Impairment of T Cell Development in $\delta$EF1 Mutant Mice

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Summary

Using the method of gene targeting in mouse embryonic stem cells, regulatory function of $\delta$EF1, a zinc finger and homeodomain-containing transcription factor, was investigated in vivo by generating the $\delta$EF1 mutant mice. The mutated allele of $\delta$EF1 produced a truncated form of the $\delta$EF1 protein lacking a zinc finger cluster proximal to COOH terminus. The homozygous $\delta$EF1 mutant mice had poorly developed thymus with no distinction of cortex and medulla. Analysis of the mutant thymocyte showed reduction of the total cell number by two orders of magnitude accompanying the impaired thymocyte development. The early stage intrathymic c-kit+ T precursor cells were largely depleted. The following thymocyte development also seemed to be affected as assessed by the distorted composition of CD4- or CD8-expressing cells. The mutant thymocyte showed elevated $\alpha_4$ integrin expression, which might be related to the T cell defect in the mutant mice. In the peripheral lymph node tissue of the mutant mice, the CD4+CD8+ single positive cells were significantly reduced relative to CD4+CD8+ single positive cells. In contrast to T cells, other hematopoietic lineages appeared to be normal. The data indicated that $\delta$EF1 is involved in regulation of T cell development at multiple stages.

Recent progress in our understanding of the T cell development clarified a major developmental pathway in thymus at cellular level: T cell precursors that originate from hematopoietic stem cells located in fetal liver and in adult bone marrow migrate and colonize in thymus. Starting from the CD4−CD8− double negative (DN)1 stage, thymocytes begin to rearrange their TCR genes and express CD3, a TCR coreceptor molecule, then proceed to the CD4−CD8− double positive (DP) stage. The DP thymocytes go through positive and negative selections depending on the specificity of the TCR. Finally, the CD4+CD8− or CD4+CD8− single positive (SP) mature thymocytes are produced, and these immunocompetent cells migrate out and populate the peripheral lymphoid organs (1).

Some of these steps have been assigned to specific genes, and mutant mice of such genes produced by gene targeting have contributed greatly in defining each regulatory step of T cell development (2). However, it is obvious that more knowledge of genetic regulation is required to understand cellular events in T cell development. The $\delta$EF1 mutant mice to be reported in this paper has a novel phenotype: only T cells are affected among hematopoietic lineages and major defects are found in early T cell precursors, thus defining a new step in T cell development.

$\delta$EF1 was originally identified as an enhancer binding factor of the chicken $\alpha_1$-crystallin gene (3). $\delta$EF1 is a unique protein in that it has multipartite DNA-binding motifs, containing two Kruppel-type C2H2 zinc finger clusters located close to NH$_2$ and COOH termini and a homeodomain in between (4). Subsequent analysis showed that $\delta$EF1 possesses repressive activity on transcription (5), and that $\delta$EF1 is expressed besides lens cells in various anlages of developing tissues, such as notochord, myotome, limb bud, and neural crest derivatives in chickens (4) and mice (Takagi T., H. Kondoh, and Y. Higashi, unpublished results), suggesting that $\delta$EF1 is involved in regulation of a number of genes other than the crystallin genes (6-8).

To clarify the regulatory function of $\delta$EF1 and to understand the functional significance of the multipartite DNA-binding motifs in vivo, we have initiated a study using $\delta$EF1 mutant mice of several different alleles generated by the gene targeting technique. So far, we have produced two $\delta$EF1 mutant alleles of mice: one, a null mutation, in which most of the coding sequence was replaced by bacterial $\beta$-galactosidase (Null–LacZ), the other coding for a truncated protein lacking only the COOH-proximal zinc finger clusters (ΔC-fin). Unexpectedly, as presented in this report,
one of the major phenotype of both homozygous mutant mice was impairment of thymus development: severe hypopcellularity in thymus without clear distinction of cortex and medulla. Since N-ull-LacZ homozygous mutant mice are perinatally lethal with skeletal defects (to be published elsewhere), while ~20% of the ΔC-fin homozygous mutant mice were born alive and grown up to adulthood, we analyzed the lymphoid tissues in detail using the surviving young adult ΔC-fin homozygous mutant mice. Here we describe the generation and analysis of ΔC-fin mutant mice and demonstrate that the defect of the thymus was ascribed to depletion of T precursor cells and to aberration of intrathymic development of T cells.

