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Gordon Chambers
Dublin Institute of Technology, Gordon.chambers@dit.ie

Sanchali Mukherjee
Dublin Institute of Technology

Alan Casey
Dublin Institute of Technology, alan.casey@dit.ie

Niall Ó Cláonadh
Dublin Institute of Technology

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Comparative in vitro cytotoxicity study of silver nanoparticle on two mammalian cell lines

Sanchali Gupta Mukherjee a,b,⁎, Niall O’Clonadha, Alan Casey a, Gordon Chambers a,b

a Nanolab Research Centre, Focas Institute, Dublin Institute of Technology, Kevin Street, Dublin 8, Ireland
b School of Physics, Dublin Institute of Technology, Kevin Street, Dublin 8, Ireland

A B S T R A C T

In this study the cytotoxic effect of commercially available silver (Ag) nanoparticle was evaluated using human dermal and cervical cancer cell lines. Prior to the cellular studies a full particle size characterisation was carried out using Dynamic Light Scattering (DLS), Transmission Electron Microscopy and Scanning Electron Microscopy in distilled water and cell culture media. The Zeta Potential (ZP) associated with the Ag nanoparticle was also determined in order to assess its stability in the solutions and its possible interaction with the media. The DLS and ZP study have suggested interaction of Ag nanoparticles with the media, which can lead to secondary toxicity. The toxic effects of Ag nanoparticles were then evaluated using different cytotoxic endpoints namely the lysosomal activity, mitochondrial metabolism, basic cellular metabolism, cellular protein content and cellular proliferative capacity. The cytotoxic effect of Ag nanoparticle was dependant on dose, exposure time and on the cell line tested. Further investigation was carried out on HeLa and HaCaT cell lines to elucidate the mechanism of its cytotoxicity. The Ag nanoparticle was noted to induce elevated levels of oxidative stress, glutathione depletion and damage to the cell membrane as found from the adenylate kinase assay and that leads to the apoptosis. Overall, significant differences were observed between the sensitivity of the two cell lines which can be understood in terms of their natural antioxidant levels.

1. Introduction

Silver (Ag) nanoparticles offer remarkably different physiochemical and biological characteristics from that of bulk silver, such as increased optical, electromagnetic, catalytic properties and antimicrobial activity (Choi et al., 2009). Ag has traditionally been used as an antimicrobial agent for many years. Ag displays an oligodynamic effect which is an inherent antimicrobial property associated with mainly heavy metals (Shuval et al., 1995). The exact mechanism by which this occurs is still largely unknown however, some literature proposes that Ag interacts with the thiol groups on proteins necessary for microbial respiration (Cho et al., 2005). Ag ions are seen to actively seek out the thiol groups of proteins and enzymes thus making them unsuitable for microbial development (Liau et al., 1997). Ag has also been shown to directly interact with the cell membranes of bacteria causing breaks in the membrane and the cell to essentially burst (Percival et al., 2005). It is this innate property that has lead to an array of potential applications of Ag nanoparticles in both the medical arena and consumer products.

In terms of medical applications Ag nanoparticles has found applications in wound dressings, contraceptive devices, surgical instruments and bone prostheses which can be coated or embedded with the Ag nanoparticles (Cheng et al., 2004; Chen et al., 2006; Muangman et al., 2006; Cohen et al., 2007; Lansdown, 2006; Zhang and Sun, 2007). In addition, consumer products containing Ag nanoparticles include antibacterial room sprays, laundry detergents, water purificants and wall paint (Cheng et al., 2004; Zhang and Sun, 2007). Ag nanoparticles are also incorporated into textiles for manufacture of clothing, undergarments and socks (Lee et al., 2007; Vigneshwaran et al., 2007).

As a result of these applications exposure of Ag nanoparticles to us and our environment are becoming increasingly widespread. Consequently, silver in the form of nanoparticles has gained an increasing access to tissues, cells and biological molecules within the human body. Once reaching nanoscale, certain materials do exhibit significant toxicity to mammalian cells even if they are bio-chemically inert and biocompatible in bulk size, e.g. carbon (Magrez et al., 2006; Shvedova et al., 2005). Several studies exist in literature showing that the release of Ag ions inhibit the normal respiratory functions of the cells and cause cell death. A study

⁎ Corresponding author at: Nanolab Research Centre, Focas Institute, Dublin Institute of Technology, Kevin Street, Dublin 8, Ireland. Tel.: +353 14027933; fax: +353 4027901.
E-mail address: sanchali.guptamukherjee@student.dit.ie (S.G. Mukherjee).
carried out by Hussain et al. (2005) have shown that Ag nanoparticles was cytotoxic to rodent liver cells causing them to change morphologically and also interfering with mitochondrial function. It has also been shown that Ag nanoparticles are highly toxic to the osteoblast cell lines (Moadab et al., 2011). One study has shown a very high toxic response of Ag nanoparticle of 5–10 nm diameter on hepatoma cell line, HepG2, by causing oxidative stress and DNA damage (Kim et al., 2009).

The human body has several semi-open interfaces for direct substance exchange with the environment, i.e. the respiratory tract, gastrointestinal tract (GIT) and skin. In view of Ag nanoparticle’s diverse forms of existence, they are also the principle routes of exposure. The irreversible pigmentation of skin and/or eye, i.e. arctact, gastrointestinal tract (GIT) and skin. In view of Ag nanoparticle’s diverse forms of existence, they are also the principle routes of exposure. The irreversible pigmentation of skin and/or eye, i.e. arctact, gastrointestinal tract (GIT) and skin. In view of Ag nanoparticles, and studies into their biological effects are of great importance. The aim of this study is to examine the effects of Ag nanoparticles on cell models of two possible routes of entry and exposure for this two human cell lines were employed – a dermal cell line and a cervical cancerous cell line. The effect has been evaluated using variety of end-points to postulate a mechanism of action of Ag nanoparticles. A battery of standard cytotoxicity assays, viz., Alamar Blue (AB) assay, Neutral Red (NR) assay, MTT assay, Coomassie Blue (CB) assay, were employed to study the effect of Ag nanoparticles on both the cell lines. AB, NR, CB and MTT assays show the effect of Ag nanoparticle on different aspects of cellular activity. Therefore the interpretation of the dose response from these assays is very important to understand the cellular response upon Ag nanoparticle exposure.

