Early T Cell Receptor β Gene Expression Is Regulated by the Pre-T Cell Receptor-CD3 Complex

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Summary

We have examined the question of whether there is an additional checkpoint in T cell development that regulates T cell receptor (TCR)-β expression in CD25−44− thymocytes by mechanisms that are independent of the pre-TCR. Our analysis in various mutant mice indicates that all changes in cytoplasmic TCR-β expression can be accounted for by pre-TCR-dependent signal mediation, putting into question the function of a putative pro-TCR.

Key words: lymphocyte development • pre-T cell receptor • cytoplasmic T cell receptor β • clonotype-independent CD3 complex

Several years ago, it was found that on the surface of immature T cells CD3e could be expressed in the apparent absence of TCR chains for antigen, and that signals could be transduced by cross-linking this complex with CD3 antibodies (1–3). Since then, it has been discussed whether CD3e surface expression in apparent association with calsenin (4) represents a biological accident without implication for physiological T cell maturation (5), or whether it is perhaps indicative of a pro-TCR complex that controls development. This would represent yet another checkpoint, in addition to those controlled by the pre-TCR (6) and the α/β TCR (7). In fact, it was recently argued that the second possibility was likely to be correct, since impaired signal transduction in p56ck and CD3e-deficient mice appeared to be associated with reduced expression of productively rearranged TCR-β genes (8, 9). However, it could not be excluded from these studies that the deficiency in TCR-β gene expression was dependent on defective signaling by the pre-TCR or the α/β TCR, since the analysis included thymocyte subsets whose generation depended either on the pre-TCR (some CD25−44− cells [10]) or the α/β TCR (CD25−44− NK T cells [11]). For that reason, we have analyzed TCR-β expression by intracellular staining in a variety of mutant mouse strains in small CD25−44− cells that are independent of the pre-TCR and the α/β TCR. Our data indicate that all differences that exist in TCR-β gene expression among various mouse strains can be attributed to signal transduction by the pre-TCR, since small CD25−44− thymocytes of pre-TCR α chain-deficient mice exhibited the same phenotype observed in CD3-deficient mice, thus arguing against a distinct checkpoint in development controlled by the CD3 complex in the absence of the pre-TCR.

Materials and Methods

Animals. CD3ε−/−, pTα−/−, and γc−/− mice have been described (6, 12, 13) and were bred in the specific pathogen-free animal facilities of the Necker Institute, Paris, C57BL/6 and Rag2−/− mice were purchased from Iffa Credo. Animals were analyzed at 6–8 wk of age. Animal care was in accordance with institutional guidelines.

Antibodies and Immunofluorescence Analysis. The following mAbs were used: CD25 (PC-61) conjugated to FITC, CD44 (Pgp-1)–PE, CD25–PE, CD4 (L3T4)–CyChrome, CD8 (Ly-2)–CyChrome (PharMingen), and anti-pan TCR-β (H57-597). Biotinylated mAbs were revealed with either streptavidin-PE (Southern Biotechnology) or streptavidin-allophycocyanin (APC; Molecular Probes Europe). Cells were stained in microtiter plates (105 cells/well) using combinations of directly conjugated mAbs. Simultaneous four-color cell analyses were performed on a FACSCalibur™ flow cytometer (Becton Dickinson). Dead cells were excluded by gating based on forward and side scatter characteristics.

Intracellular Staining for TCR-β Chains. TCR-β gene expression in CD4−8− subset by negative depletion of CD4/CD8− cells using Dynabeads (Dynal). For extracellular/intracellular double staining, cells were first incubated with culture supernatant of mAb 2.4G2 to block FcRII/III receptors. Cells were then stained with FITC-conjugated anti-CD25, PE-conjugated anti-CD44, and CyChrome-conjugated anti-CD4 and anti-CD8 at optimal concentration. After washing in PBS, cells were fixed in PBS plus 0.5% paraformaldehyde for 15 min at room temperature, followed by two washing steps in PBS. Cells were then per-
meabilized in 0.5% saponin for 10 min at room temperature and washed in PBS. Intracellular staining with biotinylated anti-pan TCR-β (H57-597) diluted in PBS/2% FCS plus 0.5% saponin was performed on 30 min at 4°C, washed twice in PBS/2% FCS, and revealed for 30 min at 4°C by streptavidin-APC diluted in PBS/2% FCS plus 0.5% saponin. Cytoplasmic staining was followed by two washing steps in PBS and 15 min on a rocking platform in PBS/2% FCS plus 0.5% saponin on ice. Finally, cells were washed in PBS/2% FCS.