Materials and Methods

Mice. C57BL/6 and ICR mice were purchased from Japan SLC Inc. (Shizuoka, Japan) or CLEA Japan Inc. (Tokyo, Japan). All mice were maintained under specific pathogen-free conditions. Construction of Targeting Vector. Cloning and structural analysis of mouse ΔEF1 has been described (9). The targeting vector (see Fig. 2 A) was constructed as follows. A 0.8-kb SalI–SalI fragment containing a Sau3AI–SalI genomic fragment of the exon 6 sequence (see Fig. 2 A) and a 12-bp SalI–BamHI adapter sequence which is derived from an EMBL3 cloning vector, was subcloned into the SalI site of pBlueScript II to give pSS. The SalI and XbaI sites at the 5’ end of the inserted genomic fragment of pSS were inactivated by digesting with Xbal, partially with SalI, blunt-ending by fill-in, and self-ligation. A XbaI linker carrying stop codons in the SalI site of exon 6 of full-length pSS were inactivated by digesting with XbaI, partially with SalI, and blunt-ending by fill-in, and self-ligation. A Xbal linker carrying stop codons in all three frames (CTAGTC TAGATCAG) was inserted in the remaining SalI site at the 3’ end of the insert to have pSSstop. A Xhol–KpnI fragment of pSTN neoB (10) containing neo' sequence was inserted in the Xhol–KpnI site of the pSSstop, to have pSSstopNEO. In parallel, the 5.4-kb SalI–ApaI genomic fragment, immediately 3’ of the Sau3AI–SalI fragment was once cloned into the pBlueScript II, and regenerated by digesting with Asp718. The resulting fragment was blunt-ended, digested with SalI and cloned into the SalI–EcorI V site of DT-A vector (11), generating pSADT-A plasmid. The pSSstopNEO was digested with the Asp718, blunt-ended and digested with Noot. The resulting Asp718 (blunt-ended by Klenow)-N ot1 fragment was cloned into the SalI (blunt-ended by Klenow)-N ot1 sites of the pSADT-A, generating a final targeting vector. The vector plasmid was linearized with Not1 and used for electroporation. The expected targeted gene product lacks the COOH-terminal zinc finger clusters which have been shown to be essential for the DNA binding of ΔEF1 protein (12). The neo' element has a promoter but lacks the termination and poly(A) addition signals, so that the neo' is expressed only when poly(A) addition signal is supplied by recombination with a host gene. A DT-A cassette (11) was placed in the 3’ end of the linearized vector for the negative selection against homologous recombination and genotyping was a 1.5-kb EcoRI–Xbal genomic fragment at 0.5 kb upstream of exon 6 (shown in Fig. 2 A). Total RNA of 12.5 d.p.c. embryos were prepared by a single step isolation procedure (16). 5 μg of the total RNA was separated by electrophoresis in a 1% formaldehyde agarose gel and blotted to a Hybond N (Amersham, Buckinghamshire, England) nylon membrane. The filter was hybridized with the 32P-labeled 2.5-kb EcoRI fragment of mouse ΔEF1 cDNA containing the sequence from exon 3 to exon 8 (9).

Expression of the ΔEF1 ΔNAs in cultured Cells. A cDNA coding for ΔC-fin ΔEF1 protein was constructed by inserting the Xbal linker with stop codons in the SalI site of exon 6 of full-length cDNA in the same way as targeting vector construction. The full-length and ΔC-fin ΔEF1 ΔNAs were inserted into the Not1 site of pCDM8 (17) and transfected to COS-7 cells by lipofection as described (18).

An Antibody against N-proximal Portion of ΔEF1. An antibody against human homologue of ΔEF1, AR E6, was used to detect the N-proximal region of mouse ΔEF1. HpaI–N hel (1041-2498) fragment of human AR E6 cDNA (6) was blunt-ended and cloned into the SmaI site of pGEX-3X to produce a GST-AR E6 fusion protein. An antibody against this fusion protein, which cross-reacts to mouse ΔEF1, was used for Western blot analysis.

Immunoprecipitation and Western Blot Analysis. Nuclear extracts were prepared according to the previous report (19). From each 12.5 d.p.c. embryo, 650 μl (2 μg protein/μl) of the extract was obtained, and a 100-μl aliquot was reacted with the anti-AR E6 antisem at room temperature for 2 h. The immunocomplexes were precipitated with protein A–Sepharose (PharMingen) and dissolved in 40 μl of the SDS-PAGE buffer. Each 10-μl sample was separated by SDS-PAGE (7.5% polyacrylamide) and blotted onto a nitrocellulose filter. The blot was treated with 5% skim-milk in T BST (20 mM Tris-HCl, 150 mM NaCl, 0.05% Tween 20, pH 8.0), incubated with the anti-AR E6 antisem, washed by T BST, reacted with HRP-labeled goat anti-rabbit IgG, and processed for ECL chemiluminescence reaction (Amersham).

