Human TCR alpha/beta+ CD4-CD8- Double-Negative T Cells in Patients with Autoimmune Lymphoproliferative Syndrome Express Restricted Vbeta TCR Diversity and Are Clonally Related to CD8+ T Cells.

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Human TCR α/β⁺ CD4⁻CD8⁻ double-negative T cells in patients with autoimmune lymphoproliferative syndrome (ALPS) express restricted VB TCR diversity and are clonally related to CD8+ T cells¹

[Running title: Repertoire Diversity and CD8 origin of DN T cells in ALPS]

Anne Bristeau-Leprince*,†, Véronique Mateo‡, Annick Lim*,†, Aude Magerus-Chatinet‡, Eric Solary§, Alain Fischer‡, Frédéric Rieux-Laucat‡ and Marie-Lise Gougeon*,†,2

* Antiviral Immunity, Biotherapy and Vaccine Unit, Infection and Epidemiology Department, Institut Pasteur, 28 rue du Dr. Roux, 75015 Paris
† INSERM U668, F-75015 Paris, France
‡ INSERM U768, Paris F-75015; Université René Descartes-Paris 5, Paris F-75006, Hôpital Necker-Enfants Malades, 149 rue de Sèvres, 75015 Paris, France
§ UMR INSERM U866, F-21079 Dijon, France; IFR-100 University of Burgundy, F-21079 Dijon, France, CHU, Dijon Cedex, France

Corresponding author:
Marie-Lise Gougeon, Tel: + 33145688907, Fax +33145688909, email :mlgougeo@pasteur.fr

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Abstract
The peripheral expansion of $\alpha/\beta^+$-CD4$^+$CD8$^-$ double negative (DN) T cells in patients with autoimmune lymphoproliferative syndrome (ALPS) is a consistent feature of this disease, and part of the diagnostic criteria of ALPS. The origin of these cells remains undetermined. They could derive from mature T cells that have lost co-receptor expression, or represent a special minor cell lineage. To investigate relationship of DN and single positive (SP) T cells in ALPS, we used Immunoscope technology to analyze the TCRV$\beta$ repertoire diversity of sorted DN and SP T cells, and we performed CDR3 sequence analyses of matching clonotypes. We show that DN T cells express all the V$\beta$ gene families that are used by their SP counterparts, though they dominantly use some V$\beta$ genes. Analysis of CDR3 length distribution revealed a diverse polyclonal TCR repertoire for sorted CD4$^+$ T cells, whereas both DN and CD8$^+$ T cells showed a skewed TCR repertoire with oligoclonal expansions throughout most of the V$\beta$ families. CDR3 sequencing of matching clonotypes revealed a significant sharing of CDR3 sequences from selected V$\beta$-J$\beta$ transcripts between DN and CD8$^+$ T cells. Altogether, these data strongly argue for a CD8 origin of DN T cells in ALPS.
Introduction

The autoimmune lymphoproliferative syndrome (ALPS)³ (also known as Canale-Smith syndrome) is one of the first human inherited disorders of apoptosis to be described (1,2). It is a rare disorder of disrupted lymphocyte homeostasis caused by an impaired Fas/CD95-mediated apoptosis, resulting in chronic lymphoproliferation and a breakdown in immunologic tolerance (3-7). The main features of this disease are chronic splenomegaly and lymphadenopathy of early onset, hypergammaglobulinemia (IgG and IgA), autoimmune manifestations, and expanded populations of T-cell receptor (TCR)-α/β⁺ CD3⁺CD4⁻CD8⁻ T cells, called double negative T cells (DN T cells) (3-7). The genetic defect responsible for this syndrome is in most cases a mutation in the Fas gene.

Fas, a member of the tumor necrosis factor (TNF) receptor superfamily, induces apoptosis of lymphocytes when triggered by its ligand (FasL). The interaction between Fas and FasL leads to the formation of the death-inducing signaling complex, which initiates a cascade of caspases that culminates in apoptosis (8). The Fas apoptotic pathway is crucial for the downregulation of the immune response, and loss-of-function mutations in the Fas (TNFRSF6) or FasL gene results in chronic lymphoid hyperplasia and autoimmunity, as seen in mice with lpr and gld mutations (9) and in patients with ALPS (2). ALPS is classified according to underlying genetic defect (10). In ALPS type I, heterozygous Fas mutations (ALPS type Ia) (11,12) or heterozygous FasL mutation (ALPS type Ib) (13) are usually associated with a partial defect in apoptosis mediated by Fas and its ligand. ALPS type II, which is characterized by resistance to Fas-mediated apoptosis, despite the presence of normal Fas and FasL, has been found in patients with caspase-10 mutations (14). Somatic heterozygous mutations of Fas were found to cause a sporadic form of ALPS (ALPS type III) by allowing lymphoid precursors to resist to normal process of cell death(15).

