Cancer Therapeutic Effects of Titanium Dioxide Nanoparticles Are Associated with Oxidative Stress and Cytokine Induction

Rina Fujiwara\textsuperscript{a} Yi Luo\textsuperscript{a} Takamitsu Sasaki\textsuperscript{b} Kiyomu Fujii\textsuperscript{a} Hitoshi Ohmori\textsuperscript{a} Hiroki Kuniyasu\textsuperscript{a}

\textsuperscript{a}Department of Molecular Pathology, Nara Medical University, Kashihara, and \textsuperscript{b}Department of Gastroenterological Surgery, Fukuoka University School of Medicine, Fukuoka, Japan

Abstract
Nanoparticles (NPs) are considered to influence the inflammatory process; however, the precise mechanism and the significance in tumors are still not clear. In this study, when CT26 and LL2 mouse cancer cells were treated with 6-nm anatase titanium dioxide NPs (TDNPs) without ultraviolet irradiation, oxidative stress and induction of inflammatory cytokines were observed. Oxidative stress was further increased by disease-associated conditions such as high glucose concentrations and hypoxia. Inhaled or orally administered TDNPs generated granulomatous lesions in the lungs and colon of the rodent models tested, with increased oxidative stress and inflammatory cytokines. Oxidative stress and inflammatory cytokines were also found in cancer cells treated with gold or carbon black NPs. Treatment of CT26 cells with 10- to 70-nm rutile TDNPs showed that smaller NPs produced more oxidative stress and inflammatory cytokines than larger ones did. To avoid diffusion of TDNPs and to minimize toxicity, 10-nm TDNPs were suspended in a collagen gel inserted into a subcutaneous tumor in a CT26 mouse. A single TDNP treatment via this method inhibited tumor growth in a size- and dose-dependent manner, and resulted in lower levels of urinary 8-OHdG when compared to systemically administered TDNPs. These findings suggest that TDNPs might be useful for the local treatment of tumors.

Introduction
The use of nanotechnology has accelerated in recent years. This technology is associated with advancements in engineering as well as industrial and economic factors [1]. Nanoparticles (NPs) are used ubiquitously in food, medicine, military, construction, cosmetics and sports [2]. Titanium dioxide NPs (TDNPs), in particular, are considered vital in the development of antimicrobial applications [3, 4]. However, there are increasing concerns that nanotechnology may threaten our health [1]. Toxicity has become the focus of extensive research. NPs evoke a natural immune response that is mediated by neutrophils and macrophages [5]. Inhalation of NPs induces persistent respiratory inflammation and plays an important role in respiratory disorders such as chronic obstructive pulmonary disease [5, 6]. These immunolog-
tical reactions are thought to be the basis for NP-induced tissue damage.

Anatase TDNPs are common NPs being used for many purposes such as antibacterial effects, sunscreens and cosmetics, making use of their photodynamic properties [4]. The toxic potential of TDNPs is unclear in the presence and absence of ultraviolet (UV) irradiation. We have already examined the effect of UV irradiation-evoked tissue damage in the presence of anatase TDNPs [7]; in those experiments, anatase TDNPs were found to be cytotoxic even in the absence of UV irradiation, suggesting a mechanism based on photodynamic-independent properties.

In this study, we examined the toxic potential and mechanisms of anatase TDNPs. We focused on the physical nature of the NPs, including particle size and chemical composition as well as the production of oxidative stress and cytokines by exposed cells. The possible antitumor application of TDNPs was explored in vivo.

**Materials and Methods**

**Cells**
The LL2 mouse lung cancer cell line was purchased from the Dainippon Sumitomo Pharma Co., Osaka, Japan. The CT26 mouse colon cancer cell line was a kind gift from Professor I.J. Fidler (MD Anderson Cancer Center, Houston, Tex., USA). Cells were cultured in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum.

**Cell Growth and Apoptosis**
Cell growth was assessed using the 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide dye assay, as described previously [8]. Apoptosis was assessed by examining 2,000 cells stained with Hoechst 33342 (Life Technologies, Carlsbad, Calif., USA) using a fluorescent microscope.

**Reagents**
TDNPs (10-nm anatase and 10- to 70-nm rutile) were obtained from TEIKA Co., Osaka, Japan. We also purchased gold colloid NPs (10 nm, BBI Solutions, Cardiff, UK), carbon black NPs (15 nm, WAKO Pure Chemicals, Osaka, Japan), glucose (WAKO) and glycinated bovine serum albumin (Sigma-Aldrich Chemical Co., St. Louis, Mo., USA).

**Ultraviolet Irradiation**
UVA irradiation at 365-nm peak length was performed with a UV lamp (xx-15 BLB, UVP Co., Tokyo, Japan) for 5 J/cm². UVB and UVC with <330 nm in length were cut with a UV cut filter (UV-33, Toshiba Co., Tokyo, Japan).

**Animals**
Four-week-old male BALB/c mice and 5-week-old male Fisher 344 rats were purchased from SLC Japan (Shizuoka, Japan). The animals were maintained according to institutional guidelines approved by the Committee for Animal Experimentation of Nara Medical University and the current regulations and standards of the Japanese Ministry of Health, Labor and Welfare (approval No. 9559).

**Animal Models**
To establish a subcutaneous tumor model, CT26 cancer cells (1 × 10⁷) were inoculated into the scapular subcutaneous tissue of BALB/c mice. