Threshold Effects of Nitric Oxide-Induced Toxicity and Cellular Responses in Wild-Type and p53-Null Human Lymphoblastoid Cells

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Toxicity induced by nitric oxide (NO•) has been extensively investigated in many in vitro and in vivo experimental models. Recently, our laboratories found that both concentration and cumulative total dose are critical determinants of cell death caused by NO•. Here, we report results of studies designed to define total dose thresholds and threshold effects for several NO•-induced toxicity and cellular responses and to determine impacts of p53 on them. We exposed human lymphoblastoid TK6 cells harboring wild-type p53 and isogenic p53-null NH32 cells to NO• delivered by a membrane delivery system. Cells were exposed at a steady state concentration of 0.6 \( \mu \text{M} \) for varying lengths of time to deliver increasing cumulative doses (expressed in units of \( \mu \text{M min} \)), and several end points of cytotoxicity and mutagenesis were quantified. Threshold doses for NO•-induced cytotoxicity were 150 \( \mu \text{M min} \) in TK6 cells and 300 \( \mu \text{M min} \) in NH32 cells, respectively. Threshold doses for NO•-induced apoptosis were identical to those for cytotoxicity, but mitochondrial depolarization thresholds were lower than those for cytotoxicity and apoptosis in both cell types. To gain insight into underlying mechanisms, cells of both types were exposed to sublethal (33% of cytotoxicity threshold), cytotoxicity threshold, or toxic (twice the cytotoxicity threshold) doses of NO•. In TK6 cells (p53), the sublethal threshold dose induced DNA double-strand breaks, but nucleobase deamination products (xanthine, hypoxanthine, and uracil) in DNA were increased only modestly (<50%) by toxic doses. Increased mutant fraction at the thymidine kinase gene (\( TK1 \)) locus was observed only at the toxic dose of NO•. Treatment of NH32 cells with NO• at the threshold or toxic dose elevated mutagenesis of the \( TK1 \) gene, but did not cause detectable levels of DNA double-strand breaks. At similar levels of cell viability, the frequency of DNA recombinational repair was higher in p53-null NH32 cells than in wild-type TK6 cells. NO• treatment induced p53-independent cell cycle arrest predominately at the S phase. Akt signaling pathway and antioxidant proteins were involved in the modulation of toxic responses of NO•. These findings indicate that exposure to doses of NO• at or above the cytotoxicity threshold dose induces DNA double-strand breaks, mutagenesis, and protective cellular responses to NO• damage. Furthermore, recombinational repair of DNA may contribute to resistance to NO• toxicity and potentially increase the risk of mutagenesis. The p53 plays a central role in these responses in human lymphoblastoid cells.

Introduction

At low concentrations, nitric oxide (NO•) acts as a signaling molecule with regulatory roles in many physiological processes. However, high levels of NO•, such as those occurring during inflammation, can damage DNA, RNA, lipids, and proteins, leading to mutations and altered cell physiology that are hallmarks of the process of carcinogenesis (1, 2). Recently, our laboratories have found that both concentration and cumulative total dose are critical in initiating NO• toxicity (3). The threshold NO• concentration for cell death in human lymphoblastoid TK6 cells harboring wild-type p53 was \( \sim 0.5 \mu \text{M} \), while the comparable cumulative dose threshold was \( \sim 150 \mu \text{M min} \). If neither or only one threshold was exceeded, the number of viable cells declined in a dose-dependent manner, indicating that both steady state NO• levels and cumulative total doses above the cytotoxic thresholds are required for induction of cell death (3, 4). Wink and co-workers also found that different concentration thresholds of NO• released from donor drugs elicited a discrete set of signal transduction pathways associated with cell growth and apoptosis in human breast cancer MCF7 cells (5). At low steady state concentrations (<50 nM), NO• induced extracellular signal-regulated kinase (ERK) phosphorylation via a guanylate cyclase-dependent mechanism; hypoxic inducible factor 1 (HIF-1) accumulation was associated with an intermediate amount of NO• (>100 nM), whereas p53 serine 15 phosphorylation occurred at considerably higher levels (>300 nM). The dose-dependent effects of synthetic NO• donors were mimicked by activated macrophages cocultured with MCF7 cells at varying ratios (5). These findings suggest that thresholds, which may be different for various cellular markers, are critical determinants in triggering physiological and toxic responses of NO•.

