OP-9 stromal cells (10, 16, 17) or normal thymic stroma (22) can both support the complete development of mature functional T cells from BM precursors in vitro, it is conceivable that either or both of these proteins function as physiological ligands for the N1 receptor during thymic T cell lineage commitment. To gain further insight into the expression pattern of *DL1* and *DL4* in the thymus, we analyzed thymi of mice in which the *lacZ* gene was knocked into either the *DL1* or *DL4* locus (23, 24). LacZ staining on thymic sections derived from *DL1*^{lacZ/+} mice was only positive for rare endothelial cells within the thymus, whereas *DL4*^{lacZ/+} mice exhibited a strong reticular expression pattern within the outer cortex (characteristic of epithelial cells) and a weaker more punctuated staining within the medulla of the thymus (Fig. 1 A). This expression pattern was confirmed at the protein level using novel mAbs generated against the extracellular domains of DL1 and DL4. The specificity of the anti-DL1 and -DL4 mAbs was tested on DL1- and DL4-expressing OP-9 cells. Anti-DL1 mAbs bound specifically to DL1- but not to DL4-expressing OP-9 cells, whereas the anti-DL4 mAbs showed a reciprocal binding pattern (Fig. 1 B). These results indicate that both mAbs specifically bind to the corresponding Notch ligands and that they do not exhibit any cross-reactivity. These mAbs were then used to stain cell preparations enriched for TECs and analyzed by flow cytometry. TECs were identified as CD45⁻PanCytokeratin (PanCyt)⁺ (25) and were further subdivided into cortical TECs (cTECs) and medullary TECs (mTECs) based on BP1 expression (26). Anti-DL1 mAbs did not stain cTECs or mTECs above background. In contrast anti-DL4 mAbs showed significant staining

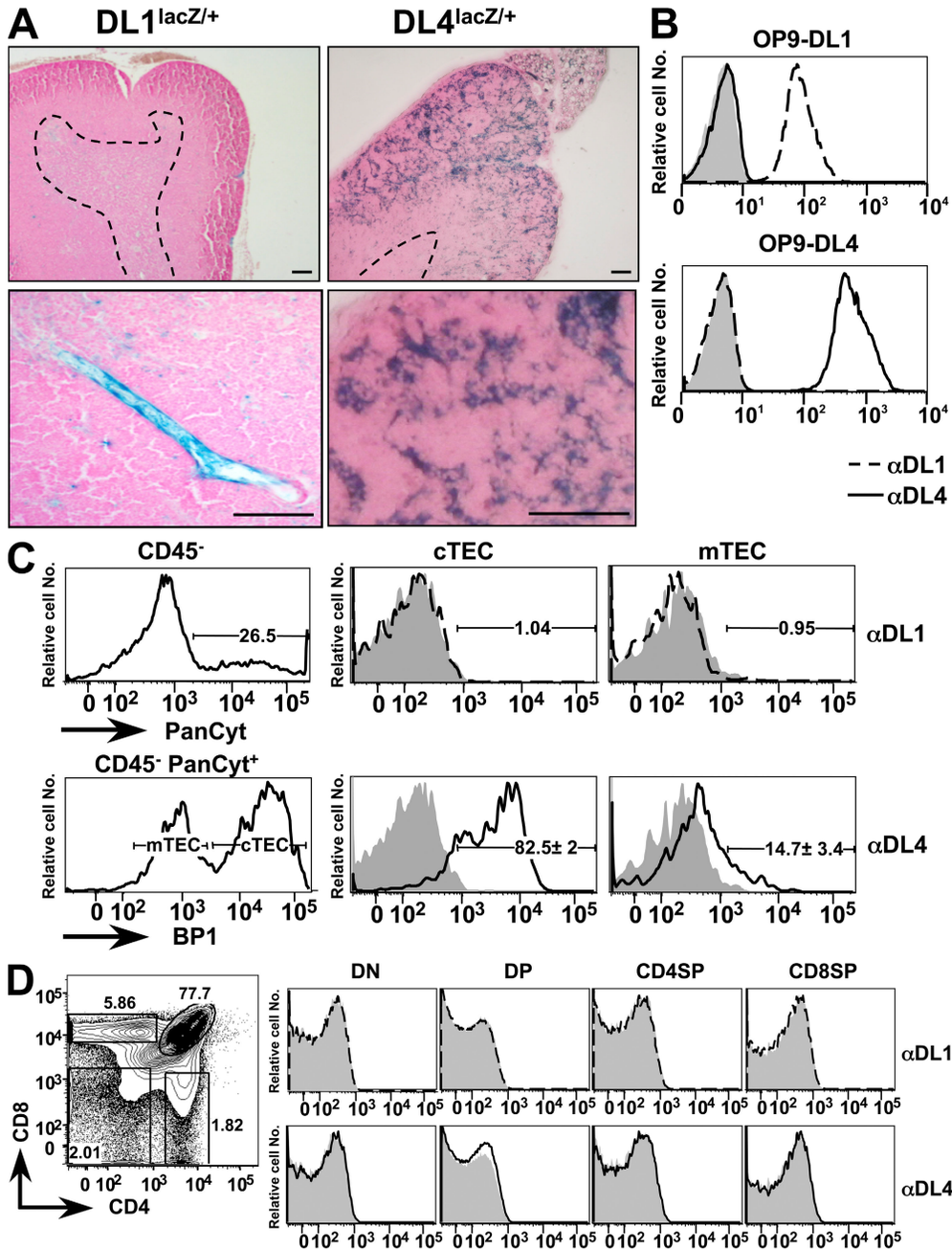


Figure 1. DL4 but not DL1 expression is detected on TECs. (A) LacZ staining on thymic sections derived from *DL1* and *DL4* lacZ knock-in mice. mRNA expression of the *DL1*-driven lacZ gene is confined to blood vessels within the thymus, whereas *DL4* drives expression preferentially within the cortical epithelium. The dashed lines indicate cortical–medullary boundaries. Bars, 100 μ m. (B) Anti-DL1 and -DL4 antibodies specifically bind their corresponding ligands without exhibiting cross-reactivity; OP-9–DL1–EGFP (top) and OP-9–DL4–EGFP (bottom) were stained with isotype Ctrl (shaded), anti-DL1 (dashed line), or anti-DL4 antibodies (continuous line; dilution, 1:100). The analysis was performed on gated EGFP-positive cells, and representative histograms are shown. (C) Anti-DL1 and -DL4 antibody staining on TECs. Enriched TECs extracted from wild-type thymi were stained for BP1 (3C6), CD45, and isotype Ctrl, anti-DL1, or anti-DL4 antibodies, followed by intracellular staining for PanCyt (C11) and flow cytometric analysis. Total TECs were gated as PanCyt⁺CD45⁻ (top left), and cTECs and mTECs were further defined as BP1⁺ and BP1⁻, respectively (bottom left); isotype Ctrl (shaded), anti-DL1 (dashed line; top right), and anti-DL4 (continuous line; bottom right) are shown. Percentages of DL4⁺ TECs represent the mean \pm SD of five mice. (D) DL1 and DL4 are not detectably expressed on thymocytes. (left) A representative flow cytometric analysis of CD4 versus CD8 of wild-type thymocytes. (right) DL1 and DL4 expression are shown as representative histograms (shaded, isotype Ctrl; dashed line, DL1; continuous line, DL4) after gating on the indicated thymic subpopulations: DN (Lin⁻, CD45⁺, CD4⁻, CD8⁻), DP (Lin⁻, CD45⁺, CD4⁺, CD8⁺), CD4SP (Lin⁻, CD45⁺, CD4⁺, CD8⁻), and CD8SP (Lin⁻, CD45⁺, CD4⁻, CD8⁺) thymocytes. Percentages of individual thymic subpopulations are indicated within the plots. Data are representative of three independent experiments.

of TECs, with brighter staining for cortical- (BP1⁺) versus medullary- (BP1⁻) derived populations (Fig. 1 C). Analysis of thymocyte subpopulations for the expression of DL1 and DL4 with these mAbs was negative (Fig. 1 D). These expression data are in general agreement with previous studies (18, 27, 28), with the notable exception of one report in which TECs were broadly stained by a commercial polyclonal antibody against DL1 (12). The specificity of this polyclonal reagent has already been challenged by others (29). In conclusion, the thymic expression pattern of DL1 and DL4 revealed by our novel mAbs confirms the results obtained with the lacZ knock-in mice, strongly suggesting that DL4 might be the physiological ligand expressed by TECs that interacts with N1-expressing T cell precursors to specify the T cell lineage.

Studying the role of DL4 in hematopoiesis and, more specifically, during T lineage commitment by a conventional loss-of-function approach is hampered by the fact that *DL4* gene-targeted mice are embryonic lethal (24, 30). We therefore generated conditional gene-targeted mice for *DL4*. These mice carry loxP sites flanking the first three coding exons of the *DL4* gene (*DL4^{lox/lox}*). Because the gene expression and antibody studies suggested that DL4 expression in TECs might be critical for T cell lineage commitment, we crossed

the *DL4^{lox/lox}* (*Ctrl*) mice to knock-in mice expressing the *Cre recombinase* under the control of the *Foxn1* gene (31), which is highly expressed in TECs, to generate *DL4^{lox/lox&Foxn1-Cre}* (*DL4^{ΔFoxn1}*) mice (Fig. 2 A). Staining of *DL4^{ΔFoxn1}* TECs with anti-DL4 and -DL1 mAbs confirmed the complete loss of DL4 protein in the gene-targeted mice compared with *Ctrl* animals (Fig. 2 B). Interestingly, DL1 expression on TECs derived from *DL4^{ΔFoxn1}* mice was not detectable, indicating that there is no functional compensation by DL1 as a consequence of loss of DL4 (Fig. 2 B).

Side-by-side analysis of littermate thymi derived from *Ctrl* and *DL4^{ΔFoxn1}* mice revealed a 14-fold decrease in absolute thymocyte numbers in *DL4^{ΔFoxn1}* compared with *Ctrl* mice (Fig. 3 A). Flow cytometric analysis of CD4 and CD8 expression on thymocytes derived from *DL4^{ΔFoxn1}* mice shows nearly complete loss (<1%) of the CD4SP, the CD8SP, and the DP thymocyte subsets, whereas all of the major subsets are present in the expected frequency in *Ctrl* mice. *DL4^{ΔFoxn1}*-derived thymi were almost exclusively populated with cells falling into the DN gate (Fig. 3 A). In absolute numbers, DP, CD4SP, and CD8SP subsets were reduced by 180–3,000-fold in *DL4^{ΔFoxn1}* mice, whereas DN cells were slightly increased (twofold). Whether the small residual population of T lineage cells in the thymus of *DL4^{ΔFoxn1}* mice results from a failure to

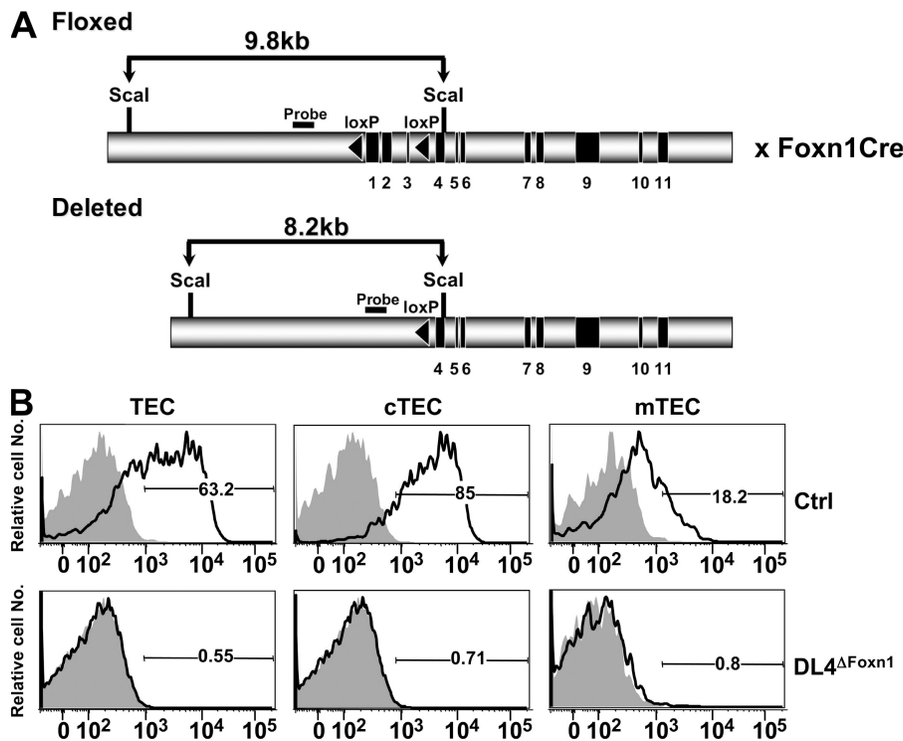


Figure 2. Specific targeting of the *DL4* gene in TECs by the *Foxn1*-*Cre* recombinase. (A) A schematic representation of the genomic organization of the *DL4^{lox/lox}* locus is shown. Exons 1–3 are flanked by loxP sequences (black triangles). *DL4^{lox/lox}* mice were crossed to mice in which the *Cre* recombinase was knocked into the *Foxn1* locus (*Foxn1-Cre*) to obtain *DL4^{lox/lox&Foxn1-Cre}* mice (*DL4^{ΔFoxn1}*). (B) Histograms of total TECs (PanCyt⁺CD45⁻), cTECs (PanCyt⁺CD45⁻BP1⁺), and mTECs (PanCyt⁺CD45⁻BP1⁻) from *Ctrl* (*DL4^{lox/lox}*) or *DL4^{ΔFoxn1}* mice stained for DL1 (shaded) and DL4 (continuous lines). DL1 staining was indistinguishable from *Ctrl* isotype staining (not depicted). Data are representative of five *Ctrl* and three *DL4^{ΔFoxn1}* mice aged 2–3 wk from three independent experiments. Percentages of cells positively staining for DL4 are indicated in the histograms.

delete *DL4* efficiently in a subset of TECs (which stain below the threshold of detection with our anti-DL4 mAb) or, alternatively, represents a minor pathway of DL4-independent (and hence, presumably Notch-independent) T cell development remains to be investigated.

Further analysis of the DN thymic compartment of *DL4^{ΔFoxn1}* mice for CD44 and CD25 expression reveals the presence of only CD44⁺CD25⁻ DN1 cells, whereas all immature thymocyte subsets (DN1–4) were present in normal numbers in *Ctrl* animals (Fig. 3 B). Thus, the thymic phenotype of *DL4^{ΔFoxn1}* mice is very similar to the one previously published for *Notch1^{lox/lox;Mx-Cre}* (*N1^{ΔMx}*) mice (4, 5). In the case of *N1^{ΔMx}* mice, accumulating DN1 cells in the thymus were found to be ectopically developing B cells (4, 5). Similarly, essentially all CD44⁺CD25⁻ DN cells in *DL4^{ΔFoxn1}* mice expressed B220, whereas only a small fraction of DN1 cells in *Ctrl* thymi were B220⁺ (Fig. 3 C). To examine this phenotypic similarity in greater detail, we studied side-by-side B cells within the thymi of 2-wk-old *Ctrl*, *DL4^{ΔFoxn1}*, and *N1^{ΔMx}*

mice 2 wk after *N1* inactivation and compared them with BM B cells.