Lymphoproliferation results from the gradual accumulation of lymphocytes that have not undergone normal programmed cell death, and expansion of an unusual population of peripheral DN T cells is striking. In the blood, DN T cells constitute less than 1% in normal individuals but can reach up to
40% in patients with ALPS (4,16). These DN T cells express αβ TCR chains, they exhibit an unusual phenotype including expression of the CD45RA isoform (B220) and altered cell surface O-glycans (16), and they are high producers of IL-10, in contrast to DN T cells from normal individuals (17,18). Besides elevated IL-10, ALPS patients exhibit a Th2 cytokine profile that may favor autoantibody production (19). The origin of DN T cells remains elusive. They could be either previously activated mature T cells that have lost CD8 or CD4 co-receptor expression, or a special minor cell lineage selectively expanded owing to defect in Fas signaling. To investigate the relationships of DN T cells and the single positive (SP) T cell populations (CD4+ and CD8+), we have analyzed the TCR diversity and characterized complementarity determining regions 3 (CDR3) of TCR Vβ chains in sorted DN and SP CD4+ and SP CD8+ T cells in patients with ALPS type 1a.
Materials and methods

Patients and samples

Blood samples were obtained from three patients with ALPS type Ia. All patients, or their parents, provided written informed consent, validated by the Ethics Committee (Comité Consultatif pour la Protection des Personnes en Recherche Biomédicale) from the Necker Hospital. Table 1 summarizes the clinical features of the patients.

Flow cytometry and sorting of double-negative T cells

Peripheral blood mononuclear cells (PBMC) were isolated from freshly drawn heparin-treated blood by means of Ficoll-hypaque density gradient centrifugation. The proportion of DN T cells was determined by flow cytometry on PBMC following staining with monoclonal antibodies (CD3-APC, CD4-PE, CD8-FITC and TCR\(\alpha\beta\)) purchased from Becton Dickinson. DN T cells (CD4\(^{-}\)CD8\(^{-}\)CD3\(^{+}\)TCR\(\alpha\beta\)) were identified as shown in Fig. 1A. Sorting of DN T CD3\(^{+}\)CD4\(^{-}\)CD8\(^{-}\) cells from the three patients was performed with a FACSstar with 98% purity. Sorted DN T cells therefore contain a majority of \(\alpha\beta\) T cells. Possible presence of \(\gamma\delta\) T cells within sorted DN T cells does not skew the data since repertoire analyses were performed with V\(\beta\)-specific primers.

Determination of V\(\beta\) gene usage and Immunoscope analysis

Determination of V\(\beta\) gene usage and Immunoscope analysis was performed as described in A. Lim et al. (20). Briefly, total RNA was extracted using the RNeasy mini-kit (Qiagen, Courtaboeuf, France) according to the manufacturers specifications. cDNA was then prepared by RNA reverse-transcription with 0.5\(\mu\)g/\(\mu\)l oligo (dT)17 and 200 U of Superscript II reverse transcriptase (Invitrogen, Cergy Pontoise, France). An aliquot of cDNA synthesis reaction was amplified with each of the 24 TCR V\(\beta\) family-specific primers, together with a TCR C\(\beta\) primer and a Minor Groove Binder (MGB)–TaqMan probe (Applied Biosystems). Real-time quantitative PCR was carried out in an ABI7300 device (Applied Biosystems, Foster City, CA). In a second approach, two microliters of each of these amplification reactions were used as template in run-off reactions.
using a nested fluorescent primer specific for the Cβ segment. In this reaction, all PCR products were copied into fluorescently labeled single-stranded DNA fragments, irrespective of their TCR Jβ usage or CDR3 sequence. These fluorescent products were separated on an ABI-PRISM 3730 DNA analyzer (Applied Biosystems). The size and intensity of each band were analyzed with Immunoscope software (21,22). Fluorescence intensity was plotted in arbitrary units on the y-axis, whereas the x-axis corresponds to CDR3 length in amino acids. The Gaussian distribution of the different CDR3 lengths is characteristic of normal Vβ repertoire.

