The Eδ enhancer controls the generation of CD4−CD8− αβ TCR-expressing T cells that can give rise to different lineages of αβ T cells

Iannis Aifantis,1,4 Craig H. Bassing,2,3 Annette I. Garbe,1 Katie Sawai,4 Frederick W. Alt,2 and Harald von Boehmer1

1Dana-Farber Cancer Institute and 2Children’s Hospital, Harvard Medical School, Boston, MA 02115
3University of Pennsylvania School of Medicine, Philadelphia, PA 19104
4Department of Medicine, The University of Chicago, Chicago, IL 60637

It is well established that the pre–T cell receptor for antigen (TCR) is responsible for efficient expansion and differentiation of thymocytes with productive TCRβ rearrangements. However, Ptcra- as well as Tcra-targeting experiments have suggested that the early expression of Tcra in CD4−CD8− cells can partially rescue the development of αβ CD4+CD8+ cells in Ptcra−deficient mice. In this study, we show that the TCR Eδ but not Eα enhancer function is required for the cell surface expression of αβ TCR on immature CD4−CD8− T cell precursors, which play a crucial role in promoting αβ T cell development in the absence of pre-TCR. Thus, αβ TCR expression by CD4−CD8− thymocytes not only represents a transgenic artifact but occurs under physiological conditions.
DP thymocytes in the absence of pre-TCR, (8). In the case of αβ TCR in Ptcra−/−, Tcrd−/− mice, this resulted in thymocytes of which >95% harbored TCRβ chains (i.e., DP cells that were selected by an early expressed αβ TCR and therefore contained in-frame Ieα rearrangements). In the case of an early expressed γδ TCR in Ptcra−/−, Tcrd−/− mice, only 15% of DP cells contained TCRβ chains. Thus, these experiments indicated that receptors other than the pre-TCR could relieve DN3 cells from a development block resulting in the production of DP cells and that there must be a rearrangement of TCRα V gene segments at the DN3 or earlier stages of T cell development.

The temporal order of TCR V gene segment rearrangement appears especially important with regard to the Tera and Terd locus. This is where the Tera locus is embedded in the Tera locus and where the early rearrangement of Vδ gene segments results in the formation of TCRδ chains, whereas late Vα rearrangement is accompanied by the deletion of the Terd locus and generation of Tera genes (9). Two different enhancer elements have been invoked in the control of rearrangement and expression of the Tera/Terd locus: the Eδ enhancer located in the Jβ1-C8 intron and the Eα enhancer 4 kb downstream of Cα (10–13). Both enhancers have been deleted by homologous recombination. In Eδ−/− animals, the development of αβ T cells appeared to proceed normally with the exception that there was a considerable reduction of thymic and peripheral γδ T cells. In this context, Eδ−/− deficient alleles exhibited a substantial reduction of Terd gene rearrangements. Eα−/− mice contained normal numbers of DP cells but reduced numbers of CD4+/CD8− and CD4−/CD8+ single-positive (SP) cells as a result of an almost complete block in Vα to Jα rearrangements. The Eα enhancer also controls levels of αβ as well as γδ TCR expression, as evident by reduced levels of TCRα and TCRδ transcripts in Eα−/− mice. These results indicate that Eδ functions in DN thymocytes to promote Terd gene rearrangement and gene expression but not Jα accessibility to the V(D)J recombination. On the other hand, Eα functions in DP thymocytes to promote TCRα gene rearrangement via Jα accessibility and controls both Terd and Tera gene expression. However, because TCRα chains were still present in Eα−/− mice, it was hypothesized that in the absence of Eα, the Eδ enhancer or other elements might have a role in promoting rearrangement, possibly via the promotion of Jα accessibility and/or expression of some Vα segments (11). This would account for the TCRα chains with limited diversity (mostly Vα2) that were expressed in peripheral lymphoid tissue in Eα−/− deficient mice.

In this study, we have tested the hypothesis that a low level of Vα to Jα rearrangements is controlled by the Eδ enhancer and occurs in DN thymocytes earlier than the bulk of Eα-controlled Vα to Jα rearrangements, thus leading to the expression of αβ TCR in DN cells, which permits some of these cells to enter the αβ lineage of DP thymocytes.