FACS Analysis. Multicolor analysis of lymphocytes was performed by FACScan® cell sorter as described previously. The following mAb were purchased from PharMingen (San Diego, CA): fluorescein isothiocyanate (FITC)-conjugated anti-CD45R, RA3-6B2; FITC-conjugated anti-CD8α, 53-67; FITC-conjugated anti-CD3e, 145-2C11; FITC-conjugated anti-Gr-1, R B6-8C5; FITC-conjugated anti-CD25, 7D4; PE-conjugated anti–Thy 1.2, 53-2.1; PE-conjugated anti-CD4, RM4-5; PE-conjugated anti–Mac-1, MFR4.B. FITC-conjugated anti–Gr-1, 145-2C11; FITC-conjugated anti–CD44, IM7; Biotin-conjugated anti-IgM, R 6-60/2, biontin-conjugated anti–c-kit receptor, 2B8. Biotibn-conjugated anti–c-kit integrin (CD49d), MF R 4.B. FITC-conjugated anti-IgD was purchased from Nordic Immunological Laboratories (Capistrano Beach, CA). Three-color analysis for c-kit, α4 integrin, and CD44/CD25 expression was done using Cy-Chrome-labeled streptavidin (PharMingen) as the third fluorescence dye and analyzed by FACScan® cell sorter.
Histology and Immunohistology. Tissues were fixed in Bouin's fixative and paraffin sections were stained with hematoxylin and eosin. For immunofluorescent staining of thymus, a thymus was excised from an 18.5 days post coitus (d.p.c.) embryo, and rinsed in Hapes-buffered saline (HBS), quickly frozen by dipping into liquid nitrogen, and then embedded in OCT compound (Miles Inc., Elkhart, IN). 6-μm-thick cryosections were fixed in 1% paraformaldehyde in HBS for 30 min. The sections were incubated with affinity-purified rabbit anti-ΔEF1 antibodies and biotinylated rat monoclonal anti-CD4 and anti-CD8 antibodies, then with FITC-labeled anti-rabbit Ig and Texas red-conjugated streptavidin anti-rat Ig in TBST containing 10% skim milk with washings by TBS between the steps. Finally, the specimens were mounted in Gelvatol (PBS containing 20% polyvinylalcohol, 20% glycerol, and 2.5% 1,4 diazabycyclo-[2,2,2]-octane) and examined under a microscope.

Results

Expression of ΔEF1 in Embryo and in Lymphoid Organs. A cDNA coding for mouse ΔEF1 was isolated by cross-hybridization with chicken ΔEF1 cDNA that was isolated and characterized in our laboratory (9). Comparison of the mouse ΔEF1 sequence with the chicken and other species revealed that the two zinc finger clusters and the homeodomain are highly conserved (9). Expression pattern of the mouse ΔEF1 in the embryos was almost identical to that of chicken (4): mesodermal tissues (e.g., notochord, somite, limb bud mesenchyme); neural crest derivatives (e.g., dorsal root ganglia, cephalic ganglia); a part of the central nervous system (hindbrain, motor neurons in the spinal cord). In Northern blot analysis of adult tissues, ΔEF1 mRNA was detected in all solid tissues examined (data not shown).

Among the lymphoid tissues, the ΔEF1 transcripts were detected in thymocytes, but not in splenocytes consisting of only mature T and B cells (Fig. 1, A, lanes 2 and 3). Bone marrow cells contained a low but detectable level of the transcripts (Fig. 1, A, lane 1). Immunohistological analysis showed that ΔEF1 protein was expressed in most of the thymocytes including cells stained with the mixtures of anti-CD4 and anti-CD8 antibodies (Fig. 1, B). Taken together with the results of Northern blot analysis, it was indicated that ΔEF1 is expressed in some of the bone marrow cells and in most of the thymocytes including CD4- and CD8-expressing cells, but once the cells migrate out ΔEF1 expression is lost.

Generation of ΔEF1 Mutant Mice Which Lack the Zinc Finger Cluster Proximal to COOH Terminus of ΔEF1 Protein. To generate the ΔEF1 ΔC-fin mutant allele, we constructed the targeting vector shown in Fig. 2, A. It was expected that the product of the recombinant gene lacks the C-proximal zinc finger cluster required for high affinity DNA binding (Fig. 2, B and see reference 12). Germ-line male chimeras were produced from homologous recombinant E14 ES cells (e.g., A84 in Fig. 2, C), and heterozygous mutant animals were generated by crossing these chimeras with C57BL/6 or ICR female mice. The heterozygous mice appeared normal in growth, fertility, behavior, and morphology of internal organs. Southern blot analysis of the yolk sac DNA from the 12.5 d.p.c. embryos generated by crossing the heterozygous mice showed the set of the hybridizing bands expected for the homozygous embryos (Fig. 2, C). Then we examined the ΔEF1 mRNA from the homozygous mutant embryos (12.5 d.p.c.) and compared them with the littermates of other genotypes. As shown in Fig. 2, D, transcripts from the mutated allele were longer than those from wild-type allele as expected from insertion

![Figure 1](image-url)
of the neo" sequence. The heterozygous embryos had both transcripts while the homozygous mutant embryos possessed only the longer transcript. To further confirm the mutation of δEF1 gene by homologous recombination, we examined the δEF1 protein in the mutant embryos by Western blot analysis (Fig. 2 E). The mutant protein is expected to be smaller by 40 kD than the wild-type δEF1 protein. It was demonstrated that wild-type and homozygous mutant embryos had only full-length and truncated forms of δEF1, respectively, while the heterozygous embryos had both. All these observations indicated that δEF1 gene was mutated as designed, and that the truncated form of the mutant protein was synthesized no less efficiently than the wild-type form.

Severe Hypopcellularity in the Thymus of δEF1 Mutant Mice. In embryonic stages the homozygous mutant embryos developed up to 18.5 d.p.c. with the expected Mendelian frequency. In histological analysis of the embryos from 10.5 to 18.5 d.p.c., we noted that the thymus of the homozygous mutant embryos were smaller than normal embryos. Other tissues and organs were normal in morphological and histological inspections. The number of thymocytes of 18.5 d.p.c. homozygous mutant embryos was reduced 10-fold (~5 × 10⁶ cells per thymus) compared to the heterozygous and wild-type embryos (~5 × 10⁶ cells per thymus). FACS® analysis of the thymocytes using CD4 and CD8 markers indicated that the development from the DN to DP cells appeared to be partially inhibited (data not shown). We observed no difference between wild-type and heterozygous mutant embryos.

In the postnatal period ~80% of the homozygous mutant pups died within 2 d after birth, but the remaining 20% survived and some of them had offspring. After backcrossing to C57BL/6 mice for 6 generations, 11 of the homozygous mutant mice which survived and aged from 3 to 11 wk were analyzed for lymphoid tissue development. As observed in the fetuses, thymi of all the homozygous mutant mice inspected were greatly reduced in size. In histo-
logical sections of the mutant thymi, medulla and cortex were hardly distinguishable (Fig. 3, A and B); especially, the cortex which usually consists of the densely packed and actively proliferating small thymocytes seemed to be missing in the mutant thymi (Fig. 3, C and D). Accordingly, the total cell number of thymus was greatly reduced from 1/100 to 1/500 of the heterozygous littermate (Fig. 4 A). The spleens of homozygous mutant mice were not significantly different in size from those of heterozygous littermates, although the number of splenocytes was slightly lower in homozygous mutant mice (Fig. 4 C). As will be shown below, the extent of reduction of the cell number seems to be accounted for by the reduction of peripheral T cells in the spleen. Histological inspection of the spleens of the homozygous mutant mice showed the basic architecture to be normal (data not shown). The lymph node of the homozygous mutant mice was characteristic in the reduced cellularity of the deep cortex where the T cells reside (data not shown). The number of lymphocytes recovered from a pair of inguinal lymph nodes of those mice was reduced to ~1/10 of that from the heterozygous littermates (Fig. 4 B).

The heterozygous and wild-type animals were indistinguishable in their histology and cell content of all the lymphoid organs described above. In addition, ΔEF1 mutant of another allele, Null-LacZ, which lacked almost all of the coding sequence, also exhibited the reduced size of the thymus and impaired T cell development at fetal stages as observed in the ΔC-fin mutant mice (Takagi T., H. Kondoh, and Y. Higashi, unpublished results). These observations exclude the possibility that the T cell defect in the ho-
mozygous ΔC-fin mutant mice is due to the specific effect of the truncated protein.

