Comparative and Mechanistic Genotoxicity Assessment of Nanomaterials via a Quantitative Toxicogenomics Approach across Multiple Species

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ABSTRACT: This study reports a comparative and mechanistic genotoxicity assessment of four engineered nanomaterials (ENMs) across three species, including E. coli, yeast, and human cells, with the aim to reveal the distinct potential genotoxicity mechanisms among the different nanomaterials and their association with physicochemical features. Both the conventional phenotypic alkaline comet test and the newly developed quantitative toxicogenomics assay, that detects and quantifies molecular level changes in the regulation of six DNA damage repair pathways, were employed. The proposed molecular endpoints derived from the toxicogenomics assays, namely TELI (Transcriptional Effect Level Index) and PELI (Protein Effect Level Index), correlated well with the phenotypic DNA damage endpoints from comet tests, suggesting that the molecular genotoxicity assay is suitable for genotoxicity detection. Temporal altered gene or protein expression profiles revealed various potential DNA damage types and relevant genotoxic mechanisms induced by the tested ENMs. nTiO₂_a induced a wide spectrum of DNA damage consistently across three species. Three carbon-based ENMs, namely carbon black, single wall carbon nanotube (SWCNT) and fullerene, exhibited distinct, species and ENM property-dependent DNA damage mechanisms. All carbon based ENMs induced relatively weak DNA damage repair response in E. coli, but more severe DNA double strand break in eukaryotes. The differences in cellular structure and defense systems among prokaryotic and eukaryotic species lead to distinct susceptibility and mechanisms for ENM uptake and, thus, varying DNA damages and repair responses. The observation suggested that eukaryotes, especially mammalian cells, are likely more susceptible to genotoxicity than prokaryotes in the ecosystem when exposed to these ENMs.

INTRODUCTION

The production and application of engineered nanomaterials (ENMs) is exponentially increasing in various fields, and it has led to the concomitant rise in public concern of the potential toxicological impact and risks associated with these nanoparticles.¹ Nanomaterial-induced toxicity has been observed in human and other lives via mechanisms related to oxidative stress,² inflammatory response,³ protein binding,⁴ ion influx disturbance,⁵ and dissolution and release of toxic ions. It is currently accepted that oxidative stress and inflammation are the major mechanisms of nanotoxicity.⁶,⁷ While most nanotoxicity assessment has been focusing on phenotypic endpoints-based cytotoxicity at relatively higher doses, chronic and lower concentration effects, such as their genotoxic potential on long-term environmental risk and human health have not been extensively investigated.⁸ Genotoxicity is caused by agents interacting with DNA and other cellular targets that control the integrity of the genetic materials, including induction of DNA adducts, strand breaks, point mutations, and structural and numerical chromosomal changes.⁹ Genotoxicity can not only initiate cancer development, but also has an impact on fertility and the health of subsequent generations, and contributes to the potential heritable changes in eco-systems after long-term accumulation.¹⁰,¹¹

Current genotoxicity assays are generally based on detection of induced damages, including Ames test, chromosome aberration test, and comet assay in vitro, and chromosome aberration, mutation of endogenous genes, and rodent carcinogenicity bioassay in vivo.¹² On the basis of these conventional tests, many ENMs have been reported to be genotoxic, including metal nanoparticles (e.g., nanosilver and gold nanoparticles), metal-oxide nanoparticles (e.g., nanotitanium dioxide), and carbon based ENMs (e.g., carbon black (CB), carbon nanotube (CNT), and fullerene).²,³,⁵,¹⁰,¹³ However, many inconsistencies have been reported among...
different nanogenotoxicity studies, which are likely related to differences in ENMs preparation, quantification, and characterization, and inherent limitations in the genotoxicity assays, such as assay-specific target and species-dependent results. Most in vitro assays only detect one or limited types of damage, which may lead to false negative results, or can be overly sensitive. The challenges in evaluating the large number of ENMs and their complex variations using resource-intensive and time-consuming in vivo assays motivate research in developing faster, more economical, and reliable genotoxicity assays.