RESULTS
Early αβ TCR-expressing DN cells
Analyzing total thymocytes and DN thymocytes from wild-type (wt) C57BL/6 mice, we find that ∼5% express TCRβ chains on the cell surface of which ∼10% are paired with Vα2-containing TCRα chains (Fig. 1 A). Because a subset of NKT cells (14) has the TCRβ+ CD4−/CD8− phenotype, NK1.1 cells were excluded by using NK1.1 antibodies for depletion as well as by the analysis of NKT cell–deficient CD1−/− animals (Fig. 2). The results show that Vα+, TCRβ+ cells belong to a distinct, non-NKT cell subset of DN cells. In addition, we found TCRβ+ DN T cells in secondary lymphoid organs in both wt and CD1− deficient animals (Fig. 2).
To investigate whether DN αβ TCR-expressing cells are pre-TCR selected as their DP (CD4+CD8+) or SP (CD4+CD8− and CD4−CD8+) counterparts, thymocytes from Ptcra−/− mice were analyzed (Fig. 1 B). Although Ptcra−/− mice were previously shown to be deficient in NKT cells (15), we used NK1.1 antibodies in the depletion procedure.

Figure 2. CD4−CD8− αβ T cells in the thymus and lymphoid periphery. (A) CD4,CD8 versus NK1.1 staining in the thymus of wt and CD1−/− mice (top). TCRβ versus Vα2 analysis of NK1.1− CD4−CD8− thymocytes (bottom). [B and C] Identification of peripheral CD4−CD8− TCRβ+ cells in the spleen and lymph nodes using TCRβ, Vα2, CD4, and CD8 antibody labeling. Numbers in quadrants indicate the percentages of cells in that quadrant.

Figure 3. Phenotype and developmental potential of early αβ T cells. (A) Vα2 versus TCRβ staining of Ptcra−/−,Tcrd−/− thymocytes. (B) Phenotypic analysis of CD4−CD8− and CD4−CD8− TCRβ+ cells using CD44 and CD25 antibodies. (C) Embryonic day 14.5 fetal (Rag-1−/−) thymic organ culture of CD4−CD8− TCRβ+ NK1.1− Ptcra−/− cells. Cells were cultured for 7 d and were stained with Vα2, TCRβ, CD4, and CD8 antibodies. A CD4 versus CD8 staining of Vα2+ donor cells is also shown (bottom). Numbers in quadrants indicate the percentages of cells in that quadrant.

To investigate whether DN αβ TCR-expressing cells are pre-TCR selected as their DP (CD4+CD8+) or SP (CD4+CD8− and CD4−CD8+) counterparts, thymocytes from Ptcra−/− mice were analyzed (Fig. 1 B). Although Ptcra−/− mice were previously shown to be deficient in NKT cells (15), we used NK1.1 antibodies in the depletion procedure.
that yielded DN thymocytes. In Ptcra−/− mice, ~6–7% of DN thymocytes express TCRβ chains on the cell surface, and, in this particular experiment, 40–50% of DN cells with TCRβ proteins coexpressed Vα2,TCRα chains (Fig. 1 C). In several experiments, the proportion of Vα2+ cells among TCRβ+ DN cells in Ptcra−/− mice was variable and ranged between 10 and 50% (Fig. 4; see Fig. 7).

Of interest was the observation that Vα2 is not only contained in αβ but also in γδTCRs in DN thymocytes from Ptcra−/− mice (Fig. 1 C), a notion consistent with recent findings that Vα2 gene segments can join to both Jα and DδJδ sequences (16). To exclude that the existence of TCRβ+,Vα2+ DN cells is the result of Vα2 segment pairing with TCRδ diversity joining elements (16), a similar analysis was conducted in Ptcra−/−,Tcrδ−/− double-deficient mice. About 7% of CD4+CD8− cells were found to express TCRβ proteins on the cell surface, and ~25% of the TCRβ chains were paired with Vα2-containing TCRα chains, suggesting that these TCRβ-expressing cells are bona fide αβ T cells that do not require pre-TCR selection (Fig. 3 A).

Further phenotypic analysis of the TCRαβ+ thymocytes showed that almost all “early” TCRβ-expressing CD4+CD8− cells are CD25 negative and, thus, belong to the DN4 (CD25−CD44−) subset (Fig. 3 B). The developmental potential of early αβTCR-expressing DN cells was then addressed by culturing purified LY5.2+ CD4−CD8− NK1.1−