The interpretation of the action of the AB assay is somewhat varied. One report has suggested that it may indicate the early inhibition of cytoplasmic metabolism and cell growth (O’Brien et al., 2000) while another suggests that it indicates the inhibition of mitochondrial metabolism (De Fries and Mitsuhashi, 1995). But in general it is suggested that it shows basic cellular metabolism in terms of it oxido-reductase enzyme activity (Mukherjee et al., 2010b). NR is a colorimetric assay for the quantification of the membrane permeability and lysosomal activity of cells in response to pharmaceutical, chemical and environmental compounds, and nutrients (Mukherjee et al., 2010a,b). The CB assay quantifies the amount of protein present in the cells (Clare and Cormack, 1996). The MTT colorimetric assay determines the ability of viable cells to reduce the soluble, yellow tetrazolium salt [3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT)] into an insoluble, purple formazan. Tetrazolium salts accept electrons from oxidized substrates or enzymes, such as NADH and NADPH. Reduction of MTT takes place at the ubiquinone and cytochrome b and c sites of the mitochondrial electron transport chain due to succinate dehydrogenase activity (Mosmann, 1983). The reduction of MTT to formazan within cells helps in the estimation of the mitochondrial metabolism and hence the number of viable cells after exposure to nanoparticles. Apart from accessing the effect of Ag nanoparticle on different cellular organelles and metabolism, we have also assessed its effect on the cellular proliferative capacity by clonogenic assay. Intracellular reactive oxygen species (ROS) generation, glutathione (GSH) depletion, adenylate kinase (AK) release from the cells and apoptosis induction upon Ag nanoparticle exposure was also studied. Adenylate kinase (also known as ADK or myokinase) is a phosphotransferase enzyme that catalyzes the nucleotide phosphoryl exchange reaction 2ADP ↔ ATP + AMP, critical in cell life. They are found in cytosol and mitochondria of the cells. Adenylate kinase will only leak from cells whose cell membrane integrity has been compromised. Apoptosis is an energy driven process; ATP levels are maintained in metabolically active cells, but upon cell death residual ATP is degraded (Bradbury et al., 2000; Crouch et al., 1993). With the onset of apoptosis, cells show an increase in the levels of cellular ADP (adenosine diphosphate) but without an appreciable drop in ATP. As cells pass into secondary necrosis the ADP levels continue to rise as hydrolysis of ATP starts to occur. In primary necrosis ATP levels fall rapidly as a result of exposure to high concentrations of toxic agents with a corresponding increase in the cellular ADP (Bradbury et al., 2000). Finally the differences between the responses from the two cell lines are compared and the possible reason behind the difference in the responses is elucidated.

2. Materials and methods

2.1. Test materials and reagents

Silver (Ag) nanopowder of <100 nm was purchased from Sigma Aldrich Ltd. (Dublin, Ireland). 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT), Neutral Red (NR), Coomassie Blue (CB) and 2’,7’-dichlorofluorescin diacetate (DCF) as well as cell culture media, supplements, the trypsinisation solution were all purchased from Sigma Aldrich Ltd. (Dublin, Ireland). Carbol fuchsin was purchased from BDH (UK). ToxiLight™ and ApoGlow™ kit were purchased from Lonza Ltd. (Wokingham, UK).

2.2. Cell culture

HaCaT cells, an immortal non-cancerous human keratinocyte cell line (kindly provided by Prof. Dr. Boukamp, Heidelberg), HeLa cells (ATCC, CCL-2™) an epithelial adenocarcinoma cell line of the cervix were employed in this study. HaCaT cells were cultured in RPMI-1640 medium supplemented with 10% FBS, 45 IU/ml penicillin and 45 IU/ml streptomycin (HaCaT media) at 37 °C in 5% CO₂. HeLa cells were cultured in Dulbecco’s Modified Eagle’s Medium Nutrient Mixture F-12 HAM (DMEM F-12 HAM) with 2 mM L-glutamine supplemented with 10% fetal bovine serum (FBS), 45 IU/ml penicillin and 45 IU/ml streptomycin (HaCaT media) at 37 °C in 5% CO₂. HeLa cells were cultured in RPMI-1640 medium supplemented with 10% FBS, 45 IU/ml penicillin and 45 IU/ml streptomycin (HeLa media) at 37 °C in 5% CO₂.

2.3. Characterization of Ag nanoparticles

Prior to the cytotoxicity testing Ag nanoparticles were subjected to an extensive characterisation process, where possible measurements were performed on Ag nanoparticle as purchased and on test suspensions of them. The suspensions of Ag nanoparticles were prepared in distilled water (dH₂O), HaCaT media and in HeLa media using bath-sonicator (Degussa-Ney ULTRASONIK 57X, 50/60 Hz, California, USA) prior to size and zeta potential measurements. Dynamic light scattering sizing measurements and zeta potential measurements were performed with the aid of a Malvern Zeta Sizer Nano ZS (Malvern Instruments, Worcestershire, UK) operating with version 5.03 of the systems Dispersion Technology Software (DTS Nano). For size measurement, DTS0012 disposable
sizing cuvettes were used. The refractive index (RI) of Ag nanoparticles, \(d_H\)2O, HaCaT media and HeLa media were 1.6, 1.330, 1.332, 1.300, respectively. Prior to analysis, viscosity measurements were performed on all media and \(d_H\)2O with the aid of a Viscometer SV-10 (A&D Instruments Ltd., UK) at 25 °C and the recorded values used in all DLS size estimations. The viscosity of \(d_H\)2O, HaCaT media and HeLa media at 25 °C were 0.887cP, 0.82cP, 0.550cP, respectively. The samples were equilibrated at 25 °C for 3 min before each measurement. For zeta potential measurements DTS1060C clear disposable zeta cells were used and the measurements were performed at 5 V. Smoluchowski equation was used in the DTS software to calculate the zeta potential of the Ag nanoparticles. For all zeta potential measurements analysis was performed on the solutions with the automatic model setting, using an applied voltage of 5 V, to minimize artefacts and charring of media proteins during analysis. The specific surface area of the particles was estimated with a Micrometrics GEMINI BET. BET sample holders were filled with a known mass of the powdered Ag nanoparticles and measured. The sample was degassed for 2 h at room temperature with nitrogen gas prior to analysis, nitrogen gas was used as the absorptive gas, a multipoint method was used in the estimation of specific surface area and the sample was analysed without porosity analysis. transmission electron microscopy (TEM) was also used to estimate the size of the Ag nanoparticles from its images. The instrument used was a JEOl 100CX model TEM. Samples were prepared by dispersing Ag nanoparticles in ethanol to a concentration of 10 mg/l by sonication 750 watts Ultrasonic Processor tip (Branson Ultrasonics, Ultra sonic processor VCX-750 W) sonicator at 40% amplitude for a total operating time of 45 s.

