Nickel nanoparticle-induced dose-dependent cyto-genotoxicity in human breast carcinoma MCF-7 cells

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Abstract: Despite the widespread application of nickel nanoparticles (Ni NPs) in industrial, commercial, and biomedical fields, their response to human cells has not been clearly elucidated. In the study reported here, Ni NPs with a 28 nm diameter were used to study their interaction with human breast carcinoma (MCF-7) cells. Dose-dependent decreased cell viability and damaged cell membrane integrity showed the cytotoxic potential of the Ni NPs. We further found that Ni NPs induce oxidative stress in a dose-dependent manner, as evidenced by glutathione depletion and reactive oxygen species (ROS) generation. Comet assay indicated the dose-dependent induction of DNA damage due to Ni NP exposure. The level of messenger RNA, as well as activity of caspase-3 enzyme, was higher in MCF-7 cells exposed to Ni NPs than in control cells. Moreover, we observed statistically significant correlations of ROS with cell viability ($R^2=0.984$), DNA damage (% tail DNA) ($R^2=0.982$), and caspase-3 enzyme activity ($R^2=0.991$). To the best of our knowledge, this is the first study on human breast cancer cells to have shown the cyto-genotoxicity of Ni NPs, which seems to be mediated through ROS.

Keywords: cytotoxicity, genotoxicity, reactive oxygen species, glutathione, caspase-3 enzyme

Introduction

Engineered nanoparticles (NPs) possess astonishing physicochemical properties, which has led to an exponential growth in their development and production worldwide. NPs, by definition, are materials/structures that have at least one dimension in the 1–100 nm range.1,2 While many NPs are commonly used, their interactions with biological systems as well as their environmental and human health effects have not been fully explored.3,4 There is concern that research on the possible health risks of NP exposure is not keeping pace with the number of products containing NPs that are entering the industry and market. The contributing factors to NP toxicity are their size, chemical composition, shape, particle aging, and surface charge.5 As they are smaller than cellular organelles, they can penetrate basic biological structures, which may, in turn, disrupt the normal functioning of these structures.6 Further, the unique physicochemical properties of an NP cannot be simply predicted from the properties of a fine particle or bulk material with the same chemical composition. This is supported by studies which have shown that NPs are more toxic than fine particles or bulk materials with the same chemical composition.7,8

Nickel (Ni) NPs have many characteristics that include a high level of surface energy, high level of magnetism, large surface area, and low melting and burning points. Therefore, Ni NPs are being widely used as catalysts, sensors, and in
electronic applications. Recent studies indicate the great potential of Ni NPs in biomedical applications, including cancer diagnosis and therapy. For example, Hossain and Kleve showed that Ni nanowire induces ROS-mediated apoptotic cell death in human pancreatic adenocarcinoma (PANC-1) cancer cells, while Guo et al suggested that a combination of Ni NPs and the anticancer drug daunorubicin may have a synergistic effect on the efficient cytotoxicity in leukemia cancer cells. Further, Zhang et al indicated that the presence of Ni NPs could facilitate the drug accumulation of daunorubicin inside the leukemia cells. In a recent study, Chen et al found a synergistic effect of Ni NPs with the anticancer drug verapamil on the induction of apoptosis in K562 cells. This in vivo study also indicates that treatment with verapamil-Ni effectively inhibited tumor growth in mice.

However, researchers have also suggested Ni to be a potent human carcinogen with low mutational activity. Pietruska et al showed that Ni NPs cause a rapid and prolonged activation of the hypoxia-inducible factor 1-alpha pathway, which is responsible for tumor growth, in human lung cancer cells. Zhao et al observed that Ni NPs induced toxicity in mouse epidermal JB6 cells, while Ispas et al investigated the effect of different size (30, 60 & 100 nm) of Ni NPs as well as aggregated particle clusters of 60 nm having dendritic structures on mortality and developmental defects in Zebrafish embryos. Ni NPs toxicity was compared to soluble nickel (Ni	extsuperscript{2+}). All three (30, 60 & 100 nm) of Ni NPs were equal to or less toxic than Ni	extsuperscript{2+} while dendritic clusters (60 nm) were more toxic. Although the potential benefits of Ni NPs are considerable, there is a distinct need to identify any potential hazards associated with Ni NPs. Our knowledge about the interactions between Ni NPs and living cells and the effects of these NPs on human health is still in its infancy. This study was designed to examine the cytogenotoxic effect of Ni NPs using human breast carcinoma (MCF-7) cells. To achieve this goal, we determined the cell viability, cell membrane damage, ROS generation, glutathione (GSH), mitochondrial membrane potential (MMP), DNA damage, and caspase-3 enzyme in MCF-7 cells exposed to different concentrations of Ni NPs for 24 hours. MCF-7 cells, which are derived from human breast carcinoma, have been widely used in toxicological/pharmacological studies.