A small number of thymic B cells are present in *Ctrl* mice, and the majority have a mature IgM⁺B220⁺ phenotype. In contrast, a much larger population of B cells was found in *DL4^{ΔFoxn1}* and *N1^{ΔMx}* mice, and these B cells express heterogeneous levels of IgM and B220. Moreover, *DL4^{ΔFoxn1}* and *N1^{ΔMx}* B cells also express CD93 and BP1, which are typical markers for immature B cells, normally only present in the BM (Fig. 4 A). All B220⁺IgM⁻ cells within the thymi of *DL4^{ΔFoxn1}* and *N1^{ΔMx}* mice were CD19⁺, confirming that these are indeed immature B cells (unpublished data). In absolute numbers, immature thymic B cells were increased 340- and 30-fold over *Ctrl* values in *DL4^{ΔFoxn1}* and *N1^{ΔMx}* mice, respectively (Fig. 4 B). In conclusion, the immature B cell phenotype observed in the thymi of *N1^{ΔMx}* mice is largely recapitulated in *DL4^{ΔFoxn1}* mice, strongly suggesting that DL4 is the natural ligand for N1 during thymic T cell lineage commitment.

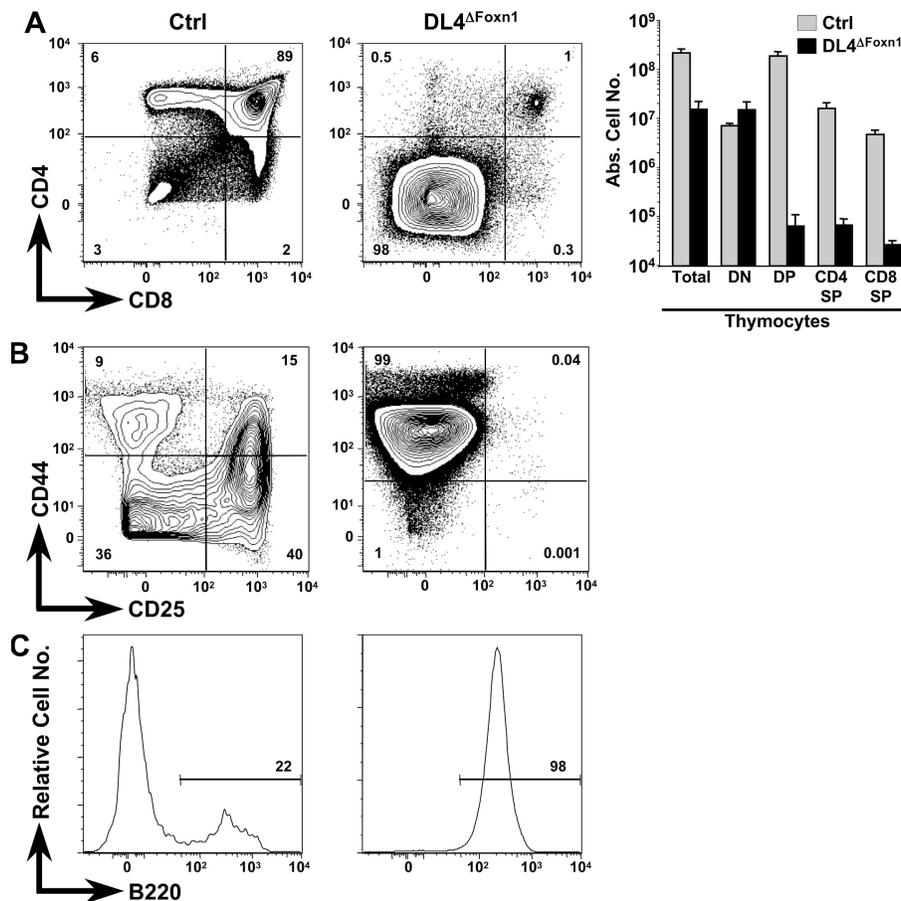


Figure 3. Complete block in T cell development and accumulation of thymic B cells in *DL4^{ΔFoxn1}* mice. (A) Representative flow cytometric analysis of thymocytes derived from *Ctrl* and *DL4^{ΔFoxn1}* 14-d-old mice. Total thymocytes were stained with anti-CD4 and -CD8 antibodies. Absolute cell numbers for total thymocytes and indicated subsets are shown as bar diagrams on a logarithmic scale. The bar diagrams represent mean values \pm SD ($n = 5$ for *Ctrl* and 6 for *DL4^{ΔFoxn1}*). (B) Representative FACS analysis of CD4⁻CD8⁻lin⁻ thymocytes stained with anti-CD44 and -CD25 antibodies. (C) Representative histograms of B220 staining gated on lin⁻CD44⁺CD25⁻ cells. Data are representative of three independent experiments. Percentages of positively stained cells are indicated within the contour plots and histograms.