Samples were then drop cast onto 3 mm FORMVAR coated Cu grids (AGAR Scientific). Once the solvent had evaporated particle size was measured at a range of magnifications up to 80,000. The accelerating voltage and probe current used for TEM analysis were 25 kV and 18 μA, respectively. SEM was also performed to analyse particle size. Hitachi SU 6600 FESEM instrument was used. The SEM was calibrated with Au on Carbon standard provided by Agar Scientific (Essex, UK). Sample preparation was identical to that of the TEM preparation. Solutions were drop cast onto a pure silicon wafer which had been thoroughly cleaned by sonication in acetone for a period of 30 min followed by boiling in propanol for 30 min. The silicon wafers were then left to air dry in a dust free environment and the nanoparticle solution was drop casted onto them 24 h prior to measurement.

2.4. Cytotoxicity evaluation

The AB, NR, CB, MTT and the clonogenic assay were performed for assessment of the cytotoxicity of Ag nanoparticles to the two cell lines. A preliminary or range finding test was conducted to determine the range of concentrations of interest for the definitive test. The definitive test used a concentration range (at least eight concentrations) in which effects was likely to occur, thereby permitting the calculation of the respective Lethal Dose (LD50), no observed effect concentration (NOEC) and lowest observed effect concentration (LOEC). In all cases, results were compared to an unexposed control, eliminating any dependence of cell line exposures on volumes, well type, seeding efficiency and numbers, exposure times, etc. A stock solution of Ag nanoparticle was prepared aseptically, from which different concentrations of Ag nanoparticles were prepared in the respective HaCaT media and HeLa cell culture media, followed by bath sonication (Degussa-Ney ULTRASONIK 57X, California, USA) for 10 min at room temperature before exposure. As a positive control, 10% dimethyl sulfoxide (DMSO) prepared in the respective media was used parallel to the Ag nanoparticle exposures.

2.4.1. Alamar Blue, Neutral Red and Coomassie Blue assays

For the AB and NR assays, cells were seeded in 96 well microplates (Nunc, Denmark) at a density of \(1 \times 10^5\) cells/ml in 100 μl of respective media containing 10% FBS. At least three independent experiments were conducted and six replicate wells were employed per concentration per plate in each independent experiment. After 24 h of cell attachment, plates were washed with 100 μl/well PBS and the cells were treated with increasing concentrations of Ag nanoparticles, prepared in 10% FBS containing media for 24, 48, 72 and 96 h. All incubations were performed at 37 °C in a 5% CO2 humidified incubator.

The AB, NR and CB assays were conducted consecutively on the same set of plates. The AB assay was performed first followed by the NR assay and then lastly the CB assay. The assays were carried out according to manufacturer’s instruction. Briefly, control media or test exposures were removed; the cells were rinsed with PBS and 100 μl of AB/NR medium (5% [v/v] solution of AB and 1.25% [v/v] of NR dye) prepared in fresh media (without FBS or supplements) were added to each well. After 3 h incubation AB fluorescence was measured at the respective excitation and emission wavelength of 531 nm and 595 nm in a VICTOR™ 1420 Multilabel Counter (Perkin Elmer, USA). Wells having only AB and media were used as blanks. After measurement, the wells were then washed with PBS and 100 μl of NR fixative (50% ethanol, 49% \(d_H\)2O and 1% glacial acetic acid) was added to each well and the plates were shaken at 240 rpm for 10 min. The NR fluorescence was then measured at the excitation and emission wavelength of 531 and 642 nm, respectively in the same instrument. Protein determinations were performed on the same plates immediately by CB assay following NR uptake study. Excess NR dye was removed from the cells by washing with 100 μl de-staining solution, COMMASSIE dye was added to each well and the plate agitated for 10 min. The dye was removed and the plate washed with an acetic acid–ethanol solution. The wash solution was discarded and the dye extracted with measuring solution (1 M potassium acetate). The plate was shaken at 240 rpm for 10 min and the absorbance of the extracted dye was read at 570 nm (reference filter 340 nm) using the microplate reader. For these three assays, mean fluorescence/absorbance units for the six replicate cultures were calculated for each exposure treatment.

2.4.2. MTT assay

A parallel set of at least 3 replicate plates was set up for the MTT assay and seeded and exposed in an identical manner as described in Section 2.4.1. After 24, 48 and 72 h of Ag nanoparticle (AgNP) exposure, the medium for the control or test exposures was removed, the cells were washed with PBS and 100 μl of freshly prepared MTT in media (5 mg/ml of MTT in media [without FBS or supplements]) were added to each well. After 3 h incubation, the medium was discarded and the cells were rinsed with PBS and 100 μl of MTT fixative solution (isopropanol with 0.04 N HCl) were added to each well and the plates were shaken at 240 rpm for 10 min. The absorbance was then measured at 595 nm in a TECAN GENios (Grofid, Austria) plate reader.

2.4.3. Clonogenic assay

The clonogenic assay was developed by Puck and Marcus (1956) and further standardized by Franken et al. (2006). The clonogenic assay was performed in 6 well plates (Nunc, Denmark) and each well was seeded with 2 ml of respective 10% FBS containing media at a cell density of 250 cells/ml. For both cell lines, the cells were allowed to attach for 12 h, which is shorter than their reported doubling time of 23 h for HaCaT cells (Boukamp et al., 1988) and 22 h for HeLa cells (York and Majerus, 1994). Therefore it is assumed that predominantly single cells were present at the time of exposure. After 12 h of cell attachment, plates were washed
with 2 ml/well PBS and the cells were treated with increasing concentrations of AgNP prepared in 10% FBS containing respective media. The plates were then incubated for 8 days. Two replicate wells were used for each control and test concentrations per 6-well plate. Following exposure, the test solution was removed; the cells were washed with PBS and finally fixed and stained with 20% carbol in formalin solution (BDH, Poole, UK). The colonies were then manually counted. The minimum size of a colony was considered to be 50 cells/colony. In all these assays 10% DMSO in 10% FBS supplemented media was used as a positive control parallel to the Ag nanoparticle test solutions.

2.5. Optical microscopic study

Interaction of Ag nanoparticles with HaCaT and HeLa cell lines was studied by optical microscopy. Both the cell lines were seeded at a density of 1 × 10^5 cells/ml in petri dishes (Nunc, Denmark) and the Petri dishes were kept in a 37 °C, 5% CO2 incubator. After 24 h of cell attachment, the cells were washed twice with PBS and were exposed to 50 mg/l, 100 mg/l and 200 mg/l Ag nanoparticle concentrations in the respective media and the Petri dishes were kept in a 37 °C, 5% CO2 incubator for 24 h. The Ag nanoparticle test solutions were prepared in the same way as stated in Section 2.4. Following exposure, the test solutions were removed; the cells were washed twice with PBS and were imaged in 37 °C PBS using a Nikon Eclipse E600 microscope. All images were recorded using a Spot RT digital camera using Spot software (version 3.2.4, Diagnostic Instruments, Inc., USA).

