Ontogeny of the immune system: γ/δ and α/β T cells migrate from thymus to the periphery in alternating waves

D. Dunon, Centre National de la Recherche Scientifique Unité de Recherche Associée
D. Courtois, Centre National de la Recherche Scientifique Unité de Recherche Associée
O. Vainio, Turku University
A. Six, University of Alabama at Birmingham
C.H. Chen, University of Alabama at Birmingham
Max Cooper, Emory University
J.P. Dangy, Basel Institute for Immunology
B.A. Imhof, Basel Institute for Immunology

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Comparative developmental studies have been informative with regard to the evolution of the immune system in vertebrates. Studies in chickens have contributed to the understanding of the hemopoietic stem cell origin of both myeloid and lymphoid T and B cell lineages (1, 2). This avian model has several advantages for the study of T and B cells undergoing differentiation in specialized central lymphoid organs, T cells in the thymus, and B cells in the bursa of Fabricius, (b) a large number of precisely staged embryos can be easily obtained, and (c) the embryos are large enough for experimental manipulation. Studies performed in chick-Quail chimeras indicate an embryonic paraaortic origin of the stem cell precursors of thymocytes, B cells, and myeloid cells, beginning around the fourth day of embryonic life (E4; 3). Embryonic stem cells native to the aortic region later migrate via the circulation to colonize the spleen, yolk sac, and, finally, the bone marrow.

The chick-Quail model has been used to show homing of the thymocyte progenitors into the embryonic epithelial thymus in three discrete waves (4-8), the first of which begins in chicken embryos on E6.5, the second on E12, and the third around E18. Each wave of progenitor cell influx lasts for 1 or 2 d, and is followed by the transient production of thymocyte progeny (7, 8). The first wave of thymus colonization involving T and B cell progenitors from the paraaortic region (7, 8), whereas the second and third waves of thymocyte progenitors come from the bone marrow and express the c-kit and the hematopoietic cell adhesion molecules (9, 10). Using congenic chicken strains that differ in the ov alloantigen expressed on hematopoietic progenitors and T lineage cells, H.B19ov+ and H.B19ov−, we have examined chimeras created by grafting thymic lobes from an ov+ donor into thymectomized ov− recipients to show the gradual replacement of donor thymocytes by ov− host thymocytes and their progeny. These experiments indicated that a series of waves or stream of thymocyte progenitors continually enter the thymus after hatching (11-13).

The ontogeny of chick T lineage cells can be monitored with anti-TCR monoclonal antibodies and molecular probes for the different TCR chains (14-16). At E12, 5 d after the initial influx of thymocyte precursors, a subpopulation of thymocytes begins to express the TCR-γδ/CD3 complex on their surface (17). These reach peak numbers on
in the second wave of thymocyte progenitors (8). TCR-\(\gamma/\delta\) cells generated sequentially in the first wave, but not faster than the H.B19ov strain was subdivided into two congenic lines, H.B19ov\(^+\) and H.B19ov\(^-\), distinguished by the ov antigen present on T lineage cells in H.B19ov\(^+\) animals. The ov antigen, which is also expressed on bone marrow cells and a B cell subset, is recognized by the 11-A-9 monoclonal antibody (9, 26, 27).

**Immunolabeling.** The ov, TCR-\(\gamma/\delta\) and TCR V\(\beta\)1 antigens were detected by the 11-A-9, TCR1, and TCR2 mAbs, respectively (17, 19, 26, 28). 11-A-9 is a mouse IgM and TCR1 and TCR2 are mouse IgG\(_1\) antibodies. Second step antibodies were fluorescein labeled, sheep anti-mouse IgM and phycoerythrin- or Texas red-coupled anti-mouse IgG1 antibodies (Southern Biotechnology Assoc., Birmingham, AL). Controls were performed using the second step antibodies alone and regular staining of tissues from noninjected individuals of the H.B19ov\(^-\) strain. Recent thymocyte emigrants, detected in blood by their FITC staining, were labeled by phycoerythrin-coupled TCR1 or TCR2 antibodies (Southern Biotechnology Assoc.). Frozen sections of embryonic organs were cut to a thickness of 5 \(\mu\)m on a cryostat (Bright, Hunkingdom, UK), fixed with acetone, and rehydrated in PBS containing 1% BSA.

Injection of Lymphoid Cells into Congenic Chickens. Adoptive transfer between H.B19ov\(^+\) and H.B19ov\(^-\) strains could be performed without complications since these strains do not differ at major histocompatibility antigens and T cell alloreactivity against a different ov antigenic determinant has not been observed in mixed lymphocyte reaction and graft versus host reactions. Bone marrow cells (2.0 \(	imes\) 10\(^7\)) from donor H.B19ov\(^+\) embryos were injected into a large vein near the aorta of recipient H.B19ov\(^-\) embryos (29). These experiments were performed with E13 and E18 age-matched donor and recipient embryos. Control injections of sorted TCR-\(\gamma/\delta\)-positive populations of E18 bone marrow cells were performed to determine that differentiated bone marrow lymphocytes were not able to colonize the thymus in this assay. For that purpose, bone marrow cells from 18-d-old H.B19ov\(^+\) embryos were suspended in PBS containing 10% FCS, filtered through a nylon sieve (mesh width of 25 \(\mu\)m; Nytal P-25 my; SST, Thal, Switzerland) and centrifuged at 225 g for 7 min. Immunofluorescence staining was performed in 96-well plates, to avoid repeated centrifugation using either the anti-TCR-\(\gamma/\delta\) antibody TCR1 or the anti-TCR V\(\beta\)1 antibody TCR2 and then fluorescein-coupled anti-mouse Ig antibody (Silenus, Hawthorn, Australia). Stained and unstained bone marrow cells were resuspended in 10% FCS/PBS and sorted using a FACSTAR Plus\(^+\) cell sorter (Becton Dickinson, Mountain View, CA). None of the recipients received irradiation or other immunosuppressive treatment. Donor ov\(^+\) cells in the thymus were analyzed by flow cytometry and by two-color immunofluorescence staining of frozen tissue sections. For analysis by FACScan\(^+\), single thymocyte suspensions were made by physical disruption in PBS and filtration through a nylon sieve.

To analyze the TCR-\(\gamma\) repertoire specifically generated by E13 and E18 bone marrow precursors, TCR-\(\gamma/\delta\) thymocytes of the donor type were sorted 9 d after injection of the precursors. Thymocytes were sorted to a two-color immunofluorescence staining using the anti-TCR-\(\gamma/\delta\) antibody TCR1 and the anti-ov antibody 11-A-9 and then FITC-coupled anti-mouse IgM and phycoerythrin-coupled anti-mouse IgG1 antibodies (Southern Biotechnology Assoc.). The cells were sorted using a FACSTAR Plus\(^+\) cell sorter. Analysis of Recent Thymocyte Emissaries. Emigration of the thymocytes into the circulation was examined in situ FITC labeling of thymocytes. Young chicks were anesthetized by intramuscular injection of 0.4 ml ketamin solution (Imalgene 500; Rhone Mérieux, Lyon, France; diluted 1:10 in PBS) followed by a short inhalation of Halothane (Hoechst, Frankfurt, Germany). The skin of the neck region was opened with scissors and each thymus lobe was injected with 10 \(\mu\)l of an FITC solution at 1

### Materials and Methods

**Animals.** Embryonated eggs from the H.B19 strain of White Leghorn chickens were produced at the Institute Chicken Facility (Gipf-Oberfrick, Switzerland). Fertilized eggs were incubated at 38\(^\circ\)C and 80% humidity in a ventilated incubator. The H.B19 strain was subdivided into two congenic lines, H.B19ov\(^+\) and H.B19ov\(^-\), distinguished by the ov antigen present on T lineage cells...
mM in DM SO. The skin was then closed using tissue clamps (Autoclip; Clay Adams, Becton Dickinson Primary Care, Sparks, MD) and the chickens were kept warm under an infrared lamp until they were fully conscious. Chickens were bled 12 h after injection and FITC-labeled lymphocytes were analyzed by flow cytometry.

