Gopalakrishnan *et al. Genome Integrity* 2010, **1**:16 http://www.genomeintegrity.com/content/1/1/16



RESEARCH Open Access

Hydrogen peroxide induced genomic instability in nucleotide excision repair-deficient lymphoblastoid cells

Kalpana Gopalakrishnan^{1†}, Grace Kah Mun Low^{1†}, Aloysius Poh Leong Ting¹, Prarthana Srikanth¹, Predrag Slijepcevic², M Prakash Hande^{1*}

Abstract

Background: The Nucleotide Excision Repair (NER) pathway specialises in UV-induced DNA damage repair. Inherited defects in the NER can predispose individuals to Xeroderma Pigmentosum (XP). UV-induced DNA damage cannot account for the manifestation of XP in organ systems not directly exposed to sunlight. While the NER has recently been implicated in the repair of oxidative DNA lesions, it is not well characterised. Therefore we sought to investigate the role of NER factors Xeroderma Pigmentosum A (XPA), XPB and XPD in oxidative DNA damage-repair by subjecting lymphoblastoid cells from patients suffering from XP-A, XP-D and XP-B with Cockayne Syndrome to hydrogen peroxide (H₂O₂).

Results: Loss of functional XPB or XPD but not XPA led to enhanced sensitivity towards H_2O_2 -induced cell death. XP-deficient lymphoblastoid cells exhibited increased susceptibility to H_2O_2 -induced DNA damage with XPD showing the highest susceptibility and lowest repair capacity. Furthermore, XPB- and XPD-deficient lymphoblastoid cells displayed enhanced DNA damage at the telomeres. XPA- and XPB-deficient lymphoblastoid cells also showed differential regulation of XPD following H_2O_2 treatment.

Conclusions: Taken together, our data implicate a role for the NER in H_2O_2 -induced oxidative stress management and further corroborates that oxidative stress is a significant contributing factor in XP symptoms. Resistance of XPA-deficient lymphoblastoid cells to H_2O_2 -induced cell death while harbouring DNA damage poses a potential cancer risk factor for XPA patients. Our data implicate XPB and XPD in the protection against oxidative stress-induced DNA damage and telomere shortening, and thus premature senescence.

Background

The nucleotide excision repair (NER) pathway is a versatile DNA repair mechanism that recognizes and efficiently removes an array of structurally diverse DNA lesions including ultraviolet (UV)-induced lesions, intrastrand crosslinks and bulky chemical adducts such as those induced by compounds in tobacco smoke. The NER comprises of more than three dozen genes working in spatial and temporal concert and is differentiated into two sub-pathways - the global genome-NER (GG-NER)

and transcription coupled repair (TCR) - that differ only in damage recognition [1,2].

Inherited defects in the NER predispose an individual to genetic disorders featuring genomic instability and segmental progeria - Xeroderma pigmentosum (XP), Cockayne syndrome (CS) and Trichothiodystrophy (TTD). XP is a rare autosomal recessive congenital disorder that arises from mutations in XP proteins, XPA - XPG, and a variant form XPV. XP patients are predisposed to suninduced cutaneous cancer incidence by more than a thousand-fold, display sunlight hypersensitivity, high frequency of internal tumours, accelerated neurodegeneration and developmental abnormalities [3,4]. XPA, XPB and XPD are three proteins that play pivotal roles in both the GGNER and TCR. XPA is involved in DNA damage recognition through site-directed binding of rigidly kinked double

Full list of author information is available at the end of the article



^{*} Correspondence: phsmph@nus.edu.sg

[†] Contributed equally

¹Department of Physiology, Yong Loo Lin School of Medicine, National University of Singapore, 117597, Singapore

stranded DNA, thereby engaging the excision of the lesion [5,6]. XPB and XPD unwind the local area of the damaged DNA; by virtue of constituting transcription factor II H (TFIIH), which is part of the RNA Polymerase II holocomplex, they are important not only for repair but also for basal transcription although the helicase activity of XPD is dispensable for transcription [7]. Mutations compromising the function of either of these XP genes can lead to specific clinical outcomes. In particular, mutations in XPA results in only XP while mutations in either XPB or XPD can result in XP, XP/CS, TTD or XP/TTD. Additionally, polymorphisms in XP genes can give rise to diseases with phenotypic heterogeneity of differing severities [8-10].

Although a common denominator for lesions repaired by the NER is the presence of significant distortion of the DNA helix [11], it has more recently been implicated in the repair of minor oxidative base damages that are not helix distorting [12]. Despite the base excision repair (BER) being the main pathway for the repair of such lesions, the NER is also important and may serve as a back-up system [13,14]. Endogenous oxidative damage occurs via the by-production of reactive oxygen species (ROS) such as hydrogen peroxide (H₂O₂) during normal cellular metabolism. Oxidative DNA damage constitutes strand breaks, helical distortions and hindrance to base pairing, all of which alter important genetic information by interfering with replication and transcription. Accumulation of oxidative lesions thus compromises DNA integrity predisposing to cancer [15,16] and ageing [17].

UV-induced damage cannot account for all the symptoms of XP and related disorders, especially those in organ systems not directly exposed to sunlight. A class of oxidative lesions has been shown to be specifically repaired by the NER and may play a role in neurodegeneration in XP patients [18,19]. In fact, oxidative base lesions have been implicated in patients suffering from XP/CS who display the skin diseases of XP together with the somatic and neurological abnormalities of CS [20]. Further, the neurological symptoms of XP/CS may have resulted from aberrant DNA repair of lesions induced endogenously during oxidative metabolism [8].

Oxidative stress through ROS generation is down-stream to various other genotoxic agents including UV-irradiation [17]. Hence oxidative stress appears to be an important player in the development of XP symptoms that do not result directly from UV exposure. In this context a recent study has suggested that TTD and XP/CS mammalian fibroblasts that lack XPD were defective in oxidative DNA lesion-repair [21]. We have previously shown that lack of functional XPA in primary human fibroblasts increased the susceptibility of sodium arsenite- and $\rm H_2O_2$ -induced genotoxicity while retaining

cell viability, posing as a risk factor for cancer in XPA patients [22]. In addition, we have found that XPB dysfunction also inclined cells to genome instability and telomere shortening when exposed to H_2O_2 [23]. This study investigates the role of XPA, XPB and XPD in maintaining the genomic stability of lymphoblastoid cells under H_2O_2 -induced oxidative stress.

Materials and methods

Cell Culture and Treatment Conditions

Epstein-Barr transformed human diploid B-lymphocytes (lymphoblastoid cells) from a normal individual (Normal-L GM03798) and patients with Xeroderma Pigmentosum (XP) Complementation Group A (XPA-L GM02344), XP Complementation Group B also exhibiting Type 2 Cockayne's Syndrome (XPB-L GM02252) and XP complementation Group D (XPD-L GM11612) were purchased from Coriell Cell Repository, U.S.A., and cultured in Roswell Park Memorial Institute (RPMI) 1640 medium (Gibco Invitrogen Corporation, U.S.A.) supplemented with 15% foetal bovine serum (FBS; Hyclone, U.S.A.) and 1% L-Glutamine (Gibco Invitrogen Corporation, U.S.A.). Cells were grown and maintained in log phase in a humidified 5% CO₂ incubator at 37°C. Unless otherwise stated, exponentially growing cells were seeded at a density of 4×10^5 per mL in 6-well plates (Nunc, Denmark) and exposed to between 20 µM and 100 µM of H₂O₂ (Kanto Chemical Co. Inc., Japan) for 2 hours, following which the medium was replaced with fresh medium for a 22-hour recovery period. Such concentrations of H₂O₂ have been shown to be of low to no cytotoxicity with some genotoxicity [22,24].

Cell Viability by 3-[4,5-Dimethylthiazol-2-yl]2,5-diphenyltetrazolium bromide (MTT) Assay

The tetrazolium salt, MTT, is reduced to a water-insoluble coloured formazan only by metabolically active cells. The relative number of viable cells can be measured following the solubilisation and quantification of the formazan salt. Suspension cells were collected and centrifuged at 260 g and resuspended in equal volume of fresh media. Fifty microlitres of the cell suspension was added in triplicates to a 96-well plate (Nunc) to an equal volume of 4 mg/mL MTT (Sigma) solution. The plate was incubated at 37°C in the dark for 2 hours before being centrifuged at 1620 g. The supernatant was carefully aspirated and the formazan formed was dissolved in 200 µL 5% v/v Sorenson's glycine buffer (0.1 M glycine (Univar, New Zealand), 0.1 M NaCl, pH 10.5) in dimethyl sulfoxide (DMSO; MP Biomedicals Inc., France), following which the dissolved salt was measured at 570 nm using µQuant Microplate Spectrophotometer. Data is represented as the percentage of absorbance of the sample over the untreated counterpart.