Genetic biomarkers-based molecular assays have been demonstrated to be promising for rapid and sensitive high throughput screening (HTS) evaluation of genotoxicants. Particularly, toxicogenomics that monitor many biomarkers simultaneously have been proven to be rapid and sensitive for genotoxicity testing with mechanism identification and adverse health effects prediction. Major challenges in application of these molecular assays are the needs to employ battery and/or tiered in vitro and in vivo genotoxicity assays in order to reliably detect all genotoxic effects and, to derive quantitative molecular endpoints that can be linked to meaningful in vivo phenotypic adverse effects. Despite the degree of biological organizational level, cells respond to potential DNA damage through a common architecture with a number of rather conserved specific response and repair pathways in an effort to repair the DNA damage to restore homeostasis. On the basis of this understanding, we recently proposed and developed a quantitative toxicogenomics-based comprehensive genotoxicity assessment approach with DNA damage and repair pathway ensemble-based cell arrays for multiple species including E. coli, yeast and human cells. Furthermore, we proposed a quantitative molecular endpoints-PELI that correlated well with conventional phenotypic genotoxicity endpoints, demonstrating that the proposed quantitative toxicogenomics-based genotoxicity assay can be a promising and alternative tool for mechanistic genotoxicity screening and evaluation in vitro (data not shown).

In this study, we performed a comprehensive and comparative genotoxicity evaluation of a number of ENMs using the newly developed quantitative toxicogenomics-based genotoxicity testing approach as well as conventional genotoxic assay (alkaline comet assay), and across three different cell lines including E. coli, yeast and human cells. The tested ENMs included nTiO$_2$_a, carbon black (CB), single wall carbon nanotube (SWCNT), and fullerene (C$_{60}$). The study aimed to detect the potential genotoxic effect of these common-used ENMs for both eco-system and human health and, to reveal differences in their genotoxic mechanisms among organisms of different taxonomic order, as well as potential association with their physicochemical properties.

**MATERIALS AND METHODS**

**Nanomaterials Used in This Study.** Four engineered nanomaterials (ENMs): nTiO$_2$_a (Sigma-Aldrich), Carbon Black N110 (CB; Cabot Corp.), Short Single Wall CNTs (SWCNT; Cheap Tubes Inc.) and Purified fullerene (C$_{60}$; M.E.R. Co.) were tested in this study (detailed information in SI Table S1). Detailed physical and chemical characterization for these ENMs was previously described by Bello et al. ENMs were prepared as 20× stock in PBS with 1% BSA (Acros, NJ, U.S.A.) as a dispersant, which is the currently common approach for bioassays. The stock solutions were sonicated at ~90 W for 15 min to maintain a better dispersion before tests, and were immediately diluted in the corresponding media (M9, SD or F12) for the following test. Cytotoxicity of ENMs, mitomycin C (MMC, a model genotoxicant as a positive control) and bisphenol A (BPA, reported to be genotoxicity negative) were evaluated by growth inhibition in E. coli, yeast and by cell counting in A549 cells (human lung cancer cells) respectively, with details in SI Figure S1 and LC5 (concentration that has 95% cell survived) in SI Table S1. Since the molecular assay intends to detect subtle and cellular level response to toxicants, three subcytotoxic doses (SLCS) for ENMs, and six for MMC and BPA were applied for E. coli and yeast, one concentration was applied for human cells for PCR and comet test (SI Table S1 for details in dose concentrations). PBS with 1%BSA was used as the vehicle control. For library tests, blank wells with medium (with or without chemicals) were used as medium controls for data processing.

**Characterization Considerations.** Extensive physico-chemical and morphological characterization of ENMs was performed as previously published. In addition, aggregation sizes of ENMs in different media for various exposure times for this study were analyzed by dynamic light scattering (DLS) analyzer (Malvern Zetasizer Nano ZS90) (SI Table S1).