cDNA Synthesis. Total cellular RNA from thymus was isolated using the guanidium isothiocyanate method (30). About 5 μg was used as template for the synthesis of randomly primed single-stranded cDNA using mouse mammary leukemia virus reverse transcriptase (Life Technologies, Inc., Gaithersburg, MD) in a reaction volume of 20 μl according to the supplier’s instructions. This cDNA was subsequently diluted to 100 μl with water and heated to 94°C for 2 min to inactivate the mouse leukemia virus reverse transcriptase.

PCR and Semiquantitative PCR of Vγ Transcripts. A PCR technique was used to amplify the TCR-γ transcripts. Transcripts deriving from rearranged TCR Vγ1, Vγ2, and Vγ3 genes were amplified independently using oligonucleotide primers specific for each Vγ family and a primer located in the Cγ region. Oligonucleotide primers CKVG1UP2, CKVG2UP3, and CKVG3UP1 were specific for the Vγ1, Vγ2, and Vγ3 regions, respectively. The CKCG1DO1 oligonucleotide primer was located at 230 nucleotides downstream the 5’ end of the Cγ region. The procedures used for semiquantitative PCR followed the detailed description given by Keller et al. (31). The amount of cDNA synthesized was calibrated by using the relative expression level of β actin as a standard. The two actin oligonucleotide primers 4611 and 4612, generated a band of 283 and 648 bp on cDNA and genomic DNA respectively (32). The following are CKVG1UP2 (Vγ1 region): GTACACGAGAGAGATC; CKVG2UP3 (Vγ2 region): CATACAGAGCCTGTATC; CKVG3UP1 (Vγ3 region): GATACTGTACATGTCTGG; CKCG1DO1 (Cγ region, antisense): CGAMMA1DD (5’ of Cγ, sense): TACCACATCTACGTGGACAAGT; CGAMMA1DD (5’ of Cγ, antisense): TCTGTGTTCTCCGTGC; 4611 (5’ of actin, from nucleotide 2057, see reference 32): TACACCAATGTACCCGGG; 4612 (3’ of actin, from nucleotide 2704, antisense, see reference 32): CTCGTCTTGTGTATGC.

PCR reactions were in 30 μl using 1 U of Taq polymerase (Perkin-Elmer Cetus, Norwalk, CT). PCR buffer was prepared as suggested by Perkin-Elmer, but with the addition of 10 mM β mercaptoethanol. Reaction mixtures were denatured by heating to 96°C for 5 min, and then subjected to 32 rounds of amplification using a Biometra T-Rio thermoblock thermocycler under the following conditions 96°C for 15 s, 50°C for 40 s, and 72°C for 1 min for cDNA amplification. Final extension was done at 72°C for 10 min.

Cloning and Sequencing of TCR-γ Transcripts. The TCR-γ V-J-C regions were specifically amplified by PCR. Amplified DNA fragments were gel purified and cloned into pCR™ II vector (Invitrogen, CA, USA). Sequences were determined from denatured double-stranded recombinant plasmid DNA (33) using Sequenase™ (Amersham Corp., Arlington Heights, IL) in the chain termination reaction (34) and the oligonucleotide primer CGAMMA1DD starting 60 nucleotides downstream the 5’ end of Cγ segment in the antisense orientation. In a number of cases where ambiguities remained, several additional nucleotide primers were used. Sequences were analyzed and assembled with the software package of the CITI2 (Paris, France). TCR-γ cDNA sequences have been submitted to the EMBL/GenBank/DDBJ database under accession numbers z97216 to z97332.