2.6. Reactive oxygen species (ROS) study

Intracellular oxidative stress was quantified with the aid of 2′,7′-dichlorofluorescin diacetate (DCFH-DA), which was reported to detect intracellular hydroperoxides (Cathcart et al. 1983) and also a probe for wide range of reactive oxygen species (Invitrogen, Ireland, website – http://www.invitrogen.com/site/us/en/home/References/Molecular-Probes-The-Handbook/tabs/Reactive-oxygen-species.html). Briefly, healthy confluent cell cultures were harvested and seeded (1000 cells/well) into black bottomed 96 well plates (Nunc, Denmark) and allowed to attach for a period of 24 h prior to exposure. For ROS quantification three independent experiments were performed for each independent experiment eight replicate wells were used for control, eight replicate wells were employed for the positive control and eight replicate wells were used for each test concentration per microplate. A working stock of 20 μM DCFH-DA in PBS was prepared and all test concentrations, unexposed negative controls and positive controls were prepared and exposed to the cells in this working stock. The negative control consisted of the working stock solely namely a 20 μM DCFH-DA solution in PBS, the positive control consisted of a 1 μM Hydrogen Peroxide (H2O2) prepared in the 20 μM DCFH-DA/PBS working stock solution and finally the test concentrations consisted of a concentration range of Ag nanoparticles (for HaCaT 7.81 to 125 mg/l and for HeLa 0.625–80 mg/l) prepared in the working stock a 20 μM DCFH-DA/PBS solution.

After the initial 24 h attachment period the media was removed, the cells were subsequently washed with 100 μl of PBS and treated with 100 μl of the negative control, positive control and the test concentrations of AgNP and incubated for 30, 60, 120, 180, 240, 300 min. The rate of intracellular oxidative stress was then monitored by monitoring the emission at 529 nm (by 504 nm excitation) of the DCFH-DA dye at various time intervals of 30–300 min (the exposure plates were re-incubated for the remaining time after each measurement had been recorded).

2.7. Intracellular reduced glutathione level measurement

A commercial kit ThiolTracker™ Violet was used to estimate the levels of intracellular reduced glutathione (GSH) in HaCaT cells upon AgNP exposure. The fluorescent dye can be used for intracellular GSH detection in flow cytometry or by fluorescent imaging in living cells (http://products.invitrogen.com/ivgn/product/T1100967C1ID=search-product). In this study, a fluorescence microscope plate reader was employed to quantify the fluorescence intensity of the dye and therefore the intracellular GSH levels in control and exposed HaCaT cultures. The study was performed in 96-well microplate format, wherein the cells were seeded at a density of 1 × 10^5 cells/ml in 100 μl of respective media containing 10% FBS. After 24 h of cell attachment, the cells were washed twice with 100 μl/well PBS and were exposed to 25–200 mg/l AgNP in 10% FBS supplemented media for 24 h. 10% DMSO in the respective media were also used as positive controls. Following exposure, the cells were washed with 100 μl/well PBS and exposed to the ThiolTracker™ Violet dye, prepared in PBS at a concentration of 20 μM in 100 μl/well and the plates were kept in a 37 °C, 5% CO2 incubator for 30 min. Following incubation, the dye solution was removed, the cells were washed thrice with 100 μl/well PBS and the fluorescence was measured at excitation and emission wavelengths of 405 and 535 nm, respectively, in a VICTOR3 V™1420 Multilabel Counter plate reader (Perkin Elmer, UK).

2.8. Adenylate kinase (AK) release assay

The bioluminescent ToxiLight™ kit (Lonza Ltd., Wokingham, UK) was employed to measure the concentration of AK present in the supernatants of test solutions after different time point exposures. The kit is based on the bioluminescent measurement of AK which is present in all cells. This kit quantitatively measures the release of AK from damaged cells. A loss of cell integrity, through damage to the plasma membrane, results in the leakage of a number of factors from cells cultured in vitro into the surrounding medium. The measurement of the release of AK from the cells allows the accurate and sensitive determination of cytotoxicity and cytolysis.

The reaction involves two steps. The first involves the addition of ADP as a substrate for AK. In the presence of the enzyme, AK, the ADP is converted to ATP for assay by bioluminescence:

\[
\text{Mg}^{2+} + \text{ADP} \rightarrow \text{Mg}^{2+} \text{AMP} + \text{PPi} + \text{CO}_2 + \text{Light}
\]

The bioluminescent method utilizes an enzyme luciferase, which catalyses the formation of light from ATP and luciferin according to the following reaction:

\[
\text{ATP} + \text{Luciferin} + \text{O}_2 \rightarrow \text{OxyLuciferin} + \text{AMP} + \text{PPi} + \text{CO}_2 + \text{Light}
\]

(Lonza Rockland, Inc., www.lonza.com.biotechserv@lonza.com). By combining the two reactions, the emitted light intensity is linearly related to the AK concentration.

To quantify AK release, the exposed test solutions from the AB, NR and CB assays were collected after their respective time point exposures and were stored at −80 °C. Upon thawing, 20 μl of each sample (which contained the pooled supernatants from six replicate treatments per plate) were removed and transferred to a 96-well white luminescence plate (Falcon, VWR Ireland) for AK measurement. Media was assayed in triplicate for AK release. AK Detection Reagent (AKDR) was then reconstituted in Assay Buffer supplied with a kit. Leave for 15 min at room temperature to ensure complete rehydration. One-hundred microlitres of AK detection reagent was then added to each well and the plate
incubated at room temperature (RT) for 5 min and a 1 s integrated luminescence reading was recorded.

2.9. Apoptosis study

Ag nanoparticulate induce apoptosis was studied using Apo-Glow™ kit of Lonza Ltd. (Wokingham, UK). In brief, 100 µl the cells were seeded in 96-well white luminescence plates (Falcon, VWR Ireland) at a density of 1 × 10⁵ cells/ml in 100 µl of respective media. Following 24 h cell attachment, the plates were washed with 100 µl/well PBS and the cells were treated with increasing concentrations of Ag nanoparticles, prepared in 10% PBS containing media for 24 h. All incubations were performed at 37 °C in a 5% CO₂ humidified incubator. Six replicate wells were used for each control and test concentrations per microplate. After the exposure the apoptosis assay was performed following the manufacturers manual. In brief, the microplates were removed from the incubator following 24 h Ag nanoparticle exposure and were allowed to cool to room temperature. The luminometer was programmed to take a 1 s integrated reading. Then 100 µl/well of nucleotide releasing agent (NRR) was added and kept for at least 5 min. After that, 20 µl/well of NMR was added and a 1 s integrated reading (Reading A) was taken in luminometric plate reader. The plates were then kept for 10 min and 1 s integrated reading (Reading B) was taken. Then finally 20 µl/well of ADP-CR was added and again held for 5 min and finally 1 s integrated reading (Reading C) was taken. Finally the ADP/ATP ratio was calculated from the readings using the following formula (Reading C – Reading B)/Reading A.