**Materials and methods**

**Ni NPs and reagents**

Ni NPs were synthesized through a solution reduction process using nickel chloride (NiCl	extsubscript{2}.6H	extsubscript{2}O) as a precursor with hydrate hydrazine solution, as described in our previous publication. The prepared Ni NPs were crystalline, spherical in shape with smooth surfaces, and had an average diameter of 28 nm. Dye-Dubelco’s Modified Eagle’s Medium (DMEM), Hank’s Balanced Salt Solution, fetal bovine serum (FBS), penicillin-streptomycin, and trypsin were purchased from Life Technologies (Carlsbad, CA, USA), while 3-(4,5-2-yl)-2,5-diphenyltetrazoliumbromide (MTT), 2,7-dichloro-dihydro-fluorescein diacetate (DCFH-DA), and Rhodamine-123 (Rh-123) were obtained from Sigma-Aldrich (St Louis, MO, USA). Kits for caspase-3 enzyme and lactate dehydrogenase (LDH) assays were bought from BioVision Inc. (Milpitas, CA, USA). All other chemicals used were of the highest purity available from commercial sources.

**Cell culture and exposure of cells to Ni NPs**

MCF-7 cells were cultured in DMEM supplemented with 10% FBS and 100 U/mL penicillin-streptomycin at 5% CO	extsubscript{2} and 37°C. At 85% confluence, cells were harvested using 0.25% trypsin and then subcultured according to the experiments. Cells were allowed to attach to the surface for 24 hours prior to NP exposure.

Ni NPs were suspended in cell-culture medium and diluted to appropriate concentrations. Dilutions of the Ni NPs were then sonicated using a sonicator bath at room temperature for 10 minutes at 40 W to avoid NP agglomeration prior to administration to the cells.

Cells not exposed to Ni NPs served as controls in each experiment.

**MTT assay**

MTT assay was carried out following the method as described by Mossman with some modifications. MTT assay assesses mitochondrial function by measuring ability of viable cells to reduce MTT into blue formazan product. In brief, 1 x 10	extsuperscript{4} cells/well were seeded in 96-well plates and exposed to different concentrations of Ni NPs (1–100 µg/mL) for 24 hours. At the end of the exposure time, culture medium was removed from each well to avoid the interference of NPs and replaced with new culture medium containing MTT solution in an amount equal to 10% of the culture volume, and incubated for 3 hours at 37°C until a purple-colored formazan product developed. The resulting formazan product was dissolved in acidified isopropanol.

Following this, each 96-well plate was centrifuged at 2,300 g for 5 minutes to settle the remaining NPs present in
the solution. Then, 100 µL of supernatant was transferred to the fresh wells of new 96-well plates and absorbance was measured at 570 nm using a Synergy™ HT microplate reader (BioTek, Winooski, VT, USA).

Neutral red uptake assay
Neutral red uptake (NRU) assay was performed following the procedure described by Borenfreund and Puerner with some modifications. In brief, 1 × 10⁴ cells/well were seeded in 96-well plates and exposed to different concentrations of Ni NPs (1–100 µg/mL) for 24 hours. At the end of the exposure time, the test solution was aspirated and cells were washed with phosphate-buffered saline (PBS) twice before being incubated for 3 hours in medium supplemented with neutral red (50 µg/mL). The medium was washed off rapidly with a solution containing 0.5% formaldehyde and 1% calcium chloride. The cells were then incubated for a further 20 minutes at 37°C in a mixture of acetic acid (1%) and ethanol (50%) to extract the dye. The 96-well plates were then centrifuged at 2,300 g for 5 minutes to settle the remaining NPs present in the solution. Following this, 100 µL of the supernatant was transferred to 96 new 96-well plates and the absorbance was measured at 540 nm using the microplate reader.

