

Monoclonal TCR-Vβ13.1⁺/CD4⁺/NKa⁺/CD8^{-/+dim} T-LGL lymphocytosis: evidence for an antigen-driven chronic T-cell stimulation origin

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Monoclonal TCR $\alpha\beta^+$ /CD4⁺ T-large granular lymphocyte (T-LGL) lymphocytosis is a T-cell disorder with a restricted TCR-V β repertoire. In the present study we explored the potential association between the expanded TCR-V β families, the CDR3 sequences of the TCR-V β gene, and the HLA genotype of patients with monoclonal TCR $\alpha\beta^+$ /CD4⁺ T-LGL lymphocytosis. For that purpose, 36 patients with monoclonal TCR $\alpha\beta^+$ /CD4⁺ T-LGL lymphocytosis (15 TCR-V β 13.1 versus 21 non–TCR-V β 13.1) were selected. For each patient,

both the HLA (class I and II) genotype and the DNA sequences of the VDJ-rearranged TCR-V β were analyzed. Our results show a clear association between the TCR-V β repertoire and the HLA genotype, all TCR-V β 13.1⁺ cases being HLA-DRB1*0701 (*P* = .004). Interestingly, the HLA-DR7/TCR-V β 13.1-restricted T-cell expansions displayed a highly homogeneous and strikingly similar TCR arising from the use of common TCR-V β gene segments, which shared (1) unique CDR3 structural features with a constantly short

length, (2) similar combinatorial gene rearrangements with frequent usage of the J β 1.1 gene, and (3) a homolog consensus protein sequence at recombination junctions. Overall, these findings strongly support the existence of a common antigendriven origin for monoclonal CD4⁺ T-LGL lymphocytosis, with the identification of the exact peptides presented to the expanded T cells deserving further investigations. (Blood. 2007;109:4890-4898)

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Introduction

Monoclonal chronic T-cell lymphocytosis and T-cell leukemias/ lymphomas are a heterogeneous group of disorders whose diagnosis and classification have been hampered by their relatively low frequency and variable clinical and histopathological behavior,¹⁻⁴ the lack of easily applicable clonality markers for T cells, and the substantial clinical overlap with nonmalignant inflammatory disorders.^{1,5,6} Although the pathogenetic mechanisms involved in the development of clonal T-cell disorders remain largely unknown, in recent years significant advances have been made in this regard.⁷⁻⁹ Among other observations, an association between chronic inflammatory and infectious processes and the occurrence of (mono)clonal expansions of lymphoid cells has recurrently been reported, particularly for chronic B-cell malignancies¹⁰ but also for mature T-cell neoplasias.^{1,8-9} Accordingly, different viruses (eg, human T-cell lymphotropic virus type I [HTLV-I], Epstein-Barr virus [EBV], and cytomegalovirus [CMV]) and bacterial superantigens (ie, staphylococci-derived superantigens) have been associated with the pathogenesis of specific mature T-cell malignancies, either because they infect tumor cells¹¹⁻¹⁴ or because they could induce an antigen-driven expansion of neoplastic T cells.9,15,16 In line with the latter hypothesis, recent reports suggest that T-cell receptor (TCR)associated signals could contribute to tumor development, particularly in T-cell large granular lymphocyte (T-LGL) leukemia.^{5,9} In

these cases, antigen-driven expansions of cytotoxic T lymphocyte (CTL) clones could precede the occurrence of oncogenic events leading to neoplastic transformation and/or dysregulation of growth/ apoptosis resulting in T-LGL leukemia9 with a restricted TCR- $V\beta/V\alpha$ usage.¹⁷ This notion is supported by the observation that TCR $\alpha\beta^+$ /CD8⁺ T-LGL leukemia often occurs in the context of specific autoimmune diseases9,16 and that in about one third of cases TCRαβ+/CD4+ T-LGL leukemia/lymphocytosis is associated with neoplasias other than the T-LGL and a preferential usage of the TCR-V β 13.1 family.¹⁸ In addition, the reactive versus neoplastic nature of some (mono)clonal expansions of T-LGL remains a matter of debate,¹⁵ particularly in cases where it is associated with viral infection, severe immune disturbances, or in the elderly.^{19,20} In this regard, the search for the potential involvement of common antigens in driving the development of monoclonal T-cell disorders through the analysis of complementary determining region 3 (CDR3) sequences of TCR genes has provided controversial findings. Accordingly, while CDR3 sequences from CD8⁺ T-LGL leukemia did not show any apparent structural homology,^{21,22} in nearly half of all TCR $\gamma\delta^+$ T-LGL neoplasias, clonal T cells express the same TCR-V δ /V γ family members (TCR-V γ 9/V δ 2) and share common TCR sequences, as reflected by the systematic presence of the antigen-selected invariant T

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Table 1. Clinical and laboratory characteristics of monoclonal TCR $\alpha\beta$ +/CD4+ LGL lymphocytosis according to the type of TCR-V β family
expressed (V eta 13.1 versus non-V eta 13.1) compared with monoclonal TCR $lphaeta^+$ /CD8 $^+$ LGL lymphocytosis