**Toxicogenomics-Based Genotoxicity Assay Library Construction.** Homologous genes specific in the DNA repair pathways that are conserved across species were selected for E. coli, yeast, and human cells based on the current knowledge of DNA damage and repair literatures as shown in Table 1.

**Measurement of DNA Damage and Repair Related Gene Expression in E. coli Library.** A library of 20 transcriptional fusions of GFP (Open Biosystem, Huntsville, AL) that includes promoters controlling the expression of genes involved in DNA damage and repair in E. coli K12, MGI1655 (Table 1) was employed in this study. A more detailed protocol description for gene expression analysis in the E. coli library is available in our previous study. Briefly, E. coli strains selected were grown with M9 medium in clear bottom black 384-well plates (Costar) for 4–6 h at 37 °C to reach early exponential growth (OD600 about 0.15 to 0.3). Freshly prepared ENM stock solutions or controls dissolved in PBS-1%BSA were added 10 μL per well to obtain the final concentrations (SI Table S1). Three dose concentrations were tested for each ENM (SI Table S1). Plates were then read in a Micro plate Reader (Synergy HT Multi-Mode, Biotech, Winoski, VT) continuously for absorbance (OD600 for cell growth) and GFP signal (with 485 nm excitation and 535 nm emission for gene expression) every 5 min for 2 h. All tests were performed in dark in triplicate.

**Measurement of DNA Damage and Repair Related Protein Expression in Yeast Library.** A library of 37 in-frame GFP fusion proteins involved in DNA damage and repair of *S. cerevisiae* (Invitrogen, no. 95702, ATCC 201388) (Table 1), constructed by oligonucleotide-directed homologous recombination to tag each open reading frame (ORF) with *Aequorea victoria* GFP (S6ST) in its chromosomal location at the 3’ end, was employed as described in our previous publications. Yeast strains selected were grown in clear bottom black 384-well plates (Costar) with SD medium for 4–6 h at 30 °C to reach early exponential growth (OD600 about 0.2 to 0.4). ENM stock solutions dissolved in PBS-1%BSA or control were added 10 μL per well to obtain the final
concentrations. Three dose concentrations were tested for each ENM (SI Table S1). The plates were then placed in a Micro plate Reader (Synergy H1Multi-Mode, Biotech, Winooski, VT) for absorbance (OD600 for cell growth) and GFP signal (filters with 485 nm excitation and 535 nm emission for protein expression) measurements every 5 min for 2 h after fast shake. The signal of medium control (with or without NM/chemical) was used as an internal control (housekeeping gene PGK143) for plate deviation range of our testing systems (data not shown).

**Gene and Protein Expression Profiling Data Processing and Quantitative Molecular Endpoints Derivation.** Gene/protein expression profiling data in *E. coli* and yeast libraries were processed as described previously.32,34-36 For the *E. coli* library that monitors promoter activities, all data were corrected for various controls, including blank with medium control (with and without NM/chemical) and promoterless bacterial controls (with and without NM/chemical). The alteration in gene expression for a given gene at each time point due to NM/chemical exposure relative to the vehicle control condition without any NM/chemical exposure, also referred as induction factor I, was represented by I = Pe/Pc, where, Pe = (GFP/OD) experiments as the normalized gene expression GFP level in the experiments condition with NM/chemical exposure, and Pc = (GFP/OD) vehicle in the vehicle control condition without any NM/chemical exposure. For the yeast library, OD and GFP raw data were corrected by background OD and GFP signal of medium control (with or without NM/chemical). The protein expression P for each measurement was then normalized by cell number (OD corrected) as P = (GFP corrected/OD corrected). The P level was corrected with vehicle internal control (housekeeping gene PGK143) for plate normalization.