Figure 4. Eα controls late but not early development of αβ T cells. (A) Absolute cell numbers of Ptcra−/− and Ptcra−/−,Eα−/− littermates. Error bars represent SD. (B) CD4 versus CD8 profiles of littermates belonging to all indicated genotypes. (C and D) Vα2 versus TCRβ antibody labeling in total DN thymocytes (C) and spleen cells (D). Numbers in quadrants indicate the percentages of cells in that quadrant.
γδTCRβ+ Ptcra−/− thymocytes together with embryonic thymus from LY5.1+ in Rag1−/− donors in fetal thymic organ cultures. After 7 d of culture, ∼40% of the donor-derived cells have up-regulated the expression of CD4,CD8 coreceptors. Moreover, >90% of the cultured thymocytes retain the surface expression of TCRβ, and ∼20% of them are Vα2+ (Fig. 3 C). Not all cells up-regulate CD4 and CD8, however, and a substantial fraction of cells expressing high levels of Vα2-containing αβTCRs remain CD4−CD8−. It is likely that these αβTCR−expressing DN cells normally exit the thymus because CD4−CD8− αβTCR− cells can be detected in the lymph nodes and spleen of adult mice (Fig. 2), and it was shown in TCRαβ transgenic mice that DN cells with the transgenic TCR can accumulate in peripheral lymphoid tissue (6). Thus, TCRαβ-expressing DN cells can give rise to both immature DP and mature SP (CD4/8) T cells as well as DN αβ T cells that do not enter the DP αβ lineage (17).

The Eα enhancer controls late but not early development of αβ T cells in Ptcra−/− mice

To address the role of the described TCR enhancers on early TCRα expression, we generated Ptcra−/− mice with a targeted deletion of the Eα locus (Ptcra−/−,Eα−/−). Ptcra−/−,Eα−/− and Ptcra−/−,Eα+/− littermate mice contain similar numbers of thymocytes (Fig. 4 A). However, there is a clear reduction of CD4+CD8− SP thymocytes in the Ptcra−/−,Eα−/− mice because of limited TCRα diversity and/or expression levels (Fig. 4 B). This is supported by the staining of either all thymocytes or only DN thymocytes with a combination of TCRβ and Va2 antibodies: although total thymocytes from Ptcra−/− single-deficient mouse contain <10% TCRβ+,Va2+ cells, among TCRβ+ cells, this proportion is much higher (>30%) in Ptcra−/−,Eα−/− double-deficient mouse. In DN thymocytes from Ptcra−/−,Eα+/− mice, TCRβ+,Va2+ cells represent 10% of all TCRβ+ cells, whereas in Ptcra−/−,Eα−/− mice, TCRβ+,Va2+ cells represent 50%. These data indicate that Eα is not required for the early rearrangement and expression of Tcrα genes (Fig. 4 C). The data show that normally Eα predominantly contributes to the rearrangement and expression of TCRα V gene segments other than Va2 gene segments. Consistent with this notion, the spleen of Ptcra−/−,Eα−/− mice contains almost exclusively Va2+,TCRβ+ cells, whereas only ∼10% of Va2+,TCRβ+ cells among TCRβ+ cells are found in the spleen of Ptcra−/−,Eα+/− mice (Fig. 4 D). These observations are in line with earlier observations in Eα−/− mice (11) showing that TCRβ+ cells in the spleen of Eα−/− mice express almost exclusively Va2 (Fig. 4 D).

DN cells but not mature T cells in Eα−/− deficient mice express Va2-negative Tcrα genes

The data in Fig. 4 suggest that DN cells in Eα−/− mice can express Va2 genes other than Va2 and that the Va2 dominance in the periphery of Eα−/− mice is established at a later developmental stage. This issue was addressed in more detail by analyzing the expression of TCR Va2 genes and other TCRα V genes (Va3, Va8, and Va11) in Ptcra−/− and Ptcra−/−,Eα−/− mice at various stages of development. As shown in Fig. 5 B, NKT cell–depleted CD4+ and CD8− (DN) cells from both Ptcra−/− and Ptcra−/−,Eα−/− mice do...
express Vα genes that stain with a cocktail of Vα3, 8, and 11 antibodies. Although the proportion of cells stained with the cocktail versus Vα2-positive cells remains about the same in CD4+CD8+ cells of Ptcra−/− mice, it drastically decreases in DP cells of Eα−/− mice such that the vast majority of cells expresses Vα2 (Fig. 5 A). This trend is also evident in peripheral T cells from Eα−/− but not wt mice in which virtually all TCRβ+ cells express Vα2 (Fig. 4 D). Thus, these data indicate that the predominance of Vα2 expression in peripheral T cells of Eα−/− mice is not caused by the fact that the Eδ enhancer only allows for Vα2 rearrangement at the DN stage of T cell development. Instead, this is likely the result of the fact that Vα2 gene segments are able to sustain a sufficiently high expression of Tcrα genes in the absence of the Eα enhancer such that only Vα2-positive αβ T cells can be positively selected and maintained in peripheral lymphoid tissue.