A aberrant T cell development in the thymus and reduced T cell population in the peripheral lymphoid organs in ΔEF1 mutant mice. Using FACS® with various cell surface markers, we first investigated the cell populations in thymi from the homozygous mutant and heterozygous control mice. In the homozygous mutant mice, the total thymocyte number was so small (see Fig. 4 A) that whole thymocytes of a mutant mouse was subjected to the FACS® analysis. A representative set of the results is shown in Fig. 5 A. The proportion of the CD4/CD8 SP or DP cell populations present was different from that found in a control heterozygous littermate: the relative proportion of cells in CD4+CD8− quadrant was reduced from 88% in normal to 57% in the mutant thymocytes while that of the SP cells was increased from 11 to 38% (Fig. 5 A). The relative frequency of αβ TCR+CD3− cells in the mutant was higher than in the control heterozygous littermate (Fig. 5 B), which was consistent with the fact that the proportion of SP cells was higher in the mutant mice compared with the control heterozygous littermate as described above.

From forward light scattering data of the FACS® analysis, which indicated the distribution pattern of cell size of a cell population examined, we found that the cell population of larger size predominantly existed in the homozygous mutant mice whereas most of the thymocytes in control mice consisted of the small cell population typical for normal mice (Fig. 5 C). The bias toward the larger size of the cell population in the mutant thymocytes seemed to be due simply to the reduced cell number of the DP thymocytes relative to that of SP cells in the mutant mice since we noticed that the most abundant cell size of the DP and SP cells in the control and mutant mice were not quite different (Fig. 5 C). However, cell size distribution pattern of DN cell population was significantly different between the control and mutant thymocytes: mutant thymocytes were not abundant in the cell population of larger size, but instead had much smaller cells compared to the control mice (Fig. 5 C).

T cells in the spleen and the lymph node were also examined by FACS®. In the mutant spleen, although decrease of total cell number in the mutant was less conspicuous than in the thymus (Fig. 4 B), the fraction of the T cells in the total splenocytes was significantly reduced from ~40% (heterozygous) to 10% (mutant) as judged by Thy1 (T cell marker) and B220 (B cell marker) expression (Fig. 6 A).

In the lymph node also, T cells were reduced in number (Fig. 7 A) and were αβ TCR+CD3+ (Fig. 7 C), indicating full maturity, but the ratio of CD4+CD8−SP to CD4−CD8+SP cells at the age of 3–8 wk was 1 in the mutant as compared to the ratio of 2 in normal (Fig. 7 B). This bias toward CD4+CD8− cell population tended to be greater in

Figure 5. FACS® analysis of thymocytes from a ΔEF1 mutant and a control heterozygous littermate. Thymocytes from 6-wk-old homozygous mutant (left) and heterozygous littermate (right) were analyzed by staining with the combination of mAbs: (A) PE-anti-CD4 vs. FITC-anti-CD8; (B) PE-anti-αβ TCR vs. FITC-anti-CD3 to assess developmental stages of thymocytes. Numbers in parentheses indicate the percentage of cells within the quadrant defined by fluorescence of cell surface markers. The thymocytes were also analyzed by the forward light scattering for the estimation of cell size (C). The histograms for whole or a portion of the thymocytes that were logically gated for the DN, DP, and SP cells in A are shown with combination of heterozygous (+/−) and mutant (−/−) thymocyte data: abscissa, forward light scattering and ordinate, relative cell number.
older mice (e.g., 20 wk, data not shown). In such old mice, the bias also became evident in the spleen though there seemed to be no such bias in the spleen of young adult mice at the age of 3–8 wk (Fig. 6B). The decrease of the T cells commonly observed in the peripheral lymphoid organs may be ascribed to the limited supply of T cells from the thymus.

Since MHC class I and II antigens on the thymic stromal cells are essential for the generation of the CD4 and CD8 SP T cells, we examined the expression of the class I (H-2K) and class II (I-A) antigens on the thymic epithelial cells by immunohistochemistry using anti H-2K and I-A antibodies. The levels of expression of these antigens were not different between ΔEF1ΔC-fin mutant and control mice, although

Figure 6. FACS® analysis of splenocytes from a ΔEF1 mutant and a control heterozygous littermate. Splenocytes from 6-wk-old homozygous mutant (left) and heterozygous littermate (right) were doubly stained with the following combinations of mAbs: (A) PE–anti-Thy1 vs. FITC–anti-B220 to assess fractions of T and B cells; (B) PE–anti-CD4 vs. FITC–anti-CD8 to assess CD4, CD8 expression in T cell population; (C) biotinylated anti-IgM plus streptavidin–PE vs. FITC–anti-B220 and; (D) biotinylated anti-IgM plus streptavidin–PE vs FITC–anti-IgD to assess development of B cells. Numbers in parentheses have the same indication as in Fig. 5.