To quantify the chemical-induced gene/protein expression level changes of a treatment, Transcriptional Efficacy Level Index (TELI)35 for *E. coli* and Protein Efficacy Level Index (PELI)36 for yeast were proposed and derived as quantitative molecular endpoints (details described in the SI, part 3). TELI and PELI quantify the cumulative alteration gene or protein expression change over the exposure period for a given gene (TELIGeno or PELIGeno), pathway (TELIPathway, PELIPathway), or for the overall DNA damage and repair pathway ensemble library (TELIGeno, PELIGeno). For a given chemical, TELIGeno or PELIGeno based dose-response patterns (SI Figure S2) were modeled using Four Parameter Logistic (4PL) nonlinear regression model. Genotoxicity positive or negative was determined based on the comparison of the maximal TELIgeno/PELIgeno value derived from the TELI or PELI-dose response curve with a threshold value, which was predetermined based on results with postive and negative genotoxic chemicals, as well as the standard deviation range of our testing systems (data not shown).

**Measurement of DNA Repair Related Gene Expression in Human Cells by RT-qPCR and Data Processing.** 2 × 10^4/well of human lung epithelial A549 cells were seeded in 6-well plates (Costar) for 24 h in F12 medium with 10% FBS. ENMs, MMC or BPA prepared in F12 medium with 1% BSA were added in PBS-washed cells (2 mL in each well). Dose concentrations were listed in SI Table S1. Cell samples were harvested at four time points after ENMS exposure (concentrations in SI Table S1) in the dark (0.5 h, 2 h, 4 h, or 24 h). DNA was extracted by RNeasy Mini Kit (QIAGEN, Valencia, CA, U.S.A.) and reverse transcribed to cDNA by iScript cDNA Synthesis Kit (Bio-Rad, Hercules, CA, U.S.A.) and reverse transcribed to cDNA by iScript cDNA Synthesis Kit (Bio-Rad, Hercules, CA, U.S.A.). Q-PCR was performed in duplicate using SYBR Green Supermix on iQ5 Multicolor Real-Time PCR detection system (Bio-Rad). PCR Primers targeting the six selected genes in Table 1 were obtained from the database at NCBI with the primer set of Primers listed in SI Table S2. Fold change that reflects relative gene expression change due to treatment compared to vehicle control, also referred to as induction factor I, was determined by comparative ΔΔCt method (2^−ΔΔCt).44 Similar to the *E. coli* library, the transcriptional effect in human cells was integrated into the Transcriptional Efficacy Level Index of genotoxicity (TELIGeno).

**DNA Damage Alkaline Comet Assay in Human A549 Cells for Phenotypic Confirmation.** The alkaline comet assay in human A549 cells upon exposure to ENMs or controls for 24 h was carried out according to the protocol of the ITRC.45 Dose concentrations were similar to those applied for molecular toxicity assay (SI Table S1). All the procedures were performed in the dark with triplicates. The slides were stained with ethidium bromide (2 μg/mL) and checked by fluorescence microscope (Olympus IX51, camera DP70). Fifty cells of each treatment were measured by software CASP randomly selected.
RESULTS

Characterization of Engineered Nanomaterials in Different Media for Genotoxicity Assays. ENMs tested in this study were generally stable along the exposure period (2 h/24 h), but had different sizes in different media (see SI Table S1). For nTiO$_2$-a and CB, larger aggregation sizes were observed in SD media for the yeast assay, while smaller sizes were observed in M9 and F12 media for E. coli and human cells, respectively. C$_{60}$ shared similar sizes in three different media. Particle size is not available for linear SWCNT.