LDH assay
LDH assay was carried out using a BioVision LDH-cytotoxicity colorimetric assay kit according to the manufacturer’s protocol. In brief, 1 × 10⁴ cells/well were seeded in 96-well plates and exposed to different concentrations of Ni NPs (1–100 µg/mL) for 24 hours. After the exposure period had elapsed, each 96-well plate was centrifuged at 2,300 g for 5 minutes to settle the NPs present in the solution. Then 100 µL of the supernatant was transferred to the wells of new 96-well plates that already contained 100 µL of the reaction mixture from the BioVision kit and incubated for 30 minutes at room temperature. At the end of incubation time, the absorbance of the solution was measured at 340 nm using the microplate reader. The LDH levels in the culture medium versus those in the cells were quantified and compared with the control values according to the manufacturer of the kit’s instructions.

ROS estimation
ROS generation was measured using DCFH-DA as described by Wang and Joseph. ROS generation was determined by two methods – fluorometric analysis and microscopic fluorescence imaging. For the fluorometric analysis, cells (1 × 10⁴ cells/well) were seeded in a 96-well, black-bottomed culture plate and allowed to adhere for 24 hours in a CO₂ incubator at 37°C. Next, the cells were exposed to different concentrations of Ni NPs (10–50 µg/mL) for 24 hours. At the end of the exposure time, the cells were washed twice with Hank’s Balanced Salt Solution (Sigma-Aldrich) before being incubated in 1 mL of working solution of DCFH-DA at 37°C for 30 minutes. Following this, the cells were lysed in alkaline solution and centrifuged at 2,300 g for 10 minutes. A 200 µL amount of the supernatant was transferred to a 96-well plate, and fluorescence was measured at 485 nm excitation and 520 nm emission using the microplate reader. The values were expressed as a percent of fluorescence intensity relative to the control wells.

A parallel set of cells (5 × 10⁴ cells/well) was analyzed for intracellular fluorescence using an Olympus CKX41 fluorescence microscope (Olympus Corporation, Tokyo, Japan), with images taken at 20× magnification.

GSH assay
For the measurement of reduced GSH, cells were cultured in a 75 cm² culture flask and exposed to 0, 10, 25, and 50 µg/mL of Ni NPs for 24 hours. After the treatment, cell extracts were prepared from the control and treated cells as described in our previous publication. GSH levels in the cell extracts were determined using Ellman’s method. Briefly, a mixture of 0.1 mL of cell extract and 0.9 mL of 5% trichloroacetic acid was centrifuged at 2,300 g for 15 minutes at 4°C. Then 0.5 mL of the supernatant was added into 1.5 mL of 0.01% 5,5-dithio-bis-2-nitrobenzoic acid (DTNB) and the reaction was monitored at 412 nm. The amount of GSH was expressed in terms of nmol/mg protein.

MMP assay
MMP was measured following the protocol of Zhang et al with some modifications. In brief, cells (5 × 10⁴ cells/well) treated with 0, 10, 25 and 50 µg/mL of Ni NPs for 24 hours, were harvested and washed twice with PBS. Cells were further exposed with 10 µg/mL of Rh-123 fluorescent dye for 1 hour at 37°C in dark. Again, cells were washed twice with PBS then the fluorescence intensity of Rh-123 dye was measured using the Olympus CKX41 upright fluorescence microscope by capturing images at 20× magnification. A parallel set of cells (1 × 10⁴ cells/well) in a 96-well plate was analyzed for quantification of Rh-123 using the microplate reader.

Comet assay
Comet assay was performed as described by Singh et al with some specific modifications. Briefly, 70,000 cells were...
seeded in each well of a 12-well plate. After 24 hours of seeding, the cells were treated with different concentrations of Ni NPs (10–50 µg/mL) for 24 hours. At the end of the exposure period, the cells were trypsinized and resuspended in DMEM supplemented with 10% FBS and the cell suspension was then centrifuged at 2,300 g for 5 minutes at 4°C. Finally, the resultant cell pellet was suspended in ice-chilled PBS for comet assay.

