The Role of Apoptosis Pathway in the Cytotoxicity Induced by Fresh and Aged Zinc Oxide Nanoparticles

Juan Wang1,2†, Lei Wang4†, Wenting Zhao1, Na Yu1, Meiling Cheng1, Mingqin Su1, Jian Hu1, Xiaoyan Wu2, Hua Du3* and Meimei Wang1,2*

Abstract

Zinc oxide nanoparticles (ZnO NPs) are used in a wide range of applications including industry, commercial products and medicine field. Numerous mechanistic studies for ZnO NPs' toxicity were performed on pristine (fresh) NPs. However, the cytotoxicity induced by the transformed (aged) ZnO NPs and the underlying mechanisms remain unclear. Here, we observed the physicochemical transformation of ZnO NPs underwent over time, followed by evaluating the cytotoxicity of fresh and aged NPs. We found that fresh ZnO NPs induced higher apoptosis level than their aged counterparts. Accordingly, RNA sequencing data from aged ZnO NP-treated human–hamster hybrid (A/L) cells showed that p53, PI3k–Akt, FoXO, Glutathione, ErbB, HIF-1, Oxytocin and Jak-STAT signaling pathways were enriched but no apoptosis pathway. Quantitative PCR results revealed the significantly higher mRNA level of IL1B and CD69 in fresh NP-treated groups compared to that of aged ZnO NP- and zinc chloride-treated groups. The above results indicated that the lower cytotoxicity of aged ZnO NPs is partially attributed to their reduced potency in inducing apoptosis. The transcriptional regulation of multiple signal pathways activated by aged NPs may help to build the cellular homeostasis. Taken together, our findings highlight the influence of aging (environmental transformation) process of ZnO NPs on their toxicities and biological consequences.

Keywords: Aged zinc oxide nanoparticles, Cytotoxicity, Mechanism, Apoptosis, Transcriptomics

Introduction

With the rapid development of nanotechnology over the past decades, nanoparticles (NPs) have been applied in various fields, including industry, human daily life and nanomedicine [1, 2]. The Nanotechnology Consumer Product Inventory (CPI) shows a 30-fold increase between 2005 and 2015 in the numbers of nano-products, including 762 health (fitness) products, 72 food (beverages) and 23 baby products [2]. The growing application of NPs in consumer products and various fields increased the possibility of NPs entering into the environment, which raises safety concerns with regard to their potential adverse impacts. Zinc oxide (ZnO) NPs are among the most commonly-used NPs, and its global annual output has reached nearly 3400 tons [3, 4]. Some substances that are previously considered as biologically inert could become toxic in their nanoparticulate state. An increasing number of studies elucidated that ZnO NPs may pose significant risks to mammalian cells and animals by inducing significant toxicity [5–7]. Various strategies including coating, surface functionalization and oxidation state modification have been used to attenuate the potential toxicity of NPs by modifying...
the physical and chemical properties of them (such as the dissolution, agglomeration and perturbation of cell membranes) [8–11]. Although these modifications of NPs weaken their toxic effects in certain degree, the uses of NPs are not always safe, especially under certain exposure conditions and environments [12–14]. Actually, many kinds of NPs are not stable and inclined to undergo “aging” or “environmental transformation” after being intentionally or unintentionally released into the natural environment [14–17]. In recent years, lots of work were carried out to explore the environmental transformation process of NPs; however, the research on the toxic effects of “transformed (aged)” NPs is still very limited, let alone their toxic mechanisms.

As the typical representative of non-persistent NPs, ZnO NPs have very high reactivity, and are prone to transform in physical and chemical properties and occurrence state after being released into the environment or ingested by animals, which could significantly affect their toxicological effects [17, 18]. For example, studies have shown that the sulfidation process of ZnO NPs changed the charge, hydrophobicity and aggregation state, resulting in the adsorption of sulfide state NPs in human saliva, sweat and bronchoalveolar lavage fluid. Furthermore, the protein adsorbed by ZnO NPs forms a special protein crown that usually affects its biological effect [19]. Phosphates in physiological solutions could convert ZnO NPs into metastable ZnHPO₄ and Zn₃(PO₄)₂, within about 5–10 h [20]. The process of complete transformation of ZnO NPs (≤3 μg/mL) in the in vitro exposure system to human T lymphocytes (37 °C, cell culture medium RPMI1640 containing 10% FBS for 24 h) was investigated by using synchrotron radiation X-ray absorption near-edge structure spectroscopy (XANES) [21]. The above studies suggest the underestimation of ZnO NPs’ environmental and health risks by solely evaluating the biological effects of pristine (fresh) ZnO NPs. In the light of this problem, there is an urgent need to comprehensively understand the aging and environmental transformation processes of NPs [22].