The p53 tumor suppressor gene is known as a critical cellular gatekeeper for growth and division through its involvement in
damage-induced G1 arrest, DNA repair, apoptosis, and gene amplification (6). Our group and others have found that p53 status strongly affected NO*-induced cellular stress, DNA damage, mutagenesis, and apoptotic signaling pathways (7–10). The p53-null human lymphoblastoid cells NH32 cells were much more resistant to NO*-induced cell killing as compared with wild-type TK6 cells (3, 4). Preliminary evidence revealed that p53 normally suppressed homologous recombination events that might otherwise contribute to genomic instability (11). Furthermore, we showed that repair processes that involved homologous recombination were pivotal in preventing NO*-induced toxicity, at least in Escherichia coli (12, 13). Consequently, one possibility is that in the absence of normal p53, the levels of homology directed repair may rise, contributing to both resistance to toxicity and potentially increasing the risk of deleterious sequence rearrangements. Although homologous recombination induced by NO* has been observed in mouse embryonic stem cells (14), the role of p53 in modulating the susceptibility of cells to NO*-induced homologous recombination events has not been reported. Furthermore, it was not known how p53 status would affect the threshold exposures required to elicit cellular toxicity and/or homologous recombination repair.

In this study, we first determined the thresholds for NO*-induced cytotoxicity, mitochondrial depolarization, and apoptosis in human lymphoblastoid TK6 and NH32 cells. Three NO* doses for each cell type were established, namely, 33% of the threshold for cytotoxicity (designated sublethal), the cytotoxicity threshold (toxic). These doses were then used in further experiments to investigate other dose-related end points of cellular response, as well as the role of p53 in these processes. A key feature was the use of a NO* delivery system specifically designed to provide controlled, steady state concentrations of NO* and molecular oxygen (O2) mimicking the chemical environment thought to exist in inflamed tissues (3, 4, 15, 16). NO* and O2 are delivered into culture medium by diffusion through semipermeable Silastic tubing to achieve constant and physiologically relevant levels of both gases (3, 4, 15, 16). Cells of the TK6 lymphoblastoid line, derived from a human spleen nearly 30 years ago, (17, 18) have been used extensively in mutagenicity and genotoxicity studies (3, 4, 8–10) and in some studies were used in conjunction with NH32 cells, an isogenic derivative of TK6 cells in which both alleles of the p53 gene were knocked out (3, 4, 19). We found that the NO* cytotoxicity threshold is associated with DNA double-strand breaks, mutagenesis, and protective cellular responses to NO*-induced damage and that recombinational DNA repair may contribute both to resistance to NO* toxicity and potentially increase the risk of mutagenesis, in which p53 plays a central role.

Materials and Methods

Chemicals and Cells. TK6 cells were kindly provided by Dr. W. G. Thilly (Massachusetts Institute of Technology), and NH32 cells were provided by Dr. C. C. Harris (National Cancer Institute). Cell culture reagents were purchased from Cambrex (Walkersville, MD). All gases were purchased from BOC Gases (Edison, NJ), and Silastic tubing (0.058 in. i.d., 0.077 in. o.d.) was purchased from Dow Corning (Midland, MD). Propidium iodide, nucleas P1, alkaline phosphatase, trilfluorothymidine, BrdU, and demecolcine were purchased from Sigma Chemical Co.; phosphodiesterase I was purchased from USB Corp. (Cleveland, OH); the MTT assay kit and RNase were from Roche Diagnostics GmbH (Mannheim, Germany); JC-1 (5,5′,6,6′-tetracloro-1,1′,3,3′-tetraethylbenzimidazolylcarbocyanine iodide) was purchased from Molecular Probes (Eugene, OR); the annexin V-FITC apoptotic assay kit was from Clontech Laboratories (Palo Alto, CA); the Puceredene genomic DNA purification kit was from Gentra Systems, Inc. (Minneapolis, MN); and the supersignal ultrachemiluminescence kit was from Pierce (Rockford, IL). Primary monoclonal anti-human Cu/Zn-superoxide dismutase (SOD1) antibody and monoclonal anti-human glutathione peroxidase (GPX) antibody were purchased from MBL (Woburn, MA); polyclonal anti-human catalase antibody was from Calbiochem (San Diego, CA); monoclonal anti-p53 antibody (Ab-6) was from Oncogene; PTEN antibody was from Upstate (Lake Placid, NY); and anti-Akt antibody, anti-NF-xB antibody, PI3 kinase antibody, and phosphor-Akt (Ser 473) antibody were from Cell Signaling Technology (Beverly, MA). Secondary goat anti-rabbit- or anti-mouse IgG conjugated to horseradish peroxidase was provided by Bio-Rad (Hercules, CA), and anti-actin antibody was provided by Oncogene (Cambridge, MA).

Cell Culture and NO* Treatment. TK6 and NH32 cells were maintained in exponentially growing suspension cultures at 37 °C in a humified, 5% CO2 atmosphere in RPMI medium1640 supplemented with 10% heat-inactivated donor calf serum, 100 units/mL penicillin, 100 μg/mL streptomycin, and 2 mM l-glutamine. Cell suspensions at a density of 5 × 10⁶ cells/mL in 115 mL of RPMI medium 1640 with calf serum were exposed, at 37 °C, to 1% NO*, delivered by diffusion through permeable Silastic tubing at a steady state concentration of 0.6 μM, as reported previously (3, 4). A mixture of 50% O2 and 5% CO2 was delivered through a second tubing loop to maintain the liquid O2 level near air saturation. The total NO* dose delivered into the medium was controlled by varying the exposure time and expressed as μM min.