Analysis of Cell Cycle by Fluorescence Activated Cell Sorting (FACS)

Harvested cells were washed in $1 \times PBS$, fixed in 70% ethanol: $1 \times PBS$ and stained with 2 mg propidium iodide (PI): 2 mg RNase A (*Roche, U.S.A.*)/100 mL 0.1% BSA in $1 \times PBS$ for 30 minutes at 37°C in the dark. Samples were analysed by flow cytometry at 488 nm excitation λ and 610 nm emission λ . Approximately 20,000 events were collected and the data obtained was analysed using WINMDI software.

Cytokinesis Blocked Micronucleus (CBMN) Analysis

Immediately after the 2-hour H_2O_2 treatment, cells were incubated in fresh medium with 4.0 µg/mL cytochalasin B (*Sigma*) for 22 hours. The protocol used is adapted from [25] and cells were processed as described before [22,26]. One thousand binucleated cells with/without the presence of micronuclei were scored under an Axioplan 2 imaging fluorescence microscope (*Carl Zeiss, Germany*) with an appropriate triple band filter.

Chromosome Aberration (CA) Analysis

Following 2-hour H_2O_2 treatment, cells were allowed to grow in fresh medium for 24 hours, then arrested at metaphase with 10 μ L/mL karyomax colcemid solution (*Gibco*) for 4 hours before being subjected to hypotonic swelling in pre-warmed 0.075 M KCl at 37°C for 12 minutes and fixation with Carnoy's fixative (3 parts methanol: 1 part glacial acetic acid). Fluorescent *in situ* hybridisation and analysis were performed as described before [22,26] using Cy3-labelled PNA-telomere and FITC-labelled PNA-centromere probes (*Applied Biosystems, U.S.A.*) [27].

Alkaline Single Cell Gel Electrophoresis (Comet) Assay

Two sets of cells were plated for the comet assay. Following treatment, one set was allowed to undergo a recovery period of 22 hours in fresh media without H_2O_2 whilst the other was harvested immediately. Cells were processed as described before [22]. One hundred randomly selected cells were examined per sample using Comet Imager Software (*Metasystems, Germany*). Extent of DNA damage was expressed as a measure of comet tail moments, which corresponds to the fraction of DNA in the comet tail.

Western Blotting

Cells were harvested at the same time points as that of the Comet assay (see above), lysed in 10 mM Tris-HCl (pH 7.4)/1% SDS/1 mM sodium ortho-vanadate (Sigma)), homogenised with a 291/2" syringe to shear DNA and centrifuged at 16,100 g at 4°C for 6 minutes. The supernatant was collected and estimated for protein concentrations using microBCA[™] Protein Assay

Kit (Pierce) according to manufacturer's instructions. An equal amount of protein (~40 μg) was resolved in SDS-polyacrylamide gels before electro-blotting onto nitrocellulose membranes (Biorad Co.). Membranes were blocked at room temperature for 1 hour with 5% (v/w) non-fat milk powder in TBS-T (0.1% v/v Tween-20 (Sigma) in 1×TBS (10 mM NaCl, 10 mM Tris pH 7.4 in dH₂O)), incubated with primary antibodies specific for the protein of interest, washed in TBS-T to remove excess antibody, incubated in secondary antibodies specific for the primary antibody at room temperature for 1 hour, then washed in TBS-T to remove excess antibody. Primary antibodies used were the mouse monoclonal antibodies for p53 (DO-1, 1:1,000), phospho-p53 (Ser 15 16G8, 1:1000; Cell Signaling Technology, Inc., U.S.A.), p21 (F-5, 1:500), survivin (D-8, 1:200), PCNA (PC-10, 1:500) and XPD (184.7, 1:100), rabbit polyclonal antibodies for XPA (FL-273, 1:100) and XPB (S-19, 1:100), and goat polyclonal antibody for actin (I-19, 1:2000). Secondary antibodies used were goat anti-mouse Ig G (H+L)-HRP (1:5,000), sheep anti-rabbit Ig G (H+L)-HRP (1:5,000) or donkey anti-goat Ig G (H+L)-HRP (1:5,000) depending on the primary antibody used. All antibodies unless otherwise stated were purchased from Santa Cruz, U.S.A.. Protein bands were visualized against a protein marker (Biorad) after the membranes were incubated with Supersignal® West Pico Chemiluminescent Substrate (Thermo scientific, U.S.A) and exposed to X-ray films (Pierce).

Statistical Analysis

Statistical significance between and among data sets was assessed using two-way ANOVA with Graphpad Prism. The difference was considered to be statistically significant when */* $^{\#}p < 0.05$; **/* $^{\#}p < 0.01$; and ***/ $^{\#\#}p < 0.001$.

Results

Hydrogen peroxide exposure resulted in loss of viability in XPB-L and XPD-L but not XPA-L cells

While the control lymphoblastoid cells, XPB-L, and XPD-L cells exhibited a dose-dependent decrease in viability (p < 0.05; Figure 1), XPA-L cells did not exhibit a dose-dependent change in viability following treatment. Furthermore, XPA-L cells exhibited significantly higher viability (p < 0.05) than Normal-L cells at $\rm H_2O_2$ concentrations of 60 μM and above. XPB-L and XPD-L cells on the other hand, were significantly more sensitive to $\rm H_2O_2$ treatment than Normal-L cells even at the low dose of 20 μM (p < 0.05). Both XPB-L and XPD-L also exhibited a significant decrease in viability than untreated counterparts following treatment at doses of 40 μM $\rm H_2O_2$ and above (p < 0.05), while Normal-L cells

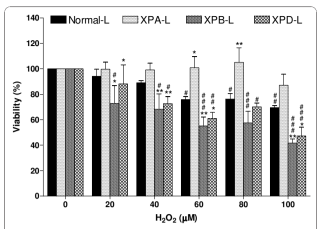


Figure 1 Cell Viability. Dose-dependent decrease in viability in Normal-L, XPB-L and XPD-L lymphoblastoid cells but not in XPA-L lymphoblastoid cells following H_2O_2 treatment. $^\#p < 0.05$; $^\#p < 0.01$; $^\#p < 0.001$ compared to untreated counterparts. At higher concentrations, XPA-L exhibited significantly higher viability than Normal-L. On the contrary, XPB-L and XPD-L both exhibited significantly lower viability than Normal-L following treatment. $^*p < 0.05$; $^*p < 0.01$ compared to Normal-L. It is interesting to note that both XPB-L and XPD-L were also significantly less viable than XPA-L at 60 μM and higher of *H_2O_2 .

exhibited significant sensitivity to H_2O_2 compared to untreated counterparts only from doses as high as 80 μ M (p < 0.05). It is also interesting to note that XPB-L and XPD-L cells were more sensitive to H_2O_2 -induced oxidative stress than XPA-L cells. These differences in viability response between XP-deficient cell types prompted our investigation of cell cycle profiles by PI staining.

XPB-L and XPD-L cells exhibited an increase in the sub-G1 population, while XPA-L cells demonstrated a dosedependent increase in the G2/M population

The primary purpose of cell cycle analysis was to observe changes in the sub-G1 population of cells, which would indicate the occurrence of early necrosis and/or apoptosis.

In response to treatment, cell cycle profiles of Normal-L cells did not change noticeably except for a slight increase in sub-G1 population at the higher concentrations (Figure 2A). XPB-L and XPD-L cells exhibited increases in the sub-G1 population with increasing concentrations of $\rm H_2O_2$ (Figure 2C and 2D respectively, and Figure 2E). It is further noteworthy that XPB-L cells displayed a corresponding decrease in percentage of cells in the G1 and G2/M phases following treatment. XPA-L cells on the other hand, did not demonstrate changes in the sub-G1 population but exhibited a dose-dependent increase in the G2/M population (Figure 2B, E and 2F).