DNA Damage and Repair Profiles Induced by Nanomaterials are Temporally Dynamic, Concentration- and Species-Dependent. The temporal altered gene/protein expression profiles provided a holistic view of the DNA damage and repair pathways and, reflected dynamic responses of E. coli, yeast, and human cells exposed to ENMs over the exposure time at different dose concentrations. Altered molecular (gene or protein) expression changes were quantified by induction factors as shown in Figure 1 for one representative dose condition and in SI Figure S3 for all other doses. The DNA-damaging chemical MMC (positive control, SI Figure S3) induced significant up-regulation of biomarkers in most of the species tested. The mean natural log of induction factor (ln I) indicates the magnitude of altered gene/protein expression (represented by a green–black–red color scale at bottom. Red spectrum colors indicate up regulation, green spectrum colors indicate down regulation. Values beyond ±2 are shown as ±2). X-axis bottom: for E. coli and yeast: testing time in minutes, the first data point shown is at 20 min after exposure due to data smoothing with moving average of every five data points; for human cells: testing time in hours. Y-axis left: clusters of genes/proteins by DNA damage repair pathways. Y-axis right for each species: list of genes or proteins (ORFs) tested, with details in Table 1. For CB exposure, corrections are made to the gene expression analysis for the physical interference of CB on the fluorescence signal reading (details in SI Figure S4).
DNA repair pathways in *E. coli*, yeast, and human cells, with time- and dose-dependent patterns. As expected, no significant up-regulation was observed in the three libraries for the negative control bisphenol A (SI Figure S3).

The DNA damage potential and mechanisms for different ENMs seemed to vary and were concentration- and species-dependent, as indicated by the distinct altered molecular expression profiles (Figure 1 and SI Figure S3). nTiO$_2$-$\alpha$ (Figure 1A) induced expression changes in biomarkers for all the three species. The three carbon based ENMs induced distinct DNA damage repair pathways activation profiles, mainly in eukaryotes, suggesting structure-dependent genotoxicity. CB (Figure 1B) seemed to exert more severe DNA damaging potential than the other two carbon nanomaterials, namely SWCNT (Figure 1C) and C$_{60}$ (Figure 1D). See more discussion in later genotoxicity mechanisms section.

**Comparison of Molecular Genotoxicity Assay Results with Other in Vitro and in Vivo Genotoxicity Assays.**

Quantitative toxicogenomics assay endpoints, namely TELI$_{geno}$ or PELI$_{geno}$ for different dose concentrations were derived for each ENM and they exhibited dose response curves for the four ENMs examined (SI Figure S2). The maximum values of TELI$_{geno}$ for *E. coli* assay and PELI$_{geno}$ for yeast assay, derived from dose response model fitting, were then compared to predetermined endpoint threshold for molecular genotoxicity assay as positive or negative (see description in methods and SI). The molecular genotoxicity assay endpoints from our assay were compared to those from conventional phenotypic genotoxicity assays, including alkaline comet assay in *E. coli* and yeast assay, (Figure 2). The phenotypic genotoxicity endpoint based on 24 h comet assay in human cells correlated well with the molecular endpoint TELI$_{geno}$ for human cell (*r*$_p$ = 0.8147, *P* = 0.0483), as well as for the endpoint PELI$_{geno}$ in yeast cell (*r*$_p$ = 0.9073, *P* = 0.0125). The higher correlation coefficient values of eukaryotes than those of prokaryotic *E. coli* cells (*r*$_p$ = 0.6102, *P* = 0.1983) suggest that eukaryotic cells shared more similar responses than prokaryotic cells. Note that yeast cell assay incorporated much more key biomarkers of DNA damage repair pathways than the human cell assay and therefore captures more comprehensive DNA damage information, which may explain the better correlation of comet assay results with yeast cells than with human cells. These results suggest that the applied toxicogenomics genotoxicity assay and proposed quantitative molecular endpoints are adequate for genotoxicity assessment across multiple species and, they can be potentially linked to phenotypic effects quantitatively.

**Genotoxic Mechanisms of ENMs Revealed by DNA Damage Repair Pathways Profiling and Their Association with ENM Properties.** To gain further insights into the detailed DNA damage repair pathway activities induced by the tested ENMs, the magnitude of expression changes in each pathway was quantified by TELI$_{pathway}$ or PELI$_{pathway}$ (see SI, part 3) and compared among ENMs at multiple concentrations (Figure 3). The model genotoxicant MMC induced severe DNA damage, as indicated by the SOS response activation in *E. coli* and strong pathway activation for double strand break and base damage repair in all three species. In addition, it seemed to cause a potential DNA cross-link, as suggested by the nucleotide excision repair biomarkers in yeast and human cells. These DNA damage responses induced by MMC were consistent with its known high alkylation reactivity to produce base alkylation and cross-link to DNA.