Cells exposed to argon gas under the same conditions served as negative controls. For sister chromatid exchange (SCE) assays, cells were exposed to 10% NO*, at a steady state concentration of 1.8 μM.

Cell Viability Analysis. Cell viability 48 h after treatment was determined by both trypan blue exclusion (8, 9) and MTT assay following the manufacturer’s instructions. After determination of the thresholds for NO*-induced cell death, three NO* doses for each cell type, namely, sublethal, threshold, and toxic as defined above, were used in further experiments to investigate other dose-associated endpoints.

Cell Cycle Analysis. Cells were harvested 20 h after NO* treatment, washed twice in PBS, and fixed in 70% cold ethanol overnight. Cells were suspended in 1% BSA (bovine serum albumin)—PBS solution containing 50 μg/mL propidium iodide and 50 μg/mL RNase incubated at 37 °C for 30 min and analyzed on a Becton Dickinson FACScan. Cell fit analysis determined the percentage of cells in a specific stage of the cell cycle.

Mitochondrial Membrane Potential (MMP) and Apoptosis Analysis. Mitochondrial depolarization and apoptosis in the cells harvested 48 h after NO* treatment were quantitatively evaluated by a Becton Dickinson FACScan equipped with CellQuest software following JC-1 or annexin V-FITC and propidium iodide staining, respectively, according to methods reported previously (8, 9).

Analysis of DNA Double-Strand Breaks (Neutral Comet Assay). Following NO* treatment, cells at a density of 1 × 10⁶ cells/mL were combined with molten LMAgarose (at 42 °C) at a ratio of 1:10, and 50 μL was immediately pipetted onto CometSlides (Trevigen, Gaithersburg, MD) (8, 9). After incubation in lysis buffer (2.5 M sodium chloride, 100 mM EDTA, pH 10, 10 mM Tris base, 1% sodium lauryl sarcosinate, and 0.01% Triton X-100), slides were electrophoresed, photographed, and analyzed with the Komet 4.2 Single Cell Gel Electrophoresis Analysis (Kinetic Imaging Limited, Liverpool, United Kingdom). Olive tail moment, defined as the product of percentage DNA in the tail and displacement between the position of the mean centers of mass in the heads and tails (20), was determined for at least 40 cells per sample. Cells treated with argon gas were used as negative controls, and cells treated with 100 μM H2O2 in RPMI medium 1640 for 20 min at 4 °C were used as positive controls.

SCE Assay. The SCE assay was performed following the method reported previously (21). Immediately after NO* treatment, aliquots
of cell suspensions were incubated with 20 μM BrdU in RPMI medium 1640 for 39 h (two cell replication cycles), followed by further incubation with 0.1 μg/mL of demecolcine for an additional 6 h. The cells were subsequently harvested, resuspended, and incubated for 15 min at 37 °C in hypotonic solution (0.2% potassium chloride, 0.2% sodium citrate, and 10% fetal bovine serum) and fixed in Carnoy’s solution. To produce “harlequin” chromosomes, slides were stained in Hoechst 33258 (5 μg/mL) for 20 min, mounted in 0.067 M Sorensen’s buffer with a coverslip, and exposed to a General Electric 15 W black light bulb at 65 °C in 20 × SSC for 20 min. After staining with 5% Giemsa solution, 20 metaphase spreads were counted for each sample.

LC/MS/MS Quantification of DNA Base Deamination Lesions. Deoxynucleoside deamination products in cells treated with NO· were quantitatively evaluated by LC/MS/MS immediately after exposure using a previously reported LC/MS method (16) adapted to LC/MS/MS. Genomic DNA was isolated using a Puregene genomic DNA purification kit according to the manufacturer’s instructions, with deaminase inhibitors cof orm ycin and tetracy d roidine added into cell lysis buffer at final concentrations of 5 and 125 μg/mL, respectively. Genomic DNA (50 μg) was hydrolyzed with nuclease P1, phosphodiesterase 1, and alkaline phos- phatase in the presence of deaminase inhibitors and appropriate amounts of iso t o-labeled internal standards. The resulting deoxy- nucleoside mixture was resolved by CI8 reversed phase HPLC, and the fractions containing 2′-deoxy-xanthosine (dX), 2′-deoxy- ximine (di), and 2′-deoxyuridine (dU) were collected for subsequent LC/MS/MS quantification in an API 3000 triple quadrupole mass spectrometer (Applied Biosystems, Foster City, CA).