This assay is based upon the bioluminescent measurement of ATP that is present in all metabolically active cells. The bioluminescent method uses luciferase, which generates light from ATP and luciferin according to the following reaction:

\[
\text{ATP} + \text{Luciferin} \xrightarrow{\text{Mg}, \text{Luciferase}} \text{Oxyluciferin} + \text{AMP} + \text{PPi} + \text{CO}_2
\]

(Lonza Rockland, Inc., www.lonza.com/biotechserv@lonza.com).

The emitted light intensity is linearly related to the ATP concentration and is measured using a luminometer. The assay is conducted at ambient temperature (18–22 °C), the optimal temperature for luciferase. This kit measures the intracellular ADP:ATP ratio which can be used to distinguish between the different modes of cell death, whether the cell dies via apoptosis or necrosis.

2.10. Statistical analysis

At least three independent experiments were conducted for each toxicity endpoint. Test results for each assay were expressed as percentage of the unexposed control ± standard deviation (SD). Control values were set as 100%. Differences between samples and the control were evaluated using the statistical analysis package SPSS 14.0. Statistically significant differences were set at \( p < 0.01 \). Normality of data was confirmed with Q–Q percentile plots and Kolmogorov–Smirnov tests. Equality of variances was evaluated using Levene tests. One-way analysis of variances (ANOVA) followed by Dunnett’s multiple comparison tests were carried out for normally distributed samples with homogeneous variances. Cytotoxicity data was fitted to a sigmoidal curve and a four parameter nonlinear logistic model used to calculate the Lethal Dose of nanomaterial that caused a 50% inhibition in comparison to untreated controls (LD₅₀). All LD₅₀ values were calculated using the average cytotoxicity data of the three independent experimental results and their associated errors. LD₅₀ values are reported ±95% confidence intervals (±95% CI). Lethal dose 50 (LD₅₀) values were estimated using Xlfit3™, a curve fitting add-on for Microsoft® Excel (ID Business Solutions, UK).

3. Results

3.1. Particle characterization

The particle size of the Ag nanoparticles was evaluated using a variety of methods namely DLS, TEM and SEM. DLS study have shown average number size-distribution peak-sizes of 100 mg/l Ag nanoparticles in dH₂O, HeLa media and in HaCaT media as 28.41, 65.45 and 68.55 nm, respectively (Fig. 1a). Size distribution plots of Ag nanoparticles in dH₂O and HaCaT media are also shown as an example in Fig. 1b and Fig. 1c, respectively. The sizes of the Ag nanoparticles were calculated from the highest peak value of size/number in DTS Nano software. The variation in size can be attributed to changes in the hydrodynamic radius of the particle in the different media due to particle and media interactions, where steric stabilization in the media was observed for the Ag nanoparticles of 65–69 nm size-range and the larger particles of around 500 nm size-range, as observed in Fig. 1c, may be due to particle agglomeration upon dispersion in the media. In fact DLS measures the hydrodynamic radius and not the physical particle size. The average diameter yielded from the TEM study was 29.57 nm (Fig. 1d) and SEM analysis was <5 nm (Fig. 1e). Both of which reflect more accurately the physical particle diameter. These combined studies confirmed the test particles had an average diameter in dH₂O of approximately 27 nm. The DLS size measurements of HaCaT and HeLa media was also performed as control and were found to be 8.14 and 9.24 nm, respectively (Fig. 1a). BET analysis was performed to estimate the surface area of the Ag nanoparticles and was found to be 2.3 ± 0.1 m²/g.

The average zeta potential measured of 100 mg/l Ag nanoparticles in dH₂O, HeLa media and in HaCaT media were −33.33, −20.67 and −15.83 mV (Fig. 1f). Zeta potential values of this nature would suggest that upon dispersion with the bath sonicator the particles form an unstable dispersion and would precipitate out with time. The average zeta potential of HaCaT and HeLa media without Ag nanoparticle were found to be 6.61 and −6.22 mV (Fig. 1d).

3.2. Standard cytotoxicity assays

For all assays, i.e., AB, NR, Coomassie Blue and MTT, a dose dependent response was observed and an LD₅₀ values were calculated. Cytotoxicity data for the Ag nanoparticles and the HaCaT and HeLa cells are presented in Table 1 and Figs. 2–5.

The LD₅₀ values presented in table suggest that MTT assay was the most sensitive assay among the four assays in both the cell lines. Coomassie Blue assay was found to be least sensitive in HaCaT cells, while in HeLa cells Neutral Red assay was found to be least sensitive. Viability was found to be decreasing with the increasing exposure time in AB, NR, CB, MTT assay in both the cell lines.

LD₅₀ values of HaCaT cells are much lower than HaCaT cells at all exposure time points in Coomassie Blue and Neutral Red assay. In Alamar Blue and MTT assay the LD₅₀ values of HeLa cells was slightly higher as compared to HaCaT cells after 24 h of exposure, but then again get lowered after 48 and 72 h exposure time.

The clonogenic assays showed decreasing colony numbers with the increasing concentration of Ag nanoparticles (Fig. 6). In the clonogenic assay the average plating efficiencies for the HaCaT and HeLa cells were 37% and 48%, respectively. The LD₅₀ values calculated from the clonogenic assay were 7.74 mg/l (confidence interval ±2.97) for the HaCaT cell line and 1.16 mg/l (confidence interval ±1.09) for that of the HeLa cells. At 2.5 and 5 mg/l Ag nanoparticle concentrations the percentage of colonies formed in HeLa cells were 13.96% and 0%, respectively. While in HaCaT cells at 2.5 mg/l and 5 mg/l Ag nanoparticle concentration there was 83.16% and 62.5% respective colonies were observed, the colonies
formed in HaCaT cells are much higher in percent than HeLa cells at same concentration. This shows that the proliferative capacity to form colonies of HaCaT cells in comparison to HeLa cells on exposure of Ag nanoparticles is much higher.

3.3. Optical microscopic study

Optical microscopic study have shown, with the increasing concentration of Ag nanoparticles the HaCaT (Fig. 7) and HeLa (Fig. 8) cells were looking unhealthy and the cells were dying out of the population. The population of HeLa cells were found to me more reduced at 200 mg/l Ag nanoparticle exposure for 24 h in comparison to HaCaT cells.