Results

Pathways of T Hymocyte Precursor Migration into the Embryonic Thymus. The initial wave of progenitor colonization and thymocyte differentiation can be examined in situ in unmanipulated animals. However, analysis of members of the succeeding waves requires a discriminating strategy. Embryonic hemopoietic precursors express the ov antigen in H.B19ov+ birds, and the antigen is maintained on T lineage cells and a subset of B cells. This expression pattern allowed us to examine successive waves of the ov+ progenitor populations in the embryonic thymus and the fate of their T cell progeny in ov− congenic recipients. To examine the second wave of thymus colonization by progenitor cells, E13 bone marrow cells of the ov+ strain were injected into E13 chicken embryos of the congenic H.B19ov− strain, and the thymocyte progenitor influx, migration, and differentiation patterns were examined in thymus sections by immunohistochemistry. By E16, donor cells of bone marrow origin accumulated in the thymic blood vessels, both in interlobular venules and in capillaries located at the corticomedullary junction (Fig. 1); relatively few donor cells were found in the parenchyma of the thymus at this time. E19 ov+ cell invasion and accumulation within the thymic cortex was evident, but by E20 the donor cells had relocated to occupy the outermost cortex of the thymic lobules. This ontogenetic pattern suggests that thymocyte progenitors entering the embryonic thymus either via the corticomedullary junction or the capsular subsequently make their way to the outer cortex of the thymus (Fig. 1 C). The donor cells were later found throughout the cortex and by day 23, mature ov+ donor T cells had begun to accumulate in the medulla. This complex intrathymic pattern of migration appears specific to bone marrow-derived thymocyte progenitors, since mature thymocytes and T cells failed to home to the thymus in other adoptive transfer experiments (not shown).

Intrathymic Differentiation Kinetics Are Consistently Accelerated for the γδ T Cell Subpopulation. The appearance of γδ T cells precedes that of Vβ1 α/β cells by a period of ~3 d during the initial wave of thymocyte development (18), but studies in chick–quail chimeras suggest this may be a one time occurrence (8). In our studies of the second wave of thymocyte differentiation, the injection of E13 H.B19ov+ bone marrow cells into E13 H.B19ov− embryos led to the appearance of donor γδ+ thymocytes 5 d later and donor α/β thymocytes ~8 d later (Figs. 2 and 3 A). The proportion of ov+ donor thymocytes expressing TCR-γ/δ peaked at 40% on day 21. The first donor α/β (Vβ1+) thymocytes were detected on E20 and these reached a peak level of 57% on E26 (Fig. 3 A). When the third wave of thymocyte differentiation was examined by injection of E18 ov+ bone marrow cells into E18 ov− recipients, the same rule held true; γδ T cells appeared 4 d before the α/β T cells (Fig. 3 B). Interestingly, for the cell transfer experiments performed during the second wave of precursor colonization, the level of chimerism was relatively greater for γδ than for α/β T cells (Fig. 3 C). However, taking into account the proliferation kinetics for the progeny of each precursor wave, the TCR-γ/δ thymocyte progeny appeared to be 12 to 16 times less numerous than the TCR-α/β (Vβ1) thymocyte progeny (Fig. 3 D, and
and a/b T Cells Migrate from Thymus to the Periphery in Alternating Waves

Figure 1. Migration pathways of thymocyte progenitors. Thymus sections from E13 H.B19ov+ embryos injected with donor E13 H.B19ov+ bone marrow cells were examined by differential immunofluorescence staining at E16, E19, E20, and E23. Donor ov+ progenitors are labeled with fluorescein; TCR-γδ and TCR-αβ-positive cells are labeled with Texas red. Original magnifications: (A) 270, (B) 170. Scale bars correspond to 100 μm. (C) Diagrammatic representation of chicken T cell progenitor migration pathways in the thymus. At E16, T cell progenitors were located either in capillaries at
The levels of donor $\gamma/\delta$ and $\alpha/\beta$ thymocytes peaked at days 23 and 26, respectively, corresponding to the main period of second wave emigration to the periphery. The differential chimerism of $\gamma/\delta$ and $\alpha/\beta$ T cells thus may reflect the differential emigration kinetics of $\gamma/\delta$ and $\alpha/\beta$ T cells.