Determination of Mutant Fraction (MF). After NO· treatment, cells were grown for 6–10 days to allow phenotypic expression and then plated in selective medium to determine MF at the thymidine kinase gene (TK1) locus (19, 22). A total of 24 × 10^6 TK6 cells from each treatment group were transferred to ten 96 well plates at densities of 40000 cells per well in medium containing 2 μg/mL of trifluorothymidine (TFT) to select TK1 mutants. In the pilot experiments, TK1 mutants in NH32 cells were not measurable due to the large number of surviving cells under the experimental conditions used for TK6 cells, and optimal conditions for studies with NH32 cells were determined to be plating at a density of 20000 cells per well in medium containing 4 μg/mL of TFT. Cells from each culture were also plated at 1 cell per 100 μL per well in the absence of TFT to determine plating efficiency. After 2 weeks of incubation, colonies were counted and MF was calculated with the Poisson distribution (19, 22). The spontaneous mutation rate was estimated from the argon-treated cells, and the cells treated with 4-nitroquinoline 1-oxide (4-NQO; 140 ng/mL for 1.5 h) were used as a positive control.

Western Blot Analysis. Cells were harvested at the indicated times after NO· treatment, and 20 μg aliquots of whole cell lysate proteins were denatured, resolved on 15% SDS–PAGE gels, and electrotransferred at 180 mA for 1 h onto a poly(vinylidene difluoride) membrane (Bio-Rad). Blots were probed with primary antibodies overnight at 4 °C, followed by a secondary goat anti-rabbit or anti-mouse IgG conjugated to horseradish peroxidase (1:3000 dilution) and determination of supersignal ultrachemilumi- nescence by exposure to Hyperfilm ECL (Amersham Pharmacia, Piscataway, NJ). To control for protein loading, membranes were stripped and reprobed with anti-actin antibody (1:3000 dilution) and determination of supersignal ultrachemiluminescence by exposure to Hyperfilm ECL (Amersham Pharmacia, Piscataway, NJ). To control for protein loading, membranes were stripped and reprobed with anti-actin antibody (1:3000 dilution) and determination of supersignal ultrachemiluminescence by exposure to Hyperfilm ECL (Amersham Pharmacia, Piscataway, NJ). To control for protein loading, membranes were stripped and reprobed with anti-actin antibody (1:3000 dilution) and determination of supersignal ultrachemiluminescence by exposure to Hyperfilm ECL (Amersham Pharmacia, Piscataway, NJ).

Statistical Analysis. All experiments were repeated 2–4 times after experimental conditions were optimized. Statistical analysis was performed using a two-tailed Student’s t-test, and p < 0.05 was considered to be statistically significant.

Results

Using a recently developed controlled NO· delivery system (3, 4, 15, 16), TK6 and NH32 cells were exposed to a steady state concentration of 0.6 μM NO· and 180 μM O3 for 1–24 h, which resulted in cumulative total NO· doses ranging from 35 to 920 μM min. As shown in Figure 1, NO· induced cell death in a dose-dependent manner in both cell lines; both the trypan blue exclusion and the MTT assays produced similar results with 48 h cytotoxicity (data not shown). However, NH32 cells were more resistant to NO·-induced growth inhibition or killing than TK6 cells. For example, when treated with 600 μM min of NO·, cell viability 48 h after exposure was approximately 5% in TK6 cells, whereas it was nearly 50% in NH32 cells, as compared to argon-treated controls. Thresholds for NO·-induced cell death were shown to be approximately 150 μM min in TK6 cells and 225 and 300 μM min in NH32 cells. Data represent the means ± SDs of three independent experiments; each was done in duplicate.

Figure 1. Cell viability in TK6 and NH32 cells 48 h after NO· treatment, as determined by MTT assay. Data represent the mean of 2–4 duplicate experiments. Standard deviations were less than 15% (not shown).

Figure 2. NO· dose-dependent MMP loss and apoptosis in TK6 and NH32 cells 48 h after exposure. The threshold doses for NO-induced MMP loss and apoptosis were approximately 35 and 150 μM min in TK6 cells and 225 and 300 μM min in NH32 cells. Data represent the means ± SDs of three independent experiments; each was done in duplicate.

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Next, we assessed the relationship of mitochondrial depolarization and apoptosis to the NO· cytotoxicity threshold. As shown in Figure 2, NO· treatment of TK6 and NH32 cells resulted in a dose-dependent MMP loss and apoptosis in both cell lines 48 h after exposure, although TK6 cells were comparatively more sensitive to mitochondrial damage and apoptosis. The threshold doses for NO·-induced apoptosis were identical to the cytotoxicity thresholds in both cell lines (150 μM min in TK6 cells and 300 μM min in NH32 cells). Thresholds for mitochondrial depolarization (35 μM min in TK6
Table 1. Effects of NO* on Cell Cycle in TK6 and NH32 Cells 20 h after Treatment (N = 4)