Eδ controls Vα expression in DN cells

Because the analysis of Ptcra−/−,Eα−/− mice has shown that Eα is not an enhancer element required for early TCRαβ expression and pre-TCR–independent T cell development (8, 11), we have focused our attention on the Eδ enhancer by generating and studying animals that lack both Ptcra as well as the Eδ enhancer element (Ptcra−/−,Eδ−/− mice). When compared with Ptcra−/−,Eδ+ thymi, it was noted that Ptcra−/−,Eδ−/− thymi contain much-reduced numbers of thymocytes (Fig. 6 A) that are severely deficient in DP and SP cells (Fig. 6 B). Also, the number of γδ TCR-expressing cells is reduced even though some γδ T cells are still present. Within the DN compartment, there is a more complete block at the DN3 stage of development in Ptcra−/−,Eδ−/− versus Ptcra−/−,Eδ+ mice (Fig. 6, C and D).

Ablation of Eδ in Ptcra−/− mice has a profound effect on the percentage and absolute number of DN αβ TCRs as well as of Vα2-expressing cells, as TCRβ+, Vα2+ thymocytes are virtually absent (<0.25%; Fig. 7, A and B). However, there was a small number of TCRβ-expressing cells among total thymocytes in some of the Ptcra−/−,Eδ−/− mice (Fig. 7), suggesting some remaining level of TCRα rearrangement in the Ptcra−/−, Eδ−/− thymi (perhaps mediated by the Eα enhancer in cells) that was “rescued” by the expression of γδ TCR at the DN developmental stage (8).

Thus, in the absence of Ptcra, the Eδ enhancer has a crucial role in the early assembly of TCRα chains as well as TCRδ chains in DN3 and/or earlier stages of T cell development (i.e., TCR δ chains that are required for the formation of γδ and αβ TCR in the absence of pre-TCR). The αβ TCRs (and γδ TCRs) can rescue some developmental progression beyond the DN3 stage in Ptcra-deficient mice, which appropriately explains the incomplete developmental block in Ptcra−/− versus CD3−/− or Tcrδ−/− × Tcrδ−/− mice (i.e., mice that cannot assemble any TCR on the cell surface).

**DISCUSSION**

The results obtained in mice with a combined deficiency of the pre-TCRα chain plus the Eα or Eδ enhancer provide an explanation for the observation that development of DN

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**Figure 6.** Eδ is essential for pre-TCR–independent T cell development. (A) Absolute cell numbers of Ptcra−/− and Ptcra−/−,Eδ−/− littermates. 4–8-wk-old mice were analyzed. Error bars represent SD. (B and C) CD4 versus CD8 (B) and TCRβ versus TCRγδ (C) profiles of littermate thymi. (D) Analysis of the DN compartment using CD25 and CD44 antibodies. Numbers in quadrants indicate the percentages of cells in that quadrant.
thymocytes into αβ lineage DP cells in Ptcra−/−,Tcrd−/− mice. Such cells were previously shown to acquire functional maturity (i.e., respond with proliferation and cytokine production to TCR ligation) and to acquire CD8α expression when antigenically stimulated. In fact, this unusual subset of T cells was analyzed in TCR transgenic mice many years ago, and it was concluded that the early expression of an αβTCR could mimic signals generated when thymocytes express γδTCR and become functionally mature (18, 20). Of interest is that such cells can express an autoreactive TCR but are not deleted because of the lack of coreceptors and can accumulate in secondary lymphoid tissue of mice, as shown in various TCR transgenic models (21). Some of these cells exhibit an activated phenotype and have an as yet undefined role in the immune system.

Recent data have, in fact, shown that the extensive accumulation of CD8α cells in the gut of TCR transgenic mice (22) to a large extent depends on the premature expression of the transgenic TCR in DN cells because “on time” expression of the same Tα transgene does not result in the strong accumulation of CD8α cells in the gut (23). Another possibility is that thymic DN αβTCR+ cells represent precursors of some peripheral regulatory T cells. Perhaps these cells are akin to both mouse and human αβTCR+CD3+ NK.1.1−CD4−CD8− DN Regulatory T cells that can suppress antigen-specific immune responses mediated by CD8+ and CD4+ T cells through a process that requires cell to cell contact and Fas–FasL interactions (24). In this regard, it is important to point out that the αβTCR+ DN studied here are different from NK1.1+ DN T cells and from αβTCR+ DN cells studied by others who concluded that all TCRβ+ DN cells were derived from DP precursors (25); either this generalization is wrong or the fate mapping approach used by the authors is not valid. However, it is clear that the TCRβ+ DN cells in Ptcra−/−, Tcrd−/− mice are involved in the rescue of development rather than being derived from DP cells (8).

