Expression of progerin does not result in an increased mutation rate

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Abstract In the premature ageing disease Hutchinson-Gilford progeria syndrome (HGPS), the underlying genetic defect in the lamin A gene leads to accumulation at the nuclear lamina of progerin—a mutant form of lamin A that cannot be correctly processed. This has been reported to result in defects in the DNA damage response and in DNA repair, leading to the hypothesis that, as in normal ageing and in other progeroid syndromes caused by mutation of genes of the DNA repair and DNA damage response pathways, increased DNA damage may be responsible for the premature ageing phenotypes in HGPS patients. However, this hypothesis is based upon the study of markers of the DNA damage response, rather than measurement of DNA damage per se or the consequences of unrepaired DNA damage—mutation. Here, using a mutation reporter cell line, we directly compared the inherent and induced mutation rates in cells expressing wild-type lamin A or progerin. We find no evidence for an elevated mutation rate in progerin-expressing cells. We conclude that the cellular defect in HGPS cells does not lie in the repair of DNA damage per se.

Keywords DNA damage · Lamin A · nuclear lamina · progeria

Introduction

Hutchinson-Gilford progeria syndrome (HGPS) is a dominant severe premature ageing genetic disease characterised by the appearance in childhood of age-related symptoms such as hair loss, thin skin, vascular defects and atherosclerosis. In the majority of cases, the underlying genetic defect is a point mutation that results in activation of a cryptic splice site in exon 11 of lamin A (LMNA) (Eriksson et al., 2003). This leads to accumulation of lamin A protein lacking the 50 amino acid (a.a.) domain toward the C-terminus (LAΔ50/progerin) that is required for the final endoproteolytic cleavage step during the normal processing of prelamin A to lamin A. As a consequence, LAΔ50 remains permanently farnesylated (Cao et al., 2007; Dechat et al., 2007). Mutation of the FACE (Zmpste 24) metalloproteinase, which catalyses the final cleavage step of lamin A processing, results in restrictive dermopathy (RD) that also has progeroid features (Navarro et al., 2004). It is the effect of the aberrantly farnesylated prelamin A proteins on nuclear architecture and nuclear function that is presumed to be responsible for the devastating phenotypes of these progeroid diseases (Mehta et al., 2011; Yang et al., 2011).

At the cellular level, progerin appears to incorporate into the lamina at the nuclear periphery (Eriksson et al., 2003; Goldman et al., 2004). In usual cell culture conditions, the resultant nuclei often develop an abnormal and irregular shape with a thickened lamina, especially as the cells reach later passages. There are also various reports of altered distribution of lamin B (Scaffidi and...
Misteli, 2005), nuclear pores and inner membrane proteins in lamin A mutant cells (Dechat et al., 2007; Scaffidi and Misteli, 2005).

A loss of peripheral electron-dense heterochromatin is seen by electron microscopy (e.m.) in later-passage (p26) HGPS fibroblasts (Goldman et al., 2004). Globally, most HGPS fibroblasts appear to have reduced levels of the heterochromatin protein HP1α and histone modifications (H3K9me3 and H3K27me3) associated with heterochromatic states (Dechat et al., 2008; Scaffidi and Misteli, 2005; Shumaker et al., 2006). But genomic profiling in HGPS cells suggests a more complex redistribution of H3K27me3 rather than a simple loss (McCord et al., 2013). H3K9me3 levels in Zmpste 24-deficient cells seem to depend on the passage number (Liu et al., 2013). These chromatin phenotypes are recapitulated in wild-type cells which ectopically express LAΔ50 (Goldman et al., 2004), but they are not rescued in HGPS cells transfected with wild-type lamin A (Scaffidi and Misteli, 2005). These data serve to underpin the idea that LAΔ50 acts as a dominant negative, or gain of function, mutation. Indeed, the presence of progerin may interfere with processing of wild-type lamin A (Goldman et al., 2004).

The presence of progerin or unprocessed prelamin A, or indeed the accumulation of farnesylated prelamin A in Zmpste24−/− cells, has been reported to lead to defects in the DNA damage response and in DNA repair and, as a consequence, an increase in DNA damage (Ghosh et al., 2015; Liu et al., 2005; Liu et al., 2006; Liu et al., 2008; Manju et al., 2006; Sedelnikova et al., 2004). This has led to the hypothesis that an accumulation of unrepaired DNA damage may be responsible for the premature ageing phenotypes in HGPS, RD and FACE/Zmpste24 mutants (Agarwal et al., 2003; Bergo et al., 2002). This is consistent with the idea that defective DNA repair and increased DNA damage are causally related to both other progeroid syndromes in which known genes of the DNA repair and DNA damage response pathways are mutated (Kudlow et al., 2007; Niedernhofer et al., 2006; Wijnhoven et al., 2005), and also to normal ageing (Garinis et al., 2008; Musich and Zou, 2009).

How lamin A mutation and lamin A processing defects might result in defective DNA repair is not clear. Because progerin seems to have an increased association with the nuclear lamina and a decreased association with internal lamin A foci, it could sequester replication proteins away from these internal sites (Barbi et al., 2004) leading to stalled and then collapsed replication forks (Barbi et al. 2004; Garinis et al., 2008; Misteli and Scaffidi, 2005; Musich and Zou, 2009). Inappropriate sequestration of repair proteins to these sites of DNA breaks may also be involved (Liu et al., 2008). Increased generation of reactive oxygen species (ROS) has also been reported in progeria fibroblasts (Richards et al., 2011). Finally, organisation at the nuclear periphery, especially of heterochromatin, may be important to physically shield the genome from incoming mutagens (Hsu, 1975).

There is a clear need to better understand the nature of any defect in DNA repair in HGPS. However, with the exception of some comet assays, previous studies have relied upon markers of the DNA damage response (e.g. γH2A.X), rather than DNA damage per se, to address this question. The consequence of unrepaired DNA damage is mutation. Here, using a mutation reporter cell line derived from Muta™Mouse (White et al., 2003), we have directly compared the inherent mutation rate in cells expressing wild-type or LAΔ50 lamin A and also the mutation rate induced in these cells by exposure to exogenous mutagens. We find no significant elevation of the mutation rate, scored at the mutation reporter, in LAΔ50/progerin-expressing cells and therefore suggest that underlying the cellular defect in HGPS cells does not lie in defective DNA repair per se.

**Results**

**Expression of wild-type and mutant lamin A in MutaMouse cells**

To investigate direct effects of progerin on the repair of DNA damage we took advantage of a well-established mutation reporter system. In the Muta™Mouse, there are ~30 copies of λgt10lacZ inserted into the mouse genome (Gossen et al., 1989; Shwed et al., 2010) and lacZ then functions as the main target sequence for scoring mutations (Myhr, 1991). A stable epithelial cell line, FE1, was established from these animals and is suitable for measurement of endogenous mutation rates, as well as the rates induced in response to a variety of mutagens (White et al., 2003). To characterise the genomic context of the mutation reporter, we used fluorescence in situ hybridisation (FISH) with a λgt10lacZ probe on metaphase chromosomes from FE1 cells. In each spread, three chromosomes were labelled by the
probe, and analysis of their DAPI-banding pattern suggested that these might be *Mus musculus* chromosomes 3 (MMU3). Combined analysis with a chromosome paint for MMU3 confirmed this (Fig. 1a). We conclude that in the aneuploid FE1 cell line (*n = 78* chromosomes/spread), there has been a triplication of the original λgt10lacZ-containing chromosome.

To determine the effects of progerin expression on the mutation rate, we created FE1 cell lines stably expressing GFP-tagged human lamin A—either wild type (wt) or ΔLA50/progerin (Goldman et al., 2004). Ectopic expression of N-terminally GFP-tagged lamin A has been shown to result in the stable integration of the tagged protein into the lamina at the nuclear periphery (Gilchrist et al., 2004). To ensure stable expression from the transgene, we FACs-sorted GFP-expressing cells. Fluorescence microscopy confirmed that both the wt and ΔLA50 mutant GFP-tagged lamin A concentrate at the nuclear lamina. Optical sectioning along the *z* axis showed that any bright, apparently
Reduced levels of heterochromatic histone modifications, especially H3K9me3, and the heterochromatin protein 1 (HP1α) that binds to this mark, have been reported in HGPS cells (Scaffidi and Misteli, 2005) and in human cells ectopically expressing ΔLA50 (Shumaker et al., 2006). By immunoblotting, we saw a small reduction of H3K9me3 and HP1α levels in late-passage FE1 cells expressing ΔLA50 as compared to cells expressing wild-type lamin A (Fig. 2a), though we did not detect loss of H3K27me3 in the presence of ΔLA50.