<table>
<thead>
<tr>
<th>total NO* dose (µM min)</th>
<th>G1/G0</th>
<th>S</th>
<th>G2/M</th>
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<td>TK6 cells</td>
<td></td>
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<tr>
<td>argon</td>
<td>32.0 ± 4.1</td>
<td>50.5 ± 7.9</td>
<td>17.6 ± 4.6</td>
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<td>50</td>
<td>37.0 ± 0.9</td>
<td>42.3 ± 0.4</td>
<td>20.7 ± 0.6</td>
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<td>150 (threshold)</td>
<td>40.6 ± 0.5</td>
<td>40.6 ± 1.0</td>
<td>18.8 ± 1.4</td>
</tr>
<tr>
<td>300</td>
<td>30.4 ± 2.3</td>
<td>59.8 ± 1.6</td>
<td>9.9 ± 3.8</td>
</tr>
<tr>
<td>NH32 cells</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>argon</td>
<td>43.5 ± 3.3</td>
<td>42.6 ± 4.8</td>
<td>13.9 ± 3.2</td>
</tr>
<tr>
<td>150</td>
<td>27.2 ± 1.7</td>
<td>53.2 ± 2.3</td>
<td>19.6 ± 1.4</td>
</tr>
<tr>
<td>300 (threshold)</td>
<td>37.1 ± 3.5</td>
<td>52.2 ± 3.1</td>
<td>10.8 ± 2.7</td>
</tr>
<tr>
<td>600</td>
<td>9.6 ± 1.0</td>
<td>57.4 ± 2.0</td>
<td>33.0 ± 1.8***</td>
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</table>

*p < 0.05 and **p < 0.01, as compared with argon-treated controls.

Figure 3. Box-and-whisker plots of Olive tail moments from neutral comet assays of TK6 and NH32 cells right after NO* treatment. At least 40 cells were analyzed in each sample. Cells treated with argon (Ar) gas or with H2O2 and were used as negative and positive controls, respectively. Data from H2O2 treatment were not shown. *p < 0.01, as compared with argon-treated control.

Table 2. SCE Induced by NO* in TK6 and NH32 Cells (N = 4)

<table>
<thead>
<tr>
<th>NO* dose (µM min)</th>
<th>TK6 cells</th>
<th>NH32 cells</th>
</tr>
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<tbody>
<tr>
<td>argon</td>
<td>7.3 ± 0.8</td>
<td>7.9 ± 0.5</td>
</tr>
<tr>
<td>300</td>
<td>8.4 ± 0.4</td>
<td>8.8 ± 0.2</td>
</tr>
<tr>
<td>600</td>
<td>19.8 ± 3.7*</td>
<td>24.7 ± 1.9**</td>
</tr>
</tbody>
</table>

*p < 0.01 and **p < 0.05, as compared with argon controls.

Figure 4. Formation of dU, dX, and dl in DNA of TK6 (solid) and NH32 (open) cells exposed to NO* at the sublethal, threshold, and toxic doses (50, 150, and 300 µM min for TK6 cells and 150, 300, and 600 µM min for NH32 cells), respectively. The only statistically significant increase in a DNA deamination product over control occurred with dU in NH32 cells treated with 600 µM min NO* (p < 0.05). Data represent the mean ± SD for N = 4.

Table 3. Mutagenesis in the TK1 Gene of TK6 and NH32 Cells Exposed to NO* (N = 4)

<table>
<thead>
<tr>
<th>NO* dose (µM min)</th>
<th>TK6 cells</th>
<th>NH32 cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>argon</td>
<td>16.6 ± 2.3</td>
<td>18.9 ± 0.5</td>
</tr>
<tr>
<td>50</td>
<td>18.9 ± 0.5</td>
<td>19.8 ± 0.5</td>
</tr>
<tr>
<td>150 (threshold)</td>
<td>30.6 ± 2.5**</td>
<td>28.6 ± 2.5**</td>
</tr>
<tr>
<td>300</td>
<td>28.4 ± 1.8**</td>
<td>71.2 ± 7.8 *</td>
</tr>
<tr>
<td>4-NQO</td>
<td>26.1 ± 3.7*</td>
<td>4-NQO</td>
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</table>

*p < 0.01 and **p < 0.05, as compared with argon controls.

To test the hypothesis that p53-regulated DNA recombination repair is responsible for the differences in DNA double-strand breaks and mutagenesis between TK6 and NH32 cells, SCE was measured in both cell types treated with NO* at 300 and 600 µM min (Table 2). Exposure to 300 µM min NO* did not increase SCE frequency in either cell type, but it was significantly elevated in both TK6 and NH32 cells treated with 600 µM min of NO*, as compared with argon-treated cells. TK6 and NH32 cells showed similar SCE frequency when treated with argon gas. As seen in Figure 1, TK6 cells treated with 300 µM min of NO* and NH32 cells treated with 600 µM min of NO* had a similar cell viability of about 50%; however, SCE frequency was greater in NH32 cells than in TK6 cells under the equivalently cytotoxic conditions (p < 0.01).