	Monoclona	I TCRαβ ⁺ /CD4 ⁺ LGL lyı	Monoclonal TCRαβ ⁺ /CD8 ⁺ LGL		
Characteristic	TCR-Vβ13.1	Non-TCR-V _β 13.1	Total cases	lymphocytosis	Р
No. of patients	15	21	36	24	
Mean age \pm 1 SD, y (range)	64 ± 8 (52-80)	61 ± 13 (36-81)	63 ± 12 (36-81)	56 ± 15 (31-79)	NS
% male/% female	25/75	61/39	47/53	37/63	NS
Reason for consulting, %					
Routine blood analysis	100	83	90	83	NS
Skin lesions	0	11	7	0	NS
Abdominal distension	0	6	3	0	NS
Fever	0	0	0	4	NS
General symptoms	0	0	0	13	NS
Physical examination, %					
Adenomegalies	11	6	7	0	NS
Hepatomegaly	0	0	0	4	NS
Splenomegaly	0	6	3	4	NS
Skin lesions	0	11	7	0	NS
Associated neoplasias, %	25	22	23	26	NS
Associated autoimmune diseases, %	18	0	13	33	.05
Laboratory parameters, %					
Leukocytosis, WBC count more than 10 $ imes$ 10 ⁹ /L	64	61	63	25	.01
Lymphocytosis, lymphocyte count more than 5 \times 10 ⁹ /L	70	67	68	42	.09
Neutropenia, neutrophil count less than 1.5 $ imes$ 10 ⁹ /L	14	0	3	61	< .001
Anemia, hemoglobin level less than 10 g/dL	0	0	0	29	.003
Thrombocytopenia, platelet count less than 100 $ imes$ 10 9 /L	10	0	3	8	NS
Increased lactic dehydrogenase level, more than 460 U/L	10	0	3	22	NS
Increased β 2-microglobulin level, more than 2 mg/dL	0	7	3	63	.001
Cases requiring treatment because of lymphocytosis or the					
associated autoimmune disease, %	0	0	0	37	.001
Outcome: stable disease, %	100	100	100	87	NS
Total deaths, %	10	18	15	8	NS
Deaths related to the TCR $lphaeta^+$ LGL lymphocytosis, %	0	0	0	4	NS
Mean follow-up \pm 1 SD, mo (range)	68 ± 48 (13-136)	74 ± 56 (10-233)	72 ± 53 (10-233)	44 ± 40 (1-152)	.03

P value corresponds to comparisons between monoclonal TCR $\alpha\beta^+$ /CD4⁺ LGL and TCR $\alpha\beta^+$ /CD8⁺ LGL lymphocytosis; NS indicates no statistically significant differences: *P* > .1. No statistically significant differences (*P* > .05) were found between TCR-Vβ13.1 and non-TCR-Vβ13.1 clonal CD4⁺ LGL lymphocytosis. A detailed description of the clinical characteristics of patients with monoclonal TCR $\alpha\beta^+$ /CD4⁺ LGL lymphocytosis is provided by Lima et al.¹⁸

nucleotide in the first codon of the V δ 2-J δ 1 junctional region from all patients.²³ Such apparent discrepancy could be related to the fact that antigen-driven TCR $\alpha\beta^+$ /CD8⁺ T-LGL leukemias would depend not only on the complementary sequences and specific binding of the TCR to the antigen, but also on the individual HLA haplotypes, while for TCR $\gamma\delta^+$ T cells this HLA restriction would not apply.

In the present study, we have analyzed a large series of 36 patients with monoclonal TCR $\alpha\beta^+$ /CD4⁺ T-LGL lymphocytosis grouped according to TCR-V β 13.1⁺ usage versus other TCR-V β families. Our aim was to explore the potential existence of an association in these patients between the expanded TCR-V β families, the CDR3 sequences of the TCR-V β gene, and the HLA genotype. Our results indicate that all patients with monoclonal expansions of TCR-V β 13.1⁺/CD4⁺ T cells display a common HLA-DRB0701⁺ genotype and express identical motifs in the CDR3-TCR-V β sequence, suggesting a common antigen-driven origin.

Patients, materials, and methods

Peripheral blood (PB) samples from human patients were obtained after informed consent was given by the patients (all of them more than 18 years old), in accordance with the local ethics committee of the University Hospital of Salamanca and the Declaration of Helsinki.

Patients and samples

A total of 161 T-LGL cases were referred to the Cytometry Service of the University Hospital of Salamanca (63 TCRαβ⁺/CD8⁺⁺/CD4⁻, 55 TCRαβ⁺/ CD4⁺/NKa⁺/CD8^{-/+dim}, 40 TCR $\gamma\delta^+$, and 3 TCR $\alpha\beta^+$ /CD8⁻/CD4⁻ cases) between September 1999 and March 2006. From them, 36 individuals with monoclonal TCR $\alpha\beta^+$ /CD4+/NKa+/CD8-/+dim lymphocytosis (19 males and 17 females; mean age, 63 \pm 12 years, ranging from 36 to 81 years) were selected and included in this study. In all these latter cases, PB samples were collected into tubes containing K3-EDTA, according to the local ethics committee's recommendations. The major clinical and laboratory features of this group of patients, according to the type of TCR-VB family expressed (TCR-VB13.1 versus non-TCR-VB13.1), are shown in Table 1 in comparison with those of a randomly selected series of 24 patients with monoclonal TCR $\alpha\beta^+$ /CD 8^+ /CD 4^- lymphocytosis. At the close of the study, the median follow-up of the patients with monoclonal TCRαβ+/CD4+/NKa+/CD8-/+dim lymphocytosis was 72 months (range, 10 to 233 months).

A total of 930 PB samples from unrelated healthy subjects were used as controls to establish the frequency of the different HLA haplotypes in the healthy population, while PB samples from 15 adult individuals (older than 50 years) were used as controls to establish the TCR-V β repertoire usage in TCR $\alpha\beta^+$ /CD4⁺ T cells.

Immunophenotypic studies

For the analysis of the TCR-V β repertoire of CD4⁺/CD8^{-/+dim} LGL T lymphocytes, a panel of 24 monoclonal antibodies (MAbs) directed against

an identical number of members of 21 different TCR-Vβ families (TCR-Vβ repertoire Kit; Immunotech, Marseille, France) was used in 4-color stainings. Further phenotypic characterization of CD4⁺/CD8^{-/+dim} LGL T cells was performed using the following 4-color combinations of MAbs: fluorescein isothiocyanate (FITC)/phycoerythrin (PE)/PE–cyanin 5 (PC5) or peridinin chlorophyll protein (PerCP)/allophycocyanin (APC): CD2/CD7/CD4/CD8, CD5/CD7/CD4/CD8, CD122/CD25/CD4/CD8, CD57/CD11c/CD4/CD8, CD16/CD56/CD4/CD8, CD122/CD25/CD4/CD8, CD45RA/CD45RO/CD4/CD8, CD62L/CD28/CD4/CD8, CD11a/HLA-DR/CD4/CD8, CD15/CD4/CD8, CD15/CD4/CD8, CD5/CD4/CD8, CD55/CD4/CD8, CD55/CD4/CD8, CD55/CD4/CD8, CD55/CD4, The source and specificity of each MAb reagent used has been previously described in detail.¹⁸

Cell staining was performed using a whole blood "stain-and-then-lyse" method (FACS lysing solution; Becton Dickinson Biosciences [BDB], San Jose, CA) and a direct immunofluorescence technique, as previously reported in detail.¹⁸ For the cytoplasmic staining, the Fix & Perm reagent kit (Invitrogen, Carlsbad, CA) was used according to the recommendations of the manufacturer.

Data acquisition was performed immediately after completion of sample preparation in a FACSCalibur flow cytometer (BDB) using the CellQUEST software program (BDB). The Paint-A-Gate Pro software program (BDB) was used for data analysis. In each case, the aberrant T-cell population was defined as CD4⁺/CD8^{-/+dim} and/or CD4⁺/CD56⁺ large granular—intermediate sideward light scatter (SSC^{intermediate})—events (Figure 1) for its further phenotypic characterization.

Preparation of DNA and HLA typing

High molecular weight DNA was prepared from 200 μ L PB using the QIAGEN bloodmicro kit (Qiagen, Hilden, Germany). HLA genotyping for HLA-ABC and both HLA-DRB1 and HLA-DQB1 was performed by sequence-specific oligonucleotide–polymerase chain reaction (SSPO-PCR) techniques using the Dynal Reli SSO kit (DYNAL Biotech, Bromborough, United Kingdom). Ambiguous results were resolved by sequence-based typing (SBT). DNA samples were amplified by PCR using the Big-Dye Terminator Cycle Sequencing Kit (Applied Biosystems, Foster City, CA). Sequencing was performed on an ABI 377 DNA sequencer (Applied Biosystems), and the data obtained were analyzed using the Match Tools v 1.0 Sequencing Analysis software program (Applied Biosystems). The ancestral haplotypes are putative because it was not possible to verify their segregation from family studies.

PCR amplification and nucleotide sequence analysis of TCR

High molecular weight DNA was prepared from freshly frozen PB samples using standard protocols including proteinase K treatment. In addition, total RNA was isolated from fluorescence-activated cell sorter (FACS)–sorted CD4⁺ and CD8⁺ T-cell populations (purity more than 95%) from pooled PB mononuclear cells (MNCs) of 5 non–HLA-DR*0701 adult healthy donors and from PB MNCs of 1 HLA-DR*0701 healthy individual and reverse transcribed into cDNA. TCR V β 13.1 gene family–specific PCR was performed using specific primers as previously described.²⁴ PCR products were cloned into pGEM-T easy vector (Promega, Madison WI), and single-colony PCR was performed on positive clones. Single-colony PCR products were directly sequenced.

DNA was amplified using a mixture of sense primers annealing to the TCR-V β 13 sequence in conjunction with a mixture of antisense primers complementary to the germ-line J regions as previously reported in detail.²⁴ In most samples, clonal products from the V β gene PCR were sequenced directly using the BigDye Terminator Cycle Sequencing Reaction Kit. In fact, the amplified sequences exhibited identical rearranged monoclonal TCR sequences, even in the VDJ junctional hypervariable regions, indicating that these expanded regions were clonal. To confirm the validity of the sequences obtained (thereby avoiding the possibility of either contamination or sequencing mistakes), a more detailed analysis of the sequences obtained was performed in some cases. For that purpose, PCR products were inserted into the PCR2.1-TOPO vector (Invitrogen, Barcelona, Spain),

which was followed by transformation into competent *Escherichia coli* cells; on average, 5 colonies were randomly selected for sequencing using the BigDye Terminator Cycle Sequencing Reaction Kit. A total of 98 V β 13.1 clones (69 from the non-HLA-DR*0701 donors and 29 from the HLA-DR*0701 donor) in the CD4⁺ T-cell fraction and 53 V β 13.1 clones (38 from the non-HLA-DR*0701 donors and 15 from the HLA-DR*0701 donor) in the CD8⁺ T-cell fraction were sequenced and analyzed. All sequence reactions were analyzed using an automated DNA sequencer (ABI 377; Applied Biosystems).

Database searches

Sequences obtained were aligned using the Basic Local Alignment Search Tool (BLAST) (National Center for Biotechnology Information, Bethesda, MD) and ImMunoGeneTics (IMGT) databases.

Statistical methods

For all clinical and laboratory parameters included in Table 1, mean, standard deviation, and range were calculated using the SPSS program (SPSS 12.0, Chicago, IL). To establish the statistical significance of the differences observed between groups, either the Pearson χ^2 test or Fisher exact test was used for categorical variables, and the Mann-Whitney *U* nonparametric test was used (SPSS 12.0) for continuous variables. *P* values below .05 were considered to be associated with statistical significance.