XP-deficient lymphoblastoid cells exhibited significantly higher DNA damage than Normal-L cells following H_2O_2 exposure

We were next interested to investigate whether the lymphoblastoid cells sustained DNA damage from $\rm H_2O_2$ exposure in the absence of cell cycle arrest. The extent of DNA damage accumulated in XP-deficient lymphoblastoid cells in response to $\rm H_2O_2\text{-}induced$ oxidative stress was studied using three DNA markers - CBMN, CA and comet.

Binucleated (BN) cells were scored for the presence and distribution of micronuclei (MN). The frequency of MN increased significantly in a dose-dependent manner in all cell types (p < 0.05; Table 1). XPA-L and XPB-L cells did not exhibit significant increase in percent MN and percent BN with MN compared to Normal-L cells (p > 0.05) while XPD-L cells showed significant increase in these damage markers from H₂O₂ doses as low as $20 \mu M$ (p < 0.001; Figure 3). Clearly XPD-L cells were the most sensitive as they also showed significantly more percent MN compared to their untreated counterparts even at 20 μ M H₂O₂ (p < 0.05; Table 1) whereas XPA-L and XPB-L cells exhibited such increase only at 40 μ M and 60 μ M H₂O₂ respectively (p < 0.05), and Normal-L cells displayed this increase only at 100 µM H_2O_2 (p < 0.01). It is noteworthy that XPD-L cells also exhibited the highest basal MN frequency as opposed to all other cells (Table 1; Figure 3).

Chromosomal aberrations (CA) such as fragments, breaks and fusions sustained by the lymphoblastoid cells following H_2O_2 exposure are shown in Figure 4. The scores and distribution of CA are represented in Table 2 and Figure 4S. H₂O₂ treatment led to an overall increase in CA in all cell types. Notably, all three XP-deficient cell types exhibited a much larger increase in total CA -specifically more breaks - than Normal-L cells following treatment (Table 2). This increase was significant in XPB-L (p < 0.05) and XPD-L (p < 0.01) but not in XPA-L cells (p > 0.05; Figure 4S); XPA-L cells appeared to be the least sensitive among the three XP-deficient lymphoblastoid cells although they exhibited more fragments than XPB-L cells (Table 2). XPB-L cells seemed to be the most sensitive to H₂O₂-induced CA particularly undetected telomeres (UTs) and fusions (Table 2). XPD-L cells also showed more UTs than XPA-L and Normal-L cells (Table 2).

Lack of functional XP proteins lead to reduced capacity to repair oxidative DNA lesions in XPD-L, but not XPA-L and XPB-L cells

The ability of cells to recover from oxidative genotoxic assault was compared using overall genomic integrity as a parameter via the comet assay, where cells were harvested and analyzed as previously described [22]. The

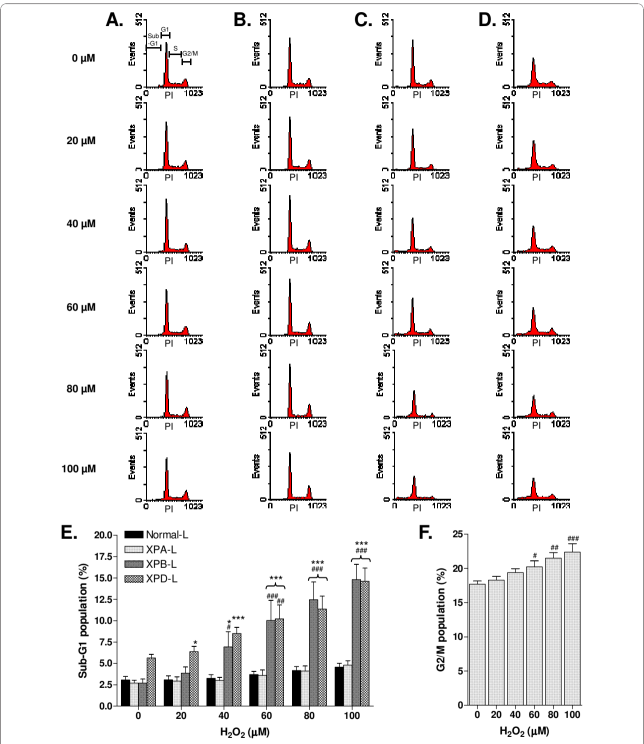


Figure 2 Cell cycle profiles of A. Normal-L, B. XPA-L, C. XPB-L, and D. XPD-L lymphoblastoid cells following H_2O_2 treatment. Pl: Propidium iodide, indicating amount of DNA stained per cell. The top profile of panel A depicts an illustration of the distribution of cells in each phase of the cell cycle (sub G1: apoptotic cells; G1-phase cells; S: S-phase cells; G2/M: G2/M-phase cells). **E.** Percentage of sub-G1 population. **F.** Percentage of G2/M population in XPA-L cells. "p < 0.05; "#p < 0.05; "#p < 0.01; "##p < 0.001 compared to untreated counterparts. "p < 0.05; "**p < 0.001 compared to Normal-L. Data is representative of at least three independent experiments.

Table 1 Cytokinesis-blocked micronucleus assay

| | H ₂ O ₂ (μΜ) | | | | | | | |
|----------|------------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Cells | | 1 | 2 | >4 | % MN | % BN with MN | | |
| | 0 | 2.80 ± 0.72 | 0.33 ± 0.12 | 0.13 ± 0.06 | 0.03 ± 0.06 | 0 | 4.00 ± 1.25 | 3.33 ± 0.86 |
| | 20 | 3.07 ± 0.60 | 0.47 ± 0.12 | 0 | 0.03 ± 0.06 | 0 | 4.13 ± 0.67 | 3.60 ± 0.62 |
| Normal-L | 40 | 3.63 ± 0.46 | 0.60 ± 0.27 | 0.13 ± 0.12 | 0.03 ± 0.06 | 0 | 5.37 ± 0.92 | 4.43 ± 0.64 |
| | 60 | 3.83 ± 1.21 | 0.53 ± 0.31 | 0.17 ± 0.06 | 0 | 0 | 5.40 ± 1.41 | 4.57 ± 1.30 |
| | 80 | 4.37 ± 0.74 | 0.57 ± 0.15 | 0.23 ± 0.06 | 0 | 0 | 6.20 ± 0.92 | 5.20 ± 0.85 |
| | 100 | 5.67 ± 0.46 | 0.87 ± 0.21 | 0.13 ± 0.06 | 0 | 0.03 ± 0.06 | 7.97 ± 0.96** | 6.70 ± 0.70** |
| | 0 | 2.33 ± 0.42 | 0.13 ± 0.15 | 0 | 0 | 0 | 2.60 ± 0.40 | 2.47 ± 0.38 |
| | 20 | 3.00 ± 0.35 | 0.30 ± 0.00 | 0 | 0 | 0 | 3.60 ± 0.35 | 3.30 ± 0.35 |
| XPA-L | 40 | 4.03 ± 1.20 | 0.80 ± 0.10 | 0.10 ± 0.10 | 0 | 0.07 ± 0.06 | 6.30 ± 1.35** | 5.00 ± 1.20* |
| | 60 | 4.00 ± 1.14 | 0.67 ± 0.25 | 0.10 ± 0.00 | 0.03 ± 0.06 | 0 | 5.77 ± 0.90* | 4.80 ± 0.95 |
| | 80 | 5.43 ± 1.92 | 0.63 ± 0.21 | 0.17 ± 0.21 | 0 | 0 | 7.20 ± 2.04*** | 6.23 ± 1.89*** |
| | 100 | 5.90 ± 0.95 | 0.83 ± 0.12 | 0.23 ± 0.21 | 0.03 ± 0.06 | 0 | 8.40 ± 0.62*** | 7.00 ± 0.72*** |
| | 0 | 2.77 ± 0.95 | 0.43 ± 0.15 | 0 | 0 | 0 | 3.70 ± 1.20 | 3.20 ± 1.01 |
| | 20 | 3.10 ± 1.15 | 0.47 ± 0.21 | 0.10 ± 0.10 | 0 | 0 | 4.43 ± 1.65 | 3.67 ± 1.26 |
| XPB-L | 40 | 3.77 ± 0.85 | 0.37 ± 0.15 | 0.10 ± 0.10 | 0 | 0 | 4.87 ± 1.47 | 4.23 ± 1.08 |
| | 60 | 4.83 ± 0.55 | 0.43 ± 0.15 | 0.23 ± 0.12 | 0 | 0 | 6.53 ± 0.74* | 5.50 ± 0.52 |
| | 80 | 5.57 ± 0.42 | 0.60 ± 0.10 | 0.23 ± 0.12 | 0 | 0 | 7.60 ± 0.35** | 6.40 ± 0.44** |
| | 100 | 6.00 ± 0.36 | 1.37 ± 0.15 | 0.37 ± 0.23 | 0 | 0 | 10.03 ± 1.00*** | 7.73 ± 0.46*** |
| | 0 | 5.10 ± 0.17 | 0.57 ± 0.12 | 0.03 ± 0.06 | 0 | 0 | 6.33 ± 0.58 | 5.70 ± 0.35 |
| | 20 | 7.07 ± 2.11 | 0.90 ± 0.17 | 0.17 ± 0.15 | 0.03 ± 0.06 | 0 | 9.50 ± 1.35* | 8.17 ± 1.82 |
| XPD-L | 40 | 7.03 ± 2.14 | 0.90 ± 0.10 | 0.23 ± 0.21 | 0.03 ± 0.06 | 0 | 9.67 ± 2.44* | 8.20 ± 2.25 |
| | 60 | 7.60 ± 0.62 | 0.90 ± 0.20 | 0.30 ± 0.17 | 0 | 0 | 10.30 ± 0.70** | 8.80 ± 0.70** |
| | 80 | 8.23 ± 1.56 | 1.20 ± 0.36 | 0.27 ± 0.21 | 0.03 ± 0.06 | 0 | 11.57 ± 0.76 | 9.73 ± 1.29*** |
| | 100 | 8.70 ± 2.19 | 1.17 ± 0.38 | 0.30 ± 0.10 | 0.07 ± 0.06 | 0.03 ± 0.00 | 12.47 ± 2.37*** | 10.27 ± 2.42*** |

