Transcriptional mutagenesis reduces splicing fidelity in mammalian cells

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ABSTRACT

Splicing fidelity is essential to the maintenance of cellular functions and viability, and mutations or natural variations in pre-mRNA sequences and consequent alteration of splicing have been implicated in the etiology and progression of numerous diseases. The extent to which transcriptional errors or lesion-induced transcriptional mutagenesis (TM) influences splicing fidelity is not currently known. To investigate this, we employed site-specific DNA lesions on the transcribed strand of a minigene splicing reporter in normal mammalian cells. These were the common mutagenic lesions O6-methylguanine (O6-meG) and 8-oxoguanine (8-oxoG). The minigene splicing reporters were derived from lamin A (LMNA) and proteolipid protein 1 (PLP1), both with known links to human diseases that result from deregulated splicing. In cells with active DNA repair, 1–4% misincorporation occurred opposite the lesions, which increased to 20–40% when repair was compromised. Furthermore, our results reveal that TM at a splice site significantly reduces in vivo splicing fidelity, thereby changing the relative expression of alternative splicing forms in mammalian cells. These findings suggest that splicing defects caused by transcriptional errors can potentially lead to phenotypic cellular changes and increased susceptibility to the development of disease.

INTRODUCTION

Splicing of pre-mRNA resulting from eukaryotic genes involves specific sequence elements both at the exon–intron boundaries and within introns, including the 5′ and 3′ splice sites preceded by the branch-point sequence and polypyrimidine tract (1). Alternative splicing results in the generation of multiple transcripts from a single gene, thereby expanding the human proteome considerably (2,3). High-throughput sequencing and genome-wide analysis have revealed that >90% of all pre-mRNAs from multi-exon human genes undergo alternative splicing (4,5) and the fidelity and regulation of this process are essential to the control of cellular and organ functions in eukaryotes. Indeed, mutations or natural variations in pre-mRNA sequences that alter splicing have been implicated in the etiology and progression of numerous pathologies, including genetic diseases, neurodegenerative disorders and cancer (6,7).

RNA polymerase II (RNA pol II) plays a central role in the synthesis of pre-mRNA and non-coding RNA in eukaryotic cells (8). Maintenance of the viability of mammalian cells, most of which are quiescent or replicate slowly (9,10), depends strongly on the fidelity of transcription. To ensure a high accuracy fidelity checkpoints are utilized at the steps of insertion, extension and proofreading during the transcriptional elongation of RNA pol II (11). In Saccharomyces cerevisiae, several alleles are associated with error-prone transcription and some of these also reduce splicing efficiency and induce phenotypic changes resembling human diseases (12–14). Notably, damaged nucleotides or lesions in DNA can also reduce the fidelity of RNA pol II significantly, thereby introducing transcriptional errors.

Lesion-specific processing of DNA damage by RNA pol II influences RNA synthesis in several ways, causing transcriptional arrest and activating transcription-coupled repair (TCR) or transcriptional bypass, which can potentially result in the incorporation of incorrect bases into nascent RNA, an event referred to as transcriptional mutagenesis (TM) (15–18). The capacity of a DNA lesion to impede transcription depends on several variables, but bulky adducts and distorting DNA cross-links are in general very efficient. Transcriptional bypass of bulky, UV-induced lesions by RNA pol II attenuates activation of TCR, allowing cells to complete ongoing mRNA synthesis and maintain protein homeostasis, thereby promoting resistance to DNA damage and cell survival (19,20). Several commonly occurring, non-bulky DNA lesions, such as O6-methylguanine (O6-meG) and 8-oxoguanine (8-oxoG), do not pose a strong hurdle to RNA pol II, allowing transcriptional bypass and misincorporation both in vitro and in vivo (16,17).

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When TM occurs, each round of transcription may produce mRNA with single-base mutations that are translated into a relatively large population of mutated or misfolded proteins with altered functions, sometimes leading to phenotypic changes with significant consequences on cellular regulation (21–25). These altered proteins could also have dominant properties leading to a more prolonged or permanent physiological change (26). Here, we tested the hypothesis that even if a DNA lesion leads to TM which does not alter coding specificity (e.g. intronic mutations, silent mutations), translation might be altered if a regulatory sequence at a splicing site is altered. The capacity of such transcriptional errors to alter splicing fidelity has been proposed before (27), but never examined. To accomplish this, we combined the well-established site-specific positioning of DNA lesions (28) and minigene splice reporters (29) designed to reflect the influence of these mutations on splicing. Our results show that when TM involves a splice site in vivo, splicing fidelity is attenuated, thereby altering the relative expression of alternatively spliced forms.

