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Exon skipping through the creation of a putative exonic splicing silencer as a consequence of the cystic fibrosis mutation R553X

Isabel Aznarez, Julian Zielenski, Johanna M Rommens, Benjamin J Blencowe, Lap-Chee Tsui

Nonsense mutations that occur more than 50 bases upstream of terminal spliced junctions are generally thought to lead to degradation of the corresponding transcripts by the process of nonsense-mediated mRNA decay. It has also been proposed that some nonsense mutations may affect splicing by the process of nonsense-associated altered splicing (NAS), or by the disruption of a splicing regulatory element. In this study, the effect of the R553X mutation on the splicing of exon 11 of the cystic fibrosis transmembrane conductance regulator gene was investigated. Evidence that R553X causes exon 11 to skip through the creation of a putative exonic splicing silencer (ESS) was provided. The putative ESS appears to be active when located immediately upstream of a 5′ splice site. These findings argue against the possibility that R553X-associated exon 11 skipping is caused by NAS. The study further suggests that aminoglycoside antibiotic treatment would not be effective for patients with the R553X mutation, owing to the skipping of exon 11, and further emphasises the need for detailed mechanistic characterisation of the consequences of nonsense disease mutations.

Cystic fibrosis is the most common autosomal recessive disease in the Caucasian population. It is a primary cause of death in young adults, and results from defects in chloride and sodium transport in the respiratory, gastrointestinal, and reproductive systems. The primary cause is a defect in chloride and sodium transport in cystic fibrosis secretory epithelial cells due to mutations in the cystic fibrosis transmembrane conductance regulator (CFTR) gene. CFTR mutations are among the most common monogenic diseases and have been associated with altered splicing patterns. Previous studies have identified sequences that promote exon skipping in the context of nonsense mutations, suggesting that ESSs can be associated with altered splicing patterns.

In addition to ESEs, other splicing elements are known to regulate the inclusion of exons. These include intronic splicing enhancers and intronic splicing silencers, which regulate the selection of splice sites and are often bound by proteins that interfere with the recruitment of the spliceosome to splice sites. Two recent studies have identified groups of exonic splicing silencers (ESSs) using bioinformatics and systematic evolution of ligands by exponential enrichment approaches. The first study identified sequences more frequently present in pseudogenes than in authentic exons and showed that these sequences promoted exon skipping in a reporter system. The second study identified sequences from random pools of hexamers and decamers that prevented the inclusion of an exon and were enriched in pseudogenes compared with constitutive exons.

Two different mechanisms have been proposed to explain how nonsense mutations may affect splicing. The first proposed mechanism, known as nonsense-associated altered splicing (NAS), could involve a scanning process that monitors the open reading frame for premature termination codons (PTCs) before splicing occurs, promoting skipping of the exon in which the PTC is located. The second proposed mechanism, which is supported by more extensive evidence, involves the inactivation of exonic splicing enhancers (ESEs), which are short and degenerate sequences that promote the selection of proximal splice sites in constitutive and alternative exons. It is of interest to note that purine-rich codons (e.g., GAA, GGA and AAG) which often overlap strong ESEs can be mutated by a single base to the PTCs TAA, TGA and TAG, respectively. Evidence that such mutations disrupt ESE function has been reported.

In addition to ESEs, other splicing elements are known to regulate the inclusion of exons. These include intronic splicing enhancers, and, exonic or intronic splicing silencers, which regulate the selection of splice sites and are often bound by proteins that interfere with the recruitment of the spliceosome to splice sites. Two recent studies have identified groups of exonic splicing silencers (ESSs) using bioinformatics and systematic evolution of ligands by exponential enrichment approaches. The first study identified sequences more frequently present in pseudogenes than in authentic exons and showed that these sequences promoted exon skipping in a reporter system. The second study identified sequences from random pools of hexamers and decamers that prevented the inclusion of an exon and were enriched in pseudogenes compared with constitutive exons.

In follow-up experiments, the mode of action of ESSs was investigated and shown to be position dependent with respect to splice sites.

Studies have shown that diseases such as cystic fibrosis caused by nonsense mutations may be treated with aminoglycoside antibiotics. Aminoglycoside antibiotics suppress translational fidelity, resulting in partial expression of the mutated allele. Thus, it is paramount to determine whether stable transcripts are generated from nonsense mutation-containing genes, and to understand the mechanism underlying the disruptive effects of these mutations. To date, distinct results about the effect of the third most common nonsense mutation, R553X, on the CFTR transcript have been obtained.
reported.\textsuperscript{22 25 26} In this report, we have carried out a detailed analysis to determine the mechanism by which the CFTR R553X mutation alters gene expression in patients with cystic fibrosis. We provide evidence that the R553X mutation results in the creation of a putative ESS element that acts on 5′ splice site selection leading to exclusion of exon 11.