Mature $\gamma/\delta$ and $\alpha/\beta$ T thymocytes migrate to the periphery in alternating waves. The colonization of the thymus in discrete waves (7, 8), and the differences in differentiation and emigration kinetics of $\gamma/\delta$ and $\alpha/\beta$ thymocytes suggest interspersed emigration of the mature $\gamma/\delta$ and $\alpha/\beta$ thymocyte subsets (11–13). To test this hypothesis, we examined the phenotype of recent thymocyte migrants at different developmental ages. In these experiments, thymocytes of chicks at 21 (hatching)–30 d were labeled in situ by intrathymic injection of FITC. Blood samples were obtained 12 h later and labeled lymphocytes in the circulation were analyzed for expression of TCR-$\gamma/\delta$ or TCR-$\alpha/\beta$ (V$\beta$1) (Fig. 4). The FITC-labeled cells represented 3–10% of the peripheral blood lymphocytes. Approximately 75% of these were $\gamma/\delta$ or $\alpha/\beta$ (V$\beta$1$^+$) T cells; of the remaining 25% approximately half were $\alpha/\beta$ (V$\beta$2$^+$) T cells and the rest were TCR$^+$. Peaks of recent $\gamma/\delta$ thymocyte emigrants were detected on days 21–23 and 27–28, and a peak of recent $\alpha/\beta$ thymocyte emigrants was observed on days 24–26. The frequency of FITC-labeled $\gamma/\delta$ thymocyte emigrants reached a maximum of 20%, whereas the peak of labeled V$\beta$1 emigrants reached a maximum of 70%. These figures reflected the fact that each precursor wave gives rise to $\sim$5% $\gamma/\delta$ and 75% $\alpha/\beta$ (V$\beta$1$^+$) thymocytes, respectively. Thymocyte progenitors in each colonization period thus gave rise to $\gamma/\delta$ T cell progeny within 9 d and $\alpha/\beta$ (V$\beta$1$^+$) T cell progeny within 12 d, and these migrated in the same sequence to the periphery.
TCR-α/β recipients were analyzed by immunofluorescence flow cytometry: ov, for T cell expression as a function of developmental age. Thymocytes of the gated thymocyte population. Hatching occurs at E21.

Each Thymocyte Progenitor Wave Contributes a Characteristic TCR-α/β Repertoire. 

Since each colonization period is followed by a refractory interval of ~4 d, the end result is alternating emigrant waves of γδ and αβ (Vβ1) T cells, with minimal overlap of migrant cells representing each of the three embryonic waves of thymocyte progenitors.

Each T-homotypic Progenitor Wave Contributes to a Characteristic TCR-γδ Repertoire. The TCR-γδ repertoire expressed by each progenitor wave was addressed in these experiments. Simple sorting of E13 thymocytes reactive with the anti-TCR-γδ antibody was sufficient to isolate first wave progeny for repertoire analysis. Since the progeny of successive waves may accumulate in the thymus during development, isolation of the second and the third wave progeny required the more complex strategy of adoptive transfer between the H.B19ov+− congenic chicken strains. The progeny of second wave progenitors was isolated after the injection of E13 bone marrow cells from H.B19ov+− donors into age-matched H.B19ov− recipients. The ov+ TCR-γδ+ thymocytes (14,000 cells) were then sampled on day 22. To purify the progeny of the third progenitor wave, a similar adoptive transfer was made into E18 embryos, and ov+ TCR-γδ+ thymocytes in the recipients were sorted (1,400 cells) on day 28.

To examine the TCR-γ repertoire expressed in each developmental wave of thymocytes, we performed a PCR using a 3′ primer specific for Cy and 5′ primers specific for Vγ1, Vγ2, or Vγ3 segments. First, we determined that Vγ1-Jγ-Cγ, Vγ2-Jγ-Cγ, and Vγ3-Jγ-Cγ transcripts were present as early as E10-11 in ov+ embryos, confirming in this congenic strain that the three different Vγ subfamilies undergo rearrangement simultaneously (Fig. 5A). A similar analysis of the second and third wave progeny indicated that all three Vγ gene subfamilies also undergo rearrangement in these waves (Fig. 5B). However, the Vγ2 subfamily representation differed among three waves. The TCR-γ repertoire of the second wave was composed mainly of Vγ2 transcripts, whereas transcripts containing the Vγ1 and

Figure 2. Differentiation kinetics of the second wave of thymocyte progenitors. After adoptive transfer of E13 H.B19ov+ bone marrow (2.0 × 10⁶ cells) into E13 H.B19ov− embryos, the donor cells were examined for T cell expression as a function of developmental age. Thymocytes of the gated thymocyte population. Hatching occurs at E21.