Although $DL4^{ΔFoxn1}$ and $N1^{ΔMx}$ mice exhibit qualitatively similar thymic phenotypes, it is clear that enhancement of immature B cell development (Fig. 4 B) is more pronounced in the $DL4^{ΔFoxn1}$ background. This quantitative difference between $DL4^{ΔFoxn1}$ and $N1^{ΔMx}$ mice most likely reflects the consequence of inactivation of the floxed $DL4$ and $N1$ genes using two different Cre systems. The $Mx-Cre$ transgene inducibly inactivates floxed genes preferentially in hematopoietic cells in the BM (32). Thus, postnatal inactivation of $N1$ will generate $N1$ -deficient BM progenitors that will migrate to the thymus and develop into B cells that will have to compete with residual thymocytes for the availability of thymic niches, growth factors, and cytokines. In contrast, constitutive $Foxn1-Cre$ -mediated $DL4$ gene inactivation in TECs occurs already during embryogenesis (31). Thus, the first wave of incoming lymphoid progenitors will encounter an empty $DL4$ -deficient thymic epithelium. Although these precursors cannot be instructed toward the T cell lineage, they can develop into the B cell lineage and

efficiently expand in the absence of competition, as all necessary cytokines and growth factors are abundantly present in the neonatal thymus.

Although $Foxn1-Cre$ expression within the thymus should be restricted to TECs (31) and wild-type thymocytes do not stain detectably with anti- $DL4$ mAbs (Fig. 1 D), it remains formally possible that rare $DL4$ -expressing hematopoietic cells might be involved in T cell lineage commitment in the thymus. In this scenario, loss of $DL4$ expression on these cells could theoretically contribute to the generation of immature B cells in the $DL4^{ΔFoxn1}$ thymus. To exclude this possibility, we generated $DL4^{lox/lox;Mx-Cre}$ ($DL4^{ΔMx}$) mice to inducibly inactivate $DL4$ in BM progenitors. Because the $Mx-Cre$ transgene is also active to some extent in TECs (16), BM from $DL4^{ΔMx}$ or $Ctrl$ mice (both $CD45.2^+$) was transplanted into $CD45.1^+$ wild-type hosts to critically evaluate whether $DL4^{ΔMx}$ hematopoietic cells could contribute to the phenotype. $Ctrl$ and $DL4^{ΔMx}$ BM chimeras were analyzed 8 wk after transplantation. The reconstitution efficiency of $CD45.2^+$ cells for