Micronuclei (MN) scores, percentage of binucleated cells (BN) with MN and percentage of MN in cytokinesis-blocked binucleated lymphoblastoids following H_2O_2 treatment. Data is represented as mean \pm SD. Statistics shown only for last two columns where *p < 0.05, **p < 0.01, ***p < 0.001, where significance is calculated to the respective cell type's untreated counterpart.

rationale for assessing damage at two time points was to elucidate susceptibility to oxidative DNA damage and capacity for damage-repair. Comet tail moments scored are a function of tail length and fluorescence intensity [28]. Figure 5 depicts a SYBR-Green stained nucleus showing apparently no damage (Figure 5A) and one with a comet tail indicative of DNA damage (Figure 5B).

All lymphoblastoid cells exhibited a dose-dependent increase in tail moments following the 2-hour treatment with $\rm H_2O_2$ and this increase was significant in XP-deficient (p < 0.01) but not in Normal-L cells (p > 0.05; Figure 5C). Amongst XP-deficient cells, XPD-L cells exhibited the most drastic increase in tail moments compared to untreated and Normal-L counterparts following the 2-hour treatment, showing significant increase from 40 μM and higher doses of $\rm H_2O_2$ (p < 0.01), while XPA-L and XPB-L lymphoblastoid cells showed significant increase in tail moment following treatment with 60 μM and higher doses of $\rm H_2O_2$ (p < 0.01). At doses of 60 μM and above, XPD-L cells were

significantly more damaged than the other two XP-deficient cell types (p < 0.05).

Post 22-hour recovery, tail moments of all lymphoblastoid cells were lower than that of the 2-hour treatment (Figure 5C), indicating repair. Notably, XPB-L cells displayed significantly lower tail moments than its untreated and Normal-L counterparts (Figure 5C) at doses as low as 20 μ M H_2O_2 , indicating capacity for repair (p < 0.01). XPD-L cells on the other hand exhibited significantly higher tail moments even after recovery compared to its untreated counterparts (p < 0.001; Figure 5A), Normal-L cells (p < 0.001) and even XPA-L and XPB-L cells (p < 0.05), indicating lowest capacity to repair damaged DNA in XPD cells as opposed to other cell types.

XPD was differentially regulated in XPA-L and XPB-L cells following H₂O₂ treatment

To gain insight into the potential role of NER in maintaining genome stability and determining cell fate upon

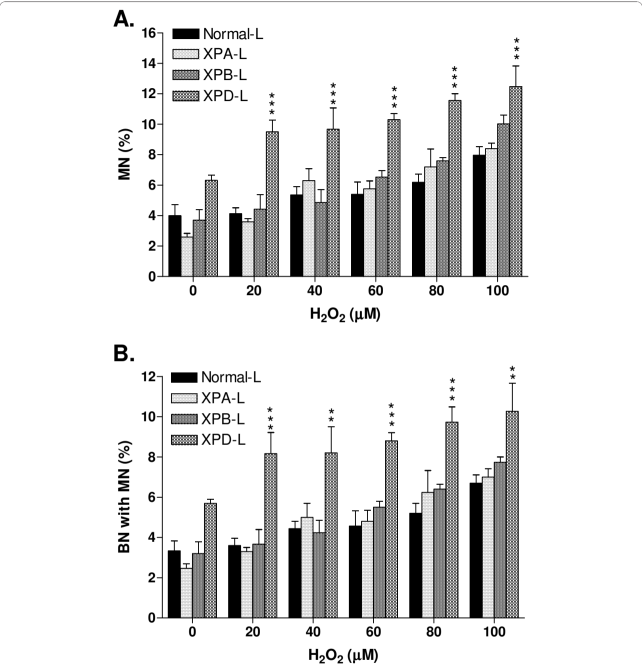


Figure 3 Cytokinesis Blocked Micronucleus (CBMN) Analysis: A. Percent MN per 1000 BN scored. B. Percent BN with presence of MN. There were dose-dependent increases in percent MN per 1000 BN as well as in percent BN with presence of MN in all lymphoblastoid cells. XPA-L and XPB-L both did not show significant increase in damage markers compared to Normal-L. XPD-L, however, exhibited significantly higher damage markers not only than Normal-L lymphoblastoid cells, but XPA-L and XPB-L as well. **p < 0.01, ***p < 0.001 compared to Normal-L. Two-way ANOVA.

 H_2O_2 exposure, we investigated the expression patterns of factors involved in cell cycle, apoptosis, survival and NER in the lymphoblastoids following exposure to 60 μM H_2O_2 (Figure 6). This dose was chosen as Normal-L cells exhibited significantly lower viability than untreated counterparts and both XPB-L and XPD-L

cells were significantly less viable than Normal-L cells at this concentration (Figure 1). Since XPB and XPD function in the same macromolecule and lymphoblastoids deficient in these proteins exhibited similar responses in our data, we explored protein expressions in only XPB-L cells.