Under the same experimental conditions, we analyzed nucleobase deamination in TK6 and NH32 cells immediately after NO* treatment. NO* treatment led to elevated formation of dU, dX, and dl in DNA of TK6 and NH32 cells in a dose-dependent fashion, but the rise was independent of p53 status (Figure 4). The only statistically significant increase in a DNA deamination product over controls occurred with dU in NH32 cells treated with 600 µM min of NO* (70.9 ± 15.2 lesions per 106 nt) (p < 0.05) (Figure 4). NO*−induced changes in MF at the TK1 locus in TK6 and NH32 cells are summarized in Table 3. MF in TK6 cells increased 2.8-fold when treated with the toxic NO* dose (300 µM min) as compared with argon-treated cells (p < 0.01),

Cells and 225 µM min in NH32 cells) were lower than the cytotoxicity and apoptosis thresholds in both cell types. Further studies were then performed using three NO* doses relative to cell death threshold for each cell type, namely, 33% of the threshold (sublethal), the threshold, and twice the threshold dose (toxic). Cell cycle changes were measured in TK6 and NH32 cells 20 h after exposure to these doses, with results shown in Table 1. While NO* treatment induced cell cycle arrest, mainly at the S phase, in TK6 cells treated with toxic dose (300 µM min), this response was observed in NH32 cells at the lower threshold (sublethal), the threshold, and twice the threshold dose (toxic). Cell cycle changes were measured in TK6 and NH32 cell death threshold for each cell type, namely, 33% of the threshold (sublethal), the threshold, and twice the threshold dose (toxic). Cell cycle changes were measured in TK6 and NH32 cells 20 h after exposure to these doses, with results shown in Table 1. While NO* treatment induced cell cycle arrest, mainly at the S phase, in TK6 cells treated with toxic dose (300 µM min), this response was observed in NH32 cells at the lower sublethal dose (150 µM min). Exposure of NH32 cells to the toxic NO* dose (600 µM min) caused arrest at both S and G2/M phases. Cell cycle behavior of argon-exposed controls was not significantly different from unexposed cells (data not shown).

Induction of DNA double-strand breaks by NO* was evaluated by neutral comet assay immediately after NO* treatment, with the results shown in Figure 3. The Olive tail moment was significantly increased in TK6 cells treated with NO* at threshold (150 µM min) (14 ± 16) or toxic (300 µM min) (26 ± 36) doses, as compared with argon-treated control (2.7 ± 2.1) (p < 0.01). In contrast, DNA double-strand breaks did not show significant changes in NH32 cells treated with NO* at any dose. As a positive control, 100 µM H2O2 treatment increased significantly DNA tail moment in TK6 cells (8.7 ± 2.9 vs 2.7 ± 2.1, p < 0.01) but not in NH32 cells (8.4 ± 6.3 vs 5.8 ± 4.1) (p > 0.05).
Doses were ineffective (p) treated cells (showed increases in MF of 1.7-fold as compared with argon ineffective (p) intervals from two to three independent experiments. *Values are mean quantitative densitometric values and 95% confidence intervals from two to three independent experiments. **Values are mean quantitative densitometric values and 95% confidence intervals from two to three independent experiments. *p < 0.05, as compared with argon-treated controls.

whereas the sublethal (50 μM min) and threshold (150 μM min) doses were ineffective (p > 0.05). NH32 cells treated with NO* at the threshold (300 μM min) or toxic doses (600 μM min) showed increases in MF of 1.7-fold as compared with argon-treated cells (p < 0.05); the sublethal dose (150 μM min) was ineffective (p > 0.05). Spontaneous as well as NO*- and 4-NQO-induced MF were significantly higher in NH32 cells than in TK6 cells (p < 0.05 or p < 0.01) when compared at the corresponding sublethal, threshold, and toxic doses, or the same dose levels.

We next assessed changes in levels of the antioxidant proteins SOD1, catalase, and GPX in response to NO* treatment. SOD1 protein in argon-treated control cells was higher in TK6 cells than in NH32 cells; after NO* treatment, levels were significantly reduced in TK6 cells treated with the toxic dose (300 μM min) (p < 0.05) but increased in NH32 cells treated with threshold (300 μM min) or toxic doses (600 μM min) (p < 0.05) (Figure 5A). Expression of catalase protein was slightly lower in TK6 cells than in NH32 cells treated with argon gas, but levels were substantially upregulated only in TK6 cells (p < 0.05) after NO* treatment (Figure 5B). Under the same conditions, GPX levels were unchanged in both cell types (Figure 5C). The prosurvival and antiapoptotic protein Akt (23) and transcription factor NF-κB p50 and p65 proteins (24) were prominently expressed in argon-treated TK6 and NH32 cells. Levels of Akt and NF-κB p65 were significantly suppressed following NO* treatment, even at the sublethal dose (50 μM min) in TK6 cells (p < 0.05 or p < 0.01), but not at any dose in NH32 cells (Figure 6).