Figure 7. FACS® analysis of lymph node lymphocytes from a ΔEF1 mutant and a control heterozygous littermate. Inguinal lymph node cells from 6-wk-old homozygous mutant (left) and heterozygous littermate (right) were doubly stained with the following combinations of mAbs: (A) PE–anti-Thy1 vs. FITC–anti-B220; (B) PE–anti-CD4 vs. FITC–anti-CD8; (C) PE–anti-β/β-TCR vs. FITC–anti-CD3. Numbers in parentheses have the same indication as Fig. 5.
the expressions of both MHC antigens were detected in the medullary region in the control mice while in the whole region in the mutant mice (data not shown). This difference may suggest and is consistent with the histological observation (Fig. 3) that the small thymi in the mutant mice lacked the typical cortex comprising the densely packed small thymocytes, most of which are DP and do not express the MHC antigens highly.

Next we examined the functional maturity of the peripheral T lymphocytes by the proliferative response to the Con A stimulation using the spleen cells as shown in Table 1. Although the degree of Δcpm of the mutant mice was about one-third of that of the control mice, it seemed that the decrease in Δcpm in the mutant mice was simply due to reduction of the number of T cells in the spleen cells of homozygous mutant mice (Fig. 6). The sensitivity to Con A stimulation and the size of colonies of the proliferating T cells were not different between the control and mutant mice (data not shown). Thus, functional maturity of the peripheral T lymphocytes accumulated in the ΔEF1 homozygous mutant mice did not seem to be affected at least in the responsiveness to the Con A stimulation.

In contrast to T cells, the development of B cells and myeloid cells was not significantly affected by the ΔEF1 mutation. B cells were comparable in number among the spleens of homozygous, heterozygous and wild-type littersmates. The level of IgM and IgD expression in splenocytes was normal in the homozygous mutant mice (Fig. 6, C and D). The population of the B cells and myeloid cells in the bone marrow was normal as analyzed using B220 (low expression), Mac1 and Gr-1 markers (data not shown). Hematocrit value of the homozygous mutant mice was also similar to the wild-type animals (data not shown). Thus, the defects in the ΔEF1 mutant mice appeared specific to T lymphocytes.

Depletion of CD4−CD8−c-kit+ thymocytes in ΔEF1 mutant mice. The FACS analysis of the mutant thymocytes using CD4 and CD8 markers indicated that, despite the severe decrease of the total cell number, intrathymic development of the T lymphocytes was not arrested at a specific stage. This is in contrast to the cases of knockout mice lacking molecules that are essential for the development and function of T lymphocytes (e.g., TCR-β [21], CD3 [22]). We suspected that the T precursor cells at a very early stage, for instance, before the rearrangement of the TCR genes, might be affected and decreased in the ΔEF1 mutant mice, and that a small fraction of the T precursor cells which escaped from the block by the ΔEF1 mutation proceeded to subsequent development.

**Table 1.** Con A–induced Proliferation of T Cells in Spleen Cells from a ΔEF1 ΔC-fin mutant and a Control Heterozygous Littermate

| Genotype | Animal* | Con A (−) | Con A (+) | Δ cpm |
|----------|---------|-----------|-----------|------|
| (+/−)    | 1       | 1,280     | 30,600    | 29,350 |
|          | 2       | 900       | 31,120    | 30,220 |
| (−/−)    | 1       | 560       | 11,140    | 10,580 |
|          | 2       | 200       | 9,110     | 8,910  |

* Numbers 1 and 2 represent different animals from different litter of 1 and 2, respectively. Animals of the same number belong to the same littermate.

1 Con A (5 μg/ml) was added to the spleen cell culture in a 96-well microtiter plate (1 × 10^6 cells/well), and incubated for 24 h. Pulse labeling with [3H]thymidine (0.5 Ci/ml) was carried out for 12 h. Incorporation of [3H]thymidine in each well was measured using Top-Counter (Packard). Values represented were average cm of the triplicate samples.

