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Re-assignment of the murine 3' TRDD1 recombination signal sequence

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Abstract

T cell receptor genes are assembled in developing T lymphocytes from discrete *V*, *D* and *J* genes by a site-specific somatic rearrangement mechanism. A flanking recombination signal, composed of a conserved heptamer and a semi-conserved nonamer separated by 12 or 23 variable nucleotides, targets the activity of the rearrangement machinery to the adjoining *V*, *D* and *J* genes. Following rearrangement of *V*, *D* or *J* genes, their respective recombination signals are ligated together. Although these signal joints are allegedly invariant, created by the head-to-head abuttal of the heptamers, some do exhibit junctional diversity. Recombination signals were initially identified by comparison and alignment of germ-line sequences with the sequence of rearranged genes. However, their overall low level of sequence conservation makes their characterization solely from sequence data difficult. Recently, computational analysis unravelled correlations between nucleotides at several positions scattered within the spacer and recombination activity, so that it is now possible to identify putative recombination signals and determine and predict their recombination efficiency. In this paper, we analyzed the variability introduced in signal joints generated after rearrangement of the *TRDD1* and *TRDD2* genes in murine thymocytes. The recurrent presence of identical nucleotides inserted in these signal joints led us to reconsider the location and sequence of the *TRDD1* recombination signal. By combining molecular characterization and computational analysis, we show that the functional *TRDD1* recombination signal is shifted inside the putative coding sequence of the *TRDD1* gene, and consequently that this gene is shorter than indicated in the databases.

Keywords:

T cell receptor gene; V(D)J recombination; recombination signal; signal joint; junctional diversity

Introduction

T cell receptor (TR) and immunoglobulin (Ig) *V*, *D* and *J* genes need to be rearranged by a site-specific recombinase before they are expressed in T and B lymphocytes, respectively. This recombinase complex is composed of two lymphoid specific proteins, the products of the recombination activating genes (RAG)-1 and -2, and ubiquitous DNA repair factors involved in DNA repair by non-homologous end joining (Jung and Alt 2004). The recombination activity is specifically targeted to TR and Ig *V*, *D* and *J* genes by the presence at their borders of a short DNA motif called recombination signal (RS) (Tonegawa 1983). After synapsis of the genes to be rearranged (Hiom and Gellert 1998), the introduction of a double strand break by the RAG proteins exactly at the border of the genes and their RS generates hairpin sealed coding ends and blunt, phosphorylated signal ends (McBlane et al. 1995). Processing of the coding ends requires opening of the hairpin structure, which can generate P nucleotides, and includes the eventual addition of non-templated N nucleotides by the terminal nucleotidyl transferase (TdT) and/or the removal of a few nucleotides from the coding ends before they are ligated together to create a novel exon (Jung and Alt 2004). The result is a highly diverse repertoire of coding joints. In sharp contrast, it is generally held that signal ends are ligated together without any processing, so that signal joints result from a perfect head to head juxtaposition of the RSs.

RSs were first identified by sequence alignment of Ig genes (Tonegawa 1983), and later functionally characterized in recombination assays using artificial extrachromosomal substrates (Akira et al. 1987; Hesse et al. 1989). They consist of a conserved heptamer (CACAGTG) separated from a semi conserved nonamer (ACAAAAAACC) by 12 or 23 bases. The first bases of the heptamer are conserved in almost all the human and murine functional RSs. The A/T rich nonamer on the other hand is less well conserved, although it acts as the docking site for RAG-1, which then recruits RAG-2 onto the RS (Difilippantonio et al. 1996).

Analysis of the spacer sequence was more complex because both 12 and 23 spacers are highly heterogeneous (Cowell et al. 2004; Glusman et al. 2001). However, the recent development of new computational tools unravelled correlations between nucleotides at several positions within the spacer, and correlations between the identity of nucleotides at key positions and the overall recombination efficiency of the RS (Cowell et al. 2002; Cowell et al. 2003). This mutual information model allows the calculation, for each RS, of a “RS information content” (RIC) score that accurately reflects and efficiently predicts the signal’s recombination activity (Cowell et al. 2004).

In spite of the generally accepted models for V(D)J recombination, a large body of evidence indicates that signal ends can be processed and modified before they are ligated to form a signal joint (Candéias et al. 1996; Kanari et al. 1998; Touvrey et al. 2006). Recently, we undertook the analysis of the junctional diversity in signal joints created when the *TRDD1* and *TRDD2* genes are rearranged together in murine thymocytes. Three quarters of the signal joints were found to exhibit variability created by the insertion of N nucleotides in between the RSs’ heptamers. The recurrence of a di-nucleotide in these modified signal joints prompted us to re-consider the location of the break point at the 3’ end of the *TRDD1* gene. Our results identified a new functional RS23 3’ of the murine *TRDD1* gene.

Materials and methods

Mice-Thymocyte DNA preparation

CD3 $\varepsilon^{\Delta 5/\Delta 5}$ mice onto a BALB/c background (Gallagher et al. 1998) were bred in the animal facility of the Commissariat à l'Energie Atomique-Grenoble, France. They were sacrificed by CO₂ inhalation and their thymus removed at 6-8 wks of age. The thymi were crushed, and thymocyte DNA was prepared from single cell suspensions with the Nucleospin Tissue Kit (Macherey-Nagel, Hoerdt, France) according to the manufacturer's instructions.

Signal Joint amplification, analysis and cloning

TR DD1/DD2 signal joints were amplified with the 3'DD1 and 5'DD2 oligonucleotide primers (Table 1) from total thymocyte DNA with the AmpliTaq Gold system (Applied Biosystems, Courtaboeuf, France) in 37 cycles consisting of 30 sec at 94°C, 30 sec at 58°C and 30 sec at 72°C followed by an incubation of 10 min at 72°C. Before the first cycle, the enzyme was activated at 94°C for 6 min. To analyze junctional diversity, one half of the product was incubated for 3 hrs with or without ApaL1 (10 U, New England Biolabs, St Quentin en Yvelines, France). Digests were then separated on an agarose gel, transferred onto a nylon membrane and hybridized with 3'DD1p (Table 1), a radiolabelled oligonucleotide specific for the signal end moiety of the amplicon. Analysis was performed on a Personnal Molecular Imager FX with the Quantity One software (Bio-Rad, Marnes la Coquette). For sequencing, after purification on an agarose gel, the PCR products were cloned into the PGEM-T Easy vector (Promega, Charbonnières, France) according to the manufacturer's instructions and transformed into competent bacteria. Following plating, signal joint-containing colonies were identified by hybridization with the 5'DD2p radio-labelled probe. Plasmid DNA was prepared from positive colonies, and digested with the restriction enzyme ApaL1 to discriminate between ApaL1-sensitive signal joints, resulting from the perfect

ligation of the TRDD1 and TRDD2 RSs (ApaL1-S) and those in which one or both of the RS ends have been processed/modified, resulting in ApaL1-resistant (ApaL1-R) signal joints, as described (Candéias et al. 1996; Touvrey et al. 2006). ApaL1-R signal joints were then sequenced with the 5'DD2p (Table 1) primer using BigDye Terminator reagent (Applied Biosystems, Courtaboeuf, France) to determine the molecular nature of the modifications.

Identification of the 3'DD1 break point

Ligation mediated-PCR (LM-PCR) was performed as described (Hempel et al. 1998) using CD3 $\epsilon^{\Delta 5/\Delta 5}$ total thymocyte DNA as template and the DR19/DR20 double stranded linker (Roth et al. 1993). Amplification was performed with the 3'DD1 and DR20 oligonucleotide primers. The resulting PCR products were digested to completion with ApaL1, separated on an agarose gel, transferred onto a nylon membrane and hybridized with 3'DD1p as described above. To determine the exact location of the double strand break, this amplicon was cloned and individual colonies sequenced.

p53 amplification.

DNA input in PCR reactions was verified by amplification of a fragment of the p53 gene with the oligonucleotides X6.5 and X7 (Table 1) for 35 cycles consisting of 30 sec at 94°C, 30 sec at 55°C and 1 min at 72°C followed by an incubation of 10 min at 72°C. Before the first cycle, DNA was denatured for 5 min at 94°C. PCR products were separated on an agarose gel in the presence of ethidium bromide and visualized on a Vistra Systems Fluorimager SI with the Image Quant 5.2 software (Molecular Dynamics, Bondoulfe, France).

Results

We analyzed the structure of TR DD1/DD2 signal joints in thymocytes prepared from CD3 ϵ -deficient CD3 $\epsilon^{\Delta 5/\Delta 5}$ mice. CD3 ϵ -deficiency prevents TR expression on developing thymocytes and arrests T cell development at an early stage, but does not impede V(D)J recombination activity. Consequently, *TRD* gene rearrangement is normal in these mice (Gallagher et al. 1998; Malissen et al. 1995). Signal joints are believed to be created by the perfect juxtaposition of RS heptamers, thereby generating a restriction site for the enzyme ApaL1 if these heptamers conform to the consensus sequence and begin by a CAC or GTG triplet. However, following amplification and ApaL1 digestion, only a minor fraction of TR DD1/DD2 signal joints contain an ApaL1 restriction site and most of the PCR product is not digested (Fig 1). Thus, it appears that in the majority of the recombination events, TR DD1/DD2 signal joints do exhibit junctional diversity. After amplification, cloning and ApaL1 digestion, only about 26% (11 out of 42) of the individual signal joints were found to contain a site for this restriction enzyme. Therefore, in almost three quarters of TR DD1/DD2 signal joints, the TRDD1 and/or the TRDD2 signal ends have been processed before ligation. To determine the exact nature of these modifications, ApaL1-R plasmids were sequenced (Fig 2C). We referred to published sequences (Chien et al. 1987) to delineate the boundaries of both *TRDD1* and *TRDD2* genes and their RSs (Fig 2A). The sequence of ApaL1-R TR DD1/DD2 signal joints were aligned with the theoretical signal and coding ends generated by RAG cleavage (Fig 2B) in order to identify the nucleotides that have been lost and/or added prior to the ligation. None of these joints were found to exhibit nucleotide loss. All were found to contain non-templated N nucleotides, as described for TRB, TRA and other TRD signal joints (Candéias et al. 1996; Kanari et al. 1998; Touvrey et al. 2006). Interestingly, each of the ApaL1-R junctions contained at least 3 added nucleotides. This feature is unusual when compared to our previous results where additions of one or two nucleotides were

frequent (Candéias et al. 1996; Touvrey et al. 2006). In addition, all the stretches of added bases contain a “CA” dinucleotide immediately adjacent to the TRDD1 heptamer (Fig 2C). As the *TRDD1* gene sequence ends with “CA”, this observation raises the possibility that the position of the double strand break introduced by the RAG-1/RAG-2 complex 3’ of the *TRDD1* gene is 2 nucleotides (nt) inside the published gene sequence.

To determine the exact location of this break point, we performed LM-PCR on DNA prepared from CD3 ϵ -deficient thymocyte. This assay allows the amplification of blunt ended DNA molecules, such as V(D)J recombination-generated signal ends, following ligation of a ds linker (Fig 3A and 3B). The product detected in RAG-2 $^{-/-}$ thymocyte DNA (Fig 3A) was considered to result from the presence of a non specific DNA break because it migrates slightly slower than the products amplified from CD3 ϵ -deficient thymocytes, is totally resistant to ApaL1 digestion (fig 3A), and because no TR DD1/DD2 signal joints could be amplified from this sample (Fig1). As expected for signal ends, (Roth et al. 1993), the PCR products obtained from CD3 ϵ -deficient DNA after linker ligation are almost fully digestible by ApaL1 (Figs 3A and 3B). The creation of a restriction site for this enzyme upon ligation with the double stranded linker indicates that most of the 3’TRDD1 signal ends end in a CAC triplet. However, this reconstitution does not allow any discrimination between the two possible cleavage sites. Therefore, the LM-PCR products were cloned and individual plasmids sequenced. All of the TRDD1 3’ signal ends identified in this experiment (20 clones from 2 mice) were found to contain an additional CA dinucleotide (Fig 3C). Because the presence of these “extra” bases would not prevent the creation of an ApaL1 restriction site upon formation of a signal joint, we then sequenced two putatively un-modified ApaL1-S signal joints identified during our initial screening procedure. They were also found to contain this CA dinucleotide (Fig 2D). We concluded from these experiments that this CA dinucleotide results

from the introduction of a double strand break 2 bases upstream of the described 3' TRDD1 RS, within the *TRDD1* gene and is present in all the TR DD1/DD2 signal joints.