Results and Discussion

Intracellular TCR-β Gene Expression in CD4-8− Subsets of Thymocytes. We have analyzed thymocytes from wild-type, γc−/− (12), pTα−/− (6), CD3ε−/− (13), and Rag2−/− mice (14) in order to analyze the effect of each mutation on TCR-β gene expression in small CD25−44− cells. The subset distribution among CD4−8− cells according to CD44 and CD25 expression is shown in Fig. 1. Wild-type and γc−/− mice exhibit a similar phenotype except for an elevated proportion of CD44+25+ cells in the latter due to a partial block at this stage of development in γc−/− mice. pTα−/− mice look similar to CD3ε−/− and Rag2−/− mice, but due to their incomplete block at the CD44−25− stage of development, contain more CD44−25− cells than the latter two strains. Of these, some 70% are γδ T cells (6). When intracellular TCR-β expression versus CD25 expression was analyzed in all CD4−8− cells, it became clear that wild-type and γc−/− thymocytes express TCR β chains in the majority of cells, but γc−/− thymocytes less so because of an early partial block before TCR-β rearrangement at the CD44−25− stage (12). In these two strains, most TCR-β expression was present in CD25− cells. In contrast, in pTα−/− and CD3ε−/− mice, most TCR-β expression was found in CD25+ cells, although less completely so in pTα−/− mice because of a partial developmental block at the CD25+44− stage resulting in a population of CD25+44− cells, of which up to 70% are γδ T cells. Of these γδ T cells, up to 25% expressed cytoplasmic TCR β chains (15), which accounts for the cytoplasmic TCR-β staining in the CD25− cells in pTα−/− mice (Fig. 2 A). There is naturally no TCR-β expression in Rag2−/− mice (Fig. 2). However, this picture changed, to some extent, when the analysis was performed on smaller cells where the proportion of TCR-β+ cells among CD25+ cells was significantly decreased in wild-type and γc−/− mice but not at all or only marginally in pTα−/− and CD3ε−/− mice (Fig. 2 B). What is also apparent in Fig. 2 B is that the proportion of TCR-β+ cells among small CD25− cells is significantly smaller in wild-type and γc−/− mice, while it is larger in pTα−/− and CD3ε−/− mice. This is due to the fact that in CD25− cells from pTα−/− and CD3ε−/− mice, TCR-β rearrangement proceeds further than in normal mice (16, 17). It is also clear from Fig. 2, A and B, that CD25+ cells in wild-type and γc−/− mice express on average higher TCR-β levels than CD25− cells from pTα−/− and CD3ε−/− mice, and that with regard to this parameter there is no significant difference between CD25+ cells from pTα−/− and CD3ε−/− cells. Actually, there is a continuous spectrum of TCR-β expression rather than a discrete peak, which would be expected from a population of cells that undergoes TCR-β rearrangement and begins to express productive genes. Nevertheless, there is no doubt that the staining is specific, since there is no staining in the same population of cells in Rag2−/− mice (Fig. 2), and also because an irrelevant control antibody of the same Ig class does not stain in all different mouse strains (data not shown). Thus, all differences that exist between wild-type and CD3ε−/− mice with regard to TCR-β expression in CD25+ cells can be attributed to defective signaling by the pre-TCR rather than to an independent control of TCR-β expression by the CD3 complex alone.

We have focused on TCR-β expression in small CD25+44− cells only, and it is clear that in this thymocyte subset the proportion of cells expressing TCR-β in their cytoplasm is much smaller than in a population that contains CD25+ as well as CD44+ cells, due to the presence of CD44+ NK T cells in the latter population that express α/β TCRs on their cell surface (11) and were included in previous analyses (8). The NK T cell population is likely to be absent in ikc−/−, ζ−/− mice because it is a highly selected population that requires signaling through the CD3 complex. Thus, this population would depend on signaling by the α/β TCR rather than just by p56cki and ζ chains, as shown.
suggested by Wurch et al. (8). The NKT cell population, for reasons so far unknown, is absent in the pTα−/− mice (18). Moreover, the CD25−/− subset contains a population of pre-TCR−dependent large cells that apparently was included in previous studies (8). If one eliminates these populations of cells from analysis, one nevertheless finds differences in the proportion of CD25+ cells expressing TCR-β, as well as in the level of TCR-β per cell between wild-type and CD3ε−/− mice: in the former, fewer CD25+ cells express TCR-β and at higher levels compared with CD3ε−/− mice. However, the same difference is also noted between wild-type and pTα−/− mice, indicating that this difference is dependent on signal transduction by the pre-TCR rather than signal transduction by p56lck and CD3ζ in the absence of the pre-TCR, as suggested previously (8, 9).

In summary, analysis of TCR-β expression in CD25+ cells argues against the notion that levels of TCR-β are upregulated by a pro-TCR rather than the pre-TCR. Rather, they demonstrate that it is the pre-TCR complex in which CD3 signal transducing molecules exert their biological function for the first time in the development of α/β lineage cells.

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