Nonsense-associated alternative splicing of T-cell receptor \( \beta \) genes: No evidence for frame dependence

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ABSTRACT

Mutations that generate premature translation-termination codons (PTCs) often result in production of alternatively spliced mRNAs. While in many cases, the PTC-causing mutation was found to affect splicing directly by disrupting an exonic splicing enhancer, induction of alternative splicing of TCR-\( \beta \) pre-mRNA has been reported to be specific for mutations that prematurely terminate the open reading frame. During testing of a cyto-nuclear feedback model that would have explained how cytoplasmic translation could influence nuclear splicing of TCR-\( \beta \) transcripts, control experiments questioned the frame dependence of the nonsense-associated altered splicing (NAS) of TCR-\( \beta \) pre-mRNA. A subsequent detailed analysis of alternatively spliced TCR-\( \beta \) mRNA expressed from different minigene constructs with nonsense, silent, or frame-shift mutations at various positions revealed no correlation between truncation of the reading frame and production of alternatively spliced mRNA. Our study thus contradicts the previously reported PTC specificity of TCR-\( \beta \) NAS and points out the need for systematically testing the PTC specificity in other cases where NAS has been observed.

Keywords: premature translation-termination codon (PTC); nonsense-associated altered splicing (NAS); exonic splicing enhancer (ESE)

INTRODUCTION

Messenger RNAs whose open reading frame (ORF) is prematurely terminated by a nonsense mutation are recognized and specifically degraded by a quality control system called nonsense-mediated mRNA decay (NMD) (Maquat 2004). In addition to triggering NMD, nonsense mutations are also frequently found to alter the splicing pattern of the nonsense codon-containing precursor mRNA (pre-mRNA), an observation that is referred to as nonsense-associated altered splicing (NAS) (Hentze and Kulozik 1999). In many cases, splice site selection is altered so that the sequence comprising the offending premature translation-termination codon (PTC) is skipped from the spliced mRNA (Li and Wilkinson 1998; Valentine 1998; Hentze and Kulozik 1999; Maquat 2001; Cartegni et al. 2002). If the alternatively spliced transcript remains in-frame, it will encode for a protein that may in some cases save the protein’s function, although an internal part of the amino acid sequence is altered or deleted.

The term NAS implies that the alternative splicing event is dependent on the presence of a PTC and thus of premature termination of the ORF. However, in most cases of NAS that were thoroughly investigated, it was found that not only nonsense, but also some silent or missense mutations cause alternative splicing, and that these mutations disrupted an exonic splicing enhancer (ESE) critical for exon inclusion (Liu et al. 2001; Buchner et al. 2003; Pagani et al. 2003). Thus, in these cases, alternative splicing is a direct effect of mutating a splicing signal and does not depend on premature termination of the ORF.

In contrast, evidence for a link between splice site selection and maintenance of an ORF within an mRNA has also been published. For example, Li and colleagues recently showed that latent 5′ splice sites present in introns of the CAD and IDUA gene are efficiently used when all in-frame termination codons between the normally selected 5′ splice site and the latent 5′ splice site are removed (Li et al. 2002). If the alternatively spliced transcript remains in-frame, it will encode for a protein that may in some cases save the protein’s function, although an internal part of the amino acid sequence is altered or deleted.

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SD-mRNA was PTC specific, since it was not observed with missense or silent mutations, and since it was suppressed in genes with a frame shift when the ORF was restored by an additional, compensating frame-shift mutation (Wang et al. 2002b). Furthermore, up-regulation of altSA/SD-mRNA was inhibited by a stem-loop structure in the 5′ UTR, by expression of suppressor tRNA or by removal of the initiator AUG codons, indicating the requirement for frame scanning (Wang et al. 2002a,b). Finally, NAS of PTC+ TCR-β transcripts was shown to require the NMD factor hUpf1/rent1, but not hUpf2/rent2, suggesting that NAS relies on a component of the nonsense surveillance machinery, but is not an indirect consequence of NMD (Mendell et al. 2002; Wang et al. 2002a).

Wang and colleagues proposed a model for frame-dependent NAS, according to which the PTC in the mature mRNA is recognized by nuclear translation at the site of transcription, affecting the local concentration of splicing factors or nucleases involved in general pre-mRNA turnover, and thus altering the production of spliced mRNAs from newly synthesized precursors in trans (Wang et al. 2002a,b). An alternative model to explain frame-dependent NAS of TCR-β transcripts that does not involve nuclear translation was proposed by Dahlberg et al. (2003). Their so-called “cyto-nuclear feedback model” postulates that a shuttling splicing factor could signal the nucleus that a cytoplasmic mRNA contained a PTC.

We set out to test the cyto-nuclear feedback model by coexpressing a PTC− and a PTC+ TCR-β minigene in HeLa cells and found that NAS was allele specific in our experimental system. But more importantly, control experiments suggested that NAS was not frame dependent in our feedback reporter system. This led us to more systematically investigate the issue of frame dependence of TCR-β NAS, including the attempt to reproduce some of the data published by Wang and colleagues. In contrast to their results, our data provide no evidence for frame-dependent up-regulation of TCR-β altSA/SD-mRNA.

RESULTS

Splicing of PTC-free TCR-β pre-mRNA is not affected in cells coexpressing both a PTC− and a PTC+ TCR-β gene

According to the model proposed by Dahlberg et al. (2003), splicing of wild-type (PTC−) TCR-β pre-mRNA should be affected by coexpression of its PTC+ counterpart, resulting in increased levels of alternatively spliced PTC− mRNA (PTC− alt-mRNA). To test this, we selected polyclonal pools of HeLa cells that constitutively expressed either PTC−, PTC+, or PTC− and PTC+ TCR-β minigenes together. To be able to distinguish PTC− from PTC+ transcripts in the subsequent RNase protection assay (RPA), a short sequence tag was inserted into the 3′ UTR of one of the constructs. Normally and alternatively spliced TCR-β mRNA levels were analyzed by RPA using a probe complementary to the final exon (including the tag) and the 3′ part of the VDJ exon (Fig. 1A). This experimental setup allowed us to determine the relative levels of all four TCR-β mRNA species at once. To avoid potential effects on mRNA levels caused by promoters competing for a limiting transcription factor, each cell pool was transfected with the same amount of the expression plasmid. Thus, for cells expressing PTC− or the PTC+ TCR-β mRNA alone, the respective TCR-β expression plasmid was cotransfected with an identical amount of the same expression vector expressing an unrelated gene. In addition, a third plasmid encoding the β-globin gene (Thermann et al. 1998) was included in each transfection for normalization.

As shown in Figure 1, B and C, the cells expressing the PTC+ TCR-β minigene produce about 3.5-fold more alt-mRNA compared to the cells expressing the PTC− construct (cf. lanes 7–9 and lanes 4–6). The cells expressing both constructs together produce fivefold more PTC+ than PTC− alt-mRNA (lanes 10–12). Thus, in contrast to the prediction made by the cyto-nuclear feedback model, alt-mRNA levels for the PTC− allele are not increased when the PTC+ TCR-β minigene is coexpressed. The difference in alt-mRNA levels between PTC+ and PTC− transcripts observed here is in the same range as reported previously from work with similar TCR-β constructs (Wang et al. 2002a,b). In summary, this result is incompatible with a feedback mechanism that relies on a freely diffusible factor and shows that NAS is allele specific. On the other hand, the result would be consistent with the proposed nuclear scanning model, under the assumption that usage of these alternative splice sites was up-regulated as a result of truncation of the reading frame, as previously published (Wang et al. 2002b). However, allele-specific effects on splicing would also be observed if the nonsense mutation in the PTC+ construct had a direct effect on splicing, independent of truncating the ORF. It was therefore important to check for PTC specificity of TCR-β NAS in our experimental system.

NAS of the TCR-β feedback reporter constructs is not PTC specific

To test whether the increased levels of TCR-β alt-mRNA were specific for the T-to-A nonsense mutation at codon 68, we also introduced the silent T-to-C mutation at this position in the untagged TCR-β minigene [PTC− (UAC); Fig. 2]. The untagged wild-type [PTC− (UAA), PTC− (UAC), PTC+ (UAA)] and the tagged wild-type [PTC− (UAA)-tag] constructs were transiently expressed in HeLa cells, and relative TCR-β mRNA and alt-mRNA levels were determined by real-time RT-PCR using splice variant-specific TaqMan probes. The probe to measure alt-mRNA spans the junction between the alternative splice donor site (altSD) in the VDJ exon and the 3′ splice site of the final exon C1/4.
normally spliced mRNA was measured over the junction between the normal 5’/H11032 splice site of the VDJ exon and the 3’/H11032 splice site of the final exon (Buhler et al. 2002). A β-globin encoding plasmid (Thermann et al. 1998) was cotransfected for normalization. As shown in Figure 2B, the normally spliced PTC+ mRNA was reduced about 100-fold compared to the wild-type mRNA, irrespective of whether the wild-type mRNA was tagged or not. Surprisingly, the T-to-C mutation resulted in twofold more mRNA. The result of the alt-mRNA levels was unexpected in two respects. First, in contrast to the 3.5- to five-fold higher alt-mRNA levels from the PTC+ TCR-β minigene observed in the stably transfected cells (Fig. 1), this difference was less than twofold in the transient transfection experiment [Fig. 2C, cf. PTC+ (UAU) and PTC− (UAU)-tag], and compared to the untagged wild-type, there was no significant difference in alt-mRNA levels [cf. PTC+ (UAU) and PTC− (UAU)]. Secondly, the T-to-C mutation resulted in a fourfold reduction of alt-mRNA, suggesting that this mutation might have strengthened VDJ exon definition and thus promotes usage of the normal splice sites (twofold elevated mRNA; see above) and inhibits alt-mRNA production. Consistent with this possibility, analysis of the sequence comprising codon 68 with ESE-RESCUE (http://genes.mit.edu/chris/), a computational tool to predict exonic splicing enhancers (Fairbrother et al. 2002, 2004), found no ESE in the wild-type (UAU) and in the PTC+ (UAA) sequence, but predicted an ESE (CGTCGC, mutated base underlined) in the PTC− (UAC) construct. In summary, these results indicated that mutations of the third position of codon 68 in our TCR-β minigene did not alter splicing by a termination codon-dependent mechanism.

TCR-β alt-mRNA levels of constructs with frame shifts at codon 58 do not correlate with truncation of the ORF

Because we did not find evidence for PTC-specific up-regulation of alt-mRNA at codon 68, we decided to more systematically investigate the question of frame dependence of
which has no PTC, while the other four constructs showed essentially the same level of alt-mRNA. In fact, the second one would expect the same result with construct FS + 2, due to the premature termination of the ORF at codon 62, leading to a polypeptide of no PTC, which has the same PTC. However, FS + 2 showed the lowest level of alt-mRNA (Fig. 3A). If this increased alternative splicing were due to the premature termination of the ORF at codon 62, one would expect the same result with construct FS + 2, which has the same PTC. However, FS + 2 showed the lowest level of alt-mRNA of all FS constructs. In fact, the second construct with an elevated alt-mRNA level was FS + 3, which has no PTC, while the other four constructs showed relatively low alt-mRNA levels. In summary, we did not find any correlation between the relative amount of alt-mRNA and the presence or absence of a PTC in this set of constructs. We conclude that also in this second set of TCR-β constructs, alt-mRNA production is not frame dependent.