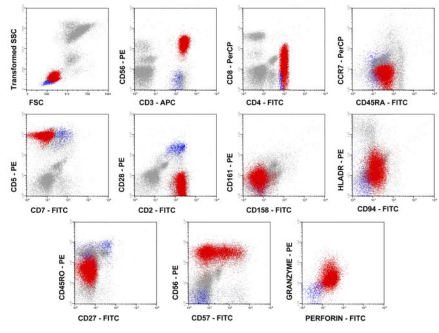
Results

Immunophenotype of the expanded CD4⁺ LGL T cells

In all cases studied, expanded CD4⁺ LGL T cells showed relatively high SSC features as compared with normal PB CD4⁺ T lymphocytes and common phenotypic characteristics consisting of TCR $\alpha\beta^+$ /CD4⁺/CD8^{-/+dim} cells with a typical cytotoxic (granzyme B⁺, CD56⁺, CD57⁺, CD11b^{+/-}) activated, memory/effector T-cell phenotype (CD2^{+bright}, CD7^{-/+dim}, CD11a^{+bright}, CD28⁻, CD62L⁻, HLA-DR⁺) (Figure 1). In about half (47%) of the cases, clonal T cells coexpressed CD45RA and CD45RO, while in the other cases they had a CD45RA⁺/CD45RO⁻ phenotype. Other NKa markers (CD11c, CD16, CD94, CD158a, CD161, and NKB1) and T-cell activation-related antigens (CD25, CD38, and CD122) were absent on the CD4⁺ LGL T cells. Overall, these cells represented 47% ± 23% of all PB lymphocytes, with a mean (±1 SD) absolute number of 6.6 × 10⁹ ± 3.3 × 10⁹ PB TCR $\alpha\beta^+$ / CD4⁺/NKa⁺/CD8^{-/+dim} T lymphocytes per liter.

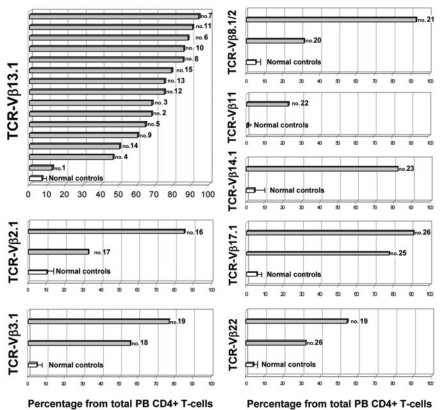
Flow cytometric analysis of the TCR-VB repertoire of CD4+/ CD8^{-/+dim} LGL T cells was consistent with a (mono)clonal expansion in all cases studied, which accounted for $72\% \pm 21\%$ of all PB CD4⁺ T cells. In 27 cases the expanded TCR-VB family was identified with the panel of TCR-VB reagents used, corresponding to TCR-VB13.1 in 15 cases (42%), TCR-VB2.1 in 2 (5.6%), TCR-Vβ3.1 in 2 (5.6%), TCR-Vβ8.1 + Vβ8.2 in 2 (5.6%), TCR-VB17.1 in 2 (5.6%), TCR-VB22 in 2 (5.6%), and TCR-VB11 or TCR-V β 14.1 in 1 case each (2.8%). In the remaining 9 patients, the expanded TCR-V β family was not identified (25%) with the panel of MAbs used. Figure 2 shows the nominal size of the TCR-V β expansion present in each patient (as percentage of the total PB CD4⁺ T cells) in comparison with the size of the corresponding TCR-VB family observed in a cohort of agematched healthy subjects; the proportion of TCR-V β 13.1⁺ cells represented 6.7% \pm 2.5% of total PB CD4⁺ T cells from healthy subjects, while in the patient group it represented between 12.5% and 94.1% (median, 75%).

Figure 1. Immunophenotypic features of monoclonal TCR $\alpha\beta^+$ /CD4+/NKa+/CD8-/+d T-LGL. Representative dot plots illustrate the phenotypic patterns shown by monoclonal TCR $\alpha\beta^+$ /CD4+/NKa+/CD8-/+d T-LGL. Red dots correspond to monoclonal TCR $\alpha\beta^+$ /CD4+/NKa+/ CD8^{-/+d} T-LGL, blue dots correspond to normal residual non-LGL CD4+ T cells, while grav dots correspond to PB leukocytes other than CD4+ T cells.



Association between the HLA haplotypes and the TCR-VB repertoire of clonal TCRαβ+/CD4+/NKa+/CD8-/+dim **T-LGL lymphocytosis**

A significant association (P = .004) was found between the TCR-VB repertoire and the HLA genotype of the studied cases (Table 2). Accordingly, all 15 patients who showed expansion of TCR-Vβ13.1⁺ CD4⁺ T cells were HLA-DRB*0701⁺ (Figure 3). In turn, the frequency of the HLA-Cw*0401 allele was slightly higher among those patients in whom the expanded TCR-V β family was not contained in the panel of MAb reagents used than among both TCR-VB13.1⁺ patients and cases expressing a known TCR-VB other than 13.1 (67% versus 33%; P = .1). In addition, the frequency of cases with a HLA-DRB1*03 and HLA-DRB1*04 genotype was lower among CD4+ T-LGL patients than in the control group (14% versus 29% and 6% versus 22%, respectively; $P \le .04$).²⁵ Interestingly, 4 of the 15 HLA-DRB*0701 cases included the 44.2 ancestral haplotype (HLA-A*2902, B*4403, C*1601, DRB*0701, and DQB1*0202) (cases 7, 9, 11, and 14)



Percentage from total PB CD4+ T-cells

Figure 2. Illustrative representation of the size of the actual identifiable TCR-VB expansion present in each patient (as percentage of total PB CD4+ T cells) in comparison with the size of the corresponding TCR-V β family observed in a cohort of age-matched healthy subjects (n = 15). Gray bars correspond to patients, each one identified by the corresponding case number, while white bars and horizontal lines correspond to the mean value and 1 SD found in healthy controls, respectively.

Case no.	Expanded TCR-Vβ family	HLA-DRB	HLA-DQB	HLA-A	HLA-B	HLA-C
1	13.1	0701/1103	0301/0303	2301/3201	3501/5002	0401/0401
2	13.1	0701/1404	0202/0503	0101/2601	1401/5701	0602/0802
3	13.1	0701/0102	0303/0501	0201/0201	5101/5701	0102/0701
4	13.1	0701/1401	0303/0503	0201/2402	4402/5701	0501/0701
5	13.1	0701/0101	0202/0501	0201/0201	3501/4901	0401/0701
6	13.1	0701/0101	0202/0501	0301/0301	1302/3501	0401/0602
7	13.1	<u>0701</u> /0401	<u>0202</u> /0301	2601/2902	4402/4403	0501/ <u>1601</u>
8	13.1	0701/0701	0202/0202	0101/2402	1801/5001	0501/0602
9	13.1	<u>0701</u> /0101	<u>0202</u> /0501	0101/2902	1401/ <u>4403</u>	0802/ <u>1601</u>
10	13.1	<u>0701</u> /1501	<u>0303</u> /0602	0201/0201	0702/ <u>5701</u>	0602/0702
11	13.1	<u>0701</u> /0301	0201/0202	0301/2902	1801/ <u>4403</u>	0501/ <u>1601</u>
12	13.1	<u>0701</u> /0301	0201/0202	0101/2301	0801/ <u>4403</u>	0401/0701
13	13.1	0701/0301	0201/0202	2301/3002	1801/5801	0501/0701
14	13.1	<u>0701</u> /0101	<u>0201</u> /0501	1101/2902	3501/ <u>4403</u>	0401/ <u>1601</u>
15	13.1	0701/1501	0202/0602	2902/3002	0702/4101	0701/1701
16	2.1	0301/1317	0201/0603	2402/2402	0801/3508	0401/0701
17	2.1	0701/1301	0202/0609	3002/6801	4403/5101	0701/1402
18	3.1	0403/1501	0302/0602	0301/3202	0702/3501	0401/0702
19	3.1	1501/1602	0502/0602	0201/0201	3701/4402	0501/0602
20	8.1 + 8.2	1101/1101	0301/0301	0201/6901	1801/3508	0701/1203
21	8.1 + 8.2	0301/1501	0201/0602	0201/0201	1518/3501	0401/0704
22	11	1101/1601	0301/0502	0101/0201	3701/4002	0602/1204
23	14.1	0701/0701	0202/0303	0103/2902	4403/5701	0701/1601
24	17.1	0802/1101	0301/0402	0102/2301	1401/3501	0701/0801
25	17.1	0701/1401	0202/0503	0201/2902	4403/5101	1502/1601
26	22	0102/0801	0402/0501	2402/3301	1402/3503	0401/0802
27	22	1103/1301	0301/0603	0201/3201	1509/4002	0202/0704
28	NI	0701/1104	0201/0301	1101/2902	4001/4403	0304/1601
29	NI	0701/1101	0201/0301	0201/0301	3503/5001	0401/0501
30	NI	0701/1301	0202/0604	3201/6801	1401/5301	0401/0802
31	NI	0701/1602	0202/0502	0301/2402	0702/3801	0702/1203
32	NI	1101/1401	0301/0503	0201/2603	4002/3501	0401/0501
33	NI	0101/1301	0501/0604	1101/1101	4004/5601	0102/0202
34	NI	0101/1101	0301/0501	0201/2402	1801/3501	0401/1203
35	NI	0101/1301	0501/0604	1101/3101	3501/5101	0401/0501
36	NI	1302/1305	0301/0604	0201/0205	3508/4101	0401/0701

Extended/ancestral haplotypes are underlined.

NI indicates TCR-V β family not identified by immunophenotyping.

(Table 2). Another 2 cases (cases 2 and 10) were related to the 57.1 ancestral haplotype (HLA-A*01, B*5701, C*0602, DRB1*0701, and DQB1*0303). The ancestral haplotypes 35.2 (HLA-A*1101, B*3501, C*0401, DRB*0101, DQB1*0501) and 35.1 (HLA-C*0401, B*3501, DRB1*11, DQB1*0301) were also found in 3

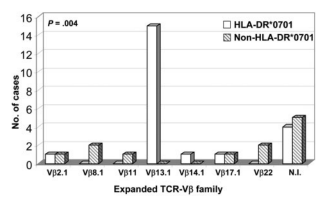


Figure 3. Frequency of the HLA-DRB1*0701 genotype in patients with monoclonal TCR $\alpha\beta^+$ /CD4⁺/NKa⁺/CD8^{-/+d} T-LGL lymphocytosis grouped according to the expanded TCR-V β family. NI indicates that the exact TCR-V β family expanded was not identified with the panel of anti–TCR-V β MAbs used.

(cases 5, 6, and 14) and 1 patients (case 1), respectively; however, among these patients, only case 14 showed a complete 35.2 haplotype, with a 4-locus haplotype being detected in the other 3 cases. Cases 8, 13, and 15 were not considered to be related with ancestral haplotypes, although they showed haplotypes that are frequently present in the Spanish population.²⁶

Expanded clonal CD4⁺/CD8^{-/+dim} T cells from HLA-DRB1*0701 patients exhibit conserved TCR β chain motifs

Molecular analysis of the length of CDR3 of CD4⁺/CD8^{-/+dim} T cells from those cases expressing TCR-V β 13.1 showed a pattern consistent with monoclonality based on both TCR-V β usage and CDR3 length (Table 3). Further comparison of CDR3 size distribution in clonal CD4⁺/CD8^{-/+dim} T cells from the same patients showed a highly restricted usage of V_βD_βJ_β gene segments and shared CDR3 configurations. Accordingly, 11 of 13 patients used the same J1.1 and V segments, and they had highly similar CDR3 configurations (Table 3). Although 3 different J segments were used in the clonal expansions derived from CD4⁺/CD8^{-/+dim} T lymphocytes, similar VDJ junctional region sequences were found (Table 4). Remarkably, in 5 of the 11 V β 13.1-D β 1-J β 1.1 cases, the combinatorial process involved the deletion of 1 to 3 nucleotides