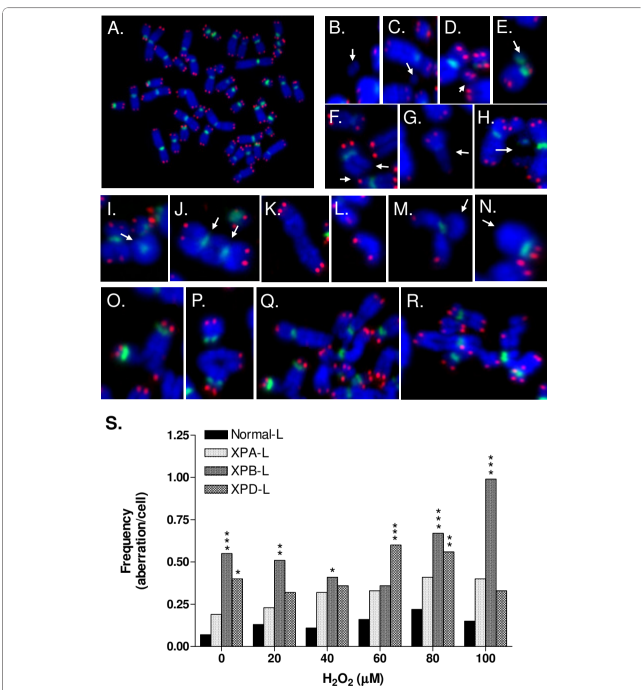


Figure 4 Telomere PNA-FISH was used to detect chromosomal aberrations (CA) in metaphases. A-R. Cy3-telomere and FITC-centromere PNA probes, depicted by the red and green signals respectively, and DAPI (blue) counterstain were used. Cell type and H_2O_2 concentrations used on the cells are stated within the parentheses. **A.** Metaphase spread with 46 chromosomes and no detected aberrations (XPA-L, 20 μM). **B-H.** Aberrations categorized as breaks. Interstitial deletion resulting in a **B.** double minute (XPD-L, 0 μM) and **C.** acentric fragment (XPD-L, 0 μM). **D.** Terminal break in chromosome resulting in an acentric fragment with telomere signals (XPA-L, 60 μM). **E.** Interstitial deletion resulting in a centric fragment (XPA-L, 80 μM). **F.** Undetected telomere signals (XPB-L, 0 μM). **G.** Chromatid break (XPD-L, 80 μM). **H.** Chromosome break (Normal-L, 80 μM). **I-N.** Aberrations categorized as simple fusions. **I.** Centric ring (XPA-L, 100 μM). **J.** Dicentric chromosome with both centromeres depicted by white arrows (XPB-L, 100 μM). **K.** Fusion of two terminally broken chromosomal fragments forming a large chromosome-like structure (XPD-L, 40 μM). **L.** Fusion of two terminally broken chromatid fragments (Normal-L, 20 μM). **M-N.** Sister chromatid fusions (Normal-L, 20 μM and XPD-L, 60 μM respectively). **O-R.** Aberrations categorized as complex fusions. **O-P.** Triradial-like structures (XPA-L, 60 μM respectively). **Q-R.** Quadiradial-like structures (both from XPA-L, 60 μM). **S.** Frequency of total aberration per spread following H₂O₂ treatment. *p < 0.05; **p < 0.01, ***p < 0.001 compared to Normal-L.

Table 2 Analysis of metaphase spreads of lymphoblastoids following H2O2 treatment

| Cell Type | H ₂ O ₂ (μΜ) | | Fusions | | | | | | | | Breaks | | <u></u> | |
|--------------|---------------------------------------|------------------------------|-----------|-----|---------|---|-----|------------------|----|----|--------|-----------------|----------------------|-----------------------|
| | | Chromosomes per metaphase | T/ QLS | SCF | D/ T | R | EAF | Total Fusions | СВ | F | UT | Total Breaks | Total Aberrations | Aberrant cells (%) |
| | 0 | 45.90 ± 0.68 (44-48) | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 6 | 7 | 7 | 7 |
| | 20 | 46.04 ± 0.49 (44-48) | 0 | 1 | 0 | 0 | 1 | 2 | 0 | 1 | 10 | 11 | 13 | 10 |
| Normal- L | 40 | 45.82 ± 0.73 (44-48) | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 8 | 11 | 11 | 9 |
| | 60 | 45.89 ± 0.77 (44-48) | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 | 11 | 16 | 16 | 14 |
| | 80 | 45.79 ± 0.86 (44-48) | 0 | 0 | 0 | 1 | 0 | 1 | 2 | 11 | 8 | 21 | 22 | 14 |
| | 100 | 46.03 ±0.76 (44-48) | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 3 | 11 | 14 | 15 | 13 |
| | 0 | 45.83 ± 0.67 (44-48) | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 10 | 9 | 19 | 19 | 14 |
| | 20 | 45.79 ± 0.64 (44-48) | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 15 | 6 | 22 | 23 | 14 |
| XPA-L | 40 | 45.69 ± 0.99 (43-48) | 0 | 1 | 2 | 0 | 1 | 4 | 2 | 13 | 13 | 28 | 32 | 22 |
| | 60 | 45.65 ± 0.79 (43-48) | 3 | 0 | 1 | 1 | 0 | 5 | 2 | 12 | 14 | 28 | 33 | 22 |
| | 80 | 45.35 ± 1.46 (41-48) | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 22 | 18 | 40 | 41 | 26 |
| | 100 | 45.77 ± 0.67 (44-48) | 0 | 0 | 3 | 1 | 0 | 4 | 5 | 16 | 15 | 36 | 40 | 28 |
| | 0 | 46.49 ± 0.97 (44-48) | 0 | 1 | 0 | 0 | 0 | 1 | 1 | 5 | 48 | 54 | 55 | 39 |
| | 20 | 46.28 ± 1.08 (44-48) | 0 | 1 | 1 | 0 | 0 | 2 | 0 | 5 | 44 | 49 | 51 | 37 |
| XPB-L | 40 | 46.61 ± 0.89 (44-48) | 0 | 1 | 0 | 0 | 0 | 1 | 0 | 6 | 34 | 40 | 41 | 29 |
| | 60 | 46.53 ± 0.87 (44-48) | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 34 | 36 | 36 | 30 |
| | 80 | 46.43 ± 0.96 (44-48) | 0 | 1 | 4 | 0 | 1 | 7 | 1 | 8 | 51 | 60 | 67 | 40 |
| | 100 | 46.42 ± 1.01 (44-48) | 0 | 2 | 4 | 0 | 5 | 11 | 0 | 11 | 77 | 88 | 99 | 45 |
| | 0 | 45.86 ± 0.87 (44-48) | 0 | 0 | 0 | 0 | 0 | 0 | 3 | 21 | 16 | 40 | 40 | 30 |
| | 20 | 45.86 ± 0.87 (44-48) | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 13 | 19 | 32 | 32 | 23 |
| XPD-L | 40 | 45.75 ± 0.86 (44-48) | 0 | 0 | 1 | 0 | 1 | 2 | 0 | 19 | 15 | 34 | 36 | 19 |
| | 60 | 45.95 ± 0.86 (44-48) | 2 | 1 | 1 | 0 | 0 | 4 | 6 | 19 | 31 | 56 | 60 | 37 |
| | 80 | 45.90 ± 0.91 (44-48) | 0 | 0 | 1 | 0 | 0 | 1 | 1 | 20 | 34 | 55 | 56 | 31 |
| | 100 | 45.80 ± 1.10 (44-48) | 0 | 1 | 1 | 1 | 0 | 3 | 3 | 10 | 17 | 30 | 33 | 25 |

T/QLS-Triradial/Quadiradial-like structures; SCF-Sister Chromatid Fusions; D/T-Di/Tricentrics; R-Rings; EAF - End-to-end fusions of acentric fragments; CB-Chromsosome/Chromatid breaks; F-fragments including double minutes, terminal acentric fragments, interstitial acentric fragments and centric fragments; UT-Undetected telomeres. Data is represented as total number of aberrations. A total of 100 spreads were analyzed in two independent experiments. *p < 0.05, **p < 0.01, ***p < 0.001, where significance is calculated to the respective cell type's untreated counterpart.

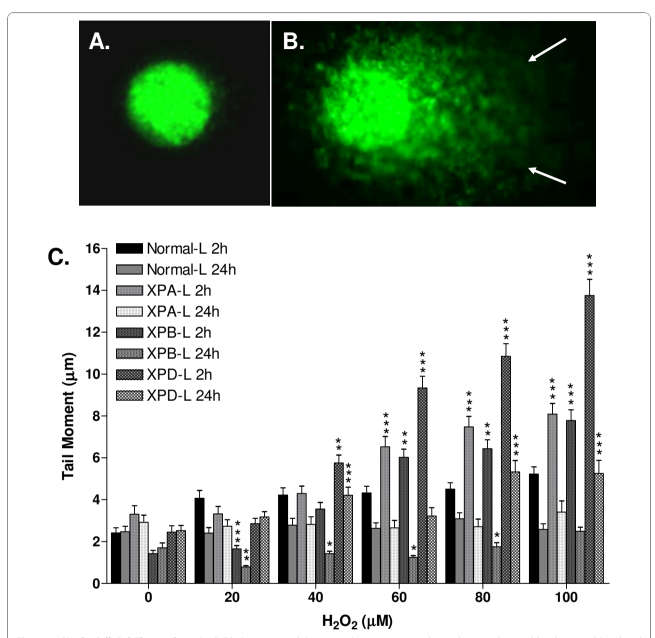


Figure 5 Single Cell Gel Electrophoresis. SYBR-Green stained **A**. comet showing apparently no damage depicted by the round head and **B**. comet with tail depicted by arrows indicating presence of damage. **C**. Tail moments immediately following 2-hour H_2O_2 treatment and 22-hour recovery in fresh medium. H_2O_2 concentrations of 60 μM and above resulted in significantly increased tail moments at 2 hours (p < 0.001) which decreased significantly following recovery (p < 0.05) in all lymphoblastoid cells. While there was no significant difference in moments between untreated Normal-L and XP-deficient lymphoblastoid cells (p > 0.05), XP-deficient lymphoblastoid cells exhibited increased tail moments when compared to Normal-L at 2 hours of H_2O_2 exposure. Following recovery, however, only XPD-L showed significantly higher moments than Normal-L (p < 0.001), while XPB-L showed significantly lower moments. *p < 0.05, **p < 0.01 and ***p < 0.001 indicate significantly greater tail moments when comparing XP-deficient lymphoblastoid cells to Normal-L counterparts. Data is represented as mean ± SE.