Figure 3. Comparative differentiation kinetics of the second and third waves of thymocyte progenitors: appearance of TCR-γδ- and TCR-α/β (Vβ1) T cell progeny. (A and B) Proportion of thymocytes expressing TCR-γδ or TCR-α/β (Vβ1) among donor ov+ thymocytes was determined by immunofluorescence flow cytometry. Each point corresponds to the mean value for three to five animals in four independent experiments. The differentiation of second wave T cell progenitors was analyzed by adoptive transfer of E13 H.B19ov+ bone marrow into E13 H.B19ov− embryos, and for analysis of the third wave, E18 H.B19ov+ bone marrow cells were injected into E18 H.B19ov− embryos (injections of 2.0 × 10⁶ cells). (C) Proportions of donor ov+ thymocytes that express TCR-γδ or TCR-α/β (Vβ1) during the differentiation of the second wave of progenitors. (D) Proportions of donor ov+ TCR-γδ- and ov+ TCR-α/β (Vβ1) thymocytes derived from second wave thymocyte progenitors among the total thymocyte population in the recipient. The realistic efficiency of the γδ and Vβ1 T cell differentiation pathways followed by the progeny of the second wave of thymocyte progenitors can be estimated, since the areas under these curves are roughly proportional to the numbers of γδ and Vβ1 thymocytes produced by the donor second wave progenitors. This assessment suggests that the donor progenitors produced ~15 times more αβ (Vβ1) than γδ thymocytes.
Vγ3 gene segments were weakly represented. In contrast, Vγ1, Vγ2, and Vγ3 transcripts appeared to be equally well represented in the first and the third thymus colonization (Fig. 5 B).

A more detailed analysis of the representative TCR-γ repertoires was performed by cloning of the PCR products and sequencing ~30 clones for each Vγ gene subfamily/thymocyte wave (Figs. 6 and 7). In the H.B19ov chicken strain, six Vγ1, 12 Vγ2, and 10 Vγ3 members were detected. The Vγ2 subfamily was divided into eight Vγ2a members, three Vγ2b members, and a new member, Vγ2c, that differs significantly from the Vγ2a and Vγ2b subfamilies. In contrast to the differences in Vγ subfamily usage (Fig. 5 B), significant differences were not seen in the representation of members within a given Vγ subfamily for the three thymocyte waves (Fig. 7). Different members of the three Vγ families were found to rearrange with all three different Jγ segments. However, preferential pairings of Vγ and Jγ segments were observed. In each of developmental waves, Vγ2 segments were rearranged with Jγ2 segments (Table 1). In addition, variable Jγ usage patterns were seen in Vγ1- and Vγ3- containing transcripts for the three thymocyte waves. A high frequency of Vγ1/Jγ2 rearrangements in the first thymocyte wave was rearranged with the Jγ1 segment in the first wave and with the Jγ3 segment in the following waves.