Our previous study revealed that ZnO NPs aged for 40–120 days in ultrapure water undergo physicochemical transformation and turn into Zn₅(CO₃)₂(OH)₆, Zn(OH)₂, and Zn²⁺ [23]. Interestingly, aged ZnO NPs exhibited lower cytotoxicity than the fresh counterparts [23], yet the toxicity mechanisms of such kind of variation are unclear. In the present study, we set out to explore the underlying reasons of different cytotoxicities between fresh and aged ZnO NPs. ZnO NPs with two different particle sizes (20 nm and 90–200 nm) were applied systematically. The cytotoxicity assays demonstrated that aged ZnO NPs induced less pronounced morphological abnormalities and relatively higher cell viabilities than their fresh counterparts. RNA sequencing data revealed that apoptotic genes were enriched in fresh ZnO NP-treated cells, whereas these genes were much less affected by aged ZnO NP-treatments. In addition, the cells exposed to aged ZnO NPs showed reduced level of cleaved Caspase-3 protein, further indicating the higher potency of fresh ZnO NPs in eliciting apoptosis in cultured cells. Combined with our previous findings, this study suggested that the decreased cytotoxicity of aged ZnO NPs is attributed to their attenuated ability in triggering cell apoptosis.

Materials and Methods

Nanoparticles and Reagents

The commercially available ZnO nanopowders (ZnO NPs), with manufacturer’s reported average size 20 nm (99.5% purity, nearly spherical) and 90–200 nm (99.9% purity, irregular morphology), were purchased from Nanostructured & Amorphous Materials (Houston, TX). Except for otherwise noted, all the reagents and chemicals used in this study were purchased from Sigma-Aldrich (Shanghai, China).

Nanoparticle Dispersion, Aging and Characterization

ZnO NPs stock suspensions (1 mg/mL) were prepared by suspending dry nanopowders in Milli-Q H₂O (Millipore, 18 MΩ cm) and sterilized by autoclaving (120 °C, 30 min) and then stored at 25 °C for natural aging period ranging from 0 to 60 days. The 0- and 60-days’ naturally transformed ZnO NPs were designated as fresh and aged NPs, respectively. To ensure proper dispersion, the fresh and aged suspensions were sonicated (100 W) for 30 min in an ultrasonic bath before characterization or incubation with cells. The morphology, particle size and aggregation of fresh and aged ZnO NPs were characterized by using transmission electron microscopy (TEM, JEOL JEM-2010, Tokyo, Japan). The crystal structure of fresh and aged ZnO NPs was determined using powder X-ray diffraction (XRD, PANalytical B. V., Shanghai, China) by comparing to authentic standards. The details of natural aging process and characterization on ZnO NPs have been described previously [23].

Cell Culture and Treatment with ZnO NPs

A₁, cell line, a kind of human–hamster hybrid cells formed by fusion of the gly24A mutant of Chinese hamster ovary (CHO) and human fibroblasts was used in this study. These hybrid cells contained a standard set of CHO-K1 chromosomes and a single copy of human chromosome 11 and were cultured in Ham’s F12 medium (HyClone, Grand Island, NY) supplemented with fetal bovine serum (8%, HyClone, Grand Island, NY), gentamicin (25 g/mL) and glycine (2 × 10⁻⁴ M) at 37 °C in a humidified 5% CO₂
Aspirated, and cells were washed 3 times with PBS. Subsequently, 1 mL of Trizol reagent (Invitrogen, Carlsbad, CA, USA) was added to each dish to extract total RNA. The total RNA samples of A549 cells from control group, aged ZnO NP-treated group and ZnCl2 treated group were sequenced by BangFei Bioscience (Beijing, China). The stock suspensions of fresh and aged ZnO NPs were dispersed by 30 min of ultrasonication (100 W) to prevent agglomeration, subsequently diluted to appropriate concentrations with cell culture media for the exposure of concentrations. Cells maintained in cell culture media without NPs were served as control in each experiment.