We observed up-regulation of p53 and phospho-p53 (p-p53), and no change in PCNA and Survivin expressions in Normal-L, XPA-L and XPB-L lymphoblastoids following 2 hours of $\rm H_2O_2$ treatment and 22 hours of recovery. p21 was up-regulated at both time points in XPA-L cells whereas it was not differentially expressed

in XPB-L cells. While Normal-L cells did not show any changes in expression in the NER factors XPA, XPB and XPD, XPA-L and XPB-L cells exhibited differential regulation of XPD, where XPA-L showed up-regulation of XPD at both time points and XPB-L displayed down-regulation at 2 hours but up-regulation at recovery. As

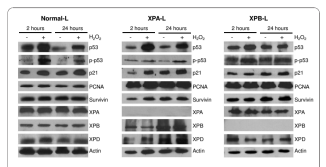


Figure 6 Protein expression profiles of Normal-L, XPA-L and XPB-L cells. p53, phospho-p53 and p21 proteins were differentially regulated in all three cell types, while PCNA, survivin, XPA, XPB and XPD proteins were not, except in XPA-L, whereby XPD exhibited differential regulation.

expected, XPA and XPB were not expressed in XPA-L and XPB-L cells respectively.

Discussion

The NER has been shown to be involved in oxidative damage repair but much is still yet to be elucidated. Through the Fenton and Haber-Weiss reactions, H₂O₂ generates OH radicals [29,30] which inflict more than 100 different DNA modifications including 8,5'-cyclopurine-2'-deoxynucleosides (CyPudNs) [31], an oxidative lesion repaired by the NER but not the BER [18,19]. These findings suggest that unrepaired oxidative lesions may contribute to XP symptoms which cannot be accounted for by UV exposure alone such as internal cancers, developmental defects and neurodegeneration. Moreover, the NER has been suggested to be functionally linked to the BER pathway, possibly explaining compromised BER functions observed in some XP patients. Specifically, NER factors XPG and XPC have been shown to interact with and activate BER DNA glycosylases [32,33]. Here we study the possible involvement of other NER factors, XPA, XPB and XPD in oxidative lesion management.

Our cell viability and cell cycle data indicate that loss of functional XPB or XPD but not XPA leads to enhanced sensitivity towards H₂O₂-induced cell death in the lymphoblastoids. The dose-dependent increase in the G2/M population and lack of cell death in H₂O₂-exposed XPA-L cells is a phenomenon that we have previously shown in XPA-deficient primary fibroblasts [22]. XPA has been shown to interact with and to be modulated by the cell cycle checkpoint proteins ATM and ATR following irradiation [34,35]. Therefore, lack of XPA may result in aberrant signalling leading to cell cycle dysfunction and failure of apoptosis. Alternatively, since the NER pathway is not the major pathway for H₂O₂-induced DNA lesions, cell cycle arrest may be triggered to allow time for repair by the BER instead.

The observed increased sensitivity of XPB-L and XPD-L cells to H₂O₂-induced cell death may possibly be attributed to the requirement of these proteins in both repair and basal transcription [36,37]. In this light, XPB and XPD are indispensible for cellular viability and only subtle mutations are found in XP, XP/CS and TTD [38,39]. However, we have previously noted that fibroblasts deficient in XPB, similar to the XPA-L lymphoblastoid cells in this study and XPA fibroblasts in our previous study, sustained genotoxic effects of H₂O₂ while retaining cell viability [23]. The differential results between XPB-deficient fibroblasts and lymphoblastoid cells could be due to cell type-specific responses, similar to an observation made between NER-deficient mouse embryonic stem cells and fibroblasts [40], or could have resulted from changes to the lymphoblastoids during transformation by Epstein-Barr virus.

Despite the observed differences in cell fate following $\rm H_2O_2$ exposure, lack of functional XPA, XPB or XPD led to increased susceptibility to $\rm H_2O_2$ -induced genome instability indicative of a role of these XP factors in maintaining genomic integrity challenged by $\rm H_2O_2$. Although lymphoblastoids deficient in XPA and XPB did not exhibit significantly more $\rm H_2O_2$ -induced MN compared to Normal-L lymphoblastoids, they demonstrated increased $\rm H_2O_2$ -induced damage susceptibility with significantly increased MN and BN per MN following treatment from lower $\rm H_2O_2$ concentrations than that of Normal-L cells (Table 1). This is consistent with our previous findings that XPA- and XPB-deficient fibroblasts were more susceptible to $\rm H_2O_2$ -induced DNA damage [22,23].

The comet assay was used to address whether the lack of a functional XP protein compromised a cell's ability to recover from H₂O₂-induced DNA damage. While the comet results also reflect that deficiency in XPA, XPB or XPD increased susceptibility to H₂O₂-induced global DNA damage, we noted that the loss of XPD resulted in the highest susceptibility to H₂O₂-induced DNA damage and the lowest capacity to repair these lesions, suggesting that XPD may have a more significant role than the other two XP factors in the repair of H₂O₂-induced lesions. On the other hand, the increased sensitivity of these NER-deficient lymphoblastoids to H₂O₂-induced comet tails may be confounded by disturbed redox homeostasis and not repair-deficiency per se as seen in XPC-deficient and XPC-silenced cells [41]. Of particular interest is the tail moment recovery of XPB-L cells to lengths shorter than that of Normal-L counterparts but comparable to that of XPB-L untreated cells, suggesting that while XPB-L may be more susceptible to H₂O₂induced damage, these cells do not lack the ability to repair the sustained damage. Similarly, while XPA-L cells sustained DNA damage, they exhibited competent

recovery. This seems to suggest that XPA and XPB may not be integral in oxidative damage-repair. While a study supports the notion that XPB in particular is not involved in the BER [42], we have previously demonstrated the importance of XPA and XPB in the protection against oxidative DNA lesions [22,23]. Moreover, XPB, XPD, and XPG deficiencies have been shown to compromise general repair [12]. The difference in the responses seen in the previous and current work may be attributed to cell-type specific responses. Nonetheless, that Normal-L lymphoblastoid cells did not sustain significant damage following $\rm H_2O_2$ exposure and exhibited competent repair mechanisms highlight the importance of XP factors in the protection against $\rm H_2O_2$ -induced oxidative DNA damage.

The marked increase in the frequency of UT in XPB-L cells suggests that XPB may have a crucial role in maintaining telomere integrity (Table 2). In line with this, we have previously shown that the lack of functional XPB leads to increased telomere attrition under oxidative stress [23]. XPD may also be integral in regulating telomere dynamics as the loss of this protein in lymphoblastoid cells also resulted in augmented frequency of UT (Table 2). The NER factor XPF was found to be implicated in telomere loss in mice overexpressing TRF2 [43], suggesting that other NER factors could play a role in telomere regulations. Moreover, XPB and XPD are part of the family of RecQ helicases which include WRN and BLM helicases shown to regulate telomere homeostasis [44-47].

Upregulation of p53 and p-p53 in Normal-L, XPA-L and XPB-L cells immediately following H₂O₂ exposure indicates proper DNA damage signalling responses were in place. H₂O₂-induced DNA damage induces the p53 pathway, which orchestrates DNA damage-repair responses, cell cycle arrest or apoptosis [48]. In response to DNA damage, p53 is upregulated and phosphorylated at Ser15 [49]. That p53 and p-p53 upregulation in XPB-L cells following 22-hour recovery is not as dramatic as that of XPA-L and Normal-L cells suggests that lack of XPB elicits only a short-lived DNA damage response, correlating with our comet data where XPB-L cells exhibited competent damage-repair. p21 is a direct downstream target of p53 and has been shown to integrate DNA damage into growth arrest or apoptotic signalling pathways [50]. XPA has shown involvement in DNA damage recognition and cell cycle checkpoints [51]. Consequently, loss of XPA may have led to aberrant signalling whereby accumulation of p53, p-p53 and p21 lead to cell cycle arrest instead of p53-dependant apoptosis. This observation correlates with our data where H2O2exposed XPA-L cells did not lose viability but exhibited a dose dependent increase in G2/M arrest.