**DNA cloning and sequencing analysis of TCR transcripts**

Vβ/Cβ PCR were carried out on cDNA with 5 U Pfu polymerase (Stratagene) for 25 cycles. PCR products were then cloned in pCR®4Blunt-TOPO vector (Invitrogen Life Technologies). For sequencing purposes, PCR was carried out directly on LacZ’ colonies with Taq polymerase (Promega, Madison, MI) (23). Sequencing was carried out with the Big Dye Terminator Reaction Kit (Applied Biosystem) and analyzed on an ABI PRISM 3730 DNA Analyzer.
Results

Clinical Features

Demographic and clinical features of the study patients are shown in Table 1. Patients presented with ALPS, characterized by a lymphoproliferative syndrome consisting in massive splenomegaly (splenectomy was performed at age 12 years in P1 because of hypersplenism), and lymphadenopathy consisting in multiple lymph nodes sometimes larger than 2 cm (P1). Autoimmune manifestations, were detected in P1 and P3, consisting in urticarial rash and arthralgia in P1, uveitis, glomerulopathy and autoimmune thrombocytopenia in P3. Hypergammaglobulinemia was found in all three patients, associated with hyper IgA in P1 and P3, and normal or low (P3) serum IgM level. This was associated with a heterozygous mutation in the Fas gene, and a defect in anti-CD95 monoclonal antibody induced-apoptosis, as previously described for such patients (12).

In all cases, an elevated proportion of TCRαβ+/CD4-/CD8- DN T cells was detected in peripheral blood, the size of this population varying from one patient to another (Table 1) (12).

Purification and immunophenotypic characteristics of DN T cells in ALPS patients

The proportion of DN T cells in the study patients ranged from 4.4% to 17.2% of peripheral blood a/ß T cells (Fig. 1A). SP and DN T cells were purified from patients’ PBMC by positive selection of CD3⁺ T cells with magnetic beads, followed by FACS sorting of three subsets: CD3⁺CD4⁺ SP, CD3⁺CD8⁺ SP, CD3⁺CD4⁺CD8⁻ DN T cells. DN T cells from these patients exhibited the phenotype of mature “end stage” cells i.e. TCR-high, CD2⁺, CD5⁺, CD27⁺, CD28⁺, CD45RA⁺RO⁻ CD31⁺, CD62Ldull, CXCR-5⁺, CD57⁺, CD11b (data not shown). Furthermore, analysis of the cytotoxic granule content showed that ex vivo DN T-cells from ALPS patients were mostly GrzA positive (for example 90.2% for P2) while GrzB was barely detected (Fig. 1B, 1C). In contrast, CD8⁺ T-cells from the same individual significantly over-expressed GrzA and GrzB, as compared to control, and granzymes were detected in CD4⁺ lymphocytes too (Fig. 1B, 1C). These data suggest that DN T cells are unable to exert a cytolytic granule-dependent activity. In addition, as it was demonstrated recently (24), cytotoxic granules-dependent cell-death in SP T cells may compensate for Fas-
deficiency in ALPS.

*The TCR Vβ-gene usage of αβ DN T cells from ALPS patients quantitatively differs from that of SP cells*

The expression of 24 Vβ gene families was examined by RT-PCR amplification of cDNA derived from sorted subpopulations. Representative data are shown for a control donor (Fig. 2A) and three patients (Fig. 3). Qualitatively, repertoire diversity was conserved in all three subsets from ALPS patients, and in particular DN T cells expressed all Vβ-gene families that were used by CD4+ and CD8+ SP cells (Fig. 3). However, quantitative differences in the expression of several Vβs were apparent in each of the three patients, with preferential expression of some Vβs by DN T cells. For example, Vβ2 and Vβ7 families were preferentially expressed in DN T cells from P1 (Vβ2: 25% in DN T cells vs 12% and 8% in SP CD8 and CD4 SP T cells respectively; Vβ7: 30% in DN T cells vs 12.1% and 2.5% in SP CD8 and CD4 SP T cells respectively). Similarly, a preferential usage of Vβ17 and Vβ7 genes was found in DN T cells from P2 and P3 respectively (Fig. 3).