MATERIALS AND METHODS
The study group consisted of five patients diagnosed with cystic fibrosis based on the presence of clinical symptoms characteristic for the disease.\textsuperscript{1} All patients were compound heterozygotic for ΔF508 (1652del3, exon 10) and R553X (1789C→T) mutations. Control samples were obtained from healthy individuals and did not carry CFTR mutations. The study was approved by the Human Subjects Review Board of The Hospital for Sick Children.

Total RNA was extracted from cultured lymphoblastoid cells treated with cycloheximide (100 μg/ml in DMSO) or DMSO only for 4 h before harvesting, using RNeasy extraction kit (Qiagen). First strand cDNA was synthesised from 2–3 μg of total RNA by oligo-dT-primed reverse transcription with superscript II reverse transcriptase (Invitrogen) according to the supplier’s directions. A PCR was then performed to generate a fragment spanning exons 10, 11 and 12 of the CFTR transcript using primers X10-5, 5′-GAG GGT AAA ATT AAC CAG AC-3′ and X12-3s, 5′-AGG TAT CCA AAA GGA GAG TC-3′, and, in a fragment spanning exons 7 and 8 of the glyceroldehyde-3-phosphate dehydrogenase (GAPDH) transcript using primers XGAPD-5, 5′-TG TCT GAT GGA GAG ACT CA-3′ and XGAPD-3, 5′-GTG TCG GTG TCG AGG TCA GA-3′. Accordingly, the PCR amplification was as follows; 30 s at 94°C, 30 s at 60°C and 30 s at 72°C for 30 cycles using 0.5 U of AmpliTaq DNA polymerase (Invitrogen). The PCR products were analysed by electrophoresis on 8% polyacrylamide gel. DNA was then transferred to a nylon membrane (Hybond N+ Pharmacia, Amersham) by Southern blotting. An internal oligonucleotide probe complementary to CFTR exon 10 and GAPDH exon 8 were labelled with the radiolabelled dCTP \textsuperscript{32}P using terminal deoxynucleotide transferase (Invitrogen) according to the supplier’s conditions. The blots were probed and the bands were visualised by autoradiography. PCR products displaying altered mobility were extracted using MiniElute Gel Extraction Kit (Qiagen) and subjected to automated DNA sequencing analysis.

The minigene reporter systems CD44v5, CD44AL and CD44AM were described previously.\textsuperscript{27} Seventeen nucleotides flanking the wild-type and R553X alleles of exon 11 of the CFTR gene were introduced into exon 5 of the CD44 gene by overlapping extension using primers CD44-R, 5′-CAT GGA TAA CCA TTA GTC TCA GTG-3′ and CD44-F, 5′-CTC TGC TCA TGG ACC TTA AGA GA-3′. PCR programme consisted of 20 cycles of 30 s at 94°C, 30 s at 50°C (increasing the annealing temperature 0.5°C per cycle) and 1 min at 72°C, followed by an additional 10 cycles of 30 s at 94°C, 30 s at 60°C and 2 min at 68°C. The resulting PCR fragments of 2490 bp were digested using BglII and HpaI and cloned into the V5CD44ΔL and V5CD44AM, respectively, using T4 DNA ligase (Invitrogen). The generated minigene reporters were verified by sequencing.

Human embryonic kidney (HEK) 293 cells (from American Type Culture Collection) were grown in Dulbecco’s modified Eagle’s medium with high glucose supplemented with 10% fetal calf serum and 1% streptomycin–penicillin antibiotics. Human bronchial epithelial (IB3) cells\textsuperscript{28} were grown in LHC-8 medium with glutamine supplemented with 5% fetal calf serum and 1% streptomycin–penicillin antibiotics. The cells were grown in 10 cm tissue culture dishes. A total of 4 μg of total plasmid DNA was transfected into HEK 293 and IB3 cells in replicates using lipofectamine reagent (Invitrogen). After 18 h of transfection, the medium was replaced with fresh complete medium. Cells were harvested 48 h after transfection. Total RNA was extracted as described previously. Oligo dT-primed cDNA synthesis was performed as described previously. PCR amplification (30 s at 94°C, 30 s at 60°C and 30 s at 72°C for 30 cycles using 1.5 U of AmpliTag DNA polymerase (Boehringer Ingelheim)) of the fragment containing CD44 exon 5 was performed using previously described primers.\textsuperscript{27} The fragments were separated on a 2% agarose gel. The ethidium bromide-stained band intensities were measured by analysis of digitised images using AlphaEase V.5.1. Statistical analysis was performed using GraphPad Prism 4.