MATERIALS AND METHODS

Construction and modification of minigene splicing reporters

The LMNA and PLP1 minigene splicing reporters were constructed on the basis of previous work (30,31) and adapted to the present study. These reporters were cloned into the pNEW-GFP vector and subsequently modified in order to obtain the following final plasmids: pNEW-LMNA-Rep-GFP WT, pNEW-LMNA-Rep-GFP 1824C>T, pNEW-PLP1-Rep-GFP WT and pNEW-PLP1-Rep-GFP 347C>A. Generation of closed circular vectors containing site-specific base modification (pNEW-LMNA-Rep-GFP O6-meG and pNEW-PLP1-Rep-GFP 8-oxoG) involved production of ssDNA, in vitro synthesis of the complementary strand modified at the specific site, and purification of closed circular dsDNA, as described previously (22,28). All DNA vectors were sequenced for confirmation. For a more detailed description of this procedure, please see the Supplementary data. The splice-site scores were calculated, which showed low levels of correlation (Supplementary Table S1; the PCR products were solved on a 3% agarose gel as shown in Supplementary Figure S5; and colonies that amplified with the mutant-specific primers for LMNA and PLP1 (see Supplementary Table S1). The gel-purified PCR products (GeneJet Purification kit, Thermo Scientific) were cloned into a pcDNA3 vector, digested with HindIII-KpnI and subsequently sequenced. In the case of the LMNA reporter, 96 individual colonies (24 for each one of the four replicates) were sequenced. With the PLP1 reporter, pre-screening was performed utilizing tetra-primer ARMS-PCR (32). In brief, colony PCR was performed on 100–150 clones (from four replicates) using the primers as indicated in Supplementary Table S1; the PCR products were resolved on a 3% agarose gel as shown in Supplementary Figure S5; and colonies that amplified with the mutant-specific primers propagated for plasmid isolation and subsequent confirmation by sequencing. Differences in levels of TM were analyzed with a two-tailed unpaired t-test ($P < 0.05$) using GraphPad Prism 6.0 software.

Flow cytometry

Cells from at least three independent transfections were harvested, re-suspended in PBS and then immediately analyzed for GFP fluorescence on a BD Accuri C6 Flow Cytometer with a FL-1 filter with 99% attenuation. The GFP-positive cells were gated relative to untransfected cells and all the available measures of GFP fluorescence were included in the analysis. First, the intra-repeat correlation in GFP intensity by combination of experiment and condition was calculated, which showed low levels of correlation (Supplementary Table S2). The observed distribution of GFP signal was markedly right-skewed and thick-tailed. Because of that, log10 transformed median GFP intensities were estimated in the statistical analysis. We used quantile regression with GFP intensity as the dependent variable. The independent variables were the indicator variables for the different conditions and for the different replicates. The WT or O6-meG condition and the data from the first replicate were left out of the model as reference groups. We estimated the standard errors of the regression coefficients with the robust sandwich estimator (33). For details see Supplementary Table S3.
Figure 1. Schematic presentation of the minigene splicing reporters containing site-specific DNA lesions with their resulting pre-mRNA. (A) The lamin A (LMNA) reporter was based on the region exon 9–11—intron 11—exon 12 of LMNA. The nucleotide pair C:O6-meG indicated corresponds to the sequence of the damaged plasmid. The WT plasmid contains a C:G pair and the mutated a T:A pair in this same position. (B) The proteolipid protein 1 (PLP1) reporter was based on the region exon 3—intron 4—exon 4 of PLP1. The nucleotide pair C:8-oxoG indicated corresponds to the sequence of the damaged plasmid. The WT plasmid contains a C:G pair and the mutated a A:T pair in this same position. The coding sequence for GFP (EGFP) was associated with the minigenes in a manner such that GFP is expressed only if alternative splicing occurs. Transcription of the minigene splicing reporters was under regulation by the human cytomegalovirus immediate early promoter (PCMV) and terminated by the bovine growth hormone polyadenylation [poly-(A)] signal.