3.4. Reactive oxygen species study

The ROS study has shown that in HaCaT cells with the increasing time of Ag nanoparticles exposure more ROS was produced at their respective concentration (Fig. 9a). In HeLa cells, similarly the amount of ROS was found to be increasing with the increasing time of exposure at their respective concentrations (Fig. 9b). In HeLa cells the maximum amount of ROS was produced at 20 mg/l concentration after 5 h exposure of Ag nanoparticles, while in HaCaT cells maximum ROS production was observed at 31.25 mg/l concentration after 5 h exposure (Fig. 9). This study also shows that there was more ROS production in HeLa cells as compared to HaCaT cells.
adenylate kinase release in HaCaT (increasing concentration of Ag nanoparticles, the percentage of GSH depletion was 3.6% and 59.57% respective). The rate of intracellular GSH level depletion is much higher in HeLa cells in compare to HaCaT cells. 100 mg/l Ag nanoparticle in both HaCaT and HeLa cell lines after 24 h exposure.

### 3.5. Intracellular GSH study

The intracellular GSH measurement study have shown decrease in the intracellular GSH level with the increasing concentration of Ag nanoparticle in both HaCaT and HeLa cell lines after 24 h exposure (Fig. 10). The rate of intracellular GSH level depletion is much higher in HeLa cells in compare to HaCaT cells. 100 mg/l Ag nanoparticle exposures to HaCaT and HeLa cells for 24 h have shown 5.3% and 59.57% respective GSH depletion (Fig. 10).

### 3.6. Adenylate kinase release study

The adenylate kinase release study have shown that with the increasing concentration of Ag nanoparticles, the percentage of adenylate kinase release in HaCaT (Fig. 11a) and HeLa (Fig. 11b) cells were increased as compare to the controls. With the increasing hours of exposure, the adenylate kinase release from both the cell lines further increases (Fig. 11a and b). Upon 100 mg/l Ag nanoparticle exposure for 24, 48 and 72 h the percentage of AK release as compare to the controls from the HaCaT cells were 133.19%, 164.41% and 233.62%, respectively, while in HeLa cells the percentage of AK release at same concentration and exposure time points was found to be 156.5%, 195.33% and 226.48%, respectively. Therefore, more AK was released from HeLa cells as compared to HaCaT cell. The huge difference in AK activity was observed at the highest concentration in both the cell lines, in HeLa cells the percentage of AK activity was 373.89% at 150 mg/l while in HaCaT cells it was 384.14% at 800 mg/l after 72 h of Ag nanoparticle exposure. This shows that there was much more loss of cell integrity, through...
the damage of plasma membrane was occurred of HeLa cells as compared to HaCaT cells upon Ag nanoparticle exposure.

3.7. Apoptosis study

The apoptosis study was shown an increase in the ADP:ATP ratios over control ratios, which shows cells has entered apoptotic pathway. In HaCaT and HeLa cell lines with increasing concentration of Ag nanoparticles there was a significant increase in apoptotic cell population (Fig. 12). When both the cell lines were compared it was found HeLa cells are undergoing more apoptotic cell death than HaCaT cells at same concentration and exposure time. Upon 100 mg/l of Ag nanoparticle exposure for 24 h, ATP:ADP ratio in HeLa was 0.31 while in HaCaT it was 0.22. This shows a much lower level of ATP production in HeLa cells than HaCaT cells when compared to their respective controls upon exposure to the same concentration of Ag nanoparticle. The assay also displayed an increase in the intracellular ADP level in HeLa cells in comparison to the HaCaT cells, this resulted in an increase in the ADP:ATP ratios compared to their associated controls.

4. Discussion

Exposure of Ag nanoparticles to the body is becoming increasingly widespread and intimate. Consequently, there is an increased potential for direct contact between Ag nanoparticles, biomolecules and cells in specific outer and inner organs. This study examines the effect of Ag nanoparticles on human dermal noncancerous (HaCaT) and cervical cancer (HeLa) cell lines. The toxicity of Ag nanoparticle is well known (AshaRani et al., 2009; Moaddab et al., 2011; Kim et al., 2009) but how cells of diverse origin react with the same Ag nanoparticle in different manner and the possible reason behind this difference has been illustrated in this paper.

The physio-chemical characteristics of nanoparticles play a significant role in their effect on biological systems (Oberdörster et al., 2000; Donaldson et al., 2004). The principal parameters of nanoparticles are their shape (including aspect ratios where appropriate), size, and the morphological sub-structure of the substance. The chemical composition and the intrinsic toxicological properties of the chemical are also of importance for toxicity of particles (Renwick et al., 2004). The zeta potential of the particle has been measured by DLS technique shows its size measurement of AgNP by DLS technique shows its potential for direct contact between Ag nanoparticles, biomolecules and cells in specific outer and inner organs. This study examines the effect of Ag nanoparticles on human dermal noncancerous (HaCaT) and cervical cancer (HeLa) cell lines. The toxicity of Ag nanoparticle is well known (AshaRani et al., 2009; Moaddab et al., 2011; Kim et al., 2009) but how cells of diverse origin react with the same Ag nanoparticle in different manner and the possible reason behind this difference has been illustrated in this paper.

The size measurement of AgNP by DLS technique shows its potential for direct contact between Ag nanoparticles, biomolecules and cells in specific outer and inner organs. This study examines the effect of Ag nanoparticles on human dermal noncancerous (HaCaT) and cervical cancer (HeLa) cell lines. The toxicity of Ag nanoparticle is well known (AshaRani et al., 2009; Moaddab et al., 2011; Kim et al., 2009) but how cells of diverse origin react with the same Ag nanoparticle in different manner and the possible reason behind this difference has been illustrated in this paper.

Fig. 4. Cytotoxicity of Ag nanoparticles after 24, 48, and 72 h exposures determined by the Coomassie Blue assay – (a) in HaCaT cells, (b) in HeLa cells. Data are expressed as percent of control mean ± SD of three independent experiments. Data are expressed as percent of control mean ± SD of three independent experiments. * denotes a statistically significant (p < 0.05) difference from the unexposed control.

Fig. 5. Cytotoxicity of Ag nanoparticles after 24, 48, and 72 h exposures determined by the MTT assay – (a) in HaCaT cells, (b) in HeLa cells. Data are expressed as percent of control mean ± SD of three independent experiments. Data are expressed as percent of control mean ± SD of three independent experiments. * denotes a statistically significant (p < 0.01) difference from the unexposed control.
cell culture media components, which have been widely reported with different nanoparticles that leads to the formation of ‘protein corona’ (Lynch and Dawson, 2008; Lundqvist et al., 2008). The zeta potential study have also shown the decrease in the negative zeta potential of the AgNP upon its dispersion in the 10% FBS supplemented cell culture media, where in dH₂O it was −33.33 mV and in HeLa and HaCaT media it was −20.67 and −15.83 mV, respectively (Fig. 1f). The zeta potential study further confirms the interaction of AgNP with the cell culture media components. Such an interaction of the nanoparticles with the components of the cell culture medium has been demonstrated to elicit a secondary or indirect toxic response in the case of single walled carbon nanotubes (Casey et al., 2007, 2008) and it is probable that there are similar contributions to the toxic response observed here. Therefore, in an independent experiment to investigate the cytotoxicity of Ag nanoparticle interacted media was performed. Where different concentrations of Ag nanoparticle were first prepared in the respective cell culture media and kept in 37 °C incubator for 24 h and then following incubation Ag nanoparticles were removed from the media by centrifuging it at 14,000 rpm for 20 min. The supernatants were then collected and filtered using cellulose acetate 0.2 μm syringe filters. Both the tested cell lines were then exposed to this supernatant for 24–72 h and cellular viability assessed using the AB assay. The toxicity found from supernatant media was not comparable in magnitude to the cytotoxicity observed from direct nanoparticle exposure and was only observed at the 72 h exposure time-point. Upon exposure of 100 mg/l Ag nanoparticle supernatant in HeLa cells after 72 h exposure to the