RESULTS
Analysis of CFTR transcripts in patient cells carrying the R553X mutation
We have performed transcript analysis of five patients with cystic fibrosis who are compound heterozygotic for ΔF508 (1652del3, exon 10) and R553X (1789C→T) mutations. These patient cell lines were used as the ΔF508 transcript has been shown to be stable,\textsuperscript{11} and can be distinguished from transcripts resulting from the R553X allele. Lymphoblastoid cell lines were available, and we and others previously showed that the splicing patterns of CFTR in these cell lines reflect those observed in airway epithelia.\textsuperscript{11 29} Accordingly, total RNA was extracted and a fragment spanning CFTR exons 10, 11 and 12 was analysed by reverse transcriptase-polymerase chain reaction (RT-PCR) assays. Variable levels of an aberrantly spliced form that corresponded to the skipping of exon 11 was observed in all five patients with cystic fibrosis that was absent in all control cell lines studied (fig 1A). Two of the three previous studies have shown that no transcript is generated from the R533X allele; however, it should be noted that for one\textsuperscript{29} the assay used would not have permitted the detection of the exon 11 transcript and the second,\textsuperscript{11} did not use a sensitive assay. The results from our study are consistent with the third study\textsuperscript{29} that also showed that the R553X mutation causes skipping of CFTR exon 11.

We assessed whether the skipped transcript (–exon 11) generated from the R553X allele arises from nonsense-associated altered splicing (NAS) and whether it is targeted by the NMD pathway, as the exclusion of exon 11 results in a frameshift that introduces a new PTC in exon 12. For this purpose, the patient with cystic fibrosis and control cell lines was treated with cycloheximide for 4 h before harvesting.

Cycloheximide, a protein synthesis inhibitor, has been reported to inhibit NAS\textsuperscript{27} as well as NMD.\textsuperscript{30} If the –exon 11 transcript was generated by NAS, a relative reduction of this transcript

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would be found in the presence of cycloheximide. If there were no NAS, both the + exon 11 and – exon 11 transcripts would be expected to increase in the presence of cycloheximide owing to suppression of NMD. Results of transcript analysis showed an increase in the overall levels for both the + exon 11 and – exon 11 transcripts as compared with endogenous GAPDH transcript levels (fig 1A). Moreover, the expression of the – exon 11 transcript was apparently not reduced on cycloheximide treatment. Although it should be kept in mind that the efficiency with which cycloheximide blocks NAS and NMD processes may vary, these results indicate that NAS is not likely to be the underlying mechanism for exon skipping.

Subsequently, RT-PCR products spanning exons 10–12 from cycloheximide-treated and untreated cell lines derived from patient 2 (fig 1A) were sequenced. Results confirmed that the full-length transcript was generated from the allele carrying the ΔF508 mutation, although the transcript lacking exon 11 resulted exclusively from the R553X allele. In either cycloheximide-treated or untreated cells of patient 2 (fig 1B) the R553X allele was specifically aberrantly spliced. We cannot exclude the possibility that low amounts of the R553X allele may lead to + exon 11 transcript owing to limitations of detection by sequencing analysis. Together, these results suggest that the primary consequence of the R553X mutation is the skipping of CFTR exon 11 and that it is not likely to be due to NAS.

**Creation of a putative exonic splicing silencer element by the R553X mutation**

To elucidate more precisely the mechanism underlying R553X mutation-associated exon skipping, a heterologous splicing reporter construct was used. The reporter system was chosen such that it allowed the assessment of sequences that promoted (ESE) or hindered (ESS) exon inclusion. Briefly, it consisted of three exons: two constant exons derived from the human insulin gene that flank the alternatively spliced exon 5 (v5) of the CD44 gene. In a previous study, two ESSs (L and M) and one ESE (R) were defined by mutagenesis within the CD44 v5 exon (fig 2A). Typically, the intact exon v5 is efficiently skipped, but removal of either of the two ESSs sequences (ΔL and ΔM) increases the level of inclusion such that approximately 50% of the total CD44 reporter transcripts contain the alternatively spliced exon. Splicing reporter cassettes were generated to assess whether the R553X allele caused exon skipping by disrupting an ESE or by creating an ESS independently of other CFTR transcript features.