At this point, 15 µL of the cell suspension (containing approximately 20,000 cells) was mixed with 85 µL of low-melting-point agarose (0.5%) and layered on one end of a plain frosted glass slide, pre-coated with a layer of 200 µL normal agarose (1%). Thereafter, it was covered with a third layer of 100 µL low-melting-point agarose (0.5%). After solidification of the gel, the slide was immersed in a freshly prepared lysing solution (2.5 M NaCl, 100 mM disodium ethylenediaminetetraacetic acid [Na₂EDTA] and 10 mM Tris pH 10 with 10% dimethyl sulfoxide [DMSO] and 1% Triton™ X-100 [The Dow Chemical Co, Midland, MI, USA]) overnight at 4°C.

The slides were then placed in a horizontal gel electrophoresis unit. Fresh, cold alkaline electrophoresis buffer (300 mM NaOH, 1 mM Na₂EDTA, and 0.2% DMSO, pH 13.5) was poured into the chamber and left for 20 minutes at 4°C for DNA unwinding and the conversion of alkali-labile sites to single-strand breaks. Electrophoresis was carried out using the same solution at 4°C for 20 minutes at 15 V. The slides were neutralized gently with 0.4 M Tris buffer at pH 7.5 and stained with 75 µL ethidium bromide (20 µg/mL). The slides were stored at 4°C in a humidified slide box until scoring.

Slides were scored at a final magnification of 400× using an image analysis system (Komet-5.0, Kinetic Imaging, Liverpool, UK) attached to a fluorescent microscope (DMLB model, Leica Microsystems GmbH, Wetzlar, Germany) equipped with appropriate filters. An undamaged cell resembles an intact nucleus without a tail and a damaged cell has the appearance of a comet. The comet parameter used to measure DNA damage in the cells was % tail DNA (fraction of DNA in the tail). Images from 50 random cells (25 from each replicate slide) were analyzed for each experiment.

**Quantitative real-time polymerase chain reaction analysis of caspase-3 gene**

Cells were exposed to different concentrations (10–50 µg/mL) of Ni NPs for 24 hours. At the end of the exposure time, total RNA was extracted using an RNeasy Mini Kit (Qiagen, Valencia, CA, USA) according to the manufacturer’s instructions. The concentrations of the extracted RNAs were determined using a NanoDrop 8000 Spectrophotometer (Thermo Fisher Scientific, Waltham, MA, USA) and the integrity of the RNAs was visualized on 1% agarose gel using a Universal Hood II gel documentation system (BioRad, Hercules, CA, USA). The first-strand complementary DNA was synthesized from 1 µg of total RNA by reverse transcriptase using M-MLV (Promega, Madison, WI, USA) and oligo (dT) primers (Promega) according to the manufacturer’s protocol. Quantitative real-time polymerase chain reaction (RT-PCRq) was performed using a QuantiTect SYBR Green PCR Kit (Qiagen) with an ABI Prism® 7900HT Sequence Detection System (Applied Biosystems, Foster City, CA, USA). Two microliters of template complementary DNA were added to the final volume of 20 µL of reaction mixture. The first RT-PCRq cycle was undertaken for 10 minutes at 95°C and this was followed by 40 cycles involving denaturation at 95°C for 15 seconds, annealing at 60°C for 20 seconds, and elongation at 72°C for 20 seconds. The sequence of the specific sets of primers for caspase-3 has been given in our previous publication. The primer sequence of the GAPDH gene was 5′-GCTGCTTCTCTTGTGACAAAGT-3′, 5′-CTCAGCTTGACTGTGCCATT-3′. The expressions of selected genes were normalized to the GAPDH gene, which was used as an internal housekeeping control. All the RT-PCRq experiments were performed in triplicate and the data expressed as the mean of at least three independent experiments.