Alignment with a consensus RS23 sequence shows (Fig 4A) that this shift results in an increase in the percentage of nucleotide identity from 36% (14 bases out of 39) to 56 % (22 matches out of 39). The numbers of identical bases in the heptamer (5) and in the nonamer (6) motifs do not change, but they are distributed differently. The main difference resides in the spacer, where the number of positions in the RS matching with the consensus sequence raises from 3 to 11. However, simple sequence comparison and alignment do not provide any information regarding the RS functionality, except for a few essential positions such as the CAC triplet in the heptamer. (sentence deleted). When the spacer sequence containing the maximum mutual information (Cowell et al. 2004) was used in place of the consensus spacer sequence, the proposed 2nt shift again induces an increase in the percentage of identity (from 41% to 53%) of the 3' TRDD1 RS with this model RS23 sequence (Fig 4B). As before, this increase results from a higher number of identical bases in the 23 nt spacer sequence. More importantly, this 2 nt shift raises the RIC score of the 3' TRDD1 RS from -70 to -52. This increased RIC score indicates that the shifted RS has the higher recombination efficiency when compared to the published sequence. This finding is in agreement with the fact that the only molecular species found during the LM-PCR analysis of the 3' TRDD1 signal ends identified a breakpoint 2 nt inside the *TRDD1* gene.

Discussion

TRD gene rearrangement is complex. This locus encompasses two *D* genes that can be used concurrently. The determination of *TRDD1* and *TRDD2* gene/RS limits were initially made solely by comparing their germ line sequences with the sequence of rearranged genes and attributing RS properties to the sequence most similar to published RS sequences (Chien et al. 1987). The short length of the *TRDD1* and *TRDD2* genes, together with a high level of end processing (exonucleolytic nibbling, P and N nucleotides) makes the exact assignment of their boundaries in coding joints difficult and sometimes imprecise (Elliott et al. 1988; Lacy et al. 1989; Takagaki et al. 1989). By analyzing the diversity of the reciprocal TR DD1/DD2 signal joints, we identified a pattern suggesting that bases from the *TRDD1* gene were specifically retained in these signal joints. Imprecise cutting leading to the retention of coding flank has been reported only within the context of V(D)J recombination performed with mutated RAG-1 and/or RAG-2 proteins (Lee and Desiderio 1999; Talukder et al. 2004). Thus, it seems highly unlikely that this phenomenon can account for the systematic inclusion of the CA dinucleotide in all the TR DD1/DD2 signal joints in mice possessing normal RAG proteins and V(D)J recombination activity, as was previously suggested by (Carroll et al. 1993). We therefore considered the possibility that the location of the double-strand break generating the inclusion of these two bases in the signal joints is not random but programmed, dictated by the position of the 3' TRDD1 RS. To verify this hypothesis, we cloned and sequenced the corresponding signal ends. The only molecular species found was terminated by this “extra” CA dinucleotide. Once loaded onto a RS, the RAG-1/RAG-2 complex creates a double strand break exactly at the gene/RS boundary (McBlane et al. 1995; Roth et al. 1993). This result therefore defines a new RS23 at the 3' end of the *TRDD1* gene, where the heptamer sequence reads CACACAG in place of CACAGGT (Fig 4C). This new RS23 presents more homology with a consensus RS23 sequence and with a model RS23 in which the spacer sequence

contains the highest mutual information possible. At the very least, our results indicate that there are two overlapping RSs downstream of the murine *TRDD1* gene, which differ in their position by 2 nt. However, out of the 20 signal ends sequenced after LM-PCR, none was found to correspond to the published 3' TRDD1 RS. Thus, if it is at all functional, the published 3' TRDD1 RS sequence, ending with the CACAGGT heptamer, is not used in more than 5% of the recombination events. This experimental conclusion is strongly supported by the fact that the calculated RIC score for the new 3' TRDD1 RS23, as defined in this study, is much higher than the score for the published TRDD1 RS23. This difference denotes a difference in recombination efficiency and correlates with our observations: out of two overlapping putative RS competing for the binding of the RAG proteins and the recombination machinery, only one is found in signal joints. Thus, our work defines a new sequence for the functional RS23 downstream of the *TRDD1* gene (Fig 4C). Interestingly, in this revised version, the nucleotides in positions 5-8 of the nonamer (AAAC) now match the consensus sequence. These nucleotide positions are those that, of positions within the RS nonamer, show the lower position-wise entropy, i.e. the most conserved, in a collection of 155 physiologic RS23 elements (Cowell et al. 2002). This conservation may reflect their involvement in base-specific DNA-protein contact, as shown by interference footprinting assays (Swanson 2002). Finally, it should be noted that this proposed revision of the murine 3' TRDD1 RS strikingly increases its similarity to its human counterpart, downstream of the *TRDD2* gene. The orthologous murine and human RSs now have an identical heptamer, and they differ at only 6 positions, 5 in the spacer and 1 in the nonamer. Without this revision, the sequences of these RSs differ at 24 of the 39 position (Fig 5).