![Graph showing cytotoxicity of Ag nanoparticles after 8 days exposures determined by the Clonogenic assay in HaCaT cells and HeLa cells.](image)

![Optical micrograph of HaCaT cells after 24 h Ag nanoparticle exposure at (a) 0 mg/l, (b) 50 mg/l, (c) 100 mg/l and (d) 200 mg/l.](image)
supernatant, the cell viability observed was 86.74%, while no cytotoxicity was observed in HaCaT cells (data not shown). This data suggests that there is indeed a contribution of secondary cytotoxicity from the Ag nanoparticle and cell culture media interaction however the observed effects were only noted at longer exposure time-points.

Following physiochemical characterization a series of toxicological studies were performed using a battery of assays to assess the in vitro cytotoxic effects of Ag nanoparticle on these two cell lines of diverse tissues and to understand the difference in the response by the two cell lines.

In Table 1 dose response from AB assay shows that in HaCaT and HeLa cell lines the cytotoxic response was increasing with the increasing concentration and time points of exposure. This indicates particles have a toxic effect on the basic cellular metabolism and proliferation of the cells as there was a decrease in the reduction of the Alamar Blue dye from Resazurin to Resorufin as observed from this assay. This Redox reaction could not happen as cytosolic and mitochondrial oxidoreductase enzyme activities were decreased with the increase in Ag nanoparticle concentration and exposure time (Fig. 2). Upon 100 mg/l AgNP exposure for 72 h, the decrease in the AB dye reduction than the control in HaCaT and HeLa cells were 16.75% and 1.11%, respectively. This further indicates the higher sensitivity of HeLa cells than HaCaT cells upon Ag nanoparticle exposure.

The dose response of Ag nanoparticles to HaCaT and HeLa cells in NR assay shows that with the increasing concentration and time of exposure of AgNPs, less NR dye was uptaken by the cellular lysosomes, which indicated the cellular viability was decreasing with the increasing concentration and time of exposure of AgNPs. In dead cells, NR can no longer retain in the cytoplasmic vacuoles and the plasma membrane does not act as a barrier to retain the NR within the cells. Fig. 3 has shown the decrease in NR dye uptake in HaCaT and HeLa cells were 25.38% and 27.42%, respectively upon 100 mg/l Ag nanoparticle exposure for 72 h.

CB assay shows that with the increasing concentration and exposure time the CB dye binding was decreased in both HaCaT and HeLa cells (Fig. 4). That suggests the degradation of cellular protein or reduction of protein synthesis upon AgNP exposure (Clare and Cormack, 1996). The percentage of CB dye binding to the cells than control upon 100 mg/l AgNP exposure for 72 h in HaCaT and HeLa cells were 51.34% and 15.75%, respectively. In this assay again the HeLa cells were found to be more sensitive than HaCaT cells.

The MTT assay determines the ability of viable cell’s mitochondria to reduce the soluble, yellow MTT into an insoluble, purple formazan. The reduction of MTT to formazan indicates the decrease in mitochondrial metabolism of the cells. Therefore, the absorbance of formazan formed, directly correlates to the number of cells whose mitochondrial metabolism is intact even after the...
exposure of Ag nanoparticles. Dose response of AgNP in HaCaT and HeLa cells shows the decrease in the reduction of MTT to formazan with the increasing concentration and time of exposure. Upon 100 mg/l AgNP exposure for 72 h in HaCaT and HeLa cells, the percentage of MTT conversion to formazan decreases to 17.71% and 3.07%, respectively (Fig. 5). Therefore MTT assay also shows the higher sensitivity of HeLa cells to the Ag nanoparticles than HaCaT cells.

Clonogenic assay indicates the proliferative capacity of a single cell upon nanoparticle exposure for longer time period, which in this study was 8 days. Dose response from this assay shows the decrease in the colony number with the increasing concentration of AgNP in both the cell lines (Fig. 6). However, it also shows that the colony forming capacity in HeLa cells were much lesser than HaCaT cells upon AgNP exposure. Exposure of 2.5 mg/l AgNP to HaCaT cells can form 83% colonies compare to the control, while HeLa cells can only form 14% colonies. This is also reflected in the LD$_{50}$ values obtained from this study. The LD$_{50}$ in HaCaT cells was 7.74 mg/l, while that in HeLa cells was 1.16 mg/l. Clonogenic assay therefore also suggest the higher sensitivity of HeLa cells when compare to HaCaT cells.

The responses from all the standard cytotoxicological assays (AB, NR, CB, MTT and Clonogenic assays) employed to test the Ag nanoparticle toxicity on the HaCaT and HeLa cells were found to be both dose and time dependant. In both cell lines the LD$_{50}$ values were seen to decrease with increasing exposure time. In all cases the cytotoxic effect was noted to be greater in HeLa cells than in HaCaT cells. Among all the colorimetric based short term exposure (up to 72 h) acute toxicity assays (i.e., AB, NR, CB and MTT assays) in both the cell lines, the MTT assay was noted to be the most sensitive as indicated by the lower LD$_{50}$ values (Table 1) evaluated from this assay. The AB assay showed less sensitivity than MTT
crease in ROS was noted in exposure concentrations well below 30 min in both the cell lines. In HaCaT cell line a significant increase in all the tested concentrations started typically within 5–10 min of the unexposed controls. It was noted that the onset of oxidative stress occurs faster in Ag nanoparticles in comparison to control and was within the range of apoptotic cell population (Fig. 12). The ADP:ATP ratio in HaCaT and HeLa cells upon 24 h 100 mg/l Ag nanoparticle exposure were 0.22 and 0.31, respectively. This indicates upon Ag nanoparticle exposure the cells were entering apoptosis and HeLa cells had more apoptotic cell population than HaCaT cells. This further indicates the higher sensitivity of HeLa cells than HaCaT cells.