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repair in E. coli and, no pathway activation in yeast but oxidative damage repair (indicated by OGG1 up-regulation) in human cells, which was consistent with previous report that BPA induced oxidative stress in mammals.54

\( n\text{TiO}_2\_a \) induced wide range of pathway activation suggesting strong DNA damage potential consistently among all three species. In E. coli, \( n\text{TiO}_2\_a \) led to activation of SOS system indicating severe DNA damage. In yeast and human cells, a wide range of repair pathway activation, including severe double strand break repair, was observed and the up-regulation of OGG1 specific for oxidative damage repair \( (I_{\text{max}} = 3.4673 \text{ for yeast}, I_{\text{max}} = 1.9740 \text{ for human cells}) \). OGG1 up-regulation in both yeast and human cells suggested that oxidative damage might play an important role in DNA damage of \( n\text{TiO}_2\_a \), which was agreeable with the current understanding that oxidative damage through reactive oxygen species (ROS) production is its main toxicity mechanism.15,34 Note that in this study, the ROS production from photocatalytic reaction was not expected to be significant under our test conditions because all bioassays were performed in the dark and the wavelength of excitation used for OD and fluorescence reading (600 and 485 nm) was out of the active photoactive range for \( n\text{TiO}_2\_a \) (385 nm). Several reports were supportive of nonphoto-induced ROS generation by unknown mechanism(s).56 In addition, the dissolution of titanium ions at neutral pH within 2 h was considered to be negligible for nano TiO\(_2\),57 and the dissolved ion was reported to exhibit no damage to the cellular DNA.57

Three carbon-based ENMs showed structure-dependent and discernible DNA damage response profiles. Carbon black (CB) seemed to lead to little response in E. coli, likely due to the uptake limit in prokaryotic cells.58,59 In contrast, a wide range of DNA repair pathway activities, including nucleotide excision repair and double strand break repair, were observed in yeast and human cells, suggesting potential damage induced by CB exposure. The up-regulation of OGG1 in yeast \( (I_{\text{max}} = 3.3050) \) suggested that oxidative damage may contribute to the genotoxicity of CB.60 However, no OGG1 up-regulation was observed in human cells, suggesting that some mechanisms other than oxidative stress may contribute to the severe DNA damage of CB in human cells, which needs further investigation.

SWCNT exposure did not lead to any DNA repair pathway activation in E. coli, but seemed to induce strong double strand
DNA damage in eukaryotes. In contrast to CB, oxidative damage may not be the main mechanism since OGG1 up-regulation was not observed in either yeast or human cells. DNA double strand break in eukaryotes may be a result of membrane damage induced apoptosis/necrosis with DNA cleavage, which has been reported to be a main target for CNTs. Although membrane damage of SWCNTs has also been observed in E. coli, it may not lead to DNA damage since E. coli lacks membrane related cell death process.

For C60 exposure, specific double strand break response was observed in human cells, but multiple DNA damage pathway activities were detected in yeast. Oxidative damage may not be the main mechanism either, as OGG1 up-regulation was not observed in either yeast or human cells. Another specific repair pathway activation, NHEJ (nonhomologous end-joining) was observed in both yeast and human cells, which repairs DNA double strand break by direct ligation, suggesting some distinguishable mechanism involved in C60 induced DNA damage, such as direct cleavage on both strands. C60 has been reported as an electron-poor photosensitizer, which can induce DNA photocleavage selectively for G base by visible light. The direct cleavage may breaks with little DNA loss, which can be repaired by direct ligation of NHEJ pathway. As previously mentioned, although the bioassays were conducted in the dark, intermittent excitation during fluorescence reading at 485 nm can potentially irritate the photocleavage of C60, which has a moderate energy gap of about 1.8 eV (∼690 nm).