In conclusion, we show by a combination of molecular biology and computational analysis that the 3' *TRDD1* gene/RS boundary is shifted 2 nt inside the *TRDD1* gene when compared with the published sequence, and therefore that when used in a recombination event, the *TRDD1* gene is shortened by 2 nt. This was probably not recognized earlier because of the

very high level of diversity in TR DD1/DD2 coding joints, in which only fragments of the *TRDD1* sequence are retained and/or identifiable (Elliott et al. 1988; Lacy et al. 1989; Takagaki et al. 1989). In this context, it is formally possible that the phenomenon we are describing in this paper for the *TRDD1* gene also applies to other TR or Ig genes but was never detected because it is masked by the exonucleolytic nibbling of coding ends that takes place during V(D)J recombination. Indeed, examination of 59 murine TRAJ genes from the IMGT database revealed for 17 of them the presence, adjacent to the heptamer, of a TG dinucleotide. These 17 genes could therefore have 2 overlapping heptamers. However, RIC score calculation predicted in each case that a 2 bp shift would result in a less efficient RS element (Table 2). Thus, it appears that the mere presence of a TG or CA dinucleotide at the border of a heptamer is not sufficient to modify the functional RS element, and that the revision that we are proposing for the 3' *TRDD1* RS is specific for this gene. This study illustrates the contribution of computational modelling to the difficult task of RS identification and characterization, even in the case of already described functional genes.

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Figure legends

Fig 1: TR DD1/DD2 signal joints are diverse.

Signal joints generated following rearrangement of *TRDD1* and *TRDD2* genes were amplified from thymocyte DNA prepared from two CD3ε-deficient and one RAG-2-deficient mice. They were then migrated on an agarose gel without (-) or with (+) ApaL1 digestion and analyzed as described in Materials and Methods. The arrowhead indicates the position of the 3' TRDD1 signal end. The bottom panel shows amplification of the *p53* gene as a DNA input control.

Fig 2: Sequences of the ApaL1-R TR DD1/DD2 signal joints

(A): Germline sequences of the murine *TRDD1* and *TRDD2* genes and their RS elements. TRDD1 and TRDD2 coding regions are in bold characters. RS heptamers and nonamers, separated by 12 or 23 bases, are underlined. The identification of the different elements is based on the published sequences (Chien et al. 1987). (B): Representation of the putative coding and signal ends. The boundaries of the putative coding and signal ends (CE and SE, respectively) are depicted for both TRDD1 and TRDD2. Coding regions are in bold characters. RS heptamers and nonamers, separated by 12 or 23 bases, are underlined. (C): Sequences of the modified signal joints. The amplified signal joints were sequenced with the 5'DD2p oligonucleotide. Identification of TRDD1 and TRDD2 RS was based on the published sequences. The sequences of TRDD1 and TRDD2 RS heptamers are indicated on the first line in bold characters; the first flanking TRDD1 and TRDD2 bases are indicated in italic characters. The nucleotides listed under the “N” column represent putative non-templated nucleotides. The recurrent CA dinucleotide flanking the TRDD1 RS is in bold, underlined characters. The number 2 in the last column indicates that these particular sequences were found twice in a sample from one mouse. Identical sequences found in

different samples were considered as distinct events. The last two sequences represent non-standard rearrangement products generated through the recombination of the 5' TR DD1 and 3' TR DD2 RS (Kanari et al. 2003). In these junctions, the sequence of the *TRDD1* and *TRDD2* genes are underlined. In (D), the sequence of two putatively un-modified ApaL1-S signal joints is shown. The additional “CA” dinucleotide adjacent to the TRDD1 RS was also found.

Fig 3: LM-PCR analysis of the 3' TRDD1 RS end from murine thymocytes.

(A) Total thymocyte DNA from two CD3ε-deficient and one RAG-2-deficient mice was amplified as described in Materials and Methods after ligation of a staggered double-stranded linker. The bottom panel shows the amplification of the *p53* gene for the three DNA samples as a DNA input control. The product amplified from RAG-2^{-/-} thymocyte DNA was considered to result from non-specific amplification (see text for details) and its nature was not explored further. (B) Total thymocyte DNA from a CD3ε-deficient animal reacted with the DR19/DR20 linker in the presence (T4 Lig +) or in the absence (T4 Lig -) of T4 DNA ligase was amplified with the 3' DD1 and DR20 oligonucleotide primers. DNA input control is shown on the bottom panel for the samples incubated with (1) and without (2) ligase. In (A) and (B), one half of the resulting PCR product was digested with ApaL1. After separation on an agarose gel and blotting, un-digested (-) and digested (+) PCR products were revealed by hybridization with the 3' DD1p radio-labelled probe. (C) Sequence of the 3' RS23 TRDD1 signal end. The germline sequence of the TRDD1 gene (in bold characters) and its 3' flanking region element is shown on the top “GL” line. The published sequence of the heptamer is underlined. The arrowhead indicates the position of the RAG-induced break. The “SE” line shows the sequence of the 3' TRDD1 signal end as determined following sequencing of the LM-PCR products. The extra “CA” dinucleotide is in bold characters.

Fig 4: Comparison of the 3' TRDD1 RS elements.

The published (Chien et al. 1987) and new (this study) 3' TRDD1 RS23 sequences are aligned with a RS23 consensus sequence (**A**), a RS sequence assembled from a consensus heptamer, a consensus nonamer and the 23bp spacer with the most mutual information (**B**), and one another (**C**). Both sequences are from Cowell et al, 2004. The *TRDD1* gene and the different elements composing the RS are separated by hyphens. DD1: *TRDD1* gene; 7: heptamer; 23: 23 bp spacer; 9: nonamer. In (**A**) and (**B**), identical bases are replaced by semi columns; in (**C**), identical bases are in bold capitals and differing bases in lowercase characters.

Fig 5: Alignment of the human 3' TRDD2 and murine 3' TRDD1 RS elements

The human 3' TRDD2 and murine 3' TRDD1 RSs are aligned using either the published (**A**) or revised (**B**) sequence for the murine element. Identical bases are in bold capitals and differing bases are in lowercase characters. The different elements composing the RSs are separated by hyphens. 7: heptamer; 23: 23 bp spacer; 9: nonamer.

Table 1: Oligonucleotides used in this study

3'DD1	5'-GACAATACAGACCAAATATACAGC-3'
3'DD1p	5'-GCGCTGTAGGGAAATATGTAAGTTT-3'
5'DD2	5'-TGGCTTGACATGCAGAAAACACCTG-3'
5'DD2p	5'-GACACGTGATAAAAGCCCAGGGAA-3'
DR20	5'-GCCTATGTACTACCCGGGAATTCTG-3'
DR19	5'-CACGAATTCCC-3'
X6.5	5'-ACAGCGTGGTGGTACCTTAT-3'
X7	5'-CACATGTACTTGTAGTGGATGG-3

Table 2: RS associated with TRAJ gene segments that have TG as the most 5' nucleotides of the coding region as defined by IMGT. The second RS in each row indicates the RS for which the heptamer includes the coding region TG as its first two nucleotides.

TRAJ gene	RS sequences	RIC score IMGT RS	RIC score shifted RS
>M64239 TRAJ11*01	cattttgtggagagggttgctgtgt ttttgtggagagggttgctgtgtgt	-24.60	-36.34
>M64239 TRAJ17*01	tgttttgcgtggcttcagatcactgt ttttgcgtggcttcagatcactgtgt	-20.72	-53.66
>M64239 TRAJ2*01	ggcttcgtaaagggtgtcacctgcagt cttcgtaaagggtgtcacctgcagtgt	-27.07	-59.52
>M38103 TRAJ2*02	ggcttcgtaaagggtgtcacctgcagt cttcgtaaagggtgtcacctgcagtgt	-27.07	-59.52
>M64239 TRAJ21*01	gcttcgtataatgggtctaaccattgt tttcgtataatgggtctaaccattgtgt	-23.60	-61.65
>M64239 TRAJ23*01	tgttttgcacagggtatgttaacacagt ttttgcacagggtatgttaacacagtgt	-21.74	-53.12
>M64239 TRAJ24*01	ccattttgttagacgtgttcacagt atttgttagacgtgttcacagtgt	-28.30	-42.01
>M64239 TRAJ30*01	cgttttgggtatggcacaatcacagt tttgggtatggcacaatcacagtgt	-21.20	-61.81
>X02858 TRAJ32*01	agttattgttaaggctctgcagggtgt ttattgttaaggctctgcagggtgtgt	-22.82	-49.57
>M64239 TRAJ33*01	tgttttgttaagggtttgtctgt ttttgttaagggtttgtctgtgt	-22.46	-45.63
>M64239 TRAJ39*01	ggttttgctgagctggagatcactgt ttttgctgagctggagatcactgtgt	-19.20	-42.07
>M64239 TRAJ4*01	aattcttgtaaagcacccttctactgt ttcttgtaaagcacccttctactgtgt	-30.19	-53.24
>X02859 TRAJ45*01	ggtttatgtcaaggcttcctcagggt tttatgtcaaggcttcctcagggtgt	-19.42	-60.55
>M64239 TRAJ47*01	agttttgtcacaggagttgaggctgt ttttgtcacaggagttgaggctgtgt	-26.15	-40.71
>M64239 TRAJ56*01	agttttgttagagtcccgtgtcattgt ttttgttagagtcccgtgtcattgtgt	-23.89	-49.61
>M64239 TRAJ57*01	agtatttgttaaggcagtgtgtgggt tatttgttaaggcagtgtgtgggtgt	-26.48	-47.20
>M64239 TRAJ58*01	agttttgcaaaggcccttcagtgcagt ttttgcaaaggcccttcagtgcagtgt	-24.44	-44.54