No evidence for frame-dependent NAS with a set of TCR-β minigenes used by Wang and colleagues

In the light of the previously published evidence for frame-dependent NAS of TCR-β pre-mRNA, it was puzzling that both of our sets of TCR-β minigenes used to address this issue did not show any evidence for frame-dependent NAS. To further investigate this discrepancy, Miles Wilkinson kindly provided some of their TCR-β constructs that were reported to up-regulate alt-mRNA specifically in response to a PTC (Wang et al. 2002b). For easier comparison, the constructs are labeled with the same letters as in the previous publication (Wang et al. 2002b). Sequencing of the region between the translation initiation codon and the PTC in the C1 exon confirms the identity of each construct (data not shown). Construct I (Wang et al. 2002b) is not informative and was not used, because the deletion of one base pair in the C1 exon shifts the alt-mRNA out of frame, which leads to degradation of this alt-mRNA by NMD. Constructs A, K, N, and M were transiently expressed in HeLa cells and relative levels of TCR-β mRNA and alt-mRNA were determined by real-time RT-PCR using TaqMan chemistry as described previously (Bühler et al. 2004). Relative TCR-β mRNA and alt-mRNA levels were normalized to the relative β-globin mRNA levels, and PTC− (UAU) was set as 100%. Average values and standard deviations of three PCR runs from one typical experiment are shown.

TCR-β NAS. To this end, we generated a set of frame-shift mutants by inserting or deleting one, two, or three nucleotides in a TCR-β minigene, giving constructs FS + 1, FS + 2, FS + 3 or FS − 1, FS − 2, FS − 3, respectively (Fig. 3A). No ESE is predicted in this region for any of the constructs by ESE-RESCUE (data not shown). The structure of these TCR-β minigenes is almost identical to the constructs used in Figures 1 and 2, except that the intron in the 5′ UTR was deleted (see Materials and Methods). The frame-shift constructs were transiently expressed in HeLa cells and the relative levels of TCR-β mRNA and alt-mRNA were measured by real-time RT-PCR. As expected, in cells expressing the PTC+, in-frame constructs FS − 1, FS − 2, FS − 3, and TCR-β mRNA levels were 15- to 40-fold higher than in cells expressing frame-shift constructs with a PTC (Fig. 3B), reflecting the highly efficient NMD for TCR-β transcripts. The highest level of alt-mRNA was obtained with the PTC+ construct FS − 1. If this increased alternative splicing were due to the premature termination of the ORF at codon 62, one would expect the same result with construct FS + 2, which has the same PTC. However, FS + 2 showed the lowest level of all FS constructs. In fact, the second construct with an elevated alt-mRNA level was FS + 3, which has no PTC, while the other four constructs show differences in the cell lines used for the experiments, al-

FIGURE 2. High-alt-mRNA levels of the TCR-β feedback reporter constructs do not correlate with the presence of a PTC. (A) Schematic representation of the two major transcripts produced from the TCR-β pre-mRNA by alternative splicing. Exonic and intronic sequences are shown as boxes and lines, respectively. The dark box depicts the alternative exon, which comprises the last 22 nt of the leader intron and the first 62 nt of the VDJ exon. Alt-mRNA is produced by usage of the same alternative splice acceptor and splice donor sites as altSA/SD-mRNA (Wang et al. 2002a,b), but our minigenes contain as terminal exon a fusion between exons C1 and C4 instead of the full-length constant region of the TCR-β gene (Li et al. 1997) and a shortened JC intron (Stoecklin et al. 2001). The position of codon 68 is depicted by an asterisk, and the position of the TaqMan probe used to measure alt-mRNA is indicated by the short black line. (B,C) HeLa cells were cotransfected with plasmids expressing the indicated TCR-β minigenes and a plasmid expressing β-globin mRNA (Thermann et al. 1998), and relative levels of normally spliced (B) and alternatively spliced (C) TCR-β mRNA and β-globin mRNA were determined by real-time RT-PCR using TaqMan chemistry as described previously (Bühler et al. 2004). Relative TCR-β mRNA and alt-mRNA levels were normalized to the relative β-globin mRNA levels, and PTC− (UAU) was set as 100%. Average values and standard deviations of three PCR runs from one typical experiment are shown.
though HeLa cells were used in both cases. To test if frame-
dependent NAS could be observed in other cells, we re-
peated our experiment using the human embryonic kidney
cell line HEK293T and the human fibrosarcoma cell line
HT1080 (Fig. 4D,E). The relative TCR-β mRNA levels mea-
sured in these cell lines after transfection was similar to
the results obtained in HeLa cells (Fig. 4, cf. C and E) and
showed again no evidence for PTC-specific up-regulation of
alt-mRNA. In both cell lines, the alt-mRNA levels of con-
struct K did not differ significantly from those of construct
A, and the highest relative levels of alt-mRNA was obtained
with the PTC− construct N.