Components of the Akt signaling pathway were characterized following treatment with the toxic doses of NO* in TK6 (300 μM min) and NH32 (600 μM min) cells. p53 protein levels increased progressively in TK6 cells through 48 h after exposure, reaching a maximum elevation of 228% as compared with argon-treated cells; p53 protein was undetectable in TK6 cells (Figure 7A). PTEN tumor suppressor protein, known to be up-regulated by p53 (25), was virtually undetectable in argon-treated TK6 cells, but after NO* treatment, levels increased 130% at 24 h, declining to 117% at 48 h. As expected, while PTEN was expressed in argon-treated NH32 cells, it remained unchanged after NO* exposure (Figure 7B). PI3 kinase protein, negatively regulated by PTEN (26), was undetectable in TK6 cells treated with argon or NO*. Very low levels of PI3 kinase protein were detected in argon-treated NH32 cells, and its level increased to 115% at 24 h and 123% at 48 h after treatment (Figure 7C). Phosphorylated Akt protein was prominently expressed in argon-treated TK6 and NH32 cells, and its levels decreased 25-38% from control levels over a 48 h period in TK6 cells after NO* treatment but showed no change in NH32 cells during the same period (Figure 7D).

Discussion

The goal of this study was to determine relationships between the threshold for NO*-induced cytotoxicity and more mechanistically informative biological end points such as genotoxicity, apoptosis, DNA repair, and cell signaling parameters. A key
feature was the controlled delivery of NO\(^*\) under conditions designed to mimic the in vivo situation found in inflamed tissues. In earlier related investigations, we found that both concentration and cumulative total dose were critical in initiating NO\(^*\) toxicity (3, 4). Here, we confirmed that at a steady state concentration of 0.6 \(\mu\)M, the thresholds for NO\(^*\)-induced cell death were approximately 150 \(\mu\)M min in TK6 cells and 300 \(\mu\)M min in NH32 cells. If the thresholds were not exceeded, cells proliferated normally. Interestingly, the thresholds for NO\(^*\)-induced apoptosis were identical to the cytotoxicity thresholds, whereas the thresholds for mitochondrial depolarization were lower in both cell types, suggesting that mitochondria were relatively sensitive to damage by NO\(^*\). We measured necrosis in TK6 and NH32 cells 48 h after NO\(^*\) treatment with the sublethal, threshold, and toxic doses, by flow cytometry following propidium iodide staining, and found that necrosis did not constitute a significant part of cell loss (data not shown). Apoptosis accounted for only a small percentage of reduced cell viability in both cell lines. For example, treatment of TK6 cells with 300 \(\mu\)M min of NO\(^*\) resulted in approximately 55% cell viability as compared with argon-treated control cells; yet, apoptosis was responsible for only 9% of the cell loss seen (Figures 1 and 2). The remainder of the cell loss was accounted for by NO\(^*\)-induced p53-independent cytostasis and cell cycle arrest at the S phase or both S and G2/M phases (Table 1). Other investigators have reported that NO\(^*\) produced from DETA NONOate at a steady state concentration of 0.5 \(\mu\)M induced cytostasis and cell cycle arrest at G1 phase in human breast cancer cells and that down-regulation of cyclin D1 played an important role in these processes (27). Cell cycle arrest induced by DNA damaging agents is presumed to provide the cell time to repair potentially mutagenic and cytotoxic DNA damage (28). The S phase arrest is thought to result from damage in DNA replicons, suppressing DNA synthesis or elongation (29, 30). If the cell can repair the damage, replication resumes, with the production of replacement cells; if damage is unrepairable, apoptosis may be initiated to eliminate irreversible injured cells (28, 30). Our findings derived from cell cycle, apoptosis, and necrosis analyses, in comparison with the cell viability data, suggest that cytostasis and mitochondria-dependent apoptosis are responsible for NO\(^*\)-induced cell viability loss at biologically relevant exposure conditions.

We found that cytotoxicity threshold and higher doses of NO\(^*\) induced a significant number of DNA double-strand breaks in wild-type TK6 cells but not in p53-null NH32 cells, implying that p53 is required for most, if not all, NO\(^*\)-induced DNA double-strand breaks, a potent inducer of apoptosis (31). Substantial evidence has demonstrated that NO\(^*\)-induced DNA double-strand breaks are caused mainly by enzymatic processing of the primary lesions created by NO\(^*\) exposure (32, 33). The majority are base lesions, most of which are repaired via the base excision repair (BER) pathway (13, 34). During excision repair, single-strand breaks are formed as intermediates, which can lead to formation of double-strand breaks, possibly during DNA replication (12, 13, 21). We observed that NO\(^*\)-induced DNA base deamination in a dose-dependent fashion but independent of p53 status. It is noteworthy that these are steady state levels of dL, dX, and dU, and many DNA base lesions have probably been repaired by the BER system or removed shortly after formation, potentially by DNA glycosylase (13, 35). p53 has been shown to modulate the efficiency of BER in mammalian cells (36) and yet to interact directly with proteins and DNA intermediates that are formed during homologous recombination (37–39), which is a major pathway for DNA double-strand break repair (40, 41).