**Caspase-3 enzyme assay**

The activity of the caspase-3 enzyme was measured in both treated and control cells using a Caspase-3 Colorimetric Assay Kit (BioVision). This assay is based on the principle that activated caspases in apoptotic cells cleave the synthetic substrates to release free chromophore p-nitroanilide (pNA), which is measured at 405 nm. The pNA was a by-product after specific action of caspase-3 on tetrapeptide substrates DEVD-pNA. Briefly, the reaction mixture – consisting of 50 µL of cell extract protein (50 µg) (from control and treated cells), 50 µL of 2X Reaction Buffer (BioVision Inc.; containing 10 mM dithiothreitol), and 5 µL of 4 mM DEVD-pNA substrate in a total volume of 105 µL – was incubated at 37°C for 1 hour then the absorbance of the product was measured using the microplate reader at 405 nm according to the manufacturer’s instructions.

**Protein estimation**

The total protein content in the cell extracts was estimated by the Bradford method using bovine serum albumin as the standard.
Statistical analysis
Statistical significance was determined by one-way analysis of variance followed by Dunnett’s multiple comparison test. Significance was ascribed at \( P < 0.05 \).

Results
Dose-dependent cell viability in response to Ni NP exposure
Cells were exposed to Ni NPs at the concentrations of 0, 1, 2, 5, 10, 25, 50, and 100 \( \mu \text{g/mL} \) for 24 hours and cell viability was determined using MTT and NRU assays. Both assays demonstrated that Ni NPs up to the concentration of 5 \( \mu \text{g/mL} \) did not produce significant reduction in cell viability. As the concentration of the NPs increased from 10 to 100 \( \mu \text{g/mL} \), reduction in cell viability was observed in a dose-dependent manner.

The MTT assay indicated that cell viability decreased to 86%, 74%, 62%, and 44% when cells were exposed to Ni NPs at the concentrations of 10, 25, 50, and 100 \( \mu \text{g/mL} \), respectively (Figure 1A). Figure 1B shows the results of cell viability obtained by NRU assay, which showed that cell viability decreased to 83%, 76%, 64%, and 53% when cells

![Figure 1 A](image1.png)

**Figure 1 A** Dose-dependent reduction of viability of MCF-7 cells after nickel nanoparticle (Ni NP) exposure: (A) 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay and (B) neutral red uptake (NRU) assay.

**Notes:** Data represented are mean \( \pm \) standard deviation of three identical experiments made in triplicate. *Statistically significant difference as compared with the control \( (P<0.05) \).
were exposed to Ni NPs at the concentrations of 10, 25, 50, and 100 μg/mL, respectively.

**Dose-dependent induction of cell membrane damage due to Ni NP exposure**

LDH is an enzyme extensively present in the cytosol that converts lactate to pyruvate. When plasma membrane integrity is disrupted, LDH leaks into culture medium and its extracellular levels are elevated depending on the toxicity of the NPs. Consequently, higher LDH values in culture medium indicate cell membrane damage. In this study, we found that Ni NP increases LDH level in cell culture medium in a dose-dependent manner (Figure 2A). A significant negative correlation was also observed between LDH leakage and MTT cell viability (Figure 2B).

![Figure 2](image)

**Figure 2** Dose-dependent induction of membrane damage to MCF-7 cells after nickel nanoparticle (Ni NP) exposure. (A) Lactate dehydrogenase (LDH) leakage. (B) A significant negative correlation between the cell viability (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide [MTT] assay) and membrane damage (LDH leakage) after 24 hours’ exposure to 0, 1, 2, 5, 10, 25, 50, and 100 μg/mL of Ni NPs.

**Notes:** *Data represented are the mean ± standard deviation of three identical experiments made in triplicate. *Statistically significant difference compared with the control (P < 0.05).

**Dose-dependent induction of ROS generation due to Ni NP exposure**

The higher production of ROS in cells due to pollutants has been suggested as signaling molecules involved in the initiation and execution of DNA damage and apoptosis. Both fluorescence microscopy (Figure 3A) and quantitative data (Figure 3B) revealed that the Ni NPs (10–50 μg/mL) induced intracellular ROS generation in a dose-dependent manner.

**Dose-dependent depletion of GSH levels due to Ni NP exposure**

The depletion of GSH, which is indicative of ROS generation, has also been implicated in the oxidative damage of cellular macromolecules such as protein and DNA. We evaluated the GSH levels in MCF-7 cells treated with Ni NPs at concentrations of 10, 25, and 50 μg/mL for 24 hours. Results showed that the GSH levels were depleted with all concentrations of Ni NPs (Figure 4A). We also observed inverse correlations of ROS with GSH (R²=0.9953; Figure 4B) and MTT cell viability (R²=0.9981; Figure 4C).