**Assay for Detecting the Cytotoxicity**

A549 cells at a logarithmic phase of growth were cultured on glass slides in 35-mm Petri dishes (6 × 10^4 cells/dish) for 24 h before stimulation, followed by treatment with 2 mL of culture medium containing 1, 5, 10, 12, 15 and 20 μg/mL fresh or aged ZnO NPs 72 h. After the completion of treatment time, the images of cell morphology were obtained using a Leica DM4B microscope (Leica, Germany). ZnCl2 was included as zinc ions reference for comparing the cytotoxicity with ZnO NPs.

The cell counting kit (CCK-8) (APExBIO, Shanghai, China) was used for detecting the cell viability. In details, A549 cells were seeded into 96-well plates (4 × 10^4 cells/well) with cell culture media for 24 h and treated with medium containing various concentrations of ZnCl2, fresh and aged ZnO NPs for 24, 48 and 72 h, respectively. For working solution, the volume of added NPs from the stock suspension was less than 5% of the total volume of the culture medium in each well. After the completion of treatment time, the culture medium was aspirated, and the cells were incubated with 100 μL CCK-8 working solution for 2 h at 37 °C following the manufacturer’s instructions. Then, the absorbance was recorded at 450 nm using a Spectra Max M2 fluorescence reader (Molecular Devices, Wokingham, Berks, UK). Cell viability was calculated as a percentage of absorbance in wells, with each concentration of NPs normalized to the absorbance of control cells (100%).

**RNA Extraction, Reverse Transcription and Quantitative PCR**

A549 cells at a logarithmic phase of growth were seeded into 35-mm-diameter Petri dish (6 × 10^4 cells/dish) with cell culture media for 24 h. Then, the medium was replaced with 2 mL of culture medium containing 12 μg/mL ZnCl2, fresh and aged ZnO NPs for 72 h. After the completion of treatment time, the culture medium was aspirated, and cells were washed 3 times with PBS. Subsequently, 1 mL of Trizol reagent (Invitrogen, Carlsbad, CA, USA) was added to each dish to extract total RNA according to the manufacturer’s instructions. Concentration and purity of total RNA obtained after the extraction were quantified using a Q5000 UV-Vis Spectrophotometer (Quawell, USA). After quantification, reverse transcription was performed using TransGene RT-PCR kit (TransGene Biotech, Beijing, China) to obtain cDNA from the RNA template according to the manufacturer’s protocols. The resulting cDNA samples were quantified by using the Q5000 UV–Vis Spectrophotometer and then analyzed using SYBR-Green as fluorescence dye (TransGene Biotech, Beijing, China) on Roche RT-PCR system (Applied Biosystems) [25].

The housekeeping gene encoding Glyceraldehyde-3-phosphate Dehydrogenase (Gapdh) was used as internal control for evaluating Il-1a, Il-1β, Caspase 3, CD69, Jun and MT1 mRNA expression. The results were expressed as the relative expression ratio between the targeted gene and Gapdh. The primer sequences used in this study are provided in Table 1.

**RNA Sequencing Data Analysis**

The total RNA samples of A549 cells from control group, aged ZnO NP-treated group and ZnCl2 treated group were sequenced by BangFei Bioscience (Beijing, China). Briefly, the total RNA of A549 cells was extracted following the TRIZOL protocols, until the isopropanol precipitation. Then, the RNA samples were resuspended in the extraction buffer before sequencing. The raw count RNA sequencing data were analyzed using R package Deseq2 [Eric1]. The venn diagram was generated by R package VennDiagram [Eric1.2]. The significantly changed genes were used for further pathway enrichment analysis. Experiments were done three independent replicates. rRNA genes, mitochondrial genes and the genes detected less than 40 bp were excluded from the analysis.

The RNA sequencing data, reference series GSE97852, GSE60159 and GSE39444, were obtained from Gene Expression Omnibus.

| Name   | Primer | Sequence | Length (bp) |
|--------|--------|----------|-------------|
| Il-1a  | Forward| 5’CGTCCGTCGTAATACGAGG3’ | 178         |
|        | Reverse| 5’GACACCTGACACCTTCAG3’   |             |
| Il-1β  | Forward| 5’ACCTTCCAGGATGAGGACATG3’| 121         |
|        | Reverse| 5’GCAATTCGAGATGACACCT3’  |             |
| Caspase3| Forward| 5’TCTTCTTCTGGAATTCTAG3’ | 162         |
|        | Reverse| 5’GGAATTCTGCGAGAGTGG3’   |             |
| CD69   | Forward| 5’GCCACACGCGCTCTGTGTCA3’ | 163         |
|        | Reverse| 5’GACAGCTGCGAGACACTT3’   |             |
| Jun    | Forward| 5’TGATATCGTCGAGACCTA3’   | 177         |
|        | Reverse| 5’CTAGTTCGAGATGACACCT3’  |             |
| MT1    | Forward| 5’GCCACACGCGCTCTGTGTCA3’ | 126         |
|        | Reverse| 5’GACACCTGACACCTTCAG3’   |             |
| Gapdh  | Forward| 5’GGAATTCTGCGAGAGTGG3’   | 123         |
|        | Reverse| 5’GCAATTCGAGATGACACCT3’  |             |

**Table 1** Primers used in this study
Expression Omnibus [Eric 2, 3, 4]. The Gene Set Enrichment Analysis plot was generated by R (version 3.6.2) using package fgsea [Eric 5]. The apoptosis genes with 1.5-fold significant change & p value < 0.05 were used for further analysis. The heatmap with gene tree was generated by R package ‘ComplexHeatmap’ [Eric 6]. Average linkage was used as the clustering method, and Euclidean was used as a distance measurement method. The pathway enrichment analysis was preceded using STRING2.0 [Eric 7].

Western Blotting

A549 cells at a logarithmic phase of growth were seeded into 60-mm-diameter Petri dish (1.5 x 10⁵ cells/dish) with cell culture media for 24 h. Then, the medium was replaced with 4 mL of culture medium containing 12 µg/mL fresh or aged ZnO NPs for 24 h. At the end of exposure period, the culture medium was aspirated, and then, cells were washed 3 times with PBS and lysed on ice with RIPA lysis buffer (Beyotime, China) to collect cellular proteins. Equal amounts of cellular proteins were separated on 12% SDS-PAGE gels and then transferred to a polyvinylidene fluoride (PVDF) membrane (Roche, Swiss). Briefly, after 2 h blocking with 5% nonfat milk in TBST at 25 °C, the membranes were subsequently incubated with primary antibody at appropriate dilutions (according to the manufacturer’s protocols) at 4 °C overnight, followed by incubating with HRP-conjugated secondary antibodies (1:5000, Promega, Madison, USA) for 2 h at 25 °C. Finally, immunolabeling was detected using an enhanced chemiluminescence (ECL) (BOSTER, China) solution. The primary antibodies of anti-pro/cleaved Caspase-3 and anti-Actin were purchased from Cell Signaling Technology and ImmunoWay, respectively.

Statistics

Statistical analysis was compiled on the means of the results obtained from at least three independent experiments. All Data were presented as means ± standard deviation (SD) and statistically compared using one-way analysis of variance (ANOVA). In all plots p values < 0.05 were showed as * and considered to be statistically significant.

Results

Characterization of ZnO NPs

To determine the differences in detailed physicochemical characteristics between fresh and aged ZnO NPs, we first observed the morphology of NPs using TEM (Fig. 1A). Our results indicated that 20 nm fresh ZnO NPs were nearly spherical crystals and 90–200 nm fresh ZnO NPs were irregularly rod-like/cubical crystals. The single particle size was consistent with the size provided by the manufacturer. Obviously, both 20 nm and 90–200 nm ZnO NPs were inclined to aggregate in ultrapure water. Also, regardless of the shape and size of the original NPs, both 20 nm and 90–200 nm ZnO NPs’ microstructure was dramatically changed from a clear crystal structure to an amorphous or sheet-/needle-like state after aged for 60 days. Furthermore, the crystalline nature and phase purity of both fresh and aged NPs were determined by using X-ray diffraction (XRD) with Cu Kα radiation (λ = 0.15418 nm) approach at 25 °C, as shown in Fig. 1B. The XRD pattern of fresh ZnO NPs

Fig. 1 Physicochemical characteristics of fresh and aged ZnO NPs. A Representative micrographs of fresh and aged NPs (100 µg/mL, 20 and 90–200 nm) in Milli-Q water using low resolution TEM; B XRD patterns of fresh NPs, aged NPs, ZnO, Zn (OH)₂ and Zn₅(CO₃)₂(OH)₆ references in dried form.
indicated that the samples were comprised of crystalline wurtzite structure and no characteristic impurity peaks were identified, suggesting a high quality of fresh NPs. For aged NPs, the XRD pattern exhibited the neoformation of Zn₅(CO₃)₂(OH)₆ (card number 00-011-0287) and Zn(OH)₂ (card number 00-003-0888) solid phases, indicating the chemical transformation of ZnO NPs (20 and 90–200 nm) during the aging process.

**Morphological Observation of A₅ Cells Exposed to Fresh and Aged ZnO NPs**

NPs’ treatment results in a noticeable change in cellular shape, or morphology, in vitro [26]. Therefore, A₅ cells exposed to fresh or aged ZnO NPs at 10 & 15 μg/mL for 72 h were examined under a stereoscopic microscope. As shown in Fig. 2, cell morphology in the control group remained normal. The cells adhered well, with most attaching within 2 h. Most cells were spindle shaped or polygonal, with a few newly dividing cells showing a more transparent cytoplasm and better dispersion during the process of adhering. Treatment with 12 μg/mL fresh ZnO NPs (20 nm & 90–200 nm) for 72 h significantly changed cell morphology. Although most cells adhered within 3–5 h, they could not spread well, and some cells became rounded and lost the polygonal shape. When the concentration of ZnO NPs was increased to 15 μg/mL, the treated cells atrophied and some cells became rounded or polygonal, with a few newly dividing cells showing a more transparent cytoplasm and better dispersion during the process of adhering. Treatment with 12 μg/mL fresh ZnO NPs (20 nm & 90–200 nm) for 72 h significantly changed cell morphology. Although most cells adhered within 3–5 h, they could not spread well, and some cells became rounded and lost the polygonal shape. When the concentration of ZnO NPs was increased to 15 μg/mL, the treated cells atrophied and could not adhere, suggesting their significantly lower cell viability than that of the cells treated with 10 μg/mL. These results indicated that the LC100 for fresh ZnO NPs is probably less than 15 μg/mL by a 72-h treatment. In contrast, cell morphology in 20 nm and 90–200 nm aged NP-treated groups (15 μg/mL) was not significantly affected, and most of the surviving cells could adhere and spread, with less than half dead cells observed, unveiling that aged ZnO NPs are much lower cytotoxic than fresh ZnO NPs.

**Fresh ZnO NPs’ Treatment Activated Apoptosis Pathways and Up-Regulated the Expression of Apoptotic Genes**

To unveil the underlying mechanisms leading to the lower cytotoxicity of aged NPs, we analyzed RNA sequencing data from both fresh and aged ZnO NPs. As shown in Fig. 4A, B, after treatment with fresh ZnO NPs, apoptosis pathway was activated in Jurkat cells (p = 0.017) and HMDM cells (p = 0.041). The apoptosis genes: ANXA1, CYLD, TNFSF10, IER3, CDKN1A, JUN, SAT1, PMAIP1, CD38 and ISG20 were significantly enriched in fresh ZnO NP-treated Jurkat cells. The apoptosis genes: CD38, TNFRSF12A, CCNA1, BMP2, PPP2R5B, EREG, IFNGR1, CD44, CD14, GNA15, GCH1, TIMP1, BTG2, IL1B, IL1A, BTG3, BCL2L11, SC5D and SPTAN1 were significantly enriched in fresh ZnO NP-treated HMDM cells (Fig. 4C, D). Since Jurket cells (peripheral blood T lymphocyte cells) and HMDM cells (human monocyte-derived macrophages) are different kinds of cells, the way they trigger apoptosis might be different. In sum, these results showed that fresh ZnO NPs’ exposure could activate different apoptosis pathways in various kinds of cells.