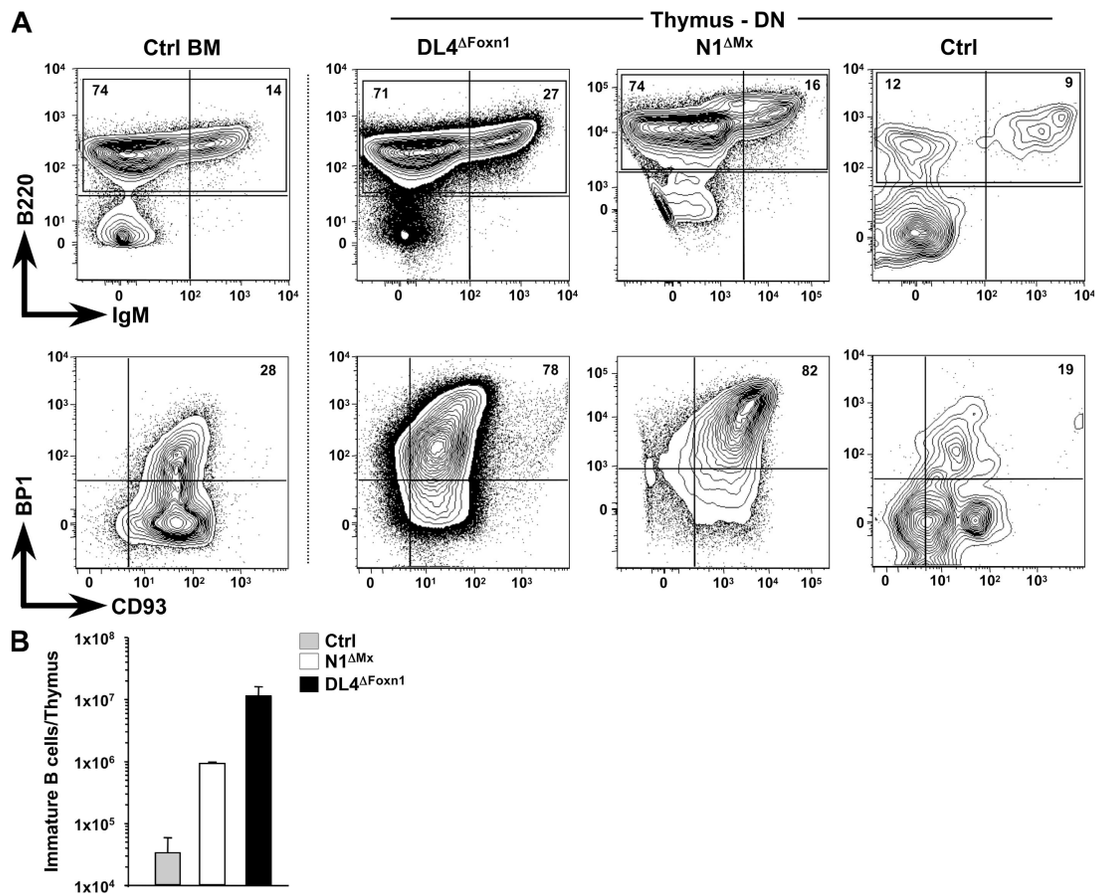


Figure 4. Comparative phenotypic analysis of B cells within thymi of $DL4^{ΔFoxn1}$ and $N1^{ΔMx}$ mice. (A) Representative flow cytometric analysis of B cells in the DN thymus compartment from $Ctrl$, $N1^{ΔMx}$, and $DL4^{ΔFoxn1}$ mice stained for B220 and IgM (top) or BP1 and CD93 (bottom). B220/IgM staining is electronically gated on lineage-negative $CD44^+CD25^-$ cells, whereas BP1/CD93 staining is gated on lineage-negative $B220^+$ cells. (left) A comparative analysis of normal BM B cells to highlight the phenotypic similarity. Percentages of populations staining positively for the indicated markers are shown in the contour plots. (B) Bar graphs show the absolute numbers \pm SD of immature ($B220^+IgM^-$) B cells per thymus derived from $Ctrl$, $Notch1^{ΔMx}$, and $DL4^{ΔFoxn1}$ mice ($n = 6$ mice per sample group). Note the logarithmic scale. All mice used were between 2–3 wk old, and data are representative of three independent experiments.

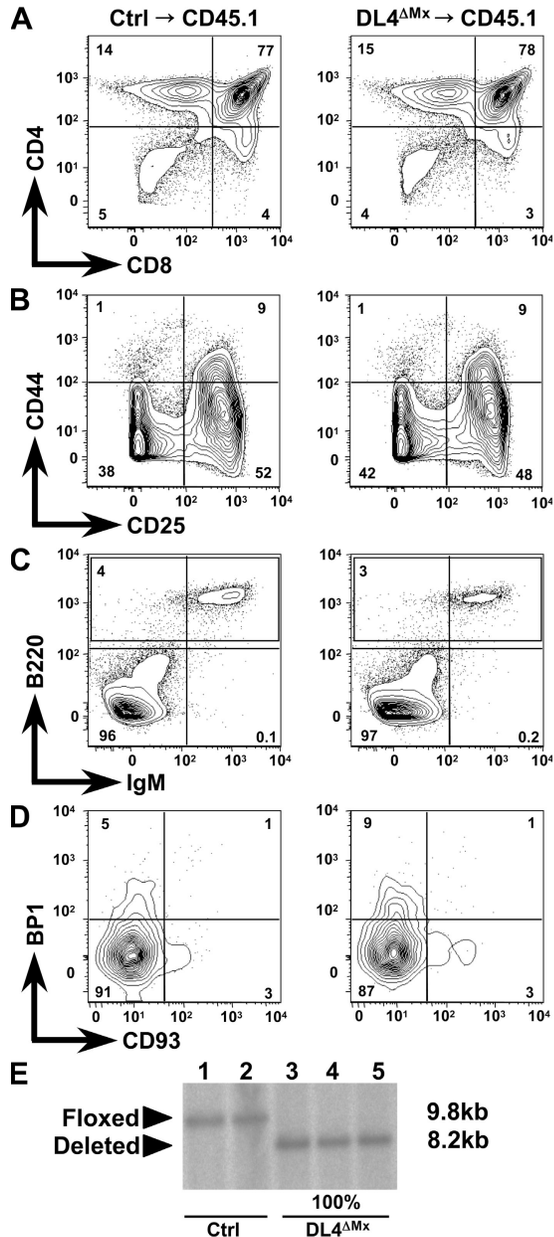


Figure 5. Normal T cell development and absence of immature thymic B cells in BM chimeras reconstituted with *DL4^{ΔMx}* BM. *Ctrl* or *DL4^{ΔMx}* CD45.2⁺ BM was transplanted into lethally irradiated CD45.1⁺ hosts and analyzed 8 wk after reconstitution. The reconstitution efficiency for both *Ctrl* and *DL4^{ΔMx}* BM chimeras was >95% (not depicted). (A) Representative flow cytometric analysis of CD45.2⁺ thymocytes stained with anti-CD4 and -CD8. (B) CD45.2⁺ lineage-negative DN thymocytes were analyzed for the expression of CD44 and CD25. (C and D) Lineage-negative thymocytes were analyzed for the presence of B cells expressing B220 and IgM, or BP1 and CD93. Data in A–D are representative of three individual chimeras per group with virtually identical results, and relative percentages are indicated in the contour plots. Two independent experiments were performed. (E) Southern blot analysis of *ScaI*-digested genomic DNA derived from BM cells of *Ctrl* and *DL4^{ΔMx}* chimeras showing the floxed alleles in *Ctrl* chimeras and the completely deleted *DL4* locus in *DL4^{ΔMx}* chimeras. The targeting strategy and size of the restriction fragments are as described in Fig. 2 A.