Our SCE analysis showed that under equivalently cytotoxic conditions, p53-null NH32 cells are susceptible to NO\(^*\) damage-induced recombination events. Because DNA double-strand breaks are potent inducers of homologous recombination (30, 31, 34, 40), it is paradoxical that the levels of DNA double-strand breaks and SCEs are opposite to the expected relationship in these data: high DNA double-strand breaks coincide with

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**Figure 7.** Immunoblot analysis of p53 (A), PTEN (B), PI3 kinase (PI3'K) (C), and phosphorylated Akt (P-Akt) (D) in TK6 (300 \(\mu\)M min) and NH32 cells (600 \(\mu\)M min) following treatment of toxic doses of NO\(^*\). Jurkat cells treated with 4 \(\mu\)M staurosporine were used as positive controls, and densitometric analysis were as indicated in Figure 5. Values are mean quantitative densitometric values and 95% confidence intervals from two to three independent experiments. *p < 0.05, as compared with argon-treated controls.
low SCEs in the wild type cells, and vice versa in the p53-null cells. A possible explanation is that p53 plays a major role in modulating cell cycle arrest and apoptosis, and suppression of either activity could potentially increase the frequency of homologous recombination—e.g., without arrest, forks hit more lesions and recombine more; without apoptosis, cells loaded with recombinogenic damage that would have been killed by damage survive instead. The latter hypothesis may apply here. Because the recombinational repair of DNA double-strand breaks is replication-dependent (41), SCEs can be analyzed only in cells that are able to go through two rounds of replication (21) after NO exposure. As noted, a large number of TK6 cells containing lethal amounts DNA double-strand breaks may go through apoptosis as a result of their functional p53 before they can complete two cycles of cell division. On the other hand, NH32 cells do not die through apoptosis but rather keep dividing due to lack of p53. As a result, SCE frequency is higher in NH32 cells relative to TK6 cells. Alternatively, p53-regulated nonhomologous end joining (42) may be a potential repair mechanism of DNA double-strand breaks in these human lymphoblasts. High recombinational activity between misaligned sequences in p53 dysfunctional cells may have led to inappropriate DNA repair and thus increased mutagenesis (43–45), as confirmed in this investigation: A higher mutagenesis at the TK1 locus was induced by NO treatment in NH32 cells than in TK6 cells.

We further probed underlying causes of the reduced toxicity and reduced DNA double-strand breaks observed in the p53-null cells and found striking differences between the wild-type and p53-null cells in the levels of key proteins involved in regulation of cell survival/death and in defending against reactive oxygen and nitrogen species. NO treatment diminished levels of the prosurvival and antiapoptotic proteins Akt and NF-κB (23, 24, 46) only in TK6 cells. Increased protein levels of catalase, which detoxifies hydrogen peroxide, may be an important cellular defense response to NO-induced oxidative stress in TK6 cells, whereas this protective mechanism is deficient in p53-null NH32 cells. SOD1 plays a fundamental role in modulating NO toxicity by acting as an antioxidant, dismutating superoxide. Cells producing an increased amount of SOD1 are less sensitive to NO-mediated toxicity (47). Conversely, decreased SOD1 levels lead to NO-mediated apoptotic cell death (48). Reduced SOD1 protein levels in TK6 cells treated with the toxic dose may therefore be a promoter of NO-induced toxicity, whereas increased expression of SOD1 protein in NH32 cells may contribute to decreased sensitivity to NO-induced cell killing in this cell type. Unexpectedly, NO treatment did not affect levels of GPX protein in either TK6 or NH32 cells, conflicting with an earlier report that GPX is a p53 down-regulated gene (49) and suggesting that GPX modulation by activated p53 may be cell type- and/or inducer-dependent.

Taking into account presently accepted models of cell survival and apoptosis, the present findings, along with our previous characterization of p53/mitochondria- and Fas-dependent apoptotic signaling pathways activated by NO treatment (8, 10), suggest that p53 status is a dominant factor in modulating NO toxicity and toxic responses in human lymphoblastoid cells. As proposed in Figure 8, NO-induced DNA damage is followed by activation of p53, which upregulates PTEN, leading to PI3K/Akt-dependent cell survival suppression (23, 46) and increased apoptosis. Phosphorylated Akt protein levels were diminished in TK6 cells following NO exposure, but expression of PI3 kinase protein was not affected, implying that PI3 kinase may not be the only kinase to phosphorylate Akt protein. Interaction between Akt and NF-κB signaling has been reported in anti-apoptotic responses by other groups (50, 51). Although p53-dependent cell cycle arrest at the G1/S checkpoint has been well-documented (6, 52), we observed a p53-independent S phase arrest after NO treatment, resulting in cells entering a program of cytostasis or apoptosis. Activated p53 inhibits DNA recombinational repair and reduces mutagenesis derived from inappropriate repair. Thus, the absence of wild-type p53 in NH32 cells leads to a higher threshold dose of NO required for cell death and fewer DNA double-strand breaks and higher mutagenesis due to increased DNA recombinational repair in this model.

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