Another characteristic of HGPS is the acquisition of aberrant nuclear morphology in late-passage cells. Indeed, under normal cell culture conditions in late-passage ΔLA50-expressing FE1 cells, we observed that the proportion of cells with aberrant nuclear morphology (evidence of nuclear blebbing, nuclear fragmentation) was >2.5-fold higher (16.4%) than in FE-1 cells expressing wild-type lamin A (6.1%) (Fig. 2b). Thus, we conclude that FE-1 transfectants exhibit many of the gross nuclear features reported for HGPS cells and other cells transfected with progerin-expressing constructs and that they therefore represent a suitable model system for studying the effects of ΔLA50 on DNA damage and mutation.

It is known that mouse cells grown in standard tissue culture conditions—i.e. under high oxygen tension (20% O2)—are subject to oxidative stress and have an elevated mutation rate (Busuttil et al., 2003). Since we wanted to study the effects of ΔLA50 on intrinsic and induced mutation rates, we did not want to work against a background of mutations introduced as a consequence of non-physiological culture conditions. Therefore, once established, we maintained the FE-1 cells and the WT and ΔLA50-expressing FE-1 lines under physiological O2 (3%) for multiple passages. The mutation rate of mouse cell lines grown under these conditions is reported to be the same as for primary cells (Busuttil et al., 2003).

Under these conditions, we noted that the proportion of ΔLA50-expressing cells exhibiting aberrant nuclear morphology was markedly reduced (10.5%) relative to the same cells grown in 20% O2 and was only slightly greater than cells expressing wild-type lamin A (Fig. 2b). Thus, it appears that the aberrant nuclear morphology of lamin A mutant cells may be due to their increased sensitivity to the cell stress of abnormally high O2 concentration. This would be consistent with the increased sensitivity to ROS reported for HGPS fibroblasts (Richards et al., 2011).

Intrinsic mutation rate in the presence of wild-type and mutant lamin A

We first assessed the intrinsic spontaneous mutation rate in lamin A FE1 transfectants by recovery and packaging of λgt10-lacZ genomes from the cellular DNAs of parental FE-1 cells, and cells stably expressing wild-type or Δ50 lamin A. Resultant phages containing lacZ mutations were selected for by infection of GalE-E. coli (BIK12001) and plating on minimal agar containing 0.3% w/v phenyl-β-d-galactosidase (PGal) (Gossen and Vignal, 1993; Ino et al., 2005). In wild-type (lacZ+) phage, release of the galactose moiety from PGal by β-galactosidase results in the accumulation of toxic UDP-galactose in GalE− strains. Therefore, only cells infected by lacZ− mutant phage survive and form plaques (Mientjes et al., 1996) (Fig. 3a).

Mutation frequency is then expressed as the ratio of mutant plaques (+PGal plates) to total plaque-forming units (pfu) on non-selective plates. The efficacy of selection was first tested using known wild-type and (L1A15) mutant stocks of λgt10-lacZ phage (Ino et al., 2005). There was a 104-fold drop in plating efficiency on PGal selective plates for the wild type over lacZ− mutant phage, comparable to previous reports using this system (Ino et al., 2005) (Fig. 3b, c).

Mutant frequency from parental FE-1 cells and lamin A transfectants was then assessed at early (<P24) and later (P48–53) passages. The intrinsic mutant frequency of all three cell lines was low (<5.5 × 10−4), and there was no evidence for an increase in transfectants stably expressing mutant lamin A (Δ50) (p = 0.67) (Fig. 3d). Indeed, there was some evidence that the mutation rate in cells expressing wild-type lamin A was slightly elevated relative to both the parental FE-1 cells and the Δ50 cell line, but this was not statistically significant.
Induced mutant rate in the presence of wild-type and mutant lamin A

To determine if cells expressing mutant lamin A were more susceptible to DNA damage induced by exogenous mutagens, we exposed FE-1 and lamin A transfectants to two different mutagens.

We first analysed the effects of exciting, but non-ionising, UV-C (254 nm) radiation (Gazave et al., 2005). This induces cyclobutane pyrimidine dimers (CPDs) and strand breaks (Rapp and Greulich, 2004; Turner et al., 2004). Cells were exposed to 10 J/m² UV 254 nm on two consecutive days and harvested 72 h after the first exposure to mutagen. Efficacy of the mutagen was assessed by measuring the frequency of mutant phage particles recovered from parental FE-1 cells before and after treatment. UV-C elevated the mutant frequency
tenfold \((p < 1 \times 10^{-6})\) (Fig. 4a). UV-C similarly elevated the mutant phage frequency from FE-1 transfectants expressing wild-type or mutant \(\Delta 50\) lamin A. However, as seen for the intrinsic mutation frequencies, and contrary to what was expected from proposed models of HGPS pathogenesis, the mutant frequency in cells expressing wild-type LMNA appeared a bit higher than in cells expressing LA\(\Delta 50\) mutant protein, although overall this was not statistically significant \((p = 0.14)\) (Fig. 4b).