Real-time quantitative and reverse-transcription PCR
RNA and cDNA was purified and synthesized as for mRNA sequencing. Real-time qPCR was performed using Maxima SYBR Green qPCR Master Mix (Thermo Fisher scientific) on an Applied Biosystems 7500 Real-time PCR. Quantification of the WT or progerin/Δ150 splicing forms of lamin A was achieved with the fw and rv primer pairs indicated with the fw and rv primer pairs indicated in Figure 3A using ribosomal protein lateral stalk subunit P0 (RPLP0) as reference gene. Differences in expression of alternative splicing products were analyzed with two-tailed unpaired t-test (P < 0.05) using GraphPad Prism 6.0 software.

These same cDNAs were used to detect splicing isoforms transcribed only from the transfected plasmids by RT-PCR. Specific detection of the WT or alternatively spliced forms of LMNA and PLP1, as well as GFP was achieved with the fw and rv primer pairs indicated in Figures 3A and 5A. RT-PCR amplification was performed with the Dream Taq PCR master mix (Thermo Fisher scientific) as follows: 2 min at 95°C, 35 cycles of 30 s at 95°C, 30 s at 54°C and 45 s at 72°C, followed by a final extension of 5 min at 72°C. The PCR products obtained were resolved on a 2% agarose gel. Densitometry was performed using ImageJ. Differences in expression of alternative splicing products were analyzed with two-way anova (P < 0.05) using GraphPad Prism 6.0 software. All primer sequences are shown in Supplementary Table S1.
Figure 2. Analysis of splicing based on the GFP fluorescence of cells transfected with the LMNA minigene reporter. (A) Representative FSC-H/FL1-H scatter plots for HEK293 cells transfected with the different plasmids as indicated (after exclusion of doublets and dead cells) with GFP-positive cells gated relative to untransfected cells. The total counted events for each condition is shown (mean ± SEM, n = 4). For representative scatter plots for WI-38 cells see Supplementary Figure S2. (B and C) Boxplots of the distributions (95% CI) of fluorescence intensity in HEK293 and WI-38 cells, respectively. Median ****P < 0.0001 in comparison to the WT reporter; median ####P < 0.0001 in comparison to the O6-meG reporter, See also Supplementary Table S3.

RESULTS

Construction of minigene splicing reporters

Two minigene splicing reporters containing site-specific DNA lesions were successfully constructed (Figure 1), as confirmed by sequencing and an Fpg nicking assay (Supplementary Figure S1). The lamin A (LMNA) reporter (Figure 1A) contained O6-meG at codon 608, which during transcription should produce RNA containing either the wild-type (WT) GGC sequence or the mutated GGU due to uridine misincorporation (22,34). This silent mutation at codon 608 in exon 11 (c.1824C>T, p.G608G) increases the usage of a natural splice donor, leading to an in-frame deletion of 150 nucleotides (Δ150) and production of the progerin protein that causes Hutchinson-Gilford progeria syndrome (35,36). The mutation in the LMNA 5′ splice site (CAG'GTGGGC to CAG'GTGGGT) increases the splice-site score from 79.04% to 84.90%.

The proteolipid protein 1 (PLP1) reporter (Figure 1B) contained 8-oxoG in codon 116, which during transcription should produce RNA containing either the WT ACG sequence or the mutated AAG due to adenine misincorporation (21,25). PLP1 can be spliced into two forms (PLP1 and DM20), due to the alternative use of donor splice sites in exon 3. Mutations that reduce the ratio of PLP1 to DM20 (e.g. c.347C>A, p.T116K) result in Pelizaeus-Merzbacher disease, an X-linked leukodystrophy (31,37). The mutation in the PLP1 5′ splice site (ACG'TAACA to
Figure 3. Analysis of splicing in cells transfected with the *LMNA* minigene reporter by qPCR and RT-PCR. (A) Schematic illustration of the primers employed. The two *LMNA* WT and progerin primer pairs were used for qPCR and the *LMNA* progerin-fw/EGFP-rv pair for RT-PCR. (B and C) Quantitative analyses of the effects of *LMNA* mRNA on splicing fidelity in HEK293 and WI-38 cells, respectively, presented as the ratio of progerin mRNA/total *LMNA* mRNA (calculated as \( \frac{R}{1 + R} \) where \( R = \text{progerin} / \text{WT *LMNA} \) (mean ± SEM, \( n = 4 \)). (D and E) Agarose gels of RT-PCR products illustrating the effects of TM on splicing fidelity of *LMNA* mRNA in HEK293 and WI-38 cells, respectively. The two splicing forms WT and Δ150 are indicated by arrows. **\( P < 0.01 \); ***\( P < 0.001 \) in comparison with the WT or \( O^6\)-meG reporter.