**DISCUSSION**

This study performed a comprehensive and comparative genotoxicity assessment for four commonly-used ENMs across multiple species using both conventional phenotypic and newly proposed quantitative toxicogenomics approaches. The results demonstrated that the toxicogenomics assay based on conserved DNA damage repair pathways across three species could detect various DNA damages induced by ENMs. And, for the first time, the molecular endpoints were quantitatively correlated to phenotypic endpoints and generally consistent with those based on conventional in vitro and in vivo genotoxicity assays, suggesting that the toxicogenomics-based assays can be promising for potential genotoxicity assessment of ENMs. Compared to current genotoxicity assay which takes several days (e.g., over two days for Ames test) or even years (two years for in vivo carcinogenesis test), our assay is faster (2 h. for E. coli and yeast, 24 h. for human cells), more sensitive (at subcytotoxic doses), more comprehensive (capture all DNA damage types instead of single or a few biomarkers or specific targets as in other assays), and more cost-effective (the gfp-fused whole cell libraries can be easily grown with less complex protocol). In addition, the molecular profiles can provide mechanistic information regarding specific DNA damage type and repair pathways.

The comparative study of the selected ENMs provided more insights for a better understanding of their genotoxic mechanisms, which was poorly understood previously. Oxidative damage has been considered the main effect of ENMs leading to the genotoxicity for nTiO2-a, CB, fullerenes, and SWCNT. In this study, oxidative damage was confirmed to play an important role in DNA damage for nTiO2-a and CB, which is consistent with previous studies. However, our results indicated that the other two carbon ENMs (SWCNT and C60) might induce DNA double strand break in eukaryotes through some other mechanisms, implying more structure-and/or property-dependent genotoxicity. Although they both lead to DNA double strand break in eukaryotes, the differences in the activation of pathway-specific genes, for example, Rad51 (for homologous recombination) and Ku70 (for nonhomologous end-joining by direct ligation) in human cells revealed different mechanisms of DSB repair. The results demonstrated the specificity and sensitivity of the gene/protein expression methods for detecting and differentiating DNA damage and related mechanisms, such as among those carbon-based yet structurally distinct ENMs.

The distinct genotoxic responses between prokaryotic and eukaryotic species revealed in this study were likely related to the different capability and mechanisms of the ENMs’ (particularly carbon based ENMs) uptake among species. For mammalian cells, nanoparticle uptake has been reported to occur through eukaryote-specific processes such as phagocytosis and endocytosis. Endocytosis in yeast also makes it possible for ENM uptake; some unclear mechanism in yeast seems to be employed to internalize nanomaterials too. However, the rigid cell wall of bacteria may avoid the direct uptake process. Moreover, bacteria lack the ability to perform endocytosis, which may make them more resistant to ENMs. Adhesion of ENMs to eukaryotic cells may also contribute to a series of membrane damage-related cell death processes, including the DNA damage response. As programed cell death is limited in bacteria, this adhesion effect may contribute little to the nanogenotoxicity of E. coli.

In conclusion, this study revealed various genotoxicity of the four commonly used ENMs by real-time gene/protein expression profiling, which seemed to be both species and ENM-property-dependent. The differences in cellular structure and defense systems among prokaryotic and eukaryotic species lead to distinct susceptibility and mechanisms for ENMs uptake and, thus varying DNA damages and repair responses. The observation suggested that eukaryotes, especially mammalian cells, are likely more susceptible to genotoxicity than prokaryotes in eco-system when exposed to these ENMs.

**ASSOCIATED CONTENT**

Supporting Information

Cytotoxicity of ENMs/chemicals in E. coli, yeast, and A549 cells; detailed physical and chemical characterization of the nanomaterials tested in this study, as well as the concentrations tested; detailed data processing and quantitative molecular endpoints derivation; the dose responses of model chemicals and ENMs; primers used for in RT q-PCR for human A549 cells; with GenBank accession number or UniSTS number; real-time gene/protein expression profiles for all concentrations tested in this study; the interference of carbon black to fluorescence signal; results of alkaline comet test in human A549 cells. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.
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