To ascertain whether a similar result could be obtained using a different mutagen with a different mode of action, we treated cells with the alkylating agent ethyl nitrosourea (ENU). ENU induces mainly A:T to T:A transversions, probably due to thymine adducts. We treated FE-1 cells with 0.85 and 1.7 mM (200 \(\mu\)g/ml in DMSO) ENU, since the latter is the dose...
reported to produce a maximal mutant rate of $2 \times 10^{-3}$, i.e. $5 \times$ spontaneous rate, at the lacZ reporter from Muta™Mouse (White et al., 2003). Indeed, we saw a similar increase in mutant frequency, compared to DMSO-treated controls, in our assay (Fig. 4c).

The mutant frequency in lamin A-expressing FE-1 stable transfectants was also elevated by ENU treatment, but as was seen for UV treatment, the ENU-induced mutant rate was higher in the cells expressing wild-type lamin A than those expressing LAΔ50, and this was statistically significant ($p = < 1 \times 10^{-4}$) (Fig. 4d). Hence, by scoring for mutant lacZ in transgenic murine cells, we are unable to find any evidence for an increased rate of unrepaired DNA damage caused by the presence of progerin. If anything, the expression of progerin suppressed the mutation frequency, at least in the case of the mutagen ENU.

No increase in γH2A.X in progerin-expressing cells

An increased appearance of γH2A.X and 53BP1 foci, and defective recruitment of 53BP1 to sites of DNA damage, has been reported in Zmpste24-deficient mouse embryonic fibroblasts (MEFs), in HGPS fibroblasts and in wild-type MEFs expressing unprocessed prelamin A (Liu et al., 2005; Liu et al., 2008), suggesting a perturbed response to DNA damage and compromised DNA repair. To investigate this further, we first showed that the appearance of γH2A.X foci could indeed be induced by exposure of FE-1 cells to oxidative damage ($H_2O_2$) and detected both by immunofluorescence (Fig. 5a) and immunoblotting (Fig. 5c). However, under low-$O_2$ growth conditions, we did not find evidence, by either immunofluorescence (Fig. 5b) or by immunoblotting (Fig. 5d), for elevated levels of γH2A.X in progerin-expressing FE-1 cells. Some
increase in γH2A.X was seen with prolonged passage in culture, but this was more pronounced in cells expressing wild-type lamin A than in those expressing the mutant protein. Finally, to assess whether our failure to detect increased levels of DNA damage was due to growth under 3% O₂, we repeated the analysis in cells grown under high (20%) O₂. As with cells under the lower [O₂], we observed no evidence for increased foci of γH2A.X (Fig. 5c), or 53BP1 (data not shown) in progerin-expressing FE-1 cells compared to cells expressing wild-type Lamin A.

Discussion

There is little doubt that the accumulation of damage to cellular components, including DNA, contributes to normal chronological ageing (Li and Vijg, 2012). Both genome instability and point mutations increase with age (Gorbunova et al., 2007). The premature ageing phenotypes seen in many genetic disorders with mutation of genes involved in DNA metabolism reaffirm that DNA damage and a loss of genome integrity is a key feature of both normal and abnormal ageing (Garinis et al., 2008).

The mechanisms that lead to a premature ageing phenotype due to the dominant LAΔ50 mutation in HGPS are a matter of ongoing debate. However, one plausible mechanism is that there are defects in DNA damage sensing and/or DNA repair (Gonzalo and Kreienkamp, 2015). Studies that have addressed this possible mechanism have examined markers of the DNA damage response and DNA repair pathways. However, compromised sensing or repair of DNA damage should lead to an increase in unrepaired DNA damage—i.e. mutation. This has been inferred (Liu et al., 2008) but not previously directly assessed, in HGPS cells or in cells expressing progerin (LAΔ50).