**AAG GTAACA** elevates the splice-site score from 72.80% to 82.01%.

To allow detection of alternative splicing, the minigenes were associated with an open reading frame GFP (EGFP) in a manner such that GFP is expressed only if alternative splicing occurs (Figure 1). In that event, a STOP codon inserted at the end of the exon containing the donor splicing site disappears from the processed mRNA, allowing for the exon encoding the acceptor splicing site and GFP to be expressed. Along with the plasmid containing a site-specific DNA lesion a control (WT sequence) and mutant (1824C>T or 347C>A) plasmid were synthesized for each reporter. These plasmids contain no known mammalian origin of replication, thereby excluding effects of replication (22).

\( O^6\)-meG and 8-oxoG induce transcriptional mutagenesis in mammalian cells

To confirm that \( O^6\)-meG and 8-oxoG induce TM, cellular transcripts of these modified reporters were analyzed by sequencing of RT-PCR products. In addition, the impact of DNA repair was assessed by treating HEK293 cells with \( O^6\)-benzylguanine (\( O^6\)-bzG), an inhibitor of the repair of methylated bases by alkylguanine transferase (AGT) (22,38) or by using primary mouse embryonic fibroblasts (MEFs) deficient in 8-oxoG DNA glycosylase (\( OGG^1\)−) (21,39).
In HEK293 cells transfected with the reporter containing O6-meG uridine misincorporation opposite the lesion was 3.6 ± 1.2% (mean ± SEM, n = 4), a value that rose to 20.1 ± 5.6% (n = 4, P < 0.05) when AGT was inhibited. In the case of MEFs transfected with the reporter containing 8-oxoG adenine misincorporation opposite the lesion in WT and OGG1−/− cells was 4.0 ± 1.2% (n = 4) and 39.5 ± 2.8% (n = 4, P < 0.0001), respectively. In HEK293 cells, presence of 8-oxoG resulted in 1.2 ± 0.9% (n = 3) TM. No other misincorporations or indels were detected. In agreement with previous studies in vivo these data demonstrate that the presence of O6-meG or 8-oxoG on the transcribed strand of a gene induces TM which, if left unrepaired, can be highly mutagenic (21–23).

Transcriptional mutagenesis reduces splicing fidelity

To determine whether TM induced by O6-meG or 8-oxoG influenced splicing fidelity, changes in GFP fluorescence were measured by flow cytometry (Figures 2A and 4A; Supplementary Figures S2 and S3). Transfection with the LMNA reporter containing O6-meG resulted in an increase of fluorescence intensity compared to WT in HEK293 cells (P < 0.0001), but not in WI-38 fibroblasts (Figure 2B and C). However, when AGT-mediated repair was inhibited, a potent increase in the fluorescence signal was detected in comparison to both the WT and O6-meG reporters and for both HEK293 and WI-38 cells (P < 0.0001), in agreement with the more frequent TM in the presence of impaired AGT. As expected, the fluorescence of both cell types was higher when transfected with the 1824C>T mutant reporter than with the WT reporter (P < 0.0001, Figure 2B and C).

The impact of TM on splicing was examined further by quantitative and qualitative PCR analyses using primers specific for each of the splicing forms of LMNA mRNA (WT and Δ150/progerin) (Figure 3A). While the qPCR primers also amplified cellular transcripts, the RT-PCR primers were specific for reporter transcripts. Both in the presence (3.6- and 4.8-fold, respectively, P < 0.01 and P < 0.001) and absence (10- and 7.4-fold, respectively, P < 0.001) of active DNA repair, HEK293 and WI-38 cells transfected with the O6-meG reporter demonstrated higher expression of the Δ150/progerin form than with the WT reporter (Figure 3B and C). Moreover, in both cell types transfected with the O6-meG this expression was higher in cells with impaired AGT than in cells with active repair (P < 0.001 and P < 0.01). This quantitative analysis was confirmed by RT-PCR (Figure 3D and E), which also showed that O6-bzG did not affect splicing of WT LMNA (Supplementary Figure S4). As in the case of the fluorescence-based assay, the more extensive alternative splicing in the absence of AGT was in accordance with the higher levels of TM. The observed expression levels of the Δ150/progerin spliced form in the untransfected cells and in the WT LMNA transfected cells (Figure 3B and C) are in agreement with previous studies (40).