A striking feature of the TCR-γ transcript analysis was the occurrence of recurrent clones exhibiting the same Vγ-Jγ junction in the second and third thymocyte waves (Fig. 7). (a) 30 identical Vγ1-Jγ1-Cγ clones (2v12) were found in the second thymocyte wave and 9 such clones (3v14) were found in the third thymocyte wave. (b) 19 identical Vγ3-Jγ3-Cγ clones were found in the second wave (2v32), and 7 clones of this type (3v32) were encountered in the third thymocyte wave. Thus, the low frequency of Vγ1 and Vγ3 subfamily usage in the second thymocyte wave was associated with an increased representation of repetitive clones.
Discussion

Three discrete waves of thymocyte progenitors enter the embryonic chick thymus to generate three successive waves of T cell progeny members which leave the thymus to colonize peripheral organs such as the spleen and the intestine. The present studies indicate that each wave of progenitors gives rise first to γ/δ thymocytes and then α/β (Vβ1) thymocytes 3 or 4 d later so that the γ/δ and α/β (Vβ1) T cells migrate asynchronously from the thymus. Progenitor colonization of the thymus in waves and an accelerated rate of γ/δ T cell differentiation thus contribute to the alternating emigration of first γ/δ and then α/β T cells to the periphery (Fig. 8). Analysis of the TCR-γ repertoire for the different thymocyte waves suggests that they differ in Vγ and Jγ usage as well as in CDR3 diversity.

Thymus colonization by waves of hemopoietic progenitors also appears to occur in mammals (35, 36), but can be examined in greater detail in the chicken because of embryonic accessibility and the opportunity to purify the progeny of individual waves of progenitors. A adoptive transfer of alloantigen-marked progenitors allowed us to elucidate the homing routes whereby these embryonic thymus T cells are guided to the cortical thymic medulla. This analysis indicates that progenitors of bone marrow origin enter the thymus either via interlobular venules or capillaries located at the corticomedullary junction. Both routes have been described in the mouse, but were not shown to be used simultaneously, the transcapsule route thought to be restricted to thymus colonization before its vascularization (37, 38). In the congenic chick chimeras, progenitors entering at the corticomedullary junction subsequently migrated to the outer cortex of the thymus, where precursors entering through the capsule were also found. This outer cortical homing pattern of thymocyte precursors has also been noted after direct needle injection in mice. As has been described in mammals (37), thymocytes then migrate from the outer cortex as they undergo T cell differentiation en route to the thymic medulla.

An interesting question concerns whether each embryonic wave of precursors generates the same or different T cell repertoires. Studies of the TCR-γ repertoire generated during mouse development indicate sequential usage of Vγ genes. The first γ/δ T cells generated during embryonic development express Vγ5-Cγ1 transcripts, the second population of γ/δ T cells express Vγ6-Cγ1 transcripts, and γ/δ T cells become more heterogeneous for Vγ usage after birth (22, 25). In similar fashion, the Vβ1 gene segments are rearranged before the Vβ2 gene segments during avian ontogeny (39). However, the Vγ gene families do not undergo sequential rearrangement during ontogeny. The chicken TCR-γ locus consists of three Vγ families with ~10 members each, 3 Jγ segments, and 1 Cγ segment (15, 16). The first wave of thymocyte progeny rearrange all three Vγ families and Jγ genes as early as E10-E11. This type of unrestricted TCR-γ rearrangement pattern has also been suggested in sheep and humans (40, 41). On the other hand, preferential Vγ Jγ pairings were observed for the three developmental waves of thymocytes, whereas preferred TCR Vβ1/D/Jβ rearrangements were not apparent during ontogeny (13). The Vγ Jγ junctional variations (CDR3) for all three embryonic thymocyte waves were more limited than in the adult (16). Such differences between embryonic and adult repertoires have also been found in mammals (42). Finally, the nonproductively rearranged TCR-γ transcripts observed in TCR-γδ+ov+ thymocytes indicate that Vγ Jγ rearrangements occur on both alleles in avian γ/δ T cells.