*CD8 SP and DN T cells from ALPS patients express a skewed TCR Vβ repertoire, in contrast to CD4 T cells*

Analysis of CDR3 length distribution in sorted CD4 T cells from ALPS patients showed for all Vβ-Cβ rearrangements Gaussian Immunoscope profiles (representative profiles are shown in Fig. 4), which is the hallmark of a polyclonal T cell repertoire observed in control donors (Fig. 2) (20). In sharp contrast, SP CD8 T cells from ALPS patients displayed skewed Immunoscope profiles with oligoclonal expansions for some Vβ families (Fig. 4), such as Vβ2, Vβ8, Vβ13a, Vβ14, Vβ17, Vβ18, and Vβ20 for P1, Vβ1, Vβ2, Vβ5, Vβ8, Vβ13b, Vβ17, Vβ22, for P2, and Vβ1 and Vβ13b for P3 (Fig. 4). This restricted diversity was not observed in sorted SP CD8 T cells from healthy donors (Fig. 2), as previously reported (20). DN T cells from ALPS patients displayed highly limited clonality with characteristic clonotypes for some Vβ families (eg Vβ2, Vβ7, Vβ13a, Vβ13b, Vβ14, Vβ20 for P1, Vβ1, Vβ5 and Vβ22 for P2) while others showed more diverse patterns with specific
T cell expansions (for example VB1, VB21, VB22 and VB23 for P1, VB2, VB8, VB13a, VB22 for P2, VB1, VB13b, VB14, VB18 for P3 (Fig. 4). Oligoclonality of both CD8 SP and DN T cells was confirmed by sequence analysis selecting within VB families matching clonotypes that had the same CDR3 length in TCR transcripts from CD8 SP and DN T cells (as exemplified for patient 2 in Fig. 5). Table 2 indicates the frequency of unique in-frame CDR3 sequences within the total analyzed selected VB TCR clonotypes from CD4 SP, CD8 SP and DN T cells from the three patients. In DN T cells from P1, as few as 14 unique CDR3 sequences out of 88 VB2 TCR, 10 out of 72 VB14 TCR, 39 out of 230 VB18 TCR and 36 out of 233 VB23 TCR were identified. A total of 682 CDR3 sequences were generated from selected transcripts, among which only 124 were unique (18%), confirming VB gene skewing among DN T cells. In contrast, paired CD4 SP T cells revealed polyclonal frequency of CDR3 sequences with almost no repeat (66 unique CDR3 sequences out of 75 VB2 TCR and 78 out of 80 VB8 TCR), as found on a total of 155 CDR3 sequences among which 144 were unique (93%) (Table 2). Oligoclonality of SP CD8 T cells suggested by Immunoscope profiles was confirmed by sequence analysis, the frequency of unique CDR3 sequences being intermediate between SP CD4 and DN T cells, 34 unique CDR3 sequences out of 80 VB2 TCR, 21 out of 84 VB14 TCR, 94 out of 196 VB18 TCR and 103 out of 216 VB23 TCR. Overall, a total of 631 CDR3 sequences were generated from selected transcripts, among which 275 were unique (44%). A similar skewed oligoclonal frequency of CDR3 sequences was found in patients 2 and 3 DN T cells (Table 2).

**CDR3 sequence analysis of TCR transcripts containing common clonotypes in DN T and SP CD8 T cells**

To address whether DN T cells may originate from SP CD8 T cells, we identified dominant clonotypes in DN T cells and selected clonotypes with similar CDR3 size in SP CD8 T cells for sequence analysis (e.g. VB7 in P2, Fig. 5). Selected TCR transcripts of common clonotypes were cloned and the resulting DNA clones were subsequently analyzed for CDR3 sequences using corresponding VB and JB primers. Fig. 6 shows that for each selected clonotype, CDR3 sequences
were detected at very high frequency in DN T cells, consistent with the oligoclonality of the Vß repertoire. For example, the Vß7 clonotype in P2 included two major CDR3 sequences, SLRQGLSTEA and SQGTGGNQPQ (76% and 6.4% of total analyzed sequences respectively). Importantly, both CDR3 sequences were also detected in SP CD8 T cells, but with a much lower frequency (4.8% and 0.4% of total analyzed sequences respectively) (Fig. 6). Similarly, the Vß13 clonotype in P3 was dominated by an 11aa CDR3 sequence (SSDSGGSYNEQ), which represented 53% of the clones in that gene family. This CDR3 sequence was also detected in SP CD8 T cells, though at a lower frequency, as expected. In P1, the 9 aa CDR3 sequence RPGLPNPEA was detected both in DN and SP CD8 T cells. Overall, the Immunoscope analysis and CDR3 sequencing experiments argue for the hypothesis that DN T cells from ALPS patients may represent, at least in part, originally SP CD8 T cells that down-modulated the co-receptor CD8. In addition, the extent of oligoclonality together with phenotypic features of DN T cells suggest recent antigenic activation by a limited set of antigens, possibly self antigens.
**Discussion**

