Intracellular accumulation of indium ions released from nanoparticles induces oxidative stress, proinflammatory response and DNA damage

Received August 3, 2015; accepted August 19, 2015; published online September 15, 2015

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Due to the widespread use of indium tin oxide (ITO), it is important to investigate its effect on human health. In this study, we evaluated the cellular effects of ITO nanoparticles (NPs), indium chloride (InCl₃) and tin chloride (SnCl₂) using human lung epithelial A549 cells. Transmission electron microscopy and inductively coupled plasma mass spectrometry were employed to study cellular ITO NP uptake. Interestingly, greater uptake of ITO NPs was observed, as compared with soluble salts. ITO NP species released could be divided into two types: ‘indium release ITO’ or ‘tin release ITO’. We incubated A549 cells with indium release ITO, tin release ITO, InCl₃ or SnCl₂ and investigated oxidative stress, proinflammatory response, cytotoxicity and DNA damage. We found that intracellular reactive oxygen species were increased in cells incubated with indium release ITO, but not tin release ITO, InCl₃ or SnCl₂. Messenger RNA and protein levels of the inflammatory marker, interleukin-8, also increased following exposure to indium release ITO. Furthermore, the alkaline comet assay revealed that intracellular accumulation of indium ions induced DNA damage. Our results demonstrate that the accumulation of ionic indium, but not ionic tin, from ITO NPs in the intracellular matrix has extensive cellular effects.

Keywords: DNA damage/indium tin oxide/nanoparticles/oxidative stress/proinflammatory response.

Abbreviations: cDNA, complementary DNA; DCFH-DA, 2',7'-dichlorodihydrofluorescin diacetate; DLS, dynamic light scattering; DMEM, Dulbecco’s modified Eagle’s medium; FBS, fetal bovine serum; FE-SEM, field emission scanning electron microscopy; HMOX-1, heme oxygenase 1; ICP-AES, inductively coupled plasma atomic emission spectrometry; ICP-MS, ICP-mass spectrometry; IL-8, interleukin-8; ITO, indium tin oxide; LDH, lactate dehydrogenase; mRNA, messenger RNA; MTIIA, metallothionein IIA; NP, nanoparticle; PCR, polymerase chain reaction; ROS, reactive oxygen species; TEM, transmission electron microscopy.

A nanoparticle (NP) is a particle with a diameter between 1 and 100 nm (ISO/TS 27678:2008). NPs have unique physicochemical properties, including high catalytic activity and distinct absorption spectra, as compared with micro-sized particles (1). These unique properties are beneficial for industrial use, and many NPs are used in consumer products, such as sunscreen, cosmetics and food additives. Additionally, several studies have focused on clinical applications of NPs, including nanomedicine and drug-delivery systems (2, 3). However, despite these potential advantages, NPs also induce several biological effects. For example, zinc oxide (ZnO) NPs are used in industrial materials, such as electronic components and sunscreen. However, several in vitro and in vivo studies have revealed severe cytotoxic effects following ZnO NP administration (4–7). Therefore, precise, case-by-case evaluation of NP-mediated cellular effects is required to avoid both the negative health effects and unnecessary prohibition of NPs.

The chemical composition of metal and metal oxide NPs is an important factor when considering cellular effects. For example, ZnO, copper oxide (CuO), nickel oxide (NiO) and antimony oxide NPs exert strong cytotoxicity, whereas titanium dioxide, cerium oxide and aluminum oxide NPs have weak cellular effects (8). Additionally, the metal ions released from NPs are an important parameter in the evaluation of the cellular effects of NPs (9). For example, ZnO, NiO and chromium oxide NPs solubilized in cell culture medium release the corresponding metal ions into the extracellular environment, exerting remarkable cytotoxicity (7, 10, 11). Internalized metal and metal oxide NPs continuously release metal ions, leading to metabolic dysfunction. For example, the cytotoxic effects of silver (Ag) and CuO NPs are mediated by cellular uptake and intracellular release of Ag and copper ions (12, 13). Therefore, to understand the mechanisms underlying NP-mediated cellular effects, the material properties, including cellular uptake and intracellular behaviour, must be considered.