A high frequency of repetitive TCR-γ transcripts was found in the second and third waves of thymocyte progeny, particularly in the second wave of thymocytes where two clones, 2v12 and 2v32, represented 97 and 66% of the Vγ1 and Vγ3 repertoires, respectively. This result could...
represent a PCR artefact or a sampling error due to the limited numbers of cells being analyzed. However, several observations suggest that the TCR-\(\gamma\) repertoire may differ for the different waves. (a) The \(V_\gamma 1\) and \(V_\gamma 3\) repertoire diversities were higher in the third wave than in second, even though the number of \(g/d\) donor-derived thymocytes examined in the second wave was ten times higher than for the third wave (\(1.4 \times 10^3\) versus \(1.4 \times 10^3\) \(g/d\) cells). (b) Repetitious clones were also encountered in the third wave repertoire, albeit at lower frequencies (31% for 2v12-3v14, 23% for 2v32-3v32). (c) The cDNA synthesis, PCR amplification, and product cloning procedures were repeated three times to confirm the findings. (d) Repetitive transcripts were not encountered in the \(V_\gamma 2\) repertoire preferentially used by the second wave. Thus, the high frequency of repetitive invariant clones in \(V_\gamma 1\) and \(V_\gamma 3\) repertoires created by cells of the second wave is in agreement with a lower usage of \(V_\gamma 1\) and \(V_\gamma 3\) families in this wave. Differences in the TCR-\(\gamma\) repertoire generated in each progenitor wave could reflect differences in the colonizing progenitors. The first wave of thymocyte progenitors are derived from multipotent hematopoietic stem cells that arose in the paraaortic region of the embryo (3), whereas the second and third waves of thymocyte progenitors were derived from the bone marrow. The generation of different TCR-\(\gamma\) repertoires could therefore reflect differences in the progenitor cells themselves. The differentiation kinetics of \(g/d\) and \(a/b\) \((V_\beta 1)\) thymocytes were conserved for the three developmental waves of thymocyte progenitors. The times required for differentiation of \(g/d\) and \(a/b\) \((V_\beta 1)\) thymocytes were \(\approx 9\) and 12 \(d\), respectively, in the chicken.
The accelerated differentiation of avian γδ versus αβ thymocytes has also been noted for mouse thymocyte precursors (43, 44). Different selection processes for the αβ and the γδ thymocytes may contribute to the differences in differentiation kinetics (45, 46). Different growth requirements during γδ and αβ T cell differentiation, such as IL-7 requirement, may also promote different differentiation kinetics (47).

The thymocyte emigration model whereby peripheral tissues are colonized by γδ T cells before αβ (Vβ1+) T cells may favor harmonious development of a strategic immune defense system mediated by interacting lymphocyte subpopulations. The interaction of γδ T cells with cells in the peripheral organs could provoke microenvironmental modification that favors the homing of αβ T cells. The location of γδ and αβ T cells in peripheral tissues may play an important role in the establishment of an immune response, in that recent evidence suggests γδ T cells may function as regulatory cells for αβ T cells (48–50).

During organogenesis, new niches for lymphocyte homing may appear throughout development. The interspersed emigration of γδ and Vβ1 thymocytes might provide a mechanism to fill these niches by γδ and αβ cells in developing organs. The observed interspersed emigration of γδ and αβ (Vβ1+) thymocytes could also affect the homing patterns of specific thymocyte subpopulations. For example, two thymocyte subpopulations are emigrating from the thymus at day 21-23, the γδ thymocytes generated by the second wave of precursors and the minor second subpopulation of αβ (Vb2+) T cells generated by the first wave of progenitors (8). Interestingly, γδ thymocytes colonize the intestine in massive numbers, whereas Vb2 T lymphocytes are rarely seen in that organ (20). These two populations may compete for homing sites, thus diverting the Vb2 population to other organs. Since an optimal immune response may require collaboration between γδ and αβ T cells, sequential migration of γδ and αβ thymocyte subpopulations may provide an efficient means to maintain a physiological balance between the two cell populations during development.
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Address correspondence to Pr. D. Dunon, UPM C, CNRS URA 1135, Equipe Adhésion et Migration Cellulaires, Bâtiment C-30, Boîte 25, 7ème étage, 9, Quai Saint-Bernard, 75252 Paris Cedex 05, France. Phone: 33-1-44-27-35-00; FAX: 33-1-44-27-34-97; E-mail: dunon@ccr.jussieu.fr. The present address of B.A. Imhof is Centre Medical Universitaire, Department of Pathology, CH-1211 Geneva, Switzerland.

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