The expansion of α/β-DN T cells in peripheral blood of ALPS patients is a consistent feature of this disease, and part of the diagnostic criteria of ALPS (1,10). DN T cell expansion is also evident histologically in secondary lymphoid organs, particularly in the paracortical areas of lymph nodes, where they accumulate and express markers of activated T cells, i.e. ki67+, CD45R0-CD45RA+ (25). Lymphoproliferation in ALPS is associated with chronic generalized activation, as demonstrated by upregulation of HLA-DR expression on peripheral CD3 T cells as well as high levels of activation markers such as soluble interleukine-2 receptor, soluble CD30 and IL10 in sera of ALPS patients (7,26,27). Consistent with their unregulated peripheral proliferation and possible recognition of self antigens, DN T cells exhibit the phenotype of mature antigen-experienced cytotoxic T cells, i.e. TCR\(^{\text{high}}\), CD2\(^+\), CD5\(^+\), CD27\(^{++}\), CD28\(^+\), CD45RA\(^-\)RO\(^-\) CD31\(^+\), CD62L\(^{\text{dull}}\), CXCR-5\(^-\), CD57\(^+\), CD11b\(^{16}\). The uniform phenotype of α/β-DN T cells from ALPS patients is similar to α/β-DN T cells from mice with defective Fas (lpr) (7,16,28) and different from the minor α/β-DN T cell compartment in healthy subjects that contains multiple subpopulations (29,30). To investigate the origin of DN T cells from ALPS patients, we analyzed with Immunoscope technology the TCR Vß repertoire diversity of sorted DN T cells and SP CD4 and CD8 T cells, and we performed sequence analyses of the CDR3 region within Vß families showing, on Immunoscope profiles, common clonotypes between DN and SP T cells. This study shows for the first time that DN T cells from ALPS patients share with CD8 T cells unique CDR3 sequences across several TCR Vß families, arguing for the CD8 origin of DN T cells.

Vß gene family usage was examined by quantitative RT-PCR amplification of cDNA derived from sorted DN and SP T cells. Though the Vß-gene usage of DN T cells did not dramatically differ from that of SP T cells in ALPS patients – i.e. DN T cells expressed all the Vß gene families that were used by the SP counterparts, preferential expression of some Vß was found in DN T cells as compared to autologous SP T cells. However, Vß usage skewing in DN T cells varied from one patient to the other. Analysis of CDR3 length distribution revealed dramatic differences between all
three subsets. A Gaussian distribution of CDR3 peaks for all Vß-Cß rearrangements, the hallmark of a polyclonal TCR repertoire, was obtained in sorted SP CD4 T cells from the ALPS patients. In contrast, DN and SP CD8 T cells displayed skewed TCR-Vß Immunoscope profiles, with oligoclonal or monoclonal expansions throughout most of the Vß families, indicating a contracted TCR-Vß repertoire. Oligoclonality of both SP CD8 and DN T cells was confirmed by sequence analysis selecting, within Vß families, matching clonotypes that had the same CDR3 length in TCR transcripts. For example, in DN T cells from P1, a total of 682 CDR3 sequences were generated from selected transcripts, among which only 18% were unique, in contrast to SP CD4 T cells that showed 93% of unique CDR3 sequences. The frequency of unique CDR3 sequences was intermediate (44%) in SP CD8 T cells from P1. Similar data were found for patients 2 and 3. That expanded DN T cells originate from SP CD8 T cells is suggested by several instances of shared CDR3 sequences from selected Vß-Jß transcripts. For example, the Vß7 clonotype in DN T cells from P2 was dominated by two major CDR3 sequences (SLRQGLSTEA and SQGTGGNQPQ) that were both detected in SP CD8 T cells. We report similar findings for Vß13 clonotype in P3, or Vß 23 clonotype in P1. Interestingly, CDR3 sequences that are shared between DN and CD8 T cells could not be detected in CD4 T cells. Therefore, from this study, we conclude that DN T cells from ALPS patients are oligoclonally expanded, and they originate from SP CD8 T cells, as suggested by shared Vß CDR3 sequences within several Vß families. Differences in Vß gene usage by DN as compared to CD8 T cells suggest that the former originate from the expansion of a minor CD8 subset. Overall, this conclusion differs from the one proposed in a recent study, focusing on CDR3 spectratyping profiles of sorted DN and SP T cells from a single ALPS patient. The authors concluded that DN are not clonally related to SP T cells because of the lack of CDR3 size matching of the oligoclonal expansions detected on spectratype profiles from the three subsets (31). However, the lack of CDR3 sequence analysis in this study precludes any valid conclusion on the origin of DN T cells.