Case no.	V-J usage	CDR2	3' end of V β	CDR3: N-D _β 1-N	5' end of J β
1	Vβ13S1-J1.1	SVGAGI	CASS	K <u>QG</u> V	TEAF <u>FG</u>
2	Vβ13S1-J1.1	SVGAGI	CASS	K <u>QG</u> A	TEAF <u>FG</u>
3	Vβ13S1-J1.1	SVGAGI	CAS	RKQGA	TEAF <u>FG</u>
4	Vβ13S1-J1.1	SVGAGI	CASS	Y <u>QG</u> A	TEAF <u>FG</u>
5	Vβ13S1-J1.1	SVGAGI	CASS	S <u>QG</u> T	TEAF <u>FG</u>
6	Vβ13S1-J1.1	SVGAGI	CAS	RH <u>QG</u> S	TEAF <u>FG</u>
7	Vβ13S1-J1.2	SVGAGI	CASS	H <u>QG</u> A	NGYT <u>FG</u>
8	Vβ13S1-J1.5	SVGAGI	CASS	Y <u>QG</u> A	QPQH <u>FG</u>
9	Vβ13S1-J1.5	SVGAGI	CASS	Y <u>QG</u> S	QPQH <u>FG</u>
10	Vβ13S1-J1.1	SVGAGI	CAS	RR <u>QG</u> Y	TEAF <u>FG</u>
12	Vβ13S1-J1.1	SVGAGI	CASS	Y <u>QGA</u>	TEAF <u>FG</u>
13	Vβ13S1-J1.1	SVGAGI	CAS	RRQGA	TEAF <u>FG</u>
14	Vβ13S1-J1.1	SVGAGI	CAS	NLQGS	TEAF <u>FG</u>
15	Vβ13S1-J1.1	SVGAGI	CASS	Y <u>QGSA</u>	EAF <u>FG</u>
10*	Vβ13S1-J1.1	ND	CASS	Y	NTEAF <u>FG</u>
52*	Vβ13S1-J1.1	ND	CASS	W <u>QG</u> VD	TEAF <u>FG</u>
29*†	Vβ13S1-J1.1	ND	CAS	NRGLY	TEAF <u>FG</u>

Comparison of the CDR2 and CDR3 of clonal TCR V β 13S1 gene rearrangements from 14 CD4⁺ T-LGL cases (this analysis could not be performed in case 11 due to sample shortage). Segments are aligned according to the conserved motifs (CASS for the V β and FG for the J β segment). Nucleotide sequences were aligned to TCR sequences according to the Basic Local Alignment Search Tool (BLAST) and ImMunoGeneTics (IMGT) databases.

ND indicates not determined. Underlining indicates the motifs shared in common.

*VDJ protein sequence of the 3 of 98 clones using the J β 1.1 segment found among PB V β 13.1+ CD4+ T cells from adult healthy donors.

†This clone corresponds to an HLA-DR*0701 donor.

from the 5' end of V β 13.1 gene and the insertion of a variable number of nontemplate nucleotides (Table 4). Accordingly, in all cases analyzed (n = 14) the mean length of CDR3 was considerably shortened (4 or 5 codons), the distance between the CASS and FG motifs constantly being of 9 codons. Interestingly, shared motifs consisting of at least 2 identical amino acids were found within the VDJ junctional regions of the expanded CD4⁺ T cells derived from different patients, a consensus XQGX motif being shared by all cases (Table 3). For 7 cases the GGG codon yielded a glycine; in contrast, in 2 other cases the glycine was generated by the GGT and in another case by a GGA codon. In turn, glutamine (Q) was yielded in all cases by the CAG codon. One of the motifs (YQGA) was identical among the clonally expanded CD4+/CD8-/+dim T cells derived from 3 patients, and the XQGA motif was detected in 7 individuals. In contrast, only 2 (2.9%), 8 (11.6%), and 2 (2.9%) of 69 clones of PB CD4⁺ T cells from non-HLA-DR*0701 healthy adults were found to use

the J β 1.1, J β 1.2, and J β 1.5 gene segments, respectively; similarly, from the 29 clones of CD4⁺ T cells sequenced from the HLA-DR*0701 adult healthy donor, only 1 (3.4%) and 3 (10%) of them were found to use J β 1.1 and J β 1.2 gene segments, respectively. Interestingly, of these 16 clones of CD4⁺ T cells, only 1 of those 3 clones using the J β 1.1 gene segment was highly similar in the CDR3 configuration to those detected in the patients analyzed (Tables 3-4); this clone was sequenced from the pooled non-HLA-DR*0701 MNCs. In addition, the XQGX configuration could not be detected in other clones of PB CD8⁺ T cells from healthy adults using the J β 1.1, J β 1.2, and J β 1.5 gene segments (0 of 9 clones from the 38 CD8⁺ T-cell clones sequenced from non-HLA-DR*0701 donors and 0 of 3 from the 15 CD8+ T-cell clones sequenced from the HLA-DR*0701 donor). Finally, we searched GenBank for VDJ rearrangements with similar XQGX amino acid sequences, but no TCR close matches were found.