In this study, we evaluated the cellular effects of indium tin oxide (ITO) NPs. ITO is a sintered material consisting of indium oxide (In₂O₃; typically 90%) and tin oxide (SnO₂; typically 10%). This material is widely used as a transparent conducting electrode in various
photoelectric devices, including liquid crystal displays and solar panels (14, 15), and is also used in protein microarrays and gas sensing devices (16, 17). The preparation and property control of nano-sized ITO particles is of interest because of their potential beneficial applications in versatile devices (18, 19). Due to the wide-spread applications of ITO, it is important to investigate their possible effects on human health. The genotoxic effects of indium and tin-containing compounds were previously reported (20, 21). Recently, Badding et al. (22) demonstrated that ITO induced robust cytokine production in both mouse macrophages and human bronchial epithelial cells. Additionally, occupational exposure in an ITO worker resulted in severe interstitial pneumonia and death due to bilateral pneumothorax in Japan (23). Furthermore, epidemiological studies and studies of fatal cases in workers exposed to ITO show that ITO induces occupational lung disease and is associated with a risk of interstitial lung damage (24–27). However, it remains unclear whether inhalation of ITO is a potential occupational health hazard in employees handling these materials.

The aim of this study was to investigate ITO NP-mediated cellular effects using human lung epithelial A549 cells. A549 cells are well-characterized in vitro model for investigating molecular and biochemical processes in airway epithelium, and are widely regarded as a valid model system for NP-induced pulmonary effects. Recently, we demonstrated that the ITO NPs possess the potential for inducing oxidative stress and DNA damage in the A549 cells (28); however, possible differences in cellular effects between ITO NPs and their corresponding metal ionic species have not been assessed yet. In this study, we focused on the importance of ITO NP cellular uptake and the subsequent release of metal ions inside the cell. Furthermore, we compared the cellular effects of ITO NPs and their corresponding metal ionic species, and examined the mechanisms underlying these effects.

**Experimental Procedure**

**Preparation of stable ITO dispersions**

ITO NPs were purchased from C. I. Kasei Co., Ltd (Tokyo, Japan) and Tomoe Works Co., Ltd (Hyogo, Japan). Physical characteristics of the ITO NPs are reported in Table I. The surface area of ITO NPs which were degassed at 120°C for 30 min was determined using a Quantachrom Autosorb-I instrument using nitrogen adsorption and the five-point Brunauer–Emmett–Teller method. Stable ITO dispersions were prepared as described previously (28). In brief, ITO NPs were dispersed in 1% bovine serum albumin at a concentration of 80 mg/ml by sonication. Subsequently, the dispersions were centrifuged at 4,000 × g for 20 min. Precipitated ITO NPs were re-dispersed in an equivalent volume of fresh culture medium. The dispersions were centrifuged at 1,000 × g for 20 min. For Sample A, the resultant 1,000 × g fraction was used for cellular response examination. Because the 1,000 × g fraction of Sample B did not contain sufficient ITO NPs, the centrifugal force was reduced from 1,000 × g to 500 × g.

**Characterization of stable ITO dispersions**

Observation of the ITO NP microstructure in stable ITO dispersions was carried out using field emission scanning electron microscopy (FE-SEM, JSM-6700F; JOEL Ltd, Tokyo, Japan). The secondary particle size of the ITO NPs in the stable ITO dispersions was determined using a Dynamic Light Scattering (DLS) Particle Size Analyzer (LB-550; HORIBA Ltd, Kyoto, Japan) at 25±0.1°C. The light source was a 650-nm laser diode of 5 mW. Approximately 20-ml aqueous ITO dispersion was measured directly without any pre-treatment.

The concentrations of In2O3 and SnO2 included in the stable ITO dispersions were measured by inductively coupled plasma atomic emission spectrometry (ICP-AES; Thermo Jarrel Ash Corp., Franklin, MA) and ICP-mass spectrometry (ICP-MS; X-Serie II; ThermoFischer Scientific, Inc., Waltham, MA). To determine the concentration of medium components, stable ITO dispersions were collected through ultrafiltration membranes (Amicon Ultra Centrifugal Filters Ultracle 30K; Merck Millipore, Billerica, MA) and centrifuged at 7,000 × g for 20 min. The concentrations of In, Sn, Na, Ca and P were determined by ICP-AES (SPS7800; Seiko Instruments, Inc., Tokyo, Japan) and ICP-MS.