**Dose-dependent depletion of MMP due to Ni NP exposure**

It is well-known that during apoptosis, the MMP of cells decreases. Ni NP-induced differences in MMP in MCF-7 cells were recorded in terms of the fluorescence intensity of the mitochondria-specific dye Rh-123. Both fluorescence microscopy (Figure 5A) and quantitative data (Figure 5B) revealed that the Ni NPs decreased the MMP of MCF-7 cells in a dose-dependent manner.

**Dose-dependent induction of DNA damage due to Ni NP exposure**

One sign of genotoxicity is the induction of DNA damage that can be determined by comet assay, a widely used method for the detection as well as measurement of DNA strand breaks. Comet assay was carried out in MCF-7 cells against Ni NPs at the concentrations of 10, 25, and 50 μg/mL for 24 hours. A dose-dependent increase in DNA damage was observed in Ni NP-treated cells, as evidenced by an increase in % tail DNA and Olive tail moment (arbitrary unit; Figure 6A and B). We also found a significant positive correlation between ROS and % tail DNA (R²=0.9816; Figure 6C).

**Dose-dependent induction of apoptosis due to Ni NP exposure**

RT-PCRq was used to analyze the messenger RNA (mRNA) levels of apoptotic gene caspase-3 in MCF-7 cells exposed...
to Ni NPs at the concentrations of 10, 25, and 50 µg/mL for 24 hours. We observed that Ni NPs significantly induced the expression of this gene in a dose-dependent manner (Figure 7A). In agreement with the mRNA results, we also found that the Ni NPs induced the activity of the caspase-3 enzyme dose dependently (Figure 7B). A significant positive correlation was also observed between ROS and the caspase-3 enzyme ($R^2=0.9911$; Figure 7C).

Figure 3 Dose-dependent induction of oxidant production in MCF-7 cells after nickel nanoparticle (Ni NP) exposure. (A) Representative microphotographs showing Ni NP-induced reactive oxygen species (ROS) generation in MCF-7 cells. (B) Percentage change in ROS generation of MCF-7 cells due to Ni NPs exposure.

Notes: *Images were captured with an Olympus CKX41 fluorescence microscope (Olympus Corporation, Tokyo, Japan); data represented are the mean ± standard deviation of three identical experiments made in triplicate. *Statistically significant difference as compared to control ($P<0.05$).
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Discussion

With the potential use of Ni NPs in the biomedical field, concerns about their toxic effects for humans and their environment have increased rapidly.

Some recent studies have shown that Ni NPs are cytotoxic to various tissues and cells. In this study, we evaluated the toxicity mechanisms of Ni NPs through which NPs generate ROS in cells are still underway. In the present study, we also found that LDH leakage increased as the concentration of Ni NPs was increased in MCF-7 cells. A significant reverse correlation between MTT and LDH was also observed.

ROS generation is likely to be one of the possible mechanisms of NP toxicity. NPs may increase ROS production and thereby cause interference in the biological antioxidant defense system. ROS induce DNA damage, including a multitude of oxidized base lesions, abasic sites, and single- and double-strand breaks, all of which can be cytotoxic and/or mutagenic.

However, studies on the exact mechanisms through which NPs generate ROS in cells are still underway. In this study, we evaluated the toxicity mechanisms of Ni NPs in MCF-7 cells. We hypothesized that the Ni NPs induced cytotoxicity in the MCF-7 cells through oxidant generation.
and antioxidant depletion. Indeed, Ni NPs were shown to generate ROS (oxidant) and GSH (antioxidant) depletion in MCF-7 cells in a dose-dependent manner. Further, the negative linear correlation between the ROS and GSH indicated that free oxygen radical species were generated by exposure to Ni NPs, which reduced intracellular antioxidant GSH levels. There was also a strong inverse correlation between cell viability (MTT) and increased ROS level. Consequently, the higher production of ROS and concomitant decrease in antioxidants such as GSH to scavenge free oxygen radicals appear to be the underlying mechanisms of Ni NP toxicity in MCF-7 cells. These results confirm the oxidant potential of Ni NPs described in previous studies.15,21,41