![Figure 8. FACS analysis of c-kit, CD25 and CD44 expression in CD4+CD8− cell population in thymocytes of a ΔEF1 mutant and a control heterozygous littermate.](image-url)
Figure 9. FACS\textsuperscript{a} analysis of $\alpha 4$ integrin expression in thymocytes of a $\delta E F 1$ mutant and a control heterozygous littermate. Thymocytes from a $\delta E F 1$ mutant (B) and a heterozygous littermate (A) were stained with FITC-anti-CD3 mAb, PE-anti-$\alpha/\beta$ TCR mAb and biotin-conjugated anti-$\alpha 4$ integrin mAb plus Cy-chrome streptavidin, and analyzed for $\alpha 4$ integrin expression in $\omega/\beta$ TCR high/CD3 low and $\omega/\beta$ TCR high/CD3 high populations marked by 1 and 2, respectively. Histograms of the $\alpha 4$ integrin expression of mutant (---) and heterozygous littermate (----) thymocytes were compared in C and D for populations 1 and 2, respectively. Each histogram was drawn so that the total number of cells sampled in each cell population was equal between the mutant and heterozygous littermate. A bscissa indicates intensity of Cy-chrome fluorescence, and ordinate represents relative cell number.

Discussion

The data presented in this report demonstrates that $\delta E F 1$ is essential for normal T cell development. In $\delta E F 1$ mutant mice, total lymphocyte number in thymus was reduced to $\sim$1% of that of normal mice. The cell populations found in the thymus of the mutant mice, though reduced in number, represented those of T cells with advanced development, such as CD4$^+$CD8$^+$ DP and CD4$^+$/CD8$^+$ SP cells. This phenotype is in contrast to the previously reported mutant mice of T cell development, e.g., RAG (34, 35), TCR-+$\alpha/\beta$ (21, 36), CD3 (22, 37). In these previous cases where the mutated genes were essential for critical steps of T cell development, advancement of thymocytes to subsequent stages was blocked resulting in accumulation of the cells arrested at the critical stages. The total thymocyte number tended to be smaller as the blocked stage became earlier. In the RAG-2-deficient mice, for instance, thymocytes were decreased 100-fold as was observed in $\delta E F 1$ mutant mice, and the existing thymocytes were mostly DN cells (34). The fact that the majority of thymocytes in $\delta E F 1$ mutant mice expressed CD4, CD8 markers despite the severe reduction in total thymocyte number led us to postulate that the major defect of this mutant lay at a very early stage, much earlier than that at which the TCRs were required, and a small fraction of the cells that somehow escaped from the $\delta E F 1$ mutation could go through the maturation stages.

The idea that the $\delta E F 1$ mutation impairs early T cell development was supported by the observation on c-kit$^+$ cells in the DN cell population that are considered to be the earliest intrathymic T precursor cells migrating from bone...
marrow (24). This particular cell population was largely depletes from the mutant thymocytes (Fig. 8 B). Consistent with this observation, the CD4\(^+\)CD25\(^-\) cells, a subset of cells in the early stage of the DN cell population expressing c-kit strongly (26), were also significantly reduced (Fig. 8 C). The cell size distribution of the mutant DN thymocytes measured by the forward light scattering was shifted to smaller than control (Fig. 5 C), which is again in agreement with the observation mentioned above because the c-kit\(^+\)DN thymocytes have been shown to have the largest cell size in the DN cell population (26). Furthermore, αEF1 was expressed in the thymocytes (Fig. 1 A, lane 2) and in the bone marrow cells (Fig. 1 A, lane 1), though at a low level in the latter. All these observations support the idea that αEF1 plays an essential role in the very early stages of T cell development so that abrogation of αEF1 results in the severe decrease of T cell populations accompanied by the significant loss of immature (CD4\(^-\)CD8\(^-\)c-kit\(^+\)) intrathymic T precursor cells.

It is interesting to note that the recently reported mutant mice of IL-7 receptor (IL-7R) (38), IL-2 receptor γ-chain (IL-2R γ) (39), and Jak3 (40-42), all showed severe reduction in the thymocyte number, and yet produced all DN, DP, and SP cells similar to αEF1 mutant mice. Since IL-7R, IL-2R γ, and Jak3 molecules are thought to be essential signaling molecules in expansion of the thymocytes at a very early stage before rearrangement of the TCR genes, it would be interesting to see if αEF1 is involved in such an signaling pathway.

Besides the early stage of T cell development, αEF1 may have additional regulatory roles in the later stage of intrathymic T cell development. The proportion of DN cells in total thymocytes was higher, and the ratio of DP to SP cells was lower in the αEF1 mutant mice. It was also noted that the mutant thymus did not develop medullo-cortical distinction, probably, due to lack of the typical cortex which usually consists of the densely packed and actively proliferating small thymocytes. The defects of αEF1 mutants are different in these two points from those of IL-7, IL-2R γ, and Jak3 mutant mice: the proportion of the four cell populations marked by CD4, CD8 expression are identical to the normal mice and the thymi of those mice develop distinct structures of cortex and medulla, though the thymi themselves were very small and the thymocyte numbers are ~1% of the normal mice as the αEF1 mutant mice (38, 39). These differences support the involvement of αEF1 in not only early but also late stages of thymocyte development. Reconstitution of the hematopoietic system in wild-type host using mutant hemopoietic tissues will clarify if the defects in early and late T cell development can be attributed to lymphocytes or stroma.