**Cell culture and treatment**

Human lung epithelial A549 cells (RCB0098) were purchased from RIKEN BioResource Center (Ibaraki, Japan). Cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Gibco Life Technologies, Gaithersburg, MD) supplemented with 10% heat-inactivated fetal bovine serum (FBS; USDA Tested FBS; ThermoFischer Scientific, Inc.), 100 U/ml penicillin, 100 μg/ml streptomycin and 250 ng/ml amphotericin B (Nacalai Tesque, Inc., Kyoto, Japan). This DMEM preparation is referred to as ‘DMEM-FBS’. Cells in DMEM-FBS were placed in 80 cm² flasks (ThermoFischer Scientific, Inc.) and cultured at 37°C in an atmosphere containing 5% CO2. For experiments, cells were seeded in different size multi-well plates (ThermoFischer Scientific, Inc.) at a density of 1 × 10⁴ cells/ml and incubated for 24 h. Subsequently, culture medium was replaced by DMEM-FBS supplemented with the stable ITO dispersion, InCl3 or SnCl2 solution (Wako Pure Chemical Ind., Osaka, Japan) and incubated at 37°C in an atmosphere of 5% CO2.

**Transmission electron microscopy observation**

A549 cells exposed to stable ITO dispersions for 24 h were fixed using 1.2% glutaraldehyde (TAAB Laboratories Equipment Ltd, Aldermaston, UK) for 1 h. The fixed samples were treated with OsO4 solution for 1 h, dehydrated in ethanol and embedded in epoxy resin. The resultant samples were cut into ultrathin sections suitable for transmission electron microscopy (TEM) observation by diamond-knife ultramicrotomy. TEM observation was then carried out using H-7600 (Hitachi Corp., Tokyo, Japan). The acceleration voltage was 80 kV.

**Table I. ITO sample characteristics**

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<tr>
<th>Names in this study</th>
<th>Primary particle size (nm)</th>
<th>Specific surface area (m²/g)</th>
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<tr>
<td>Sample A</td>
<td>26</td>
<td>35</td>
</tr>
<tr>
<td>Sample B</td>
<td>25.1</td>
<td>n/a</td>
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*According to the manufacturer’s data sheet. Other data were measured in this study. n/a = not available.*
**Cellular uptake of ITO NPs**
The delivery of ITO NPs to the cell surface and intracellular space was measured by ICP-MS. A549 cells were exposed to stable ITO dispersions for 24 h, washed with phosphate buffered saline (PBS) three times to remove loosely bound ITO NPs on the cell surface and detached by trypsinization (0.25%). The cell pellet was suspended in 1 ml PBS, and the number of cells was counted using a hemocytometer. The cells were mixed with acidic solution (water:nitric acid:hydrochloric acid = 4:1:3) and then heated at 80°C for 2 h to dissolve the cellular content. The dissolved solutions were diluted with water and used for ICP-MS measurement.

**Isolation of total RNA and quantitative real-time polymerase chain reaction**
Cells were cultured in 6-well plates and incubated in the stable ITO dispersions, InCl₃ or SnCl₂ solutions for 24 h. Subsequently, total RNA was extracted using the RNeasy mini kit (Qiagen, Valencia, CA) according to the manufacturer’s instructions. The total RNA concentrations were determined using a Nanodrop ND-1000 spectrophotometer (ThermoFisher Scientific, Inc.). First strand complementary DNA (cDNA) was synthesized from 0.5 μg total RNA using the SuperScript VILO cDNA synthesis kit (Invitrogen, Carlsbad, CA) according to the manufacturer’s instructions. The messenger RNA (mRNA) levels of metallothionein IIA (MTIIA), heme oxygenase 1 (HMOX-1) and interleukin-8 (IL-8) was analysed using the TaqMan gene expression assay (ID: MTIIA, Hs02379661_g1; HO-1, Hs01110250_m1; IL-8, Hs00174103_m1; Applied Biosystems, Foster City, CA). The housekeeping gene β-actin (ID: Hs99999903_m1) was used as an internal control. Target mRNA levels were measured using an Applied Biosystems 7300 real-time polymerase chain reaction (PCR) system. The cycling conditions were as follows: 2 min at 50°C, 10 min at 95°C and 60 cycles each of 15 s at 95°C and 1 min at 60°C. The mRNA expression levels of MTIIA, HMOX-1 and IL-8 in each sample were normalized to β-actin, and were then compared with untreated controls. The results are reported as fold change over control.

**IL-8 enzyme-linked immunossorbent assay**
Cells were cultured in 6-well plates and incubated in the stable ITO dispersions, InCl₃ or SnCl₂ solutions for 24 h. Subsequently, IL-8 protein levels in the culture supernatants were measured using an enzyme-linked immunossorbent assay kit (eBioscience, San Diego, CA) following the manufacturer’s instruction.

**Measurement of intracellular reactive oxygen species levels**
Intracellular reactive oxygen species (ROS) were detected using 2',7'-dichlorodihydrofluorescein diacetate (DCFH-DA) (Sigma-Aldrich, St. Louis, MO). After incubating the cells with stable ITO dispersions, InCl₃ or SnCl₂ solution, the medium was changed to serum-free DMEM containing 10 μM DCFH-DA. Cells were incubated for 30 min at 37°C, washed once with PBS, collected by trypsinization (0.25%), washed again with PBS and resuspended in 500 μl PBS. Cell samples in PBS were excited with a 488-nm argon ion laser in a FACScalibur flow cytometer (Becton, Dickinson and Company, Franklin Lakes, NJ). The DCF emission was recorded at 525 nm. Data were collected from at least 5,000 gated events.

**Water-soluble tetrazolium salts-1 (WST-1) assay**
Cell viability was examined using WST-1 solution (Premix WST-1 Cell Proliferation Assay System; Takara Bio, Shiga, Japan). Cells were cultured in 96-well plates and incubated in the stable ITO dispersions, InCl₃ or SnCl₂ solution for 24 h. To determine cell viability, the cells were incubated with 10-fold diluted WST-1 solution at 37°C for 1 h. The optical density of formazan was measured at 450 nm using Tcena Infinite M200 (Tecan Group Ltd, Männedorf, Switzerland).

**Lactate dehydrogenase assay**
Cells were cultured in 12-well plates and incubated in the stable ITO dispersions, InCl₃ or SnCl₂ solutions for 24 h. Lactate dehydrogenase (LDH) release was measured using the Cytotoxicity Detection Kit (LDH) (Roche Diagnostics GmbH, Mannheim, Germany) according to the manufacturer’s protocol. The cytotoxicity was calculated as follows: cytotoxicity (%) = (experimental value – low control)/(high control – low control) × 100. The low control, which refers to spontaneous LDH release, was the LDH released from untreated normal cells. The high control, which refers to the maximum LDH release, was the LDH released from cells that were lysed using the lysis solution provided in the kit.

**Alkaline comet assay**
An alkaline comet assay was performed according to the instructions provided by the manufacturer ( Comet Assay; Trevigen, Gaithersburg, MD). Cells (2 × 10⁵ cells/ml) were seeded in 6-well plates. After overnight incubation, the cells were exposed to the stable ITO dispersions, InCl₃ or SnCl₂ solutions for 24 h. Cells were then washed and resuspended in Ca²⁺- and Mg²⁺-free PBS solution. The cell suspension (1 × 10⁶ cells/ml) was mixed with L-Magroose at a ratio of 1:10 (v/v). This mixture was immediately transferred onto the CometSlide (Trevigen). After cell lysis at 4°C, CometSlides were treated with an alkaline unwinding solution (0.2 M NaOH and 1 mM ethylenediaminetetraacetic acid, pH > 13) for 60 min. The CometSlides were placed on a horizontal electrophoresis unit and the unit was filled with fresh buffer alkaline unwinding solution until the CometSlides were covered. Electrophoresis was performed at 20 V for 35 min at 4°C in the dark, and then staining was performed with an Ag-staining kit (Trevigen). The comet tail length was defined as the distance between the leading edge of the nucleus and the end of the tail. At least 50 determinations were performed for each sample.