Here, we have used transgenic reporter cell lines to directly investigate the effects of LAΔ50 mutant lamin A on the spontaneous and induced mutation rate of the phage mutation reporter integrated into the genome of these cell lines (Fig. 1). Such transgenic mutation reporters have been used to reveal the accumulation of mutation load during chronological ageing (Busuttil et al., 2007; Dolle et al., 1997; Dolle et al., 2000) but, to our knowledge, have not previously been used to assess the role of nuclear structure on DNA damage.

We found no evidence for an increase in the spontaneous mutation rate (Fig. 3) in cells expressing human progerin mutant lamin A compared with cells expressing wild-type lamin A. Many different DNA repair pathways are mutated in other progeria syndromes, and all DNA repair pathways seem to decline with chronological ageing (Gorbunova et al., 2007). The mutagens that we have investigated here (UV and ENU) likely are targeted by global-genome nucleotide excision repair (NER), base excision repair (BER) and break repair. NER is a ‘detect-excite-and-patch’ repair system for a broad class of helix-distorting lesions such as UV-induced photoproducts and many bulky chemical adducts. The global-genome NER pathway detects such lesions anywhere in the genome, whereas the transcription-coupled NER (TC-NER) pathway is selective for lesions that block transcriptional elongation. UV-induced CPD photoproduct removal is reduced in older and senescent fibroblasts (Boyle et al., 2005; Gorbunova et al., 2007), especially in the non-transcribed region (Gonzalo and Kreienkamp, 2015; Guo et al., 1998), and BER activity is reported to be decreased in old mice (Busuttil et al., 2007; Intano et al., 2003). Similarly, MEFS from lamina-associated progeroid syndromes are reported to be hypersensitive to UV (Liu et al., 2005). We did score a significantly elevated mutant rate at FE-1 cells and transfectants after their exposure to either UV or ENU; however, this induced mutant rate was no higher in FE-1 transfectants expressing LAΔ50 than in the parental cells or cells expressing wt lamin A (Fig. 4).

Because we have only scored mutation at an unexpressed reporter locus (LacZ), we cannot exclude that it is specifically transcription-coupled repair that is affected in HGPS, though no increase in the spontaneous mutation rate has been seen in progeroid TC-NER mutants.
(Wijnhoven et al., 2007). Similarly, since not all tissues seem to age prematurely in HGPS, it is possible that the FE-1 cell line is not from a suitable tissue type to reveal a defect in DNA repair caused by expression of LAΔ50 (Martin and Oshima, 2000).

As has been previously reported (Dechat et al., 2007, Goldman et al., 2004, Shumaker et al., 2006), we found an increased proportion of abnormally and irregularly shaped nuclei in cells expressing LAΔ50 mutant as compared to wt lamin A (Fig. 2). However, we noticed a decrease in this proportion when LAΔ50-expressing cells were grown at physiological oxygen levels (3%) as compared to the 20% O₂ of standard tissue culture. A similar decrease in aberrant nuclear morphology was not seen in cells expressing wt lamin A. This suggests that progerin-expressing cells may be especially sensitive to oxidative stress, and this is consistent with the elevated levels of reactive oxygen species reported in HGPS cells (Richards et al., 2011; Viteri et al., 2010).

Materials and methods

FE1 Muta™ Mouse lung epithelial cells were cultured as described by White et al. (2003). Briefly, cells were cultured in 1:1 DMEM/F12 nutrient mixture (Invitrogen) supplemented with 2% FBS (Sigma), 2 mM glutamine and 1 ng/ml murine epidermal growth factor (Invitrogen). Cells were maintained at 37 °C in hypoxic incubators (New Brunswick Galaxy 170R) with 3% O₂ or in atmospheric-O₂-level incubators when indicated. For hydrogen peroxide (H₂O₂) treatment, cells were exposed to 1 mM H₂O₂ for 25 min.

FISH

Cells were swollen in 0.56% KCl before fixation in 3:1 methanol/acetic acid. Slides were incubated with 100 μg/ml RNase A in 2× SSC for 1 h, washed in 2× SSC and dehydrated through an alcohol series. Slides were then denatured in 70% formamide/2× SSC for 1 min. λ DNA was purified using the manufacturer’s protocol (Qiagen, # 12543) and labelled by nick translation with digoxigenin-11-dUTP (Morey et al., 2007). Approximately 100 ng of labelled DNA probe and 15 μl of FITC-mouse chromosome-3 paint (Cambio) were used per slide, together with 5 μg of mouse Cot1 DNA (GIBCO BRL) and 5 μg salmon sperm DNA. Probes were denatured at 70 °C for 5 min, reannealed with Cot1 DNA for 15 min at 37 °C and hybridised to the denatured slides overnight. Digoxigenin-labelled probes were detected with Texas-Red anti-sheep (Vector). Slides were counterstained with 0.5 μg/ml 4’,6’-diamidino-2-phenylindole (DAPI) in Vectashield.