In the case of the PLP1 reporter, the effects of TM on splicing fidelity were not as evident, primarily because the two splicing forms PLP1 and DM20 are constitutively expressed, so that such effects will be reflected in the ratio between the two forms. In HEK293 cells and both WT and OGG1−/− MEFs transfected with the 347C>A mutant reporter, the increase in fluorescent intensity was only somewhat but nonetheless significantly higher than with the WT reporter (P < 0.0001, Figure 4B–D), as was the increase in the fluorescent intensity of HEK293 cells transfected with the 8-oxoG reporter (P < 0.0001, Figure 4B). An increase was also observed in OGG1−/− MEFs transfected with the 8-oxoG reporter (P < 0.0001, Figure 4D) but not in WT MEFs (Figure 4C), in agreement with the impact of OGG1 status on 8-oxoG-induced TM.

RT-PCR analyses were performed to further confirm the impact of TM on splicing fidelity of PLP1 (Figure 5A and B). Reproducible quantitative PCR analysis was not successful, although extensive method development was performed. Instead, analysis of band intensities from the PCR of the PLP1 splice variant was used to estimate changes in splicing fidelity due to the presence of 8-oxoG on the transcribed strand in MEFs (Figure 5C). In OGG1−/− MEFs a reduction of the PLP1 variant (P < 0.001) was observed but not in the WT MEFs, supporting the results based on GFP analysis and confirming the role of DNA repair in reducing TM and thereby maintaining the fidelity of splicing. Together, these results demonstrate that TM in a sequence which regulates splicing can impair splicing fidelity.

DISCUSSION

Here, we show that the presence of unrepaired DNA lesions on the transcribed strand in mammalian cells can cause TM and subsequently reduce the fidelity of mRNA splicing. Employing site-specific mutagenic DNA lesions on the transcribed strand of a reporter vector, we demonstrate that when O6-meG and 8-oxoG are bypassed by RNA pol II in vivo misincorporation opposite the lesion occurs, producing mutant transcripts. The established importance of DNA repair in preventing TM (21,22,24,41,42) was also evident here. In the case of O6-meG, the frequency of TM in HEK293 increased from 3.6% to 20.1% when AGT was inhibited and the corresponding values for 8-oxoG were from 4.0% to 39.5% when OGG1 was not expressed in MEFs, clearly showing that if left unrepaired these lesions can be highly mutagenic during transcription. Differences in the sequence flanking the DNA lesion, the relative distance between this lesion and the promoter may explain the small differences in the TM frequency observed here and in previous in vivo studies (21–23).

The occurrence, regulation and consequences of transcriptional errors, both spontaneous and due to lesions, have been receiving more and more attention, much due to the information obtainable by next-generation sequencing (11,12,14,42,43). One important aspect for understanding the potential biological significance of transcriptional errors, spontaneous or lesion-induced, is the rate at which such errors occur. In the absence of any known genotoxic insult the transcription error rate in eukaryotic cells is approximately 10−5 (14,44,45) and the frequency observed here and by others is three-to-four orders of magnitude higher, depending on the status of DNA repair. If individual transcripts can be translated between 40 and 4000 times (46,47), a single error can result in a considerable amount of erro-
neous protein, thereby changing the cellular phenotype. Indeed, since in certain cell types the average level of many mRNAs is less than one per cell (48,49), an error in this molecule may mean that all newly translated protein is abnormal.

All of the relatively few investigations that have examined the biological effects of lesion-induced transcriptional errors in mammalian systems to date reveal that TM can cause phenotypic changes. The TM caused by the presence of 8-oxoG or 5-hydroxymethyluracil at specific codons of HRAS or KRAS, respectively, in MEFs lacking OGG1 or NEIL1/2 increases the expression levels of mutated and constitutively active Ras proteins, giving rise to sustained downstream oncogenic signaling (21,50). Similarly, the presence of 8-oxoG on the transcribed strand of a gene encoding luciferase in MEFs deficient for OGG1 impaired the function of this protein due to TM (23). With a fluorescent-based reporter system, Burns and colleagues (22) found that O6-meG-induced TM in HEK293 cells led to considerably lower levels of mutated protein with altered function, an effect which was further increased when AGT was impaired. This is in agreement with our observation that during transcription in HEK293 cells with intact repair capacity O6-meG is more mutagenic than 8-oxoG. This observation might reflect dif-