**Clonogenic assay**
Colony-forming ability was detected using a clonogenic assay based on methods described by Herzog et al. (29) and Franken et al. (30). Briefly, the cells were seeded in 6-well plates (300 cells/well) and allowed to attach for 20 h. Cells were then treated with the stable ITO dispersions, InCl₃ or SnCl₂ solutions and cultured until control cells formed colonies (one colony was defined as ≤50 cells, 7 days). Subsequently, the dispersions were removed and the cells were washed with 2 ml PBS. After fixation with 100% methanol for 20 min, the cells were stained with Giemsa’s staining solution (Kanto Chemical Co., Inc., Tokyo, Japan) for 20 min and rinsed with distilled water. The number of colonies was counted.

**Statistical analysis**
Data present the mean ± standard deviation from at least three independent experiments. Statistical analyses were performed by the analysis of variance using the Dunnett test for multiple comparisons.

### Results and Discussion

**Preparation and characterization of stable ITO dispersions**
To investigate the cellular effect of NPs, the preparation of a stable NP-medium dispersion was necessary (31). When ITO NPs were dispersed in DMEM-FBS, ITO NPs formed large secondary particles and sedimented within a few minutes by gravity. Therefore, we prepared stable ITO dispersions by pre-adsorption and centrifugal fractionation methods as reported previously (28). The results of the ITO dispersion characterization are summarized in Table II. The concentrations of In₂O₃ in the ITO dispersions of Sample A and Sample B were 720 and 200 μg/ml, respectively. The concentrations of SnO₂ in Sample A and Sample B were 70 and 15 μg/ml, respectively. FE-SEM observations revealed that the ITO NPs in the prepared dispersions were globular particles (Fig. 1A). The primary particle sizes in Sample A and Sample B were within the ranges of 8–50 and 7–20 nm, respectively (Table I). Sample A contained non-uniform ITO NPs and had a broad distribution of primary particle sizes.
contained uniform ITO NPs and had a narrow particle size distribution.

Previous studies showed that dispersion stability is important to evaluate the cellular effects of NPs (32, 33). The stability of the ITO dispersions was evaluated by DLS measurement. After 3 days, the relative light scattering intensities of Sample A and Sample B were decreased to 88% and 92%, respectively, indicating that gravity sedimentation was not remarkable. The secondary particle size of ITO NPs in the dispersion was also measured by DLS (Fig. 1B). The secondary particle size histogram of Sample A revealed secondary particle sizes between 50 and 100 nm after preparation (at 0 day), which gradually increased over 3 days. However, the secondary particle size of Sample B did not change during the experimental period. These results indicate that, although slight aggregation was observed in Sample A, both ITO dispersions prepared in this study were stable for 3 days and suitable for cellular experiments.

**Cellular uptake of ITO NPs by A549 cells**

Cellular uptake is an important process that governs the effects of NPs. To investigate the uptake and intracellular localization of ITO NPs, we performed ultrastructure analysis with TEM (Fig. 2A). TEM images clearly showed the incorporation of ITO NPs into A549 cells after 24-h incubation. The ITO NPs were observed in cellular compartments, primarily in lysosome-like structures. Other cellular organelles, such as

<table>
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<tr>
<th>Names in this study</th>
<th>Metal oxide concentration (µg/ml)</th>
<th>Metal concentration in the filtrate (µM)</th>
<th>Salt composition of the dispersion</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>In2O3</td>
<td>SnO2</td>
<td>In</td>
</tr>
<tr>
<td>Sample A</td>
<td>720</td>
<td>70</td>
<td>0.03</td>
</tr>
<tr>
<td>Sample B</td>
<td>200</td>
<td>15</td>
<td>0.25</td>
</tr>
<tr>
<td>DMEM-FBS</td>
<td>0.02</td>
<td>0.00</td>
<td>0</td>
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**Fig. 1 Size distributions of ITO NPs in stable ITO dispersions.** (A) FE-SEM analysis of ITO NPs included in stable ITO dispersions. (B) Histogram of secondary ITO NP sizes based on particle number measured by DLS. The stable ITO dispersions were prepared on day 0.
nuclei and mitochondria, did not contain ITO NPs. Previous reports indicate that micro-scale ITO particles were not readily taken up by lung epithelial cells (34). This difference in results may be due to differences in ITO particle size. Size-dependent cellular uptake has been reported for gold and silica NPs (35, 36). Therefore, our results demonstrate that, similar to other NPs, ITO NPs can be readily taken up by human lung epithelial cells due to their small size.