A region comprising 17 nucleotides surrounding the wild-type (GAG GTC AAC GAG CAA GA) and R553X (GAG GTC AAT GAG CAA GA) alleles of CFTR exon 11 was introduced into the 3’ region of the CD44 reporter exon that lacked the first ESS element (ΔL). This construct maintained the position of the R553X mutation with respect to its distance from the 5’ splice site in CFTR exon 11 (fig 2B). The resulting constructs ΔL3’11WT and ΔL3’11R553X in addition to the parental construct (ΔL) were transfected into HEK 293 and IB3 cell lines. Analyses of transcripts showed that there is a relative increase in exon v5 skipping for ΔL3’11R553X transcripts, compared with ΔL3’11WT transcripts (fig 2C, compare lanes 2 and 3 with 4 and 5). The ratios of exon inclusion/exclusion from the wild-type and mutant constructs were found to be significantly different (n = 4, p = 0.014; Mann–Whitney U test; fig 2D), suggesting that the sequence carrying the R553X mutation may act as a silencer element. It is interesting to note, however, that both constructs containing CFTR sequences (wild type and R553X) showed a dramatic increase in exon skipping compared with the parental construct (ΔL; fig 2D). The 17-nucleotide wild-type sequence may act as a negative element on the splicing of the reporter exon, although the observed effect probably reflects the disruption of the ESE (R) by the CFTR sequences (fig 2B).

To evaluate these possibilities, we constructed additional reporter cassettes in which the ESE (R) was specifically deleted, maintaining the position of the CFTR wild-type or the R553X mutation-containing sequences with respect to the 3’ end of the exon (AR3’11WT or AR3’11R553X, respectively, fig 2B). These constructs together with the parental construct (ΔR) were transfected into HEK 293 cells. The results suggest that the observed increase in exon skipping resulting from the ΔL3’11WT construct with respect to the parental ΔL construct
A schematic representation of the splicing reporter construct. Analysis of the consequences of the R553X mutation in a heterologous context indicates that it confers exonic splicing silencer (ESS) activity. (A) Schematic representation of the splicing reporter construct. In, 2 and 3 = insulin exons 2 and 3, CD44V5 = exon v5 of the CD44 gene. (B) Schematic representation of the AL and AR constructs, in which 17 nucleotide sequences flanking the wild-type allele (nt C) or the R553X allele (nt T) were introduced in the 3' segment of the CD44 exon v5, in the absence of ESS1 (L) or ESE (R) segments. The resulting constructs were named ΔL3 11WT and ΔL3 11R553X and ΔR3 11R553X, respectively. The blackened AL or AR boxes denote the substitution of each of these fragments of the CD44 exon v5 with BlueScript sequences as described elsewhere.

DISCUSSION

Skipping of exon 11 caused by the R553X mutation through the creation of a position-dependent ESS

To date, more than 146 nonsense mutations identified in the CFTR gene have been associated with severe cystic fibrosis disease presentation (http://www.genet.sickkids.on.ca/cftr/). Studies have shown that PTCs elicit nonsense-mediated mRNA decay, a quality control mechanism that prevents the expression of truncated proteins. There are examples, however, where nonsense mutations are associated with exon skipping. In the present study, we have investigated the CFTR R553X mutation at the transcript level. Exon 11 skipping in patients with cystic fibrosis that are compound heterozygotic for ΔF508/R553X mutation was specifically associated with the R553X mutation. The extent of exon skipping was variable but
located in the action of ESSs results in the silencing of CD44 exon v5. The resulting constructs were named ΔLS’11WT and ΔM 11WT or ΔLS’11R553X and ΔM11R553X, respectively. ins 2 and 3 = insulin exons 2 and 3, CD44v5 = exon 5 of the CD44 gene. ΔL or ΔM in black boxes denote the substitution of the L (exonic splicing silencer (ESS)1) or M (ESS2) segments of the CD44 exon v5, respectively, with BlueScript sequences as described elsewhere. (B) Reverse transcriptase-polymerase chain reaction (RT-PCR) analysis of human embryonic kidney (HEK) 293 cell lines transfected with reporter constructs ΔL (lane 1), ΔLS’11WT (lane 2), ΔLS’11R553X (lane 3), ΔM (lane 4), ΔM11WT (lane 5) and ΔM11R553X (lane 6), respectively. PCR products were separated in a 2% agarose gel. CD44v5 = full-length transcript, –CD44v5 = transcript lacking CD44 exon 5. (C) Measurement of the resulting PCR bands was carried out as described in fig 2D. The bars correspond to the means and standard error calculated for the ratios from two independent transfection experiments. The white bar corresponds to the ΔL or ΔM, the light grey bars correspond to ΔLS’11WT and ΔM11WT and the black bars correspond to ΔLS’11R553X and ΔM11R553X transcripts.