**Aged ZnO NPs Induced Lower Cytotoxicity than Fresh NPs**

To further investigate the difference in cytotoxicity between fresh and aged ZnO NPs, we examined the cell viability by using CCK-8 kits. As shown in Fig. 3, incubation A₅ cells with gradient doses of fresh and aged ZnO NPs (ranging from 0 to 20 μg/mL, 20 nm and 90–200 nm) for 24 h, 48 h or 72 h showed a dose-dependent decrease in cell viability. No obvious cytotoxicity was observed by treating cells with ZnO NPs ≤10 μg/mL. When the dosage of fresh and aged ZnO NPs elevated to 12 and 15 μg/mL, the cell viability showed a time-dependent decrease tendency. Obviously, the cell viability in aged NP-treated groups was significantly higher than fresh NP-treated groups. In addition, ZnCl₂-treatment also compromised cell viability in a dose- and time-dependent manner, whereas the cytotoxicity of ZnCl₂ was much less than that of both fresh and aged ZnO NPs.

**Aged ZnO NPs Did Not Up-Regulate the Expression of Apoptotic Genes as Fresh ZnO NPs**

Our RNA sequencing data from aged ZnO NP-treated A₅ cells showed that p53, PI3k–Akt, FoxO, Glutathione, ErbB, HIF-1, Oxytocin and Jak-STAT signaling pathway were enriched (Fig. 5A). The apoptosis genes enriched in Jurket and HMDM cells were not significantly affected in the aged ZnO NP-treated cells (Fig. 5B). To further confirm the findings, we tested the expression of related genes by real time PCR. We found that some of the apoptosis genes: BMP2, PMAIP1,
IL1α, CD69, CCNA1, CD38 and IL1β were undetectable in aged ZnO NPs-treated A₁ cells (data not shown), probably because most of these genes are expressed in immune system cells. The other up-regulated apoptosis genes (IL1α, IL1β and CD59) observed in fresh ZnO NP-treated groups were not significantly changed in expression levels by aged ZnO NPs’ treatment. While the MT1 that serve as a positive control was significantly increased in an expression level, the expression of Caspase 3 was not significantly changed (Fig. 5C). These data suggested that aged ZnO NPs, unlike their fresh counterparts, are less potent in activating apoptosis pathway genes in A₁ cells.

**Fresh But Not Aged ZnO NPs Increased the Expression Level of the Activated Caspase 3 Protein**

Detection of gene expression of *Caspase 3* alone cannot directly indicate the activation of apoptosis pathway. To further analysis whether ZnO NPs’ treatment could change apoptotic proteins’ level, the expression of cleaved Caspase 3 protein, a commonly used biomarker to indicate the activation of cell apoptosis [27], was examined by
Western blotting analysis. As shown in Fig. 6, compared to the control group, fresh ZnO NPs (20 nm) treatment increased the cellular level of cleaved Caspase 3 protein by $1.31 \pm 0.023$-fold, which was significantly higher than that of aged 20 nm ZnO NPs-treated group ($1.12 \pm 0.039$-fold). When the particle size of fresh ZnO NPs was increased to 90–200 nm, the expression of cleaved Caspase 3 protein induced by fresh NPs was increased by $1.46 \pm 0.078$-fold, significantly greater than that of aged NPs ($1.07 \pm 0.075$-fold). These data further illustrated the higher potency of fresh ZnO NPs in inducing cell apoptosis, in comparison with their aged counterparts.

**Discussion**

ZnO NPs were reported to undergo physicochemical transformation into $\text{Zn}_5(\text{CO}_3)_2(\text{OH})_6$ with the release of $\text{Zn}^{2+}$ during the natural aging process [23, 28]. However, the cytotoxicity induced by the transformed (aged) ZnO NPs and the underlying mechanisms remain unclear. Here, to unveil the mechanism of diverse cytotoxicity between fresh and aged ZnO NPs, RNA sequencing analysis and RT-PCR test were conducted. Also, Western blotting was applied to examine the protein level of Caspase 3, the key executor in cell apoptosis.