Fig 1

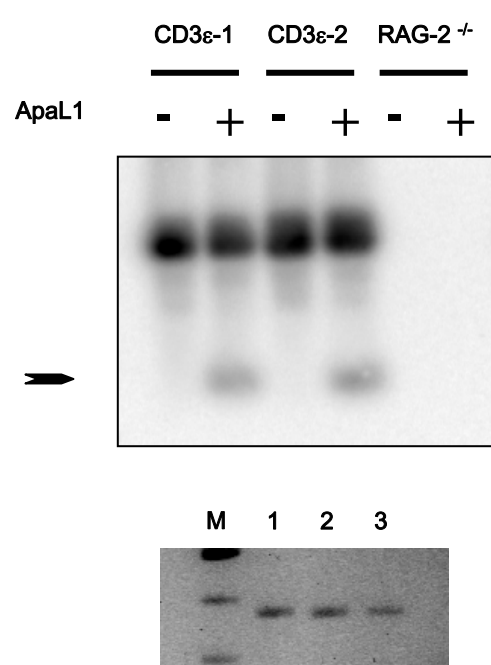


Fig2 :

A : Germline sequences

	DD1	7	23	9
TRDD1	...	/GTGGCATATCA	CACAGGTTGAAGTATATTAAACCTCTGTTCAGAACACT	
TRDD2	9	12	7	DD2
	GGTTTTGCAAAGCTCTGTAGCACCGTG	ATCGGAGGGATACGAG//..		

B: Signal and coding ends following RAG-mediated cleavage

	CE	SE	
TRDD1	.../GTGGCATATCA	CACAGGTTGAAGTATATTAAACCTCTGTTCAGAACAC	
TRDD2	GGTTTTGCAAAGCTCTGTAGCACCGTG	ATCGGAGGGATACGAG//..	

C: Sequences of the ApaL1-R TR DD1/DD2 signal joints

DD2	N	DD1
CACCGTG	ATCGGA	CACAGGT
CACCGTG	G <u>TGGGCA</u>	CACAGGT
CACCGTG	G <u>GGCA</u>	CACAGGT
CACCGTG	G <u>GTCA</u>	CACAGGT
CACCGTG	G <u>GACA</u>	CACAGGT
CACCGTG	G <u>GACA</u>	CACAGGT
CACCGTG	C <u>CCC</u> <u>CA</u>	CACAGGT
CACCGTG	C <u>ACCA</u>	CACAGGT
CACCGTG	T <u>GTC</u> <u>CA</u>	CACAGGT
CACCGTG	T <u>TCC</u> <u>CA</u>	CACAGGT
CACCGTG	G <u>GC</u> <u>CA</u>	CACAGGT
CACCGTG	C <u>CC</u> <u>CA</u>	CACAGGT
CACCGTG	A <u>CCA</u>	CACAGGT
CACCGTG	G <u>CA</u>	CACAGGT
CACCGTG	C <u>CA</u>	CACAGGT
CACCGTG	C <u>CA</u>	CACAGGT
CACCGTG	T <u>CA</u>	CACAGGT
CACCGTG	T <u>CA</u>	CACAGGT
CACCGTG	G <u>GCA</u>	CACAGGT
CACCGTG	ATCGGAGGGATACGAGGGGG <u>CA</u>	CACAGGT
CACCGTG	ATCGGAGGGATACCTGTGGCATAT <u>CA</u>	CACAGGT