In peripheral lymphoid tissues the proportion of CD4\(^-\)CD8\(^-\) T lymphocytes relative to CD4\(^+\)CD8\(^-\) cells was significantly reduced from ~0.5 (wild type) to 0.2-0.1 (mutant). In older mice, the majority of the peripheral T cell population was occupied by CD4\(^+\)CD8\^- cells (data not shown). Since the MHC class I and II antigens, which are required for generation of the functional CD4\(^-\)CD8\(^-\) and CD4\(^+\)CD8\^- SP T cells, were expressed at the normal level on thymic epithelial cells of the mutant mice (data not shown), and actually the CD4/CD8 SP T cells were produced in the mutant thymus (Fig. 5 A), survival of mature CD4\(^-\)CD8\^- cells may be affected in peripheral tissues of αEF1 mutant mice.

An interesting observation was the increase of α4 integrin (CD49d) expression in the mutant thymocyte (Fig. 9). α4β1 integrins are expressed in CD34\(^+\) hematopoietic progenitors (43) and regulate lymphocyte attachment and the homing process in the endothelial microenvironment (30, 31). Thymocytes also express α4β1 integrins (44) and stage-dependent regulation of α4 integrin expression in developing thymocytes has been demonstrated (32, 33), suggesting its possible roles in T cell development. The regulatory region of the α4 integrin promoter has been analyzed and a negative element effective in lymphocytes has been identified between positions –400 and –300 in the 5'-flanking sequence (28, 45). The αEF1 binding sequence, CACCTT, is present at –357 in a region conserved between human and mouse (45), which may be the target site of αEF1. It is possible, therefore, that derepression of α4 integrin expression caused by αEF1 mutation may affect homing of T precursor cells or intrathymic development of T cells as reported here.

nil-2-a, which seems to be a truncated form of αEF1, was reported to function as a repressor of the IL-2 gene transcriptional regulation in human T cell jarkat cell line (46). The impairment of T cell development observed in the αEF1-deficient mice could result from the incomplete regulation of the IL-2 gene expression. Deregulated expression of the IL-2 gene in transgenic mouse system, however, did not show any T cell deficiency (47, 48). Moreover, the IL-2 gene knockout mice showed no effects on thymocyte and peripheral T cell subsets (49). Thus, αEF1 mutant phenotype is not explained by altered regulation of the IL-2 gene.

It has been reported that αEF1 (and its human homologue, ZEB) is expressed in myeloma and B-lymphoma cell line (7). This is not totally consistent with our observation that αEF1 mRNA were not detected in the Northern blot analysis of the splenocytes. It is possible that expression of αEF1 might be correlated to neoplastic transformation of B cells, or, alternatively, that αEF1 mRNA in matured B or T cells are too little to be detected by our Northern blot analysis using total RNA.

Although we still do not know the reason for neonatal death observed in part of the ΔC-fin mutant mice, it was rather unexpected that homozygous mutant embryos carrying the ΔC-fin allele developed normally except for the T cell defect, given the fact that αEF1 acts as a competitive repressor against bHLH activator proteins, which are widely involved in embryogenesis (12). It should be mentioned that Sna, a mouse homologue of the Drosophila snail and es-argot, is expressed at high levels in cephalic neural crest, limb bud mesenchyme, and somites of 9.5 d.p.c. embryo and later in a variety of mesenchymal tissues (50), showing close resemblance to the expression pattern of αEF1. Moreover, consensus binding sequence of Drosophila snail and
escargot proteins contain CACCTG E2 box sequence and actually these proteins counteracts E2-box-mediated activation by heterodimer of Sna and D aughterless bHLH proteins (51, 52). It is possible that loss of normal $\delta$EF1 protein was compromised by the mouse Sna protein.

In conclusion, T cells, but not other hematopoietic lineage cells, are depleted in $\delta$EF1 mutant mice although they are thought to be derived from common stem cells. Possibly, $\delta$EF1 has an important role in growth and differentiation of early T cells in the bone marrow, in the homing process and/or in development in the thymus. To date, the early T precursor cells have been understood only poorly. $\delta$EF1 mutant mice should thus provide a unique tool to study early T cell development from the aspect of transcriptional regulation.

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