Collectively, our data from transient expression of these
TCR-β minigene constructs in three different human cell
lines indicate that the level of alt-mRNA does not alter
specifically in response to premature termination of the
ORF.

Expression of TCR-β constructs integrated into the
genome confirms data from transient transfections

In search of possible explanations for the discrepancy be-
tween our data and the data reported by Wang and col-
leagues, we also considered the possibility that high expres-
sion levels of the test genes could saturate the putative
mechanism for frame-dependent NAS. From studying
NMD of TCR-β minigenes, we know that transfecting the
cells with large quantities of plasmid DNA can easily ex-
haust the capacity of the NMD system (O. Mühlemann,
unpubl. observation). Although all transient transfection
experiments shown here were done under conditions that
do not saturate NMD (i.e., transfection of relatively small
amounts of plasmid DNA), as can be seen by the efficient
down-regulation of PTC+ mRNA levels (Figs. 2, 3, 4), the
expression levels might still saturate the putative frame-
dependent NAS mechanisms. Consistent with this idea, we
observed 3.5- to 5-fold more alt-mRNA from the PTC+
feedback reporter construct compared to the PTC− con-
struct when these constructs were stably transfected (Fig. 1),
whereas this difference was less than twofold when the same
constructs were transiently transfected (Fig. 2), which gives
a higher rate of gene expression than stable transfections.
Therefore, we selected polyclonal populations of HeLa cells
that had integrated into the genome constructs A, K, N, or
M and measured relative TCR-β mRNA and alt-mRNA
levels as before. As shown in Figure 5A, the levels of nor-
mally spliced mRNA were the same as in the transient trans-
fection (Fig. 4B), confirming that NMD is fully active under
both conditions. Likewise, there was no significant differ-
ence in alt-mRNA levels between stable (Fig. 5B) and tran-
sient (Fig. 4C) transfections. Again, the PTC+ construct K
gave only marginally more alt-mRNA than construct A,
while the PTC− construct N showed twofold more alt-
mRNA, which is inconsistent with frame-dependent regu-
lation of alt-mRNA production.

Confirmation of real-time RT-PCR data by RNase
protection assays

Finally, we wanted to rule out that the different results
obtained by us and by the Wilkinson laboratory could be
due to the different methods used to quantify RNA levels.
Although there was no reason to assume that the RPA used
by the Wilkinson laboratory or our real-time RT-PCR assay
would produce incorrect results, it should be noted that our
TaqMan assay for alt-mRNA will measure all alternatively
spliced transcripts that have altSD spliced to the 3′ splice
site of the C1 exon, while the RPA probe allows specific
quantification of the altSA/SD-mRNA. In fact, an alter-
native TCR-β transcript in which altSD is spliced to the 3′
splice site of the C1 exon, but which uses the normal 3′ splice site of the VDJ exon instead of altSA has been described (Wang et al. 2002a). However, this splice form has a PTC and hence accumulates only at minute amounts in NMD-competent cells (Wang et al. 2002a). Thus, it should influence our real-time RT-PCR measurements of alt-mRNA only marginally.

To directly compare our real-time RT-PCR data with RPA data, we checked our RNA samples from the cells stably expressing constructs A, K, N, and M by RPA. The probe used for the RPAs and the expected size of the protected fragments for normally spliced mRNA, altSA/SD-mRNA, altSD-mRNA, and pre-mRNA are shown in Figure 6A. Initial titration of probe and RNA ensured that our RPAs were performed with an excess of probe (data not shown), which is important to obtain quantitative results. As shown in Figure 6B, the RPAs confirmed the data obtained by real-time RT-PCR. Again, we could not observe a PTC-specific up-regulation of altSA/SD-mRNA. The level of altSA/SD-mRNA from construct K (PTC+) was only slightly higher than with construct A (wild-type PTC−). As in all experiments before, the highest level of altSA/SD-