Our data showed that aged ZnO NPs induced much less cytotoxicity than fresh ZnO NPs in $A_L$ cells. The $L_{C_{100}}$ of both fresh ZnO NPs (90–200 nm and 20 nm) in our present study was lower than 15 $\mu$g/mL (Fig. 3), which was consistent with previous findings that the $L_{C_{100}}$ of ZnO NPs with 19–36 nm to NIH-3T3 or MSTO cell is about 15 $\mu$g/mL [29]. We confirmed that the environmental transformations of physicochemical properties in NPs can dramatically alter their toxicity. It has been reported that the sulfidation process of ZnO NPs changes their charge, hydrophobicity and aggregation state, resulting in the adsorption of sulfide state NPs in human saliva, sweat and bronchoalveolar lavage fluid. And the protein adsorbed by ZnO NPs formed a special
Fig. 5 Apoptosis pathway was not enriched in RNA-seq data of aged ZnO NP-treated A\textsubscript{5}L cells. (A) The gene ontology analysis of enriched pathways from aged ZnO NP-treated A\textsubscript{5}L cells. (B) The heatmap of apoptotic gene expression of aged ZnO NP-treated A\textsubscript{5}L cells and control group. (C) The expression of selected apoptotic genes and control genes (MT1) in fresh and aged ZnO NP-treated A\textsubscript{5}L cells.

Fig. 6 Apoptotic levels in A\textsubscript{5}L cells incubated with fresh and aged ZnO NPs (20 and 90–200 nm). Western Blotting analysis (A) and quantification (B) of cleaved Caspase 3 protein levels when cells were incubated with 12 μg/mL fresh and aged ZnO NPs (20 and 90–200 nm) for 72 h. Data were based on ≥ 3 independent experiments and expressed as mean ± SD, *p < 0.05.
protein crown, which affected its biological effect [19]. Phosphates widely present in physiological solutions (such as saliva) could convert ZnO NPs into metastable ZnHPO₄ and Zn₅(CO₃)₂(OH)₆ within about 5–10 h and showed cytotoxicity to digestive tract epithelial cells [20]. Ivask et al. proved the occurrence of complete transformation of ZnO NPs (< 3 μg/mL) in the in vitro exposure system of human T lymphocytes (37 °C, cell culture medium RPMI1640 containing 10% FBS for 24 h) using synchrotron radiation X-ray absorption near-edge structure spectroscopy (XANES). The spectrum and cytotoxicity of the transformation products were consistent with those of ZnSO₄ [21]. Our results revealed the dose- and time-dependent toxicity of ZnCl₂ to A₁ cells, where its cytotoxicity is much lower than both fresh and aged ZnO NPs (Fig. 3). The observation further explains the finding that the cytotoxicity of fresh ZnO NP is not fully attributed to its released Zn²⁺ [30].

Our previous study also showed that aged ZnO NPs exhibit a higher potency in eliciting ROS (reactive oxygen species), as well as an attenuated ability in killing cells compared to fresh ZnO NPs [23]. We reason that the lower cytotoxicity induced by aged ZnO NPs could be more tolerable in mammalian cells. The present study of RNA sequencing data illustrated that apoptotic genes have been up-regulated in fresh ZnO NP-treated cells, where they were much less affected in aged NP-treated groups. IL1α and IL1β are members of the interleukin 1 cytokine family. The release of IL1α and IL1β activates Caspase 8 partially dependent apoptosis [31]. CD69 encodes a member of the calcium-dependent lectin superfamily of type II transmembrane receptors. Increased CD69 expression was associated with an increased expression of the apoptosis annexin V and CD95 (Fas) marker [32]. JUN is an AP-1 transcription factor subunit. Increased JUN activity proteolytically cleavages alpha-fodrin, a substrate of the interleukin 1beta-converting enzyme (ICE), and CED-3 family of cysteine proteases, which further causes programmed cell death [33]. The increased expression of these apoptotic genes revealed that fresh NPs trigger apoptosis in several different ways. After the elevation of these apoptotic gene expressions, apoptosis processes are eventually executed by apoptotic proteins (Fig. 7). Caspase 3 is the core protease for various apoptotic scenarios; cleavage of this protein is necessary to activate both extrinsic and intrinsic apoptotic pathways [34, 35]. Therefore, detection of cleaved caspase 3 is a common method for identifying apoptosis induced by a wide variety of apoptotic signals [36]. Our Western blotting data revealed that, for both 20 nm and 90–200 nm ZnO NPs, sub-thal exposure did not alter the level of Pro caspase 3 in all treatment groups. In contrast, cleaved Caspase 3 was significantly elevated by fresh NPs treatment, where aged NPs showed few (if any) effects on the level of cleaved caspase 3 (Fig. 6). Combined with RNA expression analysis, our results clearly elucidated the higher potency of fresh ZnO NPs in inducing cell apoptosis.

Conclusions

In the present study, the natural physicochemical transformation of ZnO NPs in ultrapure water was confirmed, and variations in cytotoxicity induced by fresh & aged NPs were investigated. We focused on RNA sequencing data from our aged ZnO NP-treated A₁ cells and that of fresh NPs from database. We compared those signaling pathway specifically enriched in aged NP-treated group, which are different from that of fresh NP- or ZnCl₂–treated groups. Our data indicated that the lower cytotoxicity of aged ZnO NPs is closely related to its attenuated ability in inducing apoptosis, while the transcriptional regulation of the multiple pathways activated by NPs promotes the establishment of cellular homeostasis in mammalian cells.

Abbreviations

NPs: Nanopowders; ZnO: Zinc oxide; Zn₅(CO₃)₂(OH)₆: Hydrozincite; Zn (OH)₂: Zinc hydroxide; ZnCl₂: Zinc chloride; ZnSO₄: Zinc sulfide; Zn(HPO₄)₂: Zinc hydrogen phosphate; Zn₅(PO₄)₂: Zinc phosphate; A₁ cells: Human–hamster hybrid cells; CHO cells: Chinese hamster ovary cells; Jurket cells: Peripheral blood T lymphocyte cells; HMDM cells: Human monocyte-derived macrophages; NIH-3T3 cells: Mouse embryonic cells; MTO cells: Human lung cancer cells; RPMI1640: Roswell Park Memorial Institute 1640; ICE: Interleukin 1beta-converting enzyme; CD95 (Fas) marker: CD95, Fas ligand (FasL): Membrane receptor on the surface of many cells; JUN: Jun Proto-oncogene: Transcription factor; IL1α and IL1β: Interleukins 1α and 1β; Annexin V: Phospholipid binding protein; Caspase 8 and Caspase 3: Cysteine proteases involved in apoptosis; A₁ cells: Human–hamster hybrid cells; CHO cells: Chinese hamster ovary cells; Jurkett cells: Peripheral blood T lymphocyte cells; HMDM cells: Human monocyte-derived macrophages; NIH-3T3 cells: Mouse embryonic cells; MTO cells: Human lung cancer cells; RPMI1640: Roswell Park Memorial Institute 1640; ICE: Interleukin 1beta-converting enzyme.
1beta-converting enzyme; CED-3: Caenorhabditis elegans death gene; IL1α: Interleukin 1alpha; IL1β: Interleukin 1beta; mRNA: Messenger ribonucleic acid; cDNA: Complementary deoxyribonucleic acid; FBS: Fetal bovine serum; TEM: Transmission electron microscopy; XRD: X-ray diffraction; RT-PCR: Real-time polymerase chain reaction; CPI: The Nanotechnology Consumer Product Inventory; XANES: Synchrotron radiation X-ray absorption near-edge structure spectroscopy; RIPA: Radio immunoprecipitation assay; SDS-PAGE: Polyacrylamide gel electrophoresis; PVDF: Polyvinylidine fluoride; ECL: Enhanced chemiluminescence; CCK-8: Cell counting kit-8.

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Authors’ contributions

JW, LW, NY, XYW and MMW performed the experiments. MMW, HD, JW, LW, ML, CH, MQ, SW, TZH and NY drafted the manuscript. MMW and HD reviewed the manuscript. All authors read and approved the final manuscript.

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Availability of data and materials

Not applicable.

Declaration

Competing interests

The authors declare that they have no competing financial interest.

Author details

1 Department of Pathophysiology, School of Basic Medical Science, Anhui Medical University, No. 81, Mei-Shan Road, Hefei 230032, Anhui, People's Republic of China. 2 MOE Key Laboratory of Population Health Across Life Cycle, No. 81, Mei-Shan Road, Hefei 230032, Anhui, People's Republic of China. 3 High Magnetic Field Laboratory, Key Laboratory of High Magnetic Field and Ion Beam Biological Physics, Hefei Institutes of Physical Science, Chinese Academy of Sciences, Hefei, Anhui, People's Republic of China. 4 Department of Physiology and Biophysics, Miller School of Medicine, University of Miami, Miami, FL, USA.

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