Figure 4. Analysis of splicing based on the GFP fluorescence of cells transfected with the PLP1 minigene reporter. (A) Representative FSC-H/FL1-H scatter plots obtained for HEK293 cells transfected with the different plasmids as indicated (after exclusion of doublets and dead cells) with GFP-positive cells gated relative to untransfected cells. The total counted events for each condition is shown (mean ± SEM, n = 3). For representative scatter plots for the WT and OGG1−/− MEFs see Supplementary Figure S3. (B–D) Box plots of the distributions (95% CI) of the fluorescence intensity in HEK293 cells, WT and OGG1−/− MEFs, respectively. Median ****P < 0.0001 in comparison with the WT reporter. See also Supplementary Table S3.
Figure 5. Analysis of splicing in cells transfected with the PLP1 minigene reporter by RT-PCR. (A) Schematic illustration of the primers employed. (B) Agarose gel of RT-PCR products showing the effects of TM on splicing fidelity of PLP1 mRNA in HEK293 cells, WT and OGG1−/− MEFs, respectively. The two splicing forms, PLP1 and DM20, and GFP are indicated by arrows. The top gels in B show RT-PCR with primers PLP1+DM20fw and EGFP2rv which amplifies both splice forms. Presence of PLP1 is seen as a weaker band just above the stronger DM20 band. To facilitate a semi-quantitative analysis, a PLP1-specific PCR was performed (middle gels, primers PLP1fw and EGFP2rv). Expression of GFP is shown in the bottom gels (primers EGFPfw and EGFPrv). The faint band observed in the untransfected (UT) lane in HEK293 cells is most likely an unspecific PCR product since any cross contamination would result in PCR product with the other primer pairs as well. (C) Densitometric analysis of PLP1 expression in WT and OGG1−/− MEFs using the co-expressed GFP as a loading control (mean ± SEM, n = 3–4). ***P < 0.001; ****P < 0.0001 in comparison with the WT reporter.

Genetic variation within splice sites and regulatory sequences frequently results in the aberrant splicing associated with human hereditary diseases and cancer. Single-nucleotide substitutions in the 5′ or the 3′ splice sites are the most common splicing mutations, resulting either in exon skipping, activation of a cryptic splice site or, to a lesser extent, intron retention. The pronounced fidelity of splice-site pairing by the spliceosome suggests that splicing fidelity is limited by the fidelity of transcription (51). Moreover, there is an inverse relationship in vivo between the speed of transcription elongation by RNA pol II and splicing efficiency.
Clearly, transcription and splicing fidelity are tightly coupled. Here, we demonstrate that induction of TM at a 5′ donor splice site by DNA lesions reduces splicing fidelity, leading to aberrant expression of alternative splicing forms. The impact of TM on splicing fidelity was in general more evident when analyzed on the level of transcripts compared to the GFP-based analysis which also is dependent on translation, protein folding etc., which might affect the quantitative relationship between % TM and GFP readout. This finding extends an earlier study, based on published RNA-seq data from HeLa and Huh7 cells which found enriched transcriptional errors at the 5′ donor site relative to errors at other positions in the human genome, indicating a consequent increase in intron retention during splicing (14). Our results were observed in cells with active DNA repair in response to both lesions employed here, and further emphasized when the protein mainly responsible for repair was inactivated or missing. In agreement with the overall essential role of DNA repair in maintaining genomic integrity, our results show that through its important role in protecting against TM, DNA repair status may also exert a pronounced impact on splicing fidelity.

In conclusion, bypass of mutagenic lesions by RNA pol II, proposed to be the most specific sensor of DNA damage (52), can result in considerable levels of mutant transcripts and the subsequent expression of mutant proteins can lead to phenotypic changes. Here, we show for the first time that TM can reduce splicing fidelity in vivo, thereby contributing to the production of altered proteins and/or altering the relative levels of alternatively spliced forms, potentially disturbing cellular homeostasis and triggering numerous diseases. Generation of specific splicing isoforms has recently been found to drive cancer (53,54) and the possibility that such isoforms are expressed as a consequence of lesion-induced TM emphasizes the potential significance of this phenomenon in connection with processes such as carcinogenesis and ageing.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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