D: Sequences of the ApaL1-S TR DD1/DD2 signal joints

DD2	N	DD1
CACCGTG	ATGCGA	CACAGGT
CACCGTG	<u>CA</u>	CACAGGT

Fig 3

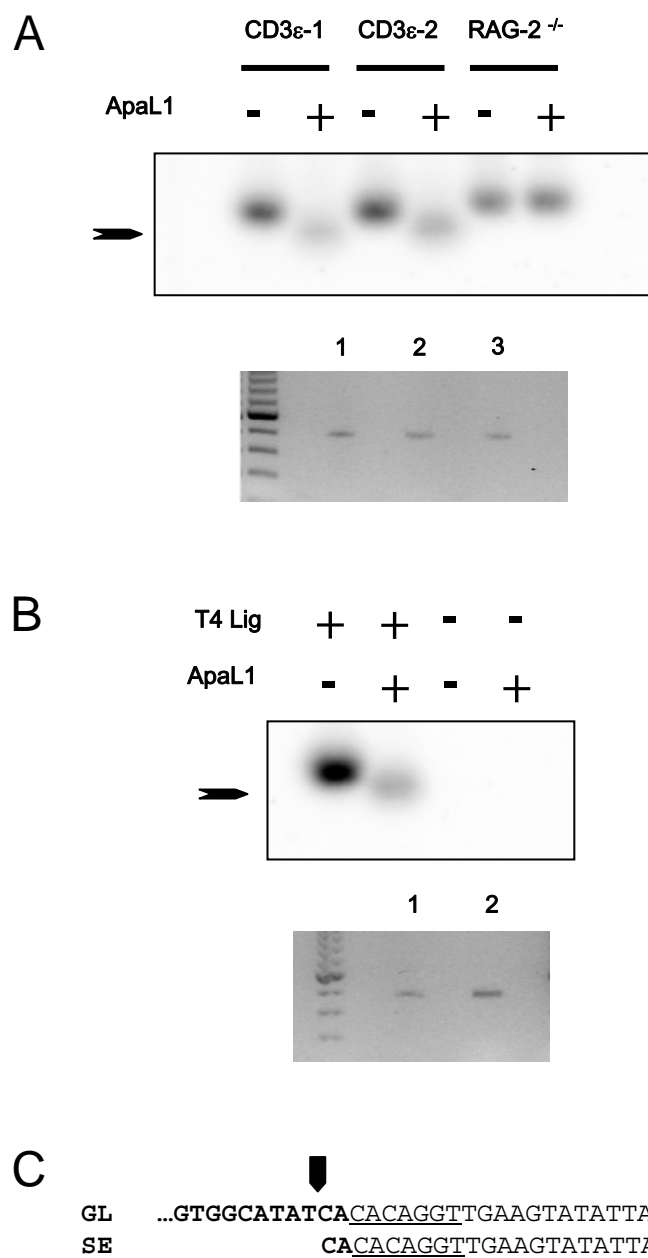


Fig 4

A

	DD1	7	23	9
RS23 Cons	GTGGCATATCA-CACAGTG-	TGGAACCACATCGGGAGCCTGT-	ACAAAAAACC	
3' DD1 (Chien)	GTGGCATATCA-:::	GT-:GAAGTAT:TTAACCTCTG:TC-:G:::C::T		
3' DD1 (this study)	GTGGCATAT -:::CA:-G:T:::GT:T::TAAACCT:::-T::G:::A			

B:

	DD1	7	23	9
RS23 MI	GTGGCATATCA-CACAGTG-	TGCAACCACATCCTGAGTGTGT-	ACAAAAAACC	
3' DD1 (Chien)	GTGGCATATCA-:::GT-:GAAGTAT:TTAACCTC:::TC-:G:::C::T			
3' DD1 (this study)	GTGGCATAT -:::CA:-G:TG::GT:T::TAAACC:C:::-T::G:::A			

C:

	DD1	7	23	9
3' DD1 (Chien)	GTGGCATATCA- CACAggt-tgaagtaTATTaaAcctCTgTtc-agAaAcACt			
3' DD1 (this study)	GTGGCATAT- CACAcag-gttgaagTATTaaacCTcTgt-tcAgAaACa			

Fig 5

A:

7 23 9
h3' TRDD2 CACAcag-gttg**GagTgcattAagcCTGTgt-gcAgAaACa**
m3' TRDD1 **CACAggt-tgaaGtaTattaaAcctCTGTtc-agAaAcAct**

B:

7 23 9
h3' TRDD2 CACACAG-GTTG**gAGTgcATTA**AgCCTtTGT-c**CAGAAACA**
m3' TRDD1 **CACACAG-GTTG**aAGTatATTAAaCCTcTGT-t**CAGAAACA**