The amount of indium and tin in the ITO NP-, InCl₃-, or SnCl₂-exposed A549 cells was quantified by ICP-MS (Fig. 2B). After exposing A549 cells for 24 h, the entire cell population was detached and analysed after lysis to determine the internalized indium and tin concentrations. A clear increase in the intracellular indium and tin concentration was observed in both Sample A and Sample B. The indium and tin concentrations in cells exposed to Sample B were significantly

![Fig. 2 Cellular uptake of ITO NPs by A549 cells. (A) TEM observation of A549 cells exposed to stable ITO dispersions for 24 h. Overview of the cell (left) and detail of the part in the white frame (right). The exposed concentrations of In₂O₃ in Sample A and Sample B were 720 and 200 µg/ml, respectively. (B) Internal concentrations of indium and tin obtained by ICP-MS measurement. The A549 cells were exposed to stable ITO dispersion, InCl₃ or SnCl₂ solutions for 24 h, and then intracellular concentrations of indium and tin were measured by ICP-MS. Inset figures depict the data for the low intracellular concentration of indium and tin (0–1.0 pg/cell).]
higher than that in cells exposed to Sample A. In contrast, InCl₃ and SnCl₂ were hardly incorporated into the A549 cells, as compared with ITO NPs. Our findings are consistent with the earlier reports on the increased cellular uptake of manganese oxide (Mn₃O₄), Ag and CuO NPs, as compared with soluble salts (MnSO₄, AgNO₃ and CuCl₂) (12, 37). These results indicated that ITO NPs were readily incorporated into A549 cells, whereas their corresponding metal ions were not easily transported into cells.

**Solubilization of ITO NPs within A549 cells**

The solubilization of micro-scale ITO particles mediated by macrophages and epithelial cells was determined by measuring the extracellular release of indium ions (38). To determine whether the ITO NPs incorporated into A549 cells were solubilized, we measured extracellular release of ionic indium and tin at 2, 4, 6 and 24 h using ICP-MS (Fig. 3). As shown in Fig. 3, the concentration of total ionic indium in the medium from A549 cells treated with Sample B was higher than cells treated with Sample A (Fig. 3A). Conversely, the concentration of total ionic tin in the medium from A549 cells treated with Sample B was lower than cells treated with Sample A (Fig. 3B). In general, ITO is insoluble at physiological pH (25). Indeed, our results show that the solubilization of ITO NPs was not observed in the absence of A549 cells. Recently, Sabella et al. (39) demonstrated that a wide variety of NPs incorporated into cells were degraded, and released corresponding ions in the acidic environment of lysosomes. Additionally, Gwinn et al. (38) reported that micro-scale ITO particles taken up by macrophages were solubilized via phagolysosomal acidification. Our results show that intracellular ITO NPs localized in lysosome-like structures were dissolved, which resulted in the release of indium or tin ions by lysosomal acidification. Furthermore, we found that when the ITO NPs were solubilized within the A549 cells, the metal ions released from Sample A were primarily tin ions (hereafter referred to as ‘tin release ITO’), whereas those released from Sample B were indium ions (hereafter referred to as ‘indium release ITO’).

**Metallothionein IIA mRNA expression**

Metallothioneins are low molecular weight, cysteine-rich metal-binding proteins found in a variety of...
**Fig. 5** Intracellular ROS levels in A549 cells exposed to stable ITO dispersion, InCl₃ or SnCl₂ solutions. Cells were exposed to stable ITO dispersions (A, B), InCl₃ (C) or SnCl₂ (D) for 6 or 24 h. Intracellular ROS levels were measured by the DCFH method using flow cytometry. The standardized DCF fluorescence value for untreated controls was 1. **P < 0.01 (versus control, Dunnett test, analysis of variance).**

**Fig. 6** The mRNA transcript levels of HMOX-1 in A549 cells exposed to stable ITO dispersion, InCl₃ or SnCl₂ solutions. The expression level of HMOX-1 was measured using real-time PCR. Each transcript level was normalized to the corresponding β-actin value, and presented as relative units compared with untreated control. **P < 0.01 (versus control, Dunnett test, analysis of variance).**
organisms, including bacteria, plants, invertebrates and vertebrates (40/42). Metallothioneins play a role in metal homeostasis and detoxification by controlling intracellular-free metal levels (40, 43). The MTIIA gene, a widely distributed isoform, is expressed under basal condition and is induced by heavy metals (44). We analysed the expression of MTIIA mRNA in A549 cells exposed to stable ITO dispersions, InCl3 or SnCl2 solutions for 24 h (Fig. 4). The mRNA levels of MTIIA dose dependently increased after incubation with indium release ITO. The expression of MTIIA was also induced after incubation with tin release ITO, InCl3 or SnCl2; however, the induction levels were not remarkable. These results indicate that the expression of MTIIA is induced by intracellular accumulation of indium ions.