The accumulation of DN T cells in murine models of ALPS, i.e. mice lacking expression of Fas
(lpr) or Fas-ligand (gld), suggests that this subset might represent a subpopulation destined for apoptosis in normal mice (9). It would arise from high affinity interactions of TCR with peptide/MHC, and would accumulate to vast numbers in the absence of Fas-induced apoptosis (32).

The SP T cell origin of DN T cells has been suggested by several findings, including their bearing of a demethylated CD8 gene (33,34), and their absence in mice lacking expression of class I MHC (35). In ALPS patients, the homogenous phenotype of DN T cells, that characterizes mature antigen-experienced cytotoxic T cells, suggest their CD8 T cell origin (16). Interestingly, the peculiar expression pattern of cytolytic proteins by DN T cells makes them unable to exert a cytolytic granule-dependent activity. Indeed, they strongly express granzyme A, whereas granzyme B and perforin are barely detected, as opposed to granule expression in SP T cells (24) (and this study). As recently discussed (24) the repressor mechanism that down-regulates expression of granzyme B and perforin while sparing granzyme A in DN T cells may involve IL-10 [highly produced by DN T-cells (17,36)] previously shown to repress granzyme B mRNA expression in cytotoxic lymphocytes (37). This repressor mechanism could be involved in the modulation of the CD4 or CD8 co-receptors (38).

The mechanisms involved in peripheral expansion of DN T cells in ALPS patients are not fully understood. Holzelova et al. have shown that resistance to Fas-mediated apoptosis would confer a selective advantage to mutant cells, leading to their accumulation and evolution towards the phenotype of DN T cells (15). The recent observation that in vivo triggering of mitochondrial apoptosis by rapamycin in murine ALPS induces a dramatic decrease in DN T cells (39) supports this hypothesis. The lack of co-receptors in DN T cells from ALPS patients may also contribute to their accumulation. Indeed, it has been reported in a TCR transgenic murine model that co-receptor engagement controls expansion of normal T cells, and in the absence of co-receptor T cells survive chronic stimulation and express B220, as seen in ALPS disease (40). In the present study, DN T cells appeared to be oligoclonally expanded, as evidenced by CDR3 length distribution. This extent of oligoclonality together with phenotypic features suggesting historical antigen encounter and
recent activation of large proportions of these cells may reflect stimulation of DN T cells by a limited yet ubiquitous set of antigens. The antigen specificities recognized by the TCR of DN T cells have not been defined, but the general assumption is that they will include self-antigens. Interestingly, this study shows for the first time that CD8 T cells from ALPS patients (children and young adults) exhibit a restricted TCR diversity, with oligoclonal and monoclonal expansions throughout most of the VB families, and redundant CDR3 sequences. This is reminiscent of repertoire contraction and oligoclonal expansions seen in aged individuals, and supposed to be a heterogeneous phenomenon including impaired homeostasis and antigen-driven stimulation (41). The conserved CDR3 sequences between CD8 T cells and DN T cells reported here suggest that they may recognize identical peptide sequences, and argue for their filiation. It has been proposed that the defect in killing of lpr CD8 T cells following stimulation by self-antigen, would lead to the modulation of CD8 co-receptor and accumulation as IL-10 secreting anergic cells (42,43). The comparative molecular analysis of CDR3 distribution and sequencing of CD8 T cells and DN T cells in the present study extends for the first time this hypothesis to the human disease.
References


_Lancet_ 348:719-723


31. Marlies, A., G. Udo, B. Juergen, S. Bernd, M. Herrmann, and J. P. Haas. 2007. The expanded double negative T cell populations of a patient with ALPS are not clonally related to CD4+ or to CD8+ T cells. _Autoimmunity_ 40:299-301


34. Wu, L., M. Pearse, M. Egerton, H. Petrie, and R. Scollay. 1990. CD4^-CD8^ thymocytes that express the T cell receptor may have previously expressed CD8. _Int. Immunol._ 2:51-56.