Cloning and expression of lamin A and LAD50

Stable FE-1 cells were prepared expressing either EGFP-myc-LA or EGFP-myc-hLAΔ50. The coding regions from pTRE-EGFP-myc-hLMNA or –hLMNAΔ50 (Goldman et al., 2004; Shumaker et al., 2006) were excised as a 2.8-kb BamHI/XbaI fragment and cloned into pcDNA3.1 (+) (Invitrogen). The resulting plasmids (pcDNA-EGFP-myc-hLMNA and pcDNA-EGFP-myc-hLMNAΔ50) were linearised with Scal and transfected into FE-1 cells by lipofectamine 2000 (Invitrogen). After 48 h, cells were trypsinised and GFP-expressing cells sorted using a BD FACSariaI SORP instrument (Becton Dickinson). The 488-nm laser was used for measuring forward scatter, side scatter and GFP fluorescence (525/50 nm bandpass filter). BD FACSDiva software (Becton Dickinson, Version 6.1.2) was used for instrument control and data analysis. The purified GFP-expressing cells were seeded in a six-well plates and expanded. Cell populations were sorted regularly for GFP fluorescence to maintain a high proportion of GFP-lamin A-expressing cells. After establishment of stably expressing cell lines, cells were maintained in a 3% O₂ incubator.

Immunofluorescence

Cells grown on coverslips were fixed in 4% paraformaldehyde in PBS and permeabilised in 0.2% Triton X-100/ PBS for 12 min. Fixed cells were incubated overnight at 4 °C with primary antibodies against Lamin A/C (1:100, Cell Signalling) or γ-H2AX phospho-[Ser139] (1:100, #05–636 Millipore). The slides were then incubated for 1 h at room temperature with Alexa-Fluor 488 or Alexa-Fluor 594 secondary antibodies (1:1000, Invitrogen). Cells were counterstained with 0.02 μg/ml DAPI in PBS and mounted in Vectashield. Slides were examined on a Nikon A1R confocal microscope equipped with a CFI Plan Fluor ×40/1.30 oil objective and NIS elements software.

Western blotting

Western blot analysis was carried out using standard protocols. Immunoblotting was performed with primary
antibodies directed against Lamin A/C (1:2000, #sc-6215, Santa-Cruz Biotechnology) or (1:1000, #4777, Cell Signalling), GFP (1:2000, #11-814-460-001 Roche), H3K27me3 (1:2000, #07-449 Millipore), H3K9me3 (1:1000, #05–1242 Millipore), γ-H2AX phospho-[Ser139] (1:500, #05-636 Millipore), HP1α (1:500, #05-689 Millipore), PCNA (1:20,000, #sc-56 Santa-Cruz Biotechnologies) and H3 (1:50,000, Millipore). Blots were detected by horseradish peroxidase (HRP)-conjugated donkey anti-rabbit, anti-mouse or anti-goat whole molecule IgG (1:10,000) and chemiluminescence using ChemiGlow West (Alpha Innotech). Signal was analysed using ImageQuantTL LAS4010 (Version 1; GE Healthcare).

Measurement of LacZ mutant frequency

3 × 10⁵ cells were cultured in a 100-mm culture dish overnight. For UV-C treatment, cells were washed with PBS and exposed to 10 J/m² (UV Stratalinker 1800, Stratagene) at 254 nm, which was repeated the following day. Cells were collected 72 h after the first exposure to mutagen. For N-ethyl-N-nitrosourea (ENU, Sigma) treatment, cells were incubated for 6 h in serum-free medium containing 200 μg/ml ENU diluted in DMSO, washed with PBS and then incubated for 72 h with complete medium. Cells were lysed for 3 h at 55 °C in 100 mM Tris pH 8, 200 mM NaCl, 5 mM EDTA, 0.2% SDS and 0.1 mg/ml proteinase K. DNA was extracted with phenol/chloroform (1:1) and chloroform/SDS and 0.1 mg/ml proteinase K. DNA was extracted with phenol/chloroform (1:1), followed by chloroform and precipitated by isopropanol, and resuspended in 100 mM Tris pH 8, 200 mM NaCl, 5 mM EDTA, 0.2%

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