![Image of Figure 4](https://example.com/image4.jpg)

**FIGURE 4.** No evidence for frame-dependent NAS with a set of TCR-β minigenes used by Wang and colleagues. (A) Schematic representation of the TCR-β minigenes used in these experiments. For easier comparison, the constructs were labeled as in the previous publication (Wang et al. 2002b). Construct K has an insertion of 10 base pairs (GCCCTCGAGG) at codon 83 of the parental TCR-β minigene A, introducing a frame-shift that generates a PTC at codon 151 in the C1 exon. Construct N has an almost identical insertion of nine base pairs (GCCCTCGAG) at codon 83 and remains in frame. Construct M is identical to construct K, except that the ORF was restored by deletion of one base pair of codon 133. After transient transfection of these constructs into HeLa cells, relative levels of TCR-β mRNA (B) and alt-mRNA (C) were determined by real-time RT-PCR and normalized to the relative mRNA levels of the neomycin resistance gene. Average values and standard deviations from four PCR runs of one typical experiment are shown, and the mRNA and alt-mRNA levels of construct A were set as 100%. (D,E) relative TCR-β mRNA and alt-mRNA levels of constructs A, K, N, and M in HEK293T and HT1080 cells. For normalization, a β-globin mRNA expressing plasmid was cotransfected. Average values and standard deviations from three PCR runs of one typical experiment are shown, and the mRNA and alt-mRNA levels of construct A were set as 100%.
mRNA was obtained with construct N. Notably, a faint band of the size expected for altSD-mRNA could also be detected, and comparison of its intensity with the intensity of the band corresponding to altSA/SD-mRNA confirms that this minor splice form does not contribute significantly to our alt-mRNA levels determined by real-time RT-PCR. In conclusion, analysis of our RNA by RPA gives results very similar to analysis by real-time RT-PCR, excluding the possibility that the discrepancy between our results and the results reported by Wang and colleagues could be due to the different methods used.

**DISCUSSION**

At the outset of the study reported here, we wanted to test the cyto-nuclear feedback model (Dahlberg et al. 2003), which was proposed as an alternative to models that postulated the involvement of nuclear ORF scanning to explain the reported PTC-specific up-regulation of an alternatively spliced TCR-β transcript (Wang et al. 2002b). In contrast to the prediction made by the cyto-nuclear feedback model, the level of PTC− TCR-β alt-mRNA did not increase when the PTC+ TCR-β construct was coexpressed in the same cells, demonstrating that NAS in this experimental system was allele specific. While at first sight, this result appeared to support the proposed nuclear scanning models, we noted that allele-specific effects on splicing would also be observed if the nonsense mutation in the PTC+ construct had a direct effect on splicing, independent of truncating the ORF. It was therefore crucial to investigate whether TCR-β NAS was PTC specific in our experimental system.

Here, our study took an unexpected turn. All our experiments that systematically addressed the issue of frame-dependent up-regulation of TCR-β alt-mRNA indicated that there was no correlation between the presence of a PTC in a TCR-β construct and an elevated level of alt-mRNA. Even with the same constructs used previously to demonstrate the PTC specificity of TCR-β NAS, we could not find any evidence for frame-dependent production of alt-mRNA. In both sets of TCR-β minigenes designed to distinguish between frame-dependent NAS and direct effects of the mutations on splicing, there was one PTC− construct that yielded higher levels of alt-mRNA than the PTC+ constructs. Based on our results, we conclude that the different levels of TCR-β alt-mRNA observed with different TCR-β minigenes is unrelated to the presence of a PTC in these genes, but that instead the sequence alterations generated by point mutations or small insertions/deletions directly affect splice site selection. The reason for the discrepancy between our data and the data of Wang and colleagues remains elusive.