PCNA plays a critical role in many DNA repair processes, including the NER [52]. Although studies have revealed that the lack of functional NER does not affect PCNA ubiquitination following UV irradiation [53], whether the protein is differentially regulated in response to $\rm H_2O_2$ treatment in an NER deficient background has not been explored. Survivin is a member of the inhibitor of apoptosis (IAP) protein family that regulates cell cycle and death, and is commonly upregulated in cancers [54]. The absence of PCNA and survivin differential expression suggests that these proteins are not involved in the processes affected by $\rm H_2O_2$ exposure.

XPA, XPB and XPD were not observed to be differentially regulated in Normal-L cells following H₂O₂ exposure, indicating that they may not be the immediate effectors of this type of stress. Indeed, the major pathway for oxidative DNA damage recognition and repair is the BER, and it would be worth investigating the modulation of BER proteins immediately following H₂O₂ exposure. Interestingly, while XPA and XPB proteins in XPB-L and XPA-L cells respectively were not modulated following H2O2 exposure, XPD was upregulated in response to H₂O₂ in XPA-L cells and in XPB-L cells following recovery, suggesting that a lack of functional XPA or XPB protein may be compensated for by upregulating XPD under H₂O₂-induced oxidative stress. Both XPB and XPD function in the TFIIH complex and have been proposed to be involved in the DNA damage verification process, whereby both helicases scan DNA strands by moving in the same direction [55,56]. XPBdeficient cells may therefore upregulate XPD to compensate for the lack of XPB function.

Conclusions

Our data shows that while all three XP-deficient lymphoblastoid cells unanimously exhibited increased susceptibility to H₂O₂-induced DNA damage, XPB and XPD, but not XPA may be involved in the repair of such lesions. While lack of functional XPA led to aberrant damage-induced cell death signalling posing a cancer risk, the lack of functional XPB and XPD led to higher damage sustenance, resulting in cell death. This could possibly explain why XPA patients develop cancer while some XPB and XPD patients are not cancer prone and manifest developmental defects of CS and TTD. Further, XPB and XPD may be involved in telomere maintenance under oxidative stress challenged by H₂O₂. Although to date telomere lengths of XP patients have not been fully explored, our findings link NER deficiency to telomere integrity, which may be associated with the progeroid symptoms of XP. Oxidative stress has been associated with premature senescence, accelerated ageing and telomere attrition [17,57] while telomere attrition and dysfunction have been linked to premature senescence and carcinogenesis [57]. Taken together, our data implicates a role for XP proteins in $\rm H_2O_2$ -induced oxidative stress management and further corroborates that oxidative stress is a significant contributing factor in XP symptoms, especially those that cannot be directly attributed to UV exposure.

Acknowledgements

MPH acknowledges the support from Defence Innovative Research Programme, Defence Science and Technology Agency, Singapore (POD: 0613592) and Academic Research Fund, Ministry of Education, Singapore (T206B3108). Supported in part by a grant from British Council, PMI2 Connect (Grant Number: RC134).

Author details

¹Department of Physiology, Yong Loo Lin School of Medicine, National University of Singapore, 117597, Singapore. ²Division of Biosciences, School of Health Sciences and Social Care, Brunel University, Uxbridge UB8 3PH, UK.

Authors' contributions

KG and GKML participated in the design of experiments, carried out the experiments, analysed the data and drafted the manuscript. APLT was involved in the experimental design, comet analysis, and manuscript preparation. PS¹ was involved in the cytogenetic analysis. PS² was involved in experimental design and manuscript preparation. MPH oversaw the overall experimental design and coordinated all research work and manuscript preparation. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

Received: 22 February 2010 Accepted: 22 December 2010 Published: 22 December 2010

References

- Hoeijmakers JH: Genome maintenance mechanisms for preventing cancer. Nature 2001, 411:366-374.
- 2. Sancar A: DNA excision repair. Annu Rev Biochem 1996, 65:43-81.
- 3. Norgauer J, Idzko M, Panther E, Hellstern O, Herouy Y: **Xeroderma pigmentosum.** *Eur J Dermatol* 2003, **13**:4-9.
- van Steeg H, Kraemer KH: Xeroderma pigmentosum and the role of UVinduced DNA damage in skin cancer. Mol Med Today 1999, 5:86-94.
- Missura M, Buterin T, Hindges R, Hubscher U, Kasparkova J, Brabec V, et al: Double-check probing of DNA bending and unwinding by XPA-RPA: an architectural function in DNA repair. EMBO J 2001, 20:3554-3564.
- Wood RD: Nucleotide excision repair in mammalian cells. J Biol Chem 1997, 272:23465-23468.
- Winkler GS, Araujo SJ, Fiedler U, Vermeulen W, Coin F, Egly JM, et al: TFIIH
 with inactive XPD helicase functions in transcription initiation but is
 defective in DNA repair. J Biol Chem 2000, 275:4258-4266.
- Kraemer KH, Patronas NJ, Schiffmann R, Brooks BP, Tamura D, DiGiovanna JJ: Xeroderma pigmentosum, trichothiodystrophy and Cockayne syndrome: a complex genotype-phenotype relationship. Neuroscience 2007, 145:1388-1396.
- Oh KS, Khan SG, Jaspers NG, Raams A, Ueda T, Lehmann A, et al: Phenotypic heterogeneity in the XPB DNA helicase gene (ERCC3): xeroderma pigmentosum without and with Cockayne syndrome. Hum Mutat 2006, 27:1092-1103.
- States JC, McDuffie ER, Myrand SP, McDowell M, Cleaver JE: Distribution of mutations in the human xeroderma pigmentosum group A gene and their relationships to the functional regions of the DNA damage recognition protein. *Hum Mutat* 1998, 12:103-113.
- de Laat WL, Jaspers NG, Hoeijmakers JH: Molecular mechanism of nucleotide excision repair. Genes Dev 1999, 13:768-785.
- Friedberg EC, Walker GC, Siede W: DNA Repair and Mutagenesis Washington, D.C.: ASM Press; 1995.