**Effects of stable ITO dispersions on oxidative stress**

Metallothioneins both regulate metal detoxification and protect against oxidative stress (45, 46). Thus, we examined the effect of ITO NPs on oxidative stress. The stable ITO dispersion, InCl3 or SnCl2 solutions were exposed to A549 cells for 6 or 24 h, and intracellular ROS levels were measured using the fluorescence dye, DCFH-DA, which shows enhanced fluorescence during oxidative stress (Fig. 5). After exposure to indium release ITO for 24 h, we observed a significant increase in intracellular ROS production (Fig. 5B). In contrast, we did not detect a significant increase in intracellular ROS when cells were exposed to tin release ITO, InCl3 or SnCl2 solutions for 24 h (Fig. 5A, C and D).

We also measured the mRNA levels of HMOX-1, a marker of oxidative stress, to determine the response of A549 cells exposed to the stable ITO dispersions, InCl3 or SnCl2 solutions for 24 h (Fig. 6). The mRNA levels of HMOX-1 were dose dependently increased after incubation with indium release ITO, consistent with ROS production. However, HMOX-1 expression was not increased in cells exposed to tin release ITO, InCl3 or SnCl2 solutions. These results indicate that oxidative stress was induced by the accumulation of intracellular indium ions, but not by intracellular tin ion or extracellular indium or tin ions.
**IL-8 mRNA and protein expression**

An increase in intracellular ROS generation can result in pro-inflammatory and cytotoxic effects (47, 48). Lung epithelial cells are important source of IL-8, and IL-8 serves as a chemoattractant for neutrophils to the site of inflammation (49, 50). Moreover, IL-8 is shown to be induced in A549 cells via a mechanism involving oxidative stress and the redox-sensitive transcription factor nuclear factor kappa-B (51, 52).

To investigate inflammatory responses in A549 cells, we examined the expression of IL-8 mRNA in A549 cells exposed to indium release ITO dispersions, InCl3 or SnCl2 solutions for 24 h (Fig. 7A).

IL-8 mRNA levels were significantly and dose-dependently induced in indium release ITO-exposed cells, whereas no effect was observed in tin release ITO-, InCl3- or SnCl2-treated cells. We also quantified IL-8 protein levels released into the cell culture medium at 24 h (Fig. 7B). Consistent with mRNA expression data, indium release ITO exposure increased IL-8 protein release from A549 cells 24 h after exposure. Taken together, these results indicate that indium release ITO exposure upregulated IL-8 expression at both the mRNA and protein level in A549 cells.

**Effect of stable ITO dispersions on cell viability and membrane integrity**

In a previous report, micro-scale ITO particles were not cytotoxic to lung-derived epithelial cells even at a high concentration level of 400 µg/ml (38). Then, the effect of ITO NPs, InCl3 and SnCl2 on cell viability was examined by WST-1 assay after 6- or 24-h treatment (Fig. 8). Cell viability decreased with increasing concentrations of indium release ITO (Fig. 8B). In contrast, no inhibition was observed using tin release ITO, InCl3 or SnCl2 (Fig. 8A, C and D). Furthermore, we measured the cell membrane integrity after incubation with stable ITO dispersions, InCl3 or SnCl2 solutions for 6 or 24 h (Fig. 9). No significant damage was observed after tin release ITO or SnCl2 treatment (Fig. 9A and D). Indium release ITO did not have a remarkable effect on cell membranes, although membrane damage was observed after 24 h at the highest dose (Fig. 9B). We also found that exposure to high InCl3 concentrations induced membrane damage in A549 cells (Fig. 9C). These results indicate that cell membrane damage is induced by extracellular indium ions, but not intracellular accumulation of indium ions.
Fig. 9 Effect of stable ITO dispersions, InCl₃ or SnCl₂ solutions on the cell membrane. The cells were exposed to stable ITO dispersions (A and B), InCl₃ (C) or SnCl₂ (D) solutions for 6 or 24 h. Cell membrane damage was determined by measuring intracellular LDH release. The method of calculating cytotoxicity is described in the Experimental procedure section. **P < 0.01 (versus control, Dunnett test, analysis of variance).