Case no.	V-J usage	3' end of V β 13.1	Ν	Dβ1	Ν	5' end of J eta
1	Vβ13S1-J1.1	AGCAGT	AA	gggACAGGGggc	AGT	aaCACTGAAGCTTTC
2	Vβ13S1-J1.1	AGCAGT	AA	gggACAGGGGGC	_	aaCACTGAAGCTTTC
3	Vβ13S1-J1.1	AGCAG	AAA	gggACAGGGGGC	_	aaCACTGAAGCTTTC
4	Vβ13S1-J1.1	AGCAGT	TAC	gggaCAGGGGGc	CG	aacACTGAAGCTTTC
5	Vβ13S1-J1.1	AGCAGT	TCC	gggaCAGGGGgc	AC	aaCACTGAAGCTTTC
6	Vβ13S1-J1.1	AGCAG	ACAT	gggaCAGGGggc	TAG	aaCACTGAAGCTTTC
10	Vβ13S1-J1.1	AGC	CGGC	ggGACAGGGGgc	AAA	aacACTGAAGCTTTC
12	Vβ13S1-J1.1	AGCAGT	TAT	gggaCAGGGGGC	_	aCACTGAAGCTTTC
13	Vβ13S1-J1.1	AGCAG	GC	ggGACAGGGGGC	_	aacACTGAAGCTTTC
14	Vβ13S1-J1.1	AGCA	ATCT	gggACAGGGggc	TAG	aaCACTGAAGCTTTC
15	Vβ13S1-J1.1	AGCAGT	—	TACCAAGGCTCGG	_	aacaCTGAAGCTTTC
10*	Vβ13S1-J1.1	AGCAGTTAC	_	_	_	AACACTGAAGCTTC
52*	Vβ13S1-J1.1	AGCAGTT	GG	gggaCAGGGGGc	TGG	aACACTGAAGCTTTC
29*†	Vβ13S1-J1.1	AGCA	_	gggACAGGGGGc	TTGT	aACACTGAAGCTTTC

- indicates not applicable.

*Sequence of the 3 clones using the J β 1.1 segment found among PB V β 13.1⁺ CD4⁺ T cells from adult healthy donors.

†This clone corresponds to an HLA-DR*0701 donor.

Discussion

Monoclonal TCRαβ+/CD4+/NKa+/CD8-/+d T-LGL lymphocytosis is a subgroup of monoclonal LGL lymphoproliferative disorders, different from both the CD8⁺ TCR $\alpha\beta^+$ T-LGL, TCR $\gamma\delta^+$ T-LGL, and natural killer (NK) cell-type LGL leukemias.18 Noteworthy, in the present study, the former subgroup of clonal T-LGL lymphocytosis was found at a higher frequency than both TCRγδ- and NK-LGL leukemias, whereas it was slightly less common than TCR $\alpha\beta^+$ CD8⁺ T-LGL. In contrast to TCR $\alpha\beta^+$ CD8+ T-LGL, monoclonal TCRaB+/CD4+/NKa+/CD8-/+d T-LGL cases have been only sporadically reported in the literature, while they were relatively frequent in our series. According to the present study, such discrepancy might be related to the fact that TCRaβ+/CD4+/NKa+/CD8-/+d T-LGL cases usually display a more indolent clinical course-although rare cases associated with aggressive disease have also been reported in the literatureassociated with a significantly lower frequency of neutropenia, anemia, and other associated autoimmune diseases, in addition to a lower percentage of cases requiring treatment, in comparison with TCR $\alpha\beta^+$ /CD8⁺ T-LGL lymphocytosis. However, the apparently high frequency of CD4 LGL cases found in our series could also be due to the fact that we actively searched for these cases. Recently, we showed that in patients with monoclonal TCR $\alpha\beta^+$ /CD4⁺/NKa⁺/ CD8^{-/+d} T-LGL lymphocytosis the expanded clonal T cells display a restricted usage of a limited number of TCR-V β families,¹⁸ from which TCR-VB13.1 was particularly overrepresented in comparison with its frequency in the PB counterpart of these cells from healthy individuals.²⁷ These observations suggest the potential involvement of a common antigen in driving the expansion of clonal T cells in these patients. In such a situation, shared HLA haplotypes, as well as common motifs in the CDR3 sequences of the TCR-VB genes, could be expected. Upon comparing TCR-VB13.1⁺ cases with all non-TCR-VB13.1 individuals, a clear association was found between the expanded TCR-VB family and the HLA genotype, all TCR-VB13.1⁺ cases displaying an HLA-DRB1*0701 allele. The random chance that both events coincide is about 2%, versus 42% in our patients. In line with these observations, it has recently been reported²⁸ that most CD4⁺ T cells from an HIV-1⁺/CMV⁺-infected patient with lytic granules containing cytotoxic proteins (such as granzymes and perforin) displayed a clear HLA class II- and not class I-restricted lytic activity. Accordingly, after specifically blocking of HLA class II, CMVspecific CD4⁺ LGL T cells from this patient resulted completely inhibited in their in vitro ability to produce cytokines. In addition to the strong association between the expanded TCR-V β and HLA class II, all (unrelated) HLA-DRB1*0701⁺ patients showing TCR-VB13.1 expansions had a common CDR3 amino acid motif (XQGX) in the expanded T lymphocytes. Interestingly, this common "XQGX" CDR3 amino acid motif could not be found among the TCR-VDJB sequences of T lymphocytes from healthy individuals deposited in GenBank, and it was detected only at very low frequencies among the few clones using the J β 1.1, J β 1.2, and J β 1.5 gene segments identified in both purified CD4⁺ (1 of 16 clones) and CD8⁺ (0 of 12 clones) PB T cells from healthy adults. In addition, in a normal T-cell repertoire, different T cells have distinct CDR3 lengths that result in a gaussian distribution, while in our series virtually all expanded monoclonal CD4⁺ T-LGL cases expressing TCR-VB13.1 showed the presence of TCRB chains characterized by a unique CDR3 length. Altogether, the association between monoclonal expansions of TCR-VB13.1 T-LGL, the