The results shown in this study also provide an adequate explanation for earlier observations in Eα-deficient mice that exhibited a TCRα repertoire limited to TCRα, Vα2 chains. In this context, on Eα-deleted alleles, other cis-acting elements such as Eβ and/or Vα2 promoters were hypothesized to promote either only Vα2 to Jα rearrangements or Vα to Jα rearrangements. These rearrangements involve a diverse array of Vα segments with assembled Vα2Jα complexes expressed in a much higher proportion in the absence of Eα. Our results indicate that in DN cells, Eδ can direct a low level of Vα to Jα rearrangements, possibly via promoting Jα accessibility that results in the expression of a variety of different Vα gene segments. The early expressed αβTCRs allow the development of some DP thymocytes with TCRs containing mostly Vα2+, TCRα chains. In the absence of Eα, only early VαJγ rearrangements with a high proportion of Vα2 would continue to be expressed in DP thymocytes and mature T cells, perhaps because their promoters do not require Eα activity to drive gene expression. This pathway of differentiation observed in Eα-deficient mice might be invisible in wt mice because of continual Tinα rearrangement in DP thymocytes that will swamp out the Eδ-initiated Vα rearrangements and, thereby, lead to a much more diverse αβTCR repertoire.

**MATERIALS AND METHODS**

**Mice.** Mice were kept in the sterile facilities of The University of Chicago, Dana-Farber Cancer Institute, and Children’s Hospital. Animal protocols
were approved by the Institutional Animal and Use Committees of these institutes. C57BL/6, Rag1"/", and Tcd"/" mice (also on the C57BL/6 background) were purchased from Jackson ImmunoResearch Laboratories. C57BL/6 CD1"/" were provided by A. Bendelac (The University of Chicago, Chicago, IL). Eca"/" and E5"/" mice were generated in the laboratory of Frederick W. Alt (10, 11). Perin"/" mice were described previously (7).

Flow cytometric analysis and cell sorting. Anti-CD4 (L3T4), CD8 (53–6–7), CD25 (3C7), CD44 (IM7), NK1.1 (PK136), TCRβ (H57–597), γδTCR (GL3), Vα2 (B20.1), Vα11 (RR8–1), and Vα3 (RR3–16) mAbs were purchased from BD Biosciences. The Vα8 (CTVA8) antibody was purchased from CALTAG. These mAbs were directly coupled to FITC, PE, Cy-crome, APC, or biotin. Surface marker expression on thymocytes and peripheral T cells was visualized using a FACScalibur (Becton Dickinson) and FACS-Aria (Becton Dickinson) sorters. Cell sorting was performed using MoFlo (DakoCytomation) and FACS-Aria (Becton Dickinson) sorters.

Fetal thymic organ culture. Thymin were cultured as described previously (26). In brief, (LY5.2-) TCRβ- CD4+ CD8+ NK1.1- cells were FACS purified from thymi of Perin"/" animals. Lineage-negative (CD3, CD8, Mac-1, NK1.1, Gr-1, Ter-119, and CD19) bone marrow progenitors were also used as a reconstitution control. Isolated embryonic day 14.5 thymus from Rag1 mouse (expressing LY5.1) were initially incubated in Terasaki plates (Nunc), subsequently applied on Transwell (Nunc) porous filters, and incubated in Iscove's modified Dulbecco's medium supplemented with 10% fetal calf serum (Sigma-Aldrich) plus penicillin, streptomycin, and mercapto-ethanol. Cultures were maintained at 37°C for 7 d, after which LYS.5.2+ cells were stained and analyzed by FACS analysis.

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CORRECTION

Please note that an error appeared in the online early release html version of this article. The final html, pdf, and print versions appear correctly. For reference, the correction appears below.

Due to a production error, two typographical errors appear in the Abstract. In the second sentence, “αγ CD4+CD8+” should have read “αβ CD4+CD8+.” In the third sentence, “Eα but not Eα” should have read “Eδ but not Eα.” The corrected sentences appear below:

“However, Ptcra- as well as Tcra-targeting experiments have suggested that the early expression of Tcra in CD4+CD8− cells can partially rescue the development of αβ CD4+CD8+ cells in Ptcra-deficient mice. In this study, we show that the TCR Eδ but not Eα enhancer function is required for the cell surface expression of αβTCR on immature CD4−CD8− T cell precursors, which play a crucial role in promoting αβ T cell development in the absence of pre-TCR.”