Fig. 10 Effect of stable ITO dispersions, InCl₃ or SnCl₂ solutions on DNA integrity. A549 cells were exposed to stable ITO dispersions, InCl₃ or SnCl₂ solutions for 24 h. The DNA tail length values were obtained by analysing at least 50 random comet images from each treatment. The upper pictures are comet images of untreated (control), ITO NPs, InCl₃ and SnCl₂-exposed cells. *P < 0.05, **P < 0.01 (versus control, Dunnett test, analysis of variance).
**DNA damage and colony-forming ability in A549 cells**

Several studies indicated that oxidative stress induces a wide variety of physiological and cellular effects, including DNA damage (33). Thus, we examined DNA single-strand breaks using alkaline comet assays. A549 cells were incubated with the stable ITO dispersions, InCl\(_3\) or SnCl\(_2\) solutions for 24 h. Representative DNA damage images are shown in Fig. 10. The ITO NPs induced DNA strand breakage over 24 h; however, only a slight difference was observed between the indium release ITO and tin release ITO. The comet tail lengths of the indium release ITO-exposed cells were longer than that of the tin release ITO-exposed cells. We also found that exposure to InCl\(_3\) and SnCl\(_2\) induced DNA damage, with SnCl\(_2\) having a greater effect. However, the DNA damage results were not correlated with intracellular ROS levels. Previous reports indicated that DNA repair enzyme activity was inhibited by exposure to indium (54). Additionally, it was suggested that tin ions have DNA damage potential via direct interactions with DNA (55). Therefore, DNA damage induced by ITO NPs is mediated by increased oxidative stress, inhibition of DNA repair enzymes and direct interaction of indium and/or tin ions with DNA.

The influence of ITO NPs, InCl\(_3\) and SnCl\(_2\) on colony-forming ability was examined using a clonogenic assay (Fig. 11). The clonogenic assay revealed that indium release ITO dose dependently inhibited colony formation. Incubation with tin release ITO induced a statistically significant, but not a dose-dependent, inhibition of colony formation. No inhibition was observed using InCl\(_3\) and SnCl\(_2\), although SnCl\(_2\) slightly inhibited colony formation at the highest concentration. These results suggest that the accumulation of intracellular indium ions induced the loss of DNA integrity and loss of colony-forming ability.

The present *in vitro* data suggest that cellular effects induced by the exposure of indium release ITO might be caused by cellular uptake of a higher number of NPs and subsequent release of indium ions. Evaluating the cellular responses, other cell types including type I and II alveolar epithelial cells and alveolar macrophages would further contribute to our understanding of the mechanisms of ITO-related lung diseases. Furthermore, future studies would be important to establish if the present studies obtained from the A549 cells also apply to the *in vivo* environment.

**Conclusion**

In this study, we investigated the effect of two types of ITO NPs, indium release ITO and tin release ITO, on human lung epithelial cells. We found that indium release ITO, but not tin release ITO, has significant cellular effects. The NPs enable high uptake and intracellular solubilization leading to high levels of intracellular indium species following cell exposure to indium release ITO. In contrast, a high uptake but few cellular effects were observed using tin release ITO, most likely due to low intracellular release of indium ions. Thus, controlling metal ion release from ITO NPs enables us to minimize and reduce NP-associated health risks, and avoid unnecessary prohibition of beneficial NPs. To our knowledge, this is the first report that describes the relationship between cellular effects and intracellular ITO NP solubility. The cellular effects induced by exposures to ITO NPs may provide insights into indium-mediated lung disease.

**Acknowledgements**

The authors are thankful to Ms. Emiko Kobayashi, Bioproduction Research Institute, National Institute of Advanced Industrial Science and Technology for carrying out the transmission electron microscopy.

**Conflict of Interest**

None declared.

**References**