In addition to TCR-β, NAS has been observed in many other genes, but only in a few cases has it been systematically investigated whether it is frame dependent. In addition to the examples mentioned in the Introduction, we have recently identified an alternatively spliced immunoglobulin μ transcript in cells expressing Ig-µ minigenes with PTCs in the VDJ exon. Systematic analysis of different mutations at two positions in the Ig-µ minigene clearly showed that production of this alternative transcript was not PTC specific. Instead, the PTC-causing mutations happened to interfere with splicing directly (Buhler and Muhlemann 2005). Collectively, our data from the TCR-β and the Ig-µ minigene systems provide strong evidence against the intellectually appealing hypothesis that cells might possess mechanisms that would allow regulation of splice site selection in response to premature termination of the ORF.

Because it is quite difficult to unambiguously demonstrate frame dependency of NAS and rule out that the altered splicing was caused by mutation of splicing signals, it would be important to carefully reexamine the cases for which NAS has been claimed to be PTC specific before embarking on further projects that build on PTC specificity of the altered splicing. The fibrillin-1 gene is an example where such a reexamination of apparently PTC-specific NAS (Dietz et al. 1993; Dietz and Kendzior 1994) has revealed that the nonsense mutations disrupted an ESE (Caputi et al. 2002).

From the point of splicing regulation, our data fit well into a growing list of examples indicating that exon definition and splice site selection involve many cis-acting signals.
distributed across the pre-mRNA (Maniatis and Tasic 2002; Zheng 2004). These signals together constitute a still poorly understood second layer of information encoded by the RNA sequence, which determines the fate of the RNA. As for the ORF, the first layer of information, single nucleotide changes in many cases are apparently sufficient to alter this second layer information, resulting for example in altered splicing of the RNA.

MATERIALS AND METHODS

Plasmids

The TCR-β feedback reporter genes used in Figures 1 and 2 are based on constructs A (pB333) and C (pB337, PTC at amino acid position 68) described by Li et al. (1997), except that the JC intron was shortened to 357 nt by deletion of a 634-nt-long Eco0109I fragment. The T-to-C mutation in construct PTC− (UAU), in which the β-actin intron in the 5′ UTR was deleted by replacing the Not1-Sal1 fragment with a double-stranded oligo (5′-GGCCGCGGCGCCCTATAAAACCCA GCGCCGCGCCGACCGCGAACCACCACCGG GGTACCG-3′), which shortened the 5′ UTR to 61 nt. One, two, and three base pair insertions and deletions were generated generated with the QuikChange Site-Directed Mutagenesis Kit (Stratagene).

Plasmids encoding constructs A (pB290), K (pB291), N (pB589), and M (pB625) were obtained from Miles Wilkinson and are described elsewhere (Muhlemann et al. 2001). pBS-B433VDI, the template for the RPA probe used in Figure 6, was generated by deleting the sequence between Nde1 and Not1 in pBS-B433-tag, pBS-neo, the template for the RPA probe detecting neomycin mRNA, was generated by inserting the neomycin resistance cassette of MSCVneoEB (Hawley et al. 1994) into Xho1–BamH1 of pBluescript II (Stratagene)

Plasmid sequences and sequences of the primers used for site-directed mutagenesis are available upon request.

Cell culture, transfection, and selection of stably transfected polyclonal cell pools

HeLa, HEK293T, and HT1080 cells were grown in Dulbecco’s modified Eagle’s medium (DMEM; Invitrogen), supplemented with 10% heat-inactivated fetal calf serum (FCS), 100 U/mL penicillin, and 100 µg/mL streptomycin (Amimed). Cells were transfected in six-well plates at about 60–80% confluency using lipofectamine (Invitrogen). For transient transfections 100–150 ng of TCR-β expressing plasmid DNA was used (Figs. 3, 4B,C) or 50