HLA-DRB1*0701 genotype, and a common XQGX motif in the CDR3 sequence strongly suggests that monoclonal TCR $\alpha\beta^+$ /CD4+/ NKa⁺/CD8^{-/+d} T-LGL from these unrelated patients has been selected by a specific common antigen and that they could be the result of a chronic, long-term, antigen-driven process, as previously reported for TCRy8 LGL leukemias23 and B-cell chronic lymphocytic leukemias using the VH3-21 gene, based on their CDR3 homology.²⁹ Because the expanded CD4+/CD8-/+d T-LGL clones expressed TCR-VB13.1 with restricted antigen-binding sites in the context of HLA-DR*0701, it could be suggested that they result from an exogenous peptide-driven T-cell stimulation. Furthermore, if this selection involves antigen binding and triggering through the TCR, the antigenic epitope would most likely be restricted in its nature and structure,30 although some differences in the amino acid sequences of the CDR3 region were noted. The overlapping phenotypes of the expanded cells between different (unrelated) patients would further reinforce an underlying common pathogenesis. As previously reported,¹⁸ monoclonal TCRαβ⁺/CD4⁺/NKa⁺/ CD8^{-/+d} T-LGL cases show a remarkably uniform cytotoxic T-cell phenotype, as reflected by a common pattern of expression of NKa surface markers and cytotoxic proteins (CD56⁺, CD57⁺, Cy granzyme B⁺) in the absence of expression of other (CD16⁻, CD94⁻, CD158a⁻, CD161⁻, NKB1⁻) NK-associated receptors.

Recent reports provide strong accumulating evidence for a role of chronic antigen stimulation in clonal selection and progression of B-cell lymphomas10 as well as T-LGL leukemias.9,22,23 Although identical TCR gene rearrangement are typically identified in LGL leukemia, indicating a (mono)clonal proliferative disease, demonstration of monoclonality does not necessarily imply either neoplastic or malignant transformation.9,23 In fact, in the present study we were unable to demonstrate the presence of any genetic alteration in the patients studied, either by conventional karyotyping or by fluorescence in situ hybridization (FISH) (data not shown). Accordingly, the most probable pathogenetic mechanism leading to an increased survival and/or proliferation of specific T-cell clones in CD4⁺ T-LGL patients could be more probably related to chronic antigenic stimulation than to a cytogenetic-associated neoplastic transformation. TCR $\alpha\beta^+$ /CD4⁺/NKa⁺/CD8^{-/+d} T cells have been found in increased proportions in humans in different disease conditions where chronic antigen stimulation may occur, such as neoplasias, chronic viral infections, autoimmune disorders, and allografts.^{28,31-34} Unfortunately, no study has been reported in which CDR3 sequences of the expanded cells have been analyzed in such disease conditions; an exception would be graft versus host disease (GVHD) after allogeneic hematopoietic stem cell transplantation (allo-HSCT), where the expanded CTL clones (including both CD4⁺ and CD8⁺ T cells) have been clonotyped.^{35,36} Accordingly, a variable but frequently high degree of CDR3 homology within a given V β family has been reported in patients undergoing allo-HSCT³⁵ whereby the extent of the alteration of the T-cell repertoire is significantly higher in PBMCs from patients with acute GVHD than it is in cases without GHVD.³⁶ Based on these results, it has been hypothesized that such abnormalities could reflect multiple antigen-driven T-cell clonal expansions against alloantigens. Altogether, the evidence of oligoclonal expansions of $TCR\alpha\beta^+/CD4^+/NKa^+/CD8^{-/+d}$ T cells in several pathological conditions, interpreted as a specific T-cell response against tumor cells, virus, and autoantigens or alloantigens, clearly suggests that clonal TCRαβ+/CD4+/NKa+/CD8-/+d T-LGL lymphocytosis represents a dysregulated reaction to exogenous antigens. As a result, a wide and complex spectrum consisting of different clinical entities (from transient immune reaction to LGL leukemia) could be expected, similar to that described for clonal TCRαβ⁺/CD8⁺⁺ T-LGL lymphocytosis.⁹ Although the exact identity of such antigen(s) remains unknown, based on our results we may conclude that monoclonal TCRαβ⁺/CD4⁺/NKa⁺/CD8^{-/+d} T-LGL lymphocytosis cases are not random, because they do not reflect the expected Vβ-J physiological frequencies. In addition, the diverse geographic origin of our patients would suggest that the potential antigen involved in these processes is widely distributed. We can also rule out the involvement of a superantigen, due to the clear major histocompatibility complex (MHC)–TCRVβ–restricted association here observed.³⁷ Finally, the HLA-II restriction found for these clonal expansions of CD4⁺ T cells supports the involvement of a peptide with an exogenous origin leading to a repetitive and chronic engagement of the TCR of the expanded CD4⁺ T-LGL.

Another interesting observation is that monoclonal expansions of CD4⁺ T-LGL have only rarely been reported in the literature¹⁸ despite the fact that HLA-DRB1*0701 is frequently observed in the Caucasian population (about 30%).²⁵ These observations further support the role of factors other than the HLA genotype in leading to the dysregulation of the immune response and clonal expansion of CD4⁺ T-LGL. In this sense, the presence of common extended haplotypes among the TCR-V β 13.1⁺ patients suggests that a genetic influence cannot be ruled out. In particular, polymorphisms in genes within the MHC (ie, MICA, cytokines) should be considered with regard to dysregulation of CD4⁺ cytotoxic T cells.

In summary, in the present study we show that patients with monoclonal expansions of TCR-V β 13.1⁺/CD4⁺ T-LGL display a common HLA-DRB1*0701 genotype and express identical motifs in a constantly shorter-length CDR3-TCR-V β sequence, supporting a common antigen-driven origin for these T-cell disorders. Further identification of the short peptides bound to HLA molecules preferentially expressed by clonal TCR $\alpha\beta^+$ /CD4⁺ T-LGL would provide new insight into the pathogenesis of the disease; at the same time it could facilitate the identification and establishment of novel preventive and/or therapeutic strategies in individuals with monoclonal CD4⁺ T-LGL lymphocytosis at risk for transformation.

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Authorship

Contribution: P.G., Y.S., M.L., A.B., M.G., M.A.L.-N., A.W.L., and A.C.G.-M. performed research and analyzed data; F.R.-C. designed research, analyzed data, and wrote the paper; P.B. performed research and collected and analyzed data; J.C. performed research; and J.A. and A.O. designed research, collected data, analyzed data, and wrote the paper.

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