35. Mixter, P.F., J. Q Russell, F. H. Durie, and R. C. Budd. 1995. Decreased CD4^-CD8^- TCR-
αβ+ cells in lpr/lpr mice lacking b2-microglobulin. *J. Immunol.* 154:2063-2067


Footnotes

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2 Correspondence: Marie-Lise Gougeon, Antiviral Immunity, Biotherapy and Vaccine Unit, Institut Pasteur, 25-28 rue du Dr. Roux, 75725 Paris Cedex 15, France, email: mlgougeo@pasteur.fr

3 Abbreviations used: ALPS, Autoimmune Lymphoproliferative Syndrome; SP, Single Positive; DN, Double Negative
**Figure legend**

**Figure 1.** *FACS analysis of cytotoxic granule content of peripheral T cells in control and ALPS patients.* Freshly drawn heparinized peripheral blood samples were stained for TCRαβ, CD3, CD4 and CD8 expression. Cells were next stained for GrzA, GrzB and perforin content.

*A:* surface CD4 and CD8 co-expression on gated TCRαβ⁺ CD3⁺ T cells from a control donor and the three ALPS patients. The percentage of cells within each quadrant among gated cells is indicated.

*B:* Intracellular GrzA and GrzB content in gated CD3⁺CD4⁺, CD3⁺CD8⁺, and CD3⁺CD4⁻CD8⁻DN T cells from a control donor and P2. *C:* Intracellular perforin and GrzB content in gated CD3⁺CD4⁺, CD3⁺CD8⁺, and CD3⁺CD4⁻CD8⁻DN T cells from a control donor and P2.

**Figure 2.** *Quantitative Vß repertoire analysis and Immunoscope profiles of SP T cells from a control donor.*

*A:* Quantitative Vß repertoire was determined by real-time PCR analysis on sorted CD4⁺ and CD8⁺ T cells from a representative healthy donor. The x axis indicates Vß families and the y axis their relative frequency of usage.

*B:* Immunoscope profiles of TCR Vß analysis on the same sorted SP T cells. Profiles for the 24 Vß families are shown. The x-axis indicates CDR3 length (amino-acid), and the y-axis displays arbitrary fluorescence intensity of the run-off products

**Figure 3.** *Quantitative Vß repertoire analysis on SP and DN T cells from ALPS patients.* Quantitative Vß repertoire was determined by real-time PCR analysis on sorted SP CD4⁺ and CD8⁺ T cells, and DN T cells from the three ALPS patients. The x axis indicates Vß families and the y axis their relative frequency of usage. Arrows point out the expansion of some Vß families detected in DN T cells as compared to SP T cells from the same individual.
**Figure 4.** *Immunoscope profiles of TCR Vβ analysis of SP and DN T cells from ALPS patients.* The diversity of TCRVβ repertoire from sorted SP and DN T cells from ALPS patients was analyzed and Immunoscope profiles are shown. The x-axis indicates CDR3 length (amino-acid), and the y-axis displays arbitrary fluorescence intensity of the run-off products. Immunoscope profiles of CD4 T cells are Gaussian for all Vβ families, indicative of a diverse repertoire, as observed in control donors (Fig. 2). Immunoscope profiles of CD8+ and DN T cells are oligoclinal in most of the Vβ families, indicative of a skewed restricted repertoire. **Vβ families targeted for further CDR3 sequence analysis are pointed out with a star.**

**Figure 5.** *Oligoclonality of both CD8+ T cells and DN T cells for indicated Vβ families confirmed by sequence analysis,* as shown in the upper part of the figure for each cell subset. For a given Vβ, the number of sequences obtained for each CDR3 length was representative of the Immunoscope profile described in Fig. 4. Identification of matching clonotypes with same CDR3 length in TCR transcripts from CD8 SP and DN T cells from ALPS P2 was performed within Vβ1, 5, 7, 17, 20 24 families, as shown by the hatched lines.