both *Ctrl* and *DL4^{ΔMx}* chimeras was >95%. A portion of the BM cells of the corresponding BM chimeras was analyzed by Southern blot analysis to ensure that reconstitution was not caused by rare cells that might have escaped deletion of the *DL4* locus (Fig. 5 E). As expected, *Ctrl* and *DL4^{ΔMx}* BM progenitors generated all major blood lineages in the BM and spleen (unpublished data). More importantly, all major mature and immature thymocyte subsets were generated normally from *Ctrl* and *DL4^{ΔMx}* BM progenitors (Fig. 5, A and B), and only very small numbers of immature B cells were found in the thymus (Fig. 5, C and D). These data exclude a significant role for putative *DL4*-expressing hematopoietic cells in thymic T cell lineage commitment.

Collectively, we have shown that *DL4* is expressed by TECs, whereas *DL1* is not detectable. Moreover, *DL4^{Foxn1}* mice exhibit a loss of *DL4* expression in TECs, which correlates with a complete block in T cell development and the accumulation of immature B cells within the thymus. *DL4* deficiency within hematopoietic progenitors seems not to contribute to this phenotype, as assessed by BM chimeras, suggesting that the loss of *DL4* on TECs is causative. A similar accumulation of immature thymic B cells is found in mice in which the *N1* receptor is inactivated in hematopoietic progenitors. Collectively, these data demonstrate that *DL4* expression on TECs is essential for thymic T cell development. Furthermore, they strongly suggest that *N1*-expressing T cell progenitors only commit to the T cell lineage after entry into the thymus.

Although in vitro *DL1* and *DL4* are functionally redundant in their ability to promote T cell development from BM precursors when expressed on OP9 cells, in vivo *DL1* and *DL4* mediate nonredundant functions within the hematopoietic system by binding to specific Notch receptors. Our data demonstrate that *DL4* and *N1* constitute within the thymus a nonredundant ligand–receptor pair essential for T cell lineage commitment. Similarly, *DL1* pairs with *N2* in the spleen to specify marginal zone B cell development in a nonredundant manner (16, 33). Differential expression of Notch ligands has also been reported to influence T helper cell differentiation in peripheral CD4 T cells (34). Thus, all of these examples are consistent with the possibility that specificity of Notch signaling within the hematopoietic system might be generally regulated by tissue-specific expression of Notch ligands.

MATERIALS AND METHODS

Mice: generation of *DL4* gene-targeted floxed mice. We generated the conditional *DL4* loss-of-function mutant mice by flanking the first three exons of the *DL4* allele with two loxP sites. Mouse *DL4* genomic clones were isolated from a 129/Sv genomic library. One loxP site was inserted into the 5' untranslated region in the *XhoI* restriction site, 75 bp downstream of the transcription initiation site. The other loxP site, together with the flipase recognition target (FRT) site–flanked *PGK-neomycin* selection cassette, was inserted in the *EcoRI* restriction site of intron 3. Upon Cre-mediated recombination, the translation initiation site and the coding sequence for the first 132 amino acids of the protein were removed, creating a null allele. Electroporation and selection of embryonic stem (ES) cells (R1) was performed using standard protocols. Homologous recombination of the targeted *DL4*

locus in ES cells was confirmed first by PCR and then by *ScaI* restriction digest of the genomic tail DNA and Southern blotting. A 400-bp *Apal-Apal* genomic fragment localized upstream of the 5' arm of the targeting vector was used as a probe (Fig. 2 A). Blastocyst injection of the targeted ES cells was performed according to standard protocols. To remove the FRT site-flanked *PGK-neomycin* selection cassette, germline chimeras were crossed with ACTB-F₁pe mice (35) in a pure C57BL6 background. *DL4* homozygous floxed mice were obtained in the expected Mendelian ratio, suggesting normal expression of a functional DL4 protein from the modified allele. Homozygous *DL4^{lox/lox}* mice were bred to *Mx-Cre* mice (32) and to *Foxn1-Cre* mice (31) to generate *DL4^{lox/+&Mx-Cre+/-}* and *DL4^{lox/+&Foxn1-Cre+/-}* mice, respectively. These mice were intercrossed to generate *DL4^{lox/lox&Mx-Cre+/-}* (*DL4^{ΔMx}*) and *DL4^{lox/lox&Foxn1-Cre+/-}* (*DL4^{ΔFoxn1}*) mice and were subsequently backcrossed for at least five generations into C57BL6 mice. Mice were genotyped for homozygosity of the loxP sites and for the presence of the *Mx-Cre* or *Foxn1-Cre* transgene by PCR.

Conditional gene-targeted *N1* mice were previously described (4). B6.SJL-Ptprca (CD45.1⁺) mice were originally purchased from the Jackson Laboratory. CD45.1⁺ recipient mice were used as BM recipients for the chimeras described in the following section. This study has been reviewed and approved by the Service Vétérinaire Cantonal of Etat de Vaud.