Figure 3 Position effect of the R553X mutation. (A) Schematic representation of the CD44V5 ΔL and CD44V5 ΔM constructs described elsewhere, in which 17 nucleotide sequences flanking the wild-type allele (nt C) or the R553X allele (nt T) were introduced at the 5’ and in the middle segments of CD44 exon v5. We further investigated the underlying cause of the R553X mutation-associated exon 11 skipping and showed that 17 nucleotides of exon 11 carrying the R553X mutation had a negative effect on splicing in a heterologous reporter system (CD44 v5 exon), compared with the wild-type allele. This effect was only apparent when the R553X mutation was positioned proximal to the 3’ end of the exon, as occurs in CFTR exon 11. The wild-type allele did not have a positive effect on the splicing of CD44 exon 5, indicating that the R553X mutation did not affect an ESE. Thus far, previous studies have shown that nonsense mutations could affect splicing by disrupting ESEs. Other reports have implicated a quality control mechanism known as nonsense-associated altered splicing, as the underlying cause of nonsense mutation-related exon skipping. Our study provides the first evidence for a nonsense mutation creating an ESS, and further shows that this putative ESS element may result in exon skipping by preventing selection of the proximal 5’ splice site. Interestingly, the sequence created by the R553X mutation partially resembles the consensus sequence of an ESS previously identified by Zhang and Chasin. A recent study has shown that the mode of action of ESSs results in the silencing of 5’ splice sites that are located 3’ of the ESS sequences. Although Wang et al also showed that several such sequences resulted in the silencing of 3’ splice sites located 5’ of the ESSs (in addition to the silencing of the 5’ splice sites), one of them (hexamer “A’A’, fig 1D) only influenced the 5’ splice site. The putative ESS sequence created by the R553X mutation seems to represent an additional example of this latter mode of action of ESSs.

Therapeutic implications

Our findings have implications for proposed treatments for nonsense mutations involving aminoglycoside antibiotics that were shown to promote read-through of PTCs. Clinical trials using gentamicin administered to patients with cystic fibrosis carrying the W1282X mutation showed improvement of nasal potential difference measurements and increased CFTR at the cell membrane, compared with patients with cystic fibrosis with non-nonsense mutations. However, our findings suggest that nonsense mutations, such as CFTR R553X, which cause exon skipping or are targeted by NMD, may not be good candidates for such treatment. The results thus emphasise the importance of determining the specific consequences of nonsense mutations at the transcript level.

Molecular treatments directed against splicing defects are being tested using various methods such as modified RNA oligonucleotides, molecular chimeras, splicing factors and chemical compounds. It was recently shown that 2’-O-methyl antisense oligonucleotides directed against the Duchenne muscular dystrophy gene systemically delivered to the dystrophic mdx mouse caused the skipping of an exon carrying a nonsense mutation, leading to an improvement in muscle function. These results suggest that 2’OMeAOs could also be used to treat patients with splicing defects. Similarly, it may be possible to treat the splicing defect caused by the R553X mutation using a 2’OMeAO directed against the region where the mutation is located in CFTR exon 11, to prevent the binding of splicing factors responsible for exon 11 skipping. The correction would lead to the presence of CFTR transcripts carrying exon 11 generated from the R553X allele, and provided it is not entirely degraded by NMD, it could then be treated using aminoglycoside antibiotics.

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Key points

- The third most common nonsense mutation in the cystic fibrosis transmembrane conductance regulator gene, R553X (1789C→T), was found to cause exon 11 skipping in cell lines from patients with cystic fibrosis.
- The R553X mutation-associated exon skipping did not seem to result from nonsense-associated altered splicing.
- The R553X mutation caused exon skipping through the creation of a position-dependent exonic splicing silencer.
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