The dose responses from these assays suggest that upon Ag nanoparticle exposure the mitochondrial metabolism was most affected followed by the basic cellular metabolism and then the lysosomal activity and finally because of the toxic effect of these nanoparticles on the cells the cellular protein contents were also decreased. Among both the cell lines, HeLa cells were found to be more sensitive than HaCaT cells. One significant difference between the two cell lines is their natural antioxidant levels. The natural glutathione (GSH) level of HaCaT cell is 7300 nmol/mg protein (Snow et al., 2005) and HeLa cell is ~220 nmol/mg protein (Tipnis et al., 1999). This huge difference in their natural GSH level could be an important reason for higher sensitivity of HeLa cells over HaCaT cells.

Following the standard cytotoxicity assays intracellular ROS generation was also monitored. In both cell lines a significant increase in the levels of ROS (Fig. 9) was observed in comparison to the unexposed controls. It was noted that the onset of oxidative stress in all the tested concentrations started typically within 30 min in both the cell lines. In HaCaT cell line a significant increase in ROS was noted in exposure concentrations well below the calculated LD_{50} values. A response of this nature would that at concentrations even were there was no observed cytotoxicity from the above standard cytotoxicity assays the presence of Ag nanoparticles can induce high levels of cellular oxidative stress. The amount of ROS generated upon AgNP exposure was higher in HeLa cells than in HaCaT cells (Fig. 9). The fold increase of ROS than control upon 10–20 mg/l AgNP exposure in HeLa cells were 3.93–4.39, whereas upon 15 mg/l AgNP exposure that in HaCaT cells was 1.77. This could be due to the much lower natural GSH level in HeLa cells (Tipnis et al., 1999) than HaCaT cells (Snow et al., 2005) as GSH is an important antioxidant and ROS scavenger of the cell. Therefore an independent GSH study has shown that the rate of depletion of intracellular GSH level was much higher in HeLa cells when compared to that of HaCaT cells (Fig. 10). Such differences would account for the higher amount of ROS production observed in HeLa cells in comparison to HaCaT cells. ROS plays an important role in triggering many cellular pathways which can lead to cellular death, including cytokine activation (Brown et al., 2004) and caspase activation (Kim et al., 2009, Stennicke et al., 1998). They can also cause damage to the nuclear DNA by altering the chemical structure of the nucleotide bases and the deoxyribosyl backbone (Cooke et al., 2003, Valko et al., 2004). Therefore, intracellular ROS generation upon AgNP exposure is an important factor to its cytotoxicity. ToxiLight™ Assay assay quantifies the release of adenylate kinase from the cells which is an indication of the damage of the cell membrane. Once the plasma membrane has been damaged by cytotoxic action the AK will be available for detection in the surrounding medium. With the increasing Ag nanoparticle concentration in both the cell lines, an increasing release of adenylate kinase from the cells was observed which also increases with the exposure time (Fig. 11). This assay therefore suggests that with the increasing concentration and exposure time the Ag nanoparticle causes more damage to the cell membrane. Upon AgNP exposure more AK was found to be released from HeLa cells than HaCaT cells at lower dose (Fig. 11).

This damage in the cell membrane also indicates that more cell population is possibly dying by either necrotic or apoptotic pathway. This was confirmed when apoptosis study was performed by ApoGlow Kit of Lonza Ltd (Wokingham, UK). ApoGlow Kit helps in the differentiation of normal healthy proliferative cells, apoptotic cells and necrotic cells from the cellular ADP:ATP ratio. If a cell undergoes apoptosis, the test gives lower levels of ATP compared to controls, but show an increase in ADP, this eventually gives an increase in the ADP:ATP ratios over control ratios. In normal healthy cells the ADP:ATP ratio was observed below 0.11, whereas in apoptotic cells the ratio was in between 0.11 and 1.0 and in cells undergoing necrosis by heat shock treatment the ADP:ATP ratio was observed in excess of 15.0 (Bradbury et al., 2000). The apoptosis study shows the ADP:ATP ratio in HaCaT and HeLa cells upon AgNP exposure were higher than that of the control and was within the range of apoptotic cell population (Fig. 12). The ADP:ATP ratio in HaCaT and HeLa cells upon 24 h 100 mg/l Ag nanoparticle exposure were 0.22 and 0.31, respectively. This indicates upon Ag nanoparticle exposure the cells were entering apoptosis and HeLa cells had more apoptotic cell population than HaCaT cells. This further indicates the higher sensitivity of HeLa cells than HaCaT cells.

Overall, the standard cytotoxicity assays shows more sensitivity of MTT assay upon short term exposure over other assays followed by AB assay, NR and CB assays were the least sensitive ones. This further indicates upon Ag nanoparticle exposure mitochondria was getting affected first followed by basic cellular metabolism and lysosomal activity was decreasing later on. Optical microscopic study had also shown the cytotoxic effect of Ag nanoparticles on HaCaT (Fig. 7) and HeLa (Fig. 8) cells with their increasing concentration. Ag nanoparticles generate reactive oxygen species upon entering the cells and that creates oxidative stress to the cells. ROS plays an important role in the mitochondrial damage, therefore its generation upon AgNP exposure causing the higher mitochondrial toxicity. Ag nanoparticles also damage the cell membrane and therefore more AK release from the cells was observed upon increasing concentration and time of exposure of Ag nanoparticles. All these effects further activate the apoptotic pathways and therefore significant apoptotic cell populations were observed upon 24 h Ag nanoparticle exposure and eventually cell dies. The interaction of AgNP with the media components, as indi-
cated from the size and zeta potential measurements study, may also lead to secondary toxicity. The difference in the sensitivity of HaCaT and HeLa cells can be understood in terms of the difference in their natural antioxidant levels.

5. Conclusions

Size measurement study of Ag nanoparticles in the cell culture media showed the interaction of AgNP with the cell culture media components that suggest the significant protein adsorption by the nanoparticles in the cell growth media. This protein adsorption from the medium is also evident in a dramatic change in the zeta potential and therefore surface properties of the particles. This could have possible effect on the toxic response of the Ag nanoparticles. HeLa cells were found to be more sensitive than HaCaT cells in all the assays and clonogenic assay showed the strongest response indicating potential chronic responses. All these different short term cytotoxicity assays show the effect of silver nanoparticles at different cellular level. Therefore proper interpretation of the dose response from these assays can help in the prediction of the mechanism of its cytotoxicity. ROS generation by Ag nanoparticles upon entering the cells is the key step which possibly initiates several pathways that leads to cell death. The difference in the sensitivity of different cell lines can be understood in terms of their natural antioxidant levels, which is a key behind their natural defence mechanism during oxidative stress. Therefore, this probably governs the sensitivity of the cell lines upon same nanoparticle exposure.

Conflict of interest

The author declares that there is no conflict of interest.

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