BM chimeras. BM chimeras were generated using T cell-depleted BM cells from polyI-polyC (pI-pC; InvivoGen) *Ctrl* (*DL4^{lox/lox}*) or *DL4^{ΔMx}* (*DL4^{lox/lox&Mx-Cre+/-}*) mice. Lethally irradiated (1,000 cGy) 10-wk-old recipients (CD45.1⁺) received 10 × 10⁶ donor BM cells (CD45.2⁺) i.v. Radiation chimeras were maintained on antibiotic water and analyzed after 8 wk.

Activation of the *Mx-Cre* recombinase. 3–4-wk-old *DL4^{ΔMx}* and *Ctrl* mice received 4 i.p. injections of 2 μg/g body weight pI-pC at 2 d intervals. Mice were killed at the time points indicated in the figures, and genomic DNA was prepared from 10 × 10⁶ BM cells. The deletion efficiency in cells from chimeric, *Ctrl*, and *DL4^{ΔMx}* mice was assessed by Southern blot analysis of *ScaI*-digested genomic DNA and hybridized with a 400-bp *Apal-Apal* probe (Figs. 2 and 5). The analysis was quantified using a phosphorimager (FLA3000; Fujifilm).

Staining for β-galactosidase activity and histology. β-Galactosidase activity was detected as previously described (36). In brief, whole thymi were isolated and fixed for 60 min in 0.2% glutaraldehyde in 100 mM of potassium phosphate buffer, pH 7.4, containing 5 mM EGTA and 2 mM MgCl₂. Thymi were washed three times in 0.01% Na-desoxycholate and 0.02% Nonidet P-40 in 100 mM of potassium phosphate buffer with 5 mM EGTA and 2 mM MgCl₂. For detection of β-galactosidase activity, thymi were incubated in washing solution containing 0.5 mg/ml X-gal, 10 mM K₃Fe(CN)₆, and 10 mM K₄Fe(CN)₆ at 37°C. After staining, thymi were washed again in washing solution and, finally, in 1 × PBS. For histology, thymi were dehydrated in isopropanol, cleared in xylene, and transferred to paraffin. Embedded thymi were sectioned at 7–10 μm and counterstained with eosin according to standard procedures.

TEC enrichment. TECs from normal and *DL4^{ΔFoxn1}* thymi were prepared using a modified protocol from Klein et al. (37). In brief, individual thymi were cut into small pieces and washed three times with DMEM supplemented with 2% FCS (PAA Laboratories) to remove thymocytes. TECs were then extracted from the thymic fragments sequentially using collagenase D and DNaseI digestions (both from Roche), followed by TrypLE Express (Invitrogen) digestion. Finally, TECs were recovered at the interface of a 38–60% Percoll gradient, stained with the mAbs indicated in the figures, and analyzed by FACS.

mAbs and flow cytometry. Single-cell suspensions from BM, thymus, and TECs were prepared and stained using standard protocols, as previously described (5). FACS analyses (6- and 8-color) were performed using a

FACSCanto flow cytometer (Becton Dickinson) and a CyAn ADP flow cytometer (Dako). Data were analyzed using FlowJo software (Tree Star, Inc.) or FACS Aria software (Becton Dickinson). The following antibodies were used to analyze hematopoietic cells and TECs, and the subsequent mAb conjugates were purchased from eBioscience: TCRβ (H57-597)-FITC; CD4 (GK1.5)-PE-Cy5, -PE-Cy5.5, and -PE-Cy7; CD8α (53.6.7)-PE and -PE-Cy7; CD11b (M1/70)-PE-Cy7; Gr1 (Ly6G, RB6-8C5)-PE-Cy7; Ter119 (Ly76)-PE-Cy7; B220 (RA3-6B2)-FITC and -PE-Cy5; CD44 (Pgp-1, IM7)-PE-Cy7; CD117 (c-kit, 2B8)-allophycocyanin (APC); CD25 (IL-2R, PC61.5)-APC-Alexa Fluor 750; BP-1 (6C3)-PE; IgM (eB121-15F9)-APC; CD93 (AA4.1, C1qRp)-FITC; CD45 (30-F11)-PE-Cy5.5; CD45.1 (A20.1)-PE and -APC-Alexa Fluor 750. CD45.2 (104)-PacificBlue was purchased from BioLegend. TCRγδ (GL3)-FITC, CD8α (53.6.7)-FITC and -Alexa Fluor 647, CD4 (GK1.5)-FITC, CD3 (145-2C11)-FITC, CD11b (M1/70)-FITC, Gr1 (Ly6G, RB6-8C5)-FITC, and Ter119 (Ly76)-FITC were purified from hybridoma supernatants and conjugated in our laboratory according to standard protocols. Alexa Fluor 647 conjugates were prepared using the appropriate Alexa Fluor protein labeling kits (Invitrogen). Intracellular staining of TECs was performed with a PanCyt mAb (C11) from Sigma-Aldrich.

Rat mAbs were generated against the extracellular domains of DL1 and DL4. Details of the immunization and screening will be presented in a future paper (unpublished data). Purified mAbs were biotinylated in our laboratory according to standard protocols and revealed with streptavidin-PE. Both mAbs were assayed for specificity on OP-9 stromal cells engineered to express DL1 or DL4 combined with enhanced GFP (OP-9-DL1-EGFP or OP-9-DL4-EGFP; Fig. 1 B).

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