**Figure 6.** *CDR3 sequence analysis of selected TCR Vβ transcripts of dominant clonotypes between CD8+ T cells and DN T cells.* Selected TCR transcripts were cloned and the resulting DNA clones were subsequently analyzed for CDR3 sequences using corresponding Vβ and Jβ primers. For each selected clonotype, the frequency of CDR3 sequences in DN and SP CD8+ T cells is indicated.
**Table 1 : Patients’ clinical and immunological characteristics**

<table>
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<th>Patient No</th>
<th>Fas Mutation</th>
<th>Sex/Age (yrs)</th>
<th>Age at presentatio n (yrs)</th>
<th>LA*</th>
<th>Spl†</th>
<th>Splenectomy Age (yrs)</th>
<th>AI‡</th>
<th>Serum Ig‡‡</th>
<th>Lymphocyte subsets¶ (%)</th>
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<td>IgA</td>
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<td>K280fs</td>
<td>M/32</td>
<td>6</td>
<td>++</td>
<td>+++</td>
<td>(12)</td>
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</tr>
<tr>
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<td>S214fs</td>
<td>M/10</td>
<td>0.3</td>
<td>+</td>
<td>+++</td>
<td>-</td>
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</tr>
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<td>3</td>
<td>S227fs</td>
<td>F/22</td>
<td>4</td>
<td>+</td>
<td>++</td>
<td>-</td>
<td></td>
<td>N</td>
<td>G,T</td>
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</tbody>
</table>

*LA : Lymph-Adenopathy + : multiple nodes, size < 2 cm, ++ : multiple nodes 2 cm < size < 5 cm; †Spl : Splenomegaly + : above umbilicus, ++ : below umbilicus, +++ : palpable in iliac fossa; ‡AI : Auto Immunity 0 : absence of autoimmune manifestation; UR : Urticarial Rash; Ar : Arthralgia; Uv : Uveitis; G : Glomerulopathy; T : Thrombocytopenia; ‡‡Serum Immunoglobulin level: N: within the normal range, > +2SD <+4SD, > +4SD; ¶ Lymphocyte counts in controls are 2000-4000 x10³/ml, with 70-80% CD3+ cells, 40-50% CD4+ cells, 20-30% CD8+ cells, 10-20% CD19+ cells and 10-20% of CD16+CD56+ cells. ** DN T cells : % CD4⁺CD8⁻ among TCR α/β T cells (normal limit : 2%); na : not available
<table>
<thead>
<tr>
<th>Donor Cells</th>
<th>Vß2</th>
<th>Vß8</th>
<th>Vß14</th>
<th>Vß18</th>
<th>Vß23</th>
<th>Total</th>
</tr>
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</tr>
<tr>
<td>DN</td>
<td>14/88 (16%)</td>
<td>25/59 (42%)</td>
<td>10/72 (14%)</td>
<td>39/230 (17%)</td>
<td>36/233 (16%)</td>
<td>124/682 (18%)</td>
</tr>
<tr>
<td>CD8+</td>
<td>34/80 (42%)</td>
<td>23/55 (42%)</td>
<td>21/84 (25%)</td>
<td>94/196 (48%)</td>
<td>103/216 (48%)</td>
<td>275/631 (44%)</td>
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<tr>
<td>CD4+</td>
<td>66/75 (88%)</td>
<td>78/80 (97.5%)</td>
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<td>144/155 (93%)</td>
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<tr>
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<td>Vß1</td>
<td>Vß5</td>
<td>Vß7</td>
<td>Vß17</td>
<td>Vß20</td>
<td>Vß24</td>
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</tr>
<tr>
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<td>40/42 (95%)</td>
<td>44/57 (77%)</td>
<td>25/186 (13%)</td>
<td>87/230 (38%)</td>
<td>43/75 (57%)</td>
<td>29/55 (53%)</td>
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<tr>
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<td>45/73 (62%)</td>
<td>9/13 (69%)</td>
<td>152/247 (61%)</td>
<td>91/142 (64%)</td>
<td>50/89 (56%)</td>
<td>66/86 (77%)</td>
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<tr>
<td>DN</td>
<td>45/94 (48%)</td>
<td>62/156 (40%)</td>
<td>56/102 (55%)</td>
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<td>163/352 (46%)</td>
</tr>
<tr>
<td>CD8+</td>
<td>57/69 (83%)</td>
<td>132/148 (89%)</td>
<td>35/64 (55%)</td>
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<td></td>
<td>224/281 (79%)</td>
</tr>
</tbody>
</table>