FIGURE 6. Confirmation of real-time RT-PCR results by RNase protection assays. (A) The RPA probe used in (B) is complementary to the last 58 nt of the leader intron and the first 154 nt of the VDJ exon of TCR-β pre-mRNA. The relevant splice sites and the alternative exon included in altSA/SD-mRNA are indicated, and the predicted sizes of protected fragments corresponding to the differently spliced TCR-β transcripts are listed. (B) Twenty-seven micrograms of total cellular RNA from cells expressing the indicated constructs were hybridized with 15 fmol 32P-labeled TCR-β and 15 fmol 32P-labeled neo probe (lanes 3–8) or with each probe alone as a control (lanes 1,2). RNA from untransfected HeLa cells was used as a specificity control of the assay (lane 7) and to show the probes when no RNases were added (lane 8). Protected fragments after RNase digestion were separated on a denaturing polyacrylamide gel and quantified by PhosphorImager scanning. The fragment sizes of the 32P-labeled probe, the template for the RPA probe used in Figure 6, was generated by deleting the sequence between Nde1 and Not1 in pBS-B433-tag, pBS-neo, the template for the RPA probe detecting neomycin mRNA, was generated by inserting the neomycin resistance cassette of MSCVneoEB (Hawley et al. 1994) into Xho1–BamH1 of pBluescript II (Stratagene)

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ng of TCR-β plasmid plus 50 ng of pCneoβglobinwt (Neu-Yilik et al. 2001; Figs. 2, 4,D,E). To generate the cell lines that stably coexpressed the tagged PTC- or (pβ433-tag) and the PTC+ (pβ434) TCR-β minigenes, 200 ng of each plasmid were transfected together with 100 ng pCneoβglobinwt. In parallel, 200 ng of pβ433-tag or pβ434 were transfected together with 200 ng pBhyb8- (Buhler et al. 2004) and 100 ng pCneoβglobinwt to generate cell lines that express only one allele of TCR-β. For generating the stably transfected cell lines used in Figures 5 and 6, 500 ng of pβ290, pβ291, p β589, or pβ625 were used. Polyclonal populations of stably transfected HeLa cells were obtained by selection with 500 µg/ml G418 (Roche) for 2–3 weeks.

RNA isolation
Total cellular RNA was purified using the Absolutely RNA RT-PCR Miniprep Kit according to the manufacturer’s protocol (Stratagene).

RNase protection assay
The TCR-β probe illustrated in Figure 1A comprises the last 199 nt of the VDJ and the exon final until the end of the tag sequence plus an unspecified 5’-part originating from pBluescript II. The probe for detecting β-globin mRNA is complementary to the AvaiI-BamHI fragment of exon 2 in pCneoβglobinwt and contains an unspecified 5’ part originating from pBluescript II. The 32P-labeled probes were generated by in vitro transcription using T7 RNA polymerase and Nde1-digested pBS-β433ΔC-TCGTAAGTGGGCGAGCACA-3′, and pBS-β-globin-exon2 as template, respectively. Nine-teen femtomoles of each gel-purified probe were hybridized with 3, 9, and 27 µg of total cellular RNA, digested with RNases A and T1, and separated on a denaturing polyacrylamide gel as previously described (Suter et al. 1999).

The TCR-β probe illustrated in Figure 6A comprises the last 58 nt of the leader intron and the first 154 nt of the VDJ exon plus an unspecified 5’ part originating from pBluescript II. The 32P-labeled probe was generated by T7 in vitro transcription from the EcoRI-digested pBS-β433VDJ. The neo probe was generated by T7 in vitro transcription from the Sap1-digested pBS-neo. Fifteen femtomoles of each gel-purified probe were hybridized with 27 µg of total cellular RNA digested with RNases A and T1, and separated on a denaturing polyacrylamide gel as previously described (Suter et al. 1999).

Real-time RT-PCR
Reverse transcription and real-time PCR was as described by Buhler et al. (2004). TCR-β alt-mRNA was measured using 800 nM om111 (5′-GCTGCACAGGCAGAGATCGC-3′), 800 nM om112 (5′-AAAGTGGGCGAGCACA-3′), and 200 nM TM111/112 (5′-FAM-TCTTGAACCTTCCAGGTGCTTGGAGTCACAAC-TAMRA-3′).

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N-terminal multimerization domain of TCR-β revisited

Nonsense-associated alternative splicing of T-cell receptor β genes: No evidence for frame dependence

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