- Rybanska I, Pirsel M: Involvement of the nucleotide excision repair proteins in the removal of oxidative DNA base damage in mammalian cells. Neoplasma 2003, 50:389-395.
- Satoh MS, Lindahl T: Enzymatic repair of oxidative DNA damage. Cancer Res 1994, 54:1899s-1901s.
- Mates JM, Segura JA, Alonso FJ, Marquez J: Intracellular redox status and oxidative stress: implications for cell proliferation, apoptosis, and carcinogenesis. Arch Toxicol 2008, 82:273-299.
- Toyokuni S: Molecular mechanisms of oxidative stress-induced carcinogenesis: from epidemiology to oxygenomics. IUBMB Life 2008, 60:441-447.
- Finkel T, Holbrook NJ: Oxidants, oxidative stress and the biology of ageing. Nature 2000, 408:239-247.
- Brooks PJ, Wise DS, Berry DA, Kosmoski JV, Smerdon MJ, Somers RL, et al: The oxidative DNA lesion 8,5'-(S)-cyclo-2'-deoxyadenosine is repaired by the nucleotide excision repair pathway and blocks gene expression in mammalian cells. J Biol Chem 2000, 275:22355-22362.
- Kuraoka I, Bender C, Romieu A, Cadet J, Wood RD, Lindahl T: Removal of oxygen free-radical-induced 5',8-purine cyclodeoxynucleosides from DNA by the nucleotide excision-repair pathway in human cells. Proc Natl Acad Sci USA 2000. 97:3832-3837.
- Rapin I, Lindenbaum Y, Dickson DW, Kraemer KH, Robbins JH: Cockayne syndrome and xeroderma pigmentosum. Neurology 2000, 55:1442-1449.
- Andressoo JO, Mitchell JR, de WJ, Hoogstraten D, Volker M, Toussaint W, et al: An Xpd mouse model for the combined xeroderma pigmentosum/ Cockayne syndrome exhibiting both cancer predisposition and segmental progeria. Cancer Cell 2006, 10:121-132.
- Low GK, Fok ED, Ting AP, Hande MP: Oxidative damage induced genotoxic effects in human fibroblasts from Xeroderma Pigmentosum group A patients. Int J Biochem Cell Biol 2008, 40:2583-2595.
- Ting AP, Low GK, Gopalakrishnan K, Hande MP: Telomere attrition and genomic instability in xeroderma pigmentosum type-B deficient fibroblasts under oxidative stress. J Cell Mol Med 2010, 14(1-2):403-16.
- Lueken A, Juhl-Strauss U, Krieger G, Witte I: Synergistic DNA damage by oxidative stress (induced by H2O2) and nongenotoxic environmental chemicals in human fibroblasts. Toxicol Lett 2004, 147:35-43.
- Hande MP, Boei JJ, Natarajan AT: Induction and persistence of cytogenetic damage in mouse splenocytes following whole-body Xirradiation analysed by fluorescence in situ hybridization. II. Micronuclei. Int J Radiat Biol 1996, 70:375-383.
- Newman JP, Banerjee B, Fang W, Poonepalli A, Balakrishnan L, Low GK, et al: Short dysfunctional telomeres impair the repair of arsenite-induced oxidative damage in mouse cells. J Cell Physiol 2008, 214:796-809.
- Hande MP, Samper E, Lansdorp P, Blasco MA: Telomere length dynamics and chromosomal instability in cells derived from telomerase null mice. J Cell Biol 1999, 144:589-601.
- Collins AR: The comet assay for DNA damage and repair: principles, applications, and limitations. Mol Biotechnol 2004, 26:249-261.
- Liu Y, Hu N: Electrochemical detection of natural DNA damage induced by ferritin/ascorbic acid/H2O2 system and amplification of DNA damage by endonuclease Fpg. Biosens Bioelectron 2009, 25:185-190.
- Shackelford RE, Kaufmann WK, Paules RS: Oxidative stress and cell cycle checkpoint function. Free Radic Biol Med 2000, 28:1387-1404.
- Michalik V, Spotheim MM, Charlier M: Calculation of hydroxyl radical attack on different forms of DNA. J Biomol Struct Dyn 1995, 13:565-575.
- Klungland A, Hoss M, Gunz D, Constantinou A, Clarkson SG, Doetsch PW, et al: Base excision repair of oxidative DNA damage activated by XPG protein. Mol Cell 1999, 3:33-42.
- Shimizu Y, Iwai S, Hanaoka F, Sugasawa K: Xeroderma pigmentosum group C protein interacts physically and functionally with thymine DNA glycosylase. EMBO J 2003, 22:164-173.
- Shell SM, Li Z, Shkriabai N, Kvaratskhelia M, Brosey C, Serrano MA, et al: Checkpoint kinase ATR promotes nucleotide excision repair of UVinduced DNA damage via physical interaction with xeroderma pigmentosum group A. J Biol Chem 2009, 284:24213-24222.
- Wu X, Shell SM, Yang Z, Zou Y: Phosphorylation of nucleotide excision repair factor xeroderma pigmentosum group A by ataxia telangiectasia mutated and Rad3-related-dependent checkpoint pathway promotes cell survival in response to UV irradiation. Cancer Res 2006, 66:2997-3005.
- Schaeffer L, Roy R, Humbert S, Moncollin V, Vermeulen W, Hoeijmakers JH, et al: DNA repair helicase: a component of BTF2 (TFIIH) basic transcription factor. Science 1993, 260:58-63.

- Schaeffer L, Moncollin V, Roy R, Staub A, Mezzina M, Sarasin A, et al: The ERCC2/DNA repair protein is associated with the class II BTF2/TFIIH transcription factor. EMBO J 1994, 13:2388-2392.
- Andressoo JO, Weeda G, de WJ, Mitchell JR, Beems RB, van SH, et al: An Xpb mouse model for combined xeroderma pigmentosum and cockayne syndrome reveals progeroid features upon further attenuation of DNA repair. Mol Cell Biol 2009, 29:1276-1290.
- de Boer J, Donker I, de WJ, Hoeijmakers JH, Weeda G: Disruption of the mouse xeroderma pigmentosum group D DNA repair/basal transcription gene results in preimplantation lethality. Cancer Res 1998, 58:89-94.
- de WH, Sonneveld E, de WJ, Esveldt-van LR, Hoeijmakers JH, Vrieling H, et al: Cell-type-specific consequences of nucleotide excision repair deficiencies: Embryonic stem cells versus fibroblasts. DNA Repair (Amst) 2008. 7:1659-1669.
- Liu SY, Wen CY, Lee YJ, Lee TC: XPC silencing sensitizes glioma cells to arsenic trioxide via increased oxidative damage. *Toxicol Sci* 2010, 116:183-193.
- Rybanska I, Gursky J, Faskova M, Salazar EP, Kimlickova-Polakovicova E, Kleibl K, et al: Newly identified CHO ERCC3/XPB mutations and phenotype characterization. Mutagenesis 2010, 25:179-185.
- Munoz P, Blanco R, Flores JM, Blasco MA: XPF nuclease-dependent telomere loss and increased DNA damage in mice overexpressing TRF2 result in premature aging and cancer. Nat Genet 2005, 37:1063-1071.
- Crabbe L, Verdun RE, Haggblom CI, Karlseder J: Defective telomere lagging strand synthesis in cells lacking WRN helicase activity. Science 2004, 306:1951-1953.
- Du X, Shen J, Kugan N, Furth EE, Lombard DB, Cheung C, et al: Telomere shortening exposes functions for the mouse Werner and Bloom syndrome genes. Mol Cell Biol 2004, 24:8437-8446.
- Opresko PL, von KC, Laine JP, Harrigan J, Hickson ID, Bohr VA: Telomerebinding protein TRF2 binds to and stimulates the Werner and Bloom syndrome helicases. J Biol Chem 2002, 277:41110-41119.
- Opresko PL, Mason PA, Podell ER, Lei M, Hickson ID, Cech TR, et al: POT1 stimulates RecQ helicases WRN and BLM to unwind telomeric DNA substrates. J Biol Chem 2005, 280:32069-32080.
- Lane DP, Midgley CA, Hupp TR, Lu X, Vojtesek B, Picksley SM: On the regulation of the p53 tumour suppressor, and its role in the cellular response to DNA damage. *Philos Trans R Soc Lond B Biol Sci* 1995, 347:83-87.
- Banin S, Moyal L, Shieh S, Taya Y, Anderson CW, Chessa L, et al: Enhanced phosphorylation of p53 by ATM in response to DNA damage. Science 1998. 281:1674-1677.
- Weinberg WC, Denning MF: P21Waf1 control of epithelial cell cycle and cell fate. Crit Rev Oral Biol Med 2002, 13:453-464.
- Zhang Y, Rohde LH, Wu H: Involvement of nucleotide excision and mismatch repair mechanisms in double strand break repair. Curr Genomics 2009, 10:250-258.
- 52. Fotedar R, Bendjennat M, Fotedar A: Role of p21WAF1 in the cellular response to UV. Cell Cycle 2004, **3**:134-137.
- Soria G, Podhajcer O, Prives C, Gottifredi V: P21Cip1/WAF1 downregulation is required for efficient PCNA ubiquitination after UV irradiation. Oncogene 2006, 25:2829-2838.
- Mita AC, Mita MM, Nawrocki ST, Giles FJ: Survivin: key regulator of mitosis and apoptosis and novel target for cancer therapeutics. Clin Cancer Res 2008, 14:5000-5005.
- Dip R, Camenisch U, Naegeli H: Mechanisms of DNA damage recognition and strand discrimination in human nucleotide excision repair. DNA Repair (Amst) 2004, 3:1409-1423.
- Wood RD: DNA damage recognition during nucleotide excision repair in mammalian cells. Biochimie 1999, 81:39-44.
- 57. Duan J, Duan J, Zhang Z, Tong T: Irreversible cellular senescence induced by prolonged exposure to H2O2 involves DNA-damage-and-repair genes and telomere shortening. Int J Biochem Cell Biol 2005, 37:1407-1420.

doi:10.1186/2041-9414-1-16

Cite this article as: Gopalakrishnan *et al.*: Hydrogen peroxide induced genomic instability in nucleotide excision repair-deficient lymphoblastoid cells. *Genome Integrity* 2010 1:16.

Submit your next manuscript to BioMed Central and take full advantage of:

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your manuscript at www.biomedcentral.com/submit