Alkaline comet assay was used to analyze the DNA damage caused by Ni NPs in MCF-7 cells. This assay is able to detect single-strand breakage or other lesions, such as alkali-labile sites, DNA cross-links, and incomplete excision repair events.52,53 This technique offers considerable advantages over other cytogenetic methods like chromosome aberrations, sister chromatid exchanges, and
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Figure 6 Dose-dependent induction of DNA damage (comet assay) in MCF-7 cells after nickel nanoparticle (Ni NP) exposure. (A) Percent tail DNA and (B) Olive tail moment (arbitrary unit). (C) Significant positive correlations between reactive oxygen species (ROS) and DNA damage (% tail DNA) in MCF-7 cells after 24 hours' exposure to 10, 25, and 50 µg/mL of Ni NPs.

Notes: Data represented are mean ± standard deviation of three identical experiments made in triplicate. *Significant difference compared with the control (P<0.05).

Abbreviation: au, arbitrary unit.

Figure 7 Dose-dependent induction of apoptosis in MCF-7 cells after nickel nanoparticle (Ni NP) exposure. (A) Ni NPs induced the expression of mRNA levels of apoptotic gene caspase-3 in MCF-7 cells. (B) Ni NPs also induced the activity of the caspase-3 enzyme in MCF-7 cells. (C) A significant positive correlation between reactive oxygen species (ROS) and the caspase-3 enzyme in MCF-7 cells after 24 hours' exposure to 10, 25, and 50 µg/mL of Ni NPs.

Notes: Data represented are mean ± standard deviation of three identical experiments made in triplicate. *Statistically significant difference compared with the control (P<0.05).

Abbreviation: au, arbitrary unit.
micronucleus tests used to detect DNA damage, because the cells do not need to be mitotically active for comet assay.\textsuperscript{14} Therefore, this assay has been widely used in the field of genetic toxicology.\textsuperscript{13} Investigators have utilized the comet assay to examine the genotoxic potential of different types of NPs.\textsuperscript{31,55} In this study, we observed that Ni NPs induce DNA damage, as evidenced by \% tail DNA and Olive tail moment. We also found a strong positive correlation between DNA damage (\% tail DNA) and ROS, suggesting that oxidative stress may be one of the causes of DNA damage in Ni NP toxicity. We further analyzed the mRNA expression of an apoptotic gene caspase-3 in response to Ni NP exposure in MCF-7 cells. Our RT-PCR results showed that the Ni NPs upregulated the mRNA level of this gene. Further, the activity of the caspase-3 enzyme was also higher in Ni NPs treated MCF-7 cells. A positive correlation between caspase-3 enzyme activity and ROS level was also observed. Studies have shown that ROS work as signaling molecules for the initiation and execution of the apoptotic pathways.\textsuperscript{56,57}

The recent literature has shown that Ni ions release from the surface of Ni NPs when they are suspended in aqueous solutions such as water and cell-culture media. However, in our study, we did not examine the degree of ionization of Ni NPs in water or culture media and the effects of this on MCF-7 cells. A recent study reported that Ni ions and nickel(II) oxide (NiO) NPs were equally toxic to human lung epithelial cells, followed by metallic Ni NPs but not by metallic Ni microparticles.\textsuperscript{15} Therefore, toxicity due to the dissolution of metallic Ni NPs in cells or tissues needs further study.

Conclusion
We observed that Ni NPs induce cytotoxicity in MCF-7 cells in a dose-dependent manner in the concentration range of 10–100 µg/mL. This NP was also found to induce oxidative stress, as evidenced by ROS production and GSH depletion. The mRNA level, as well as the activity of the caspase-3 enzyme, was increased due to Ni NP exposure. Taken together, these results suggest that Ni NPs induce cytogenotoxicity in MCF-7 cells, which is likely to be mediated through ROS. Further studies are underway examining the molecular mechanism of the toxicity induced by Ni NPs in human breast cancer cells.

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Disclosure
The authors declare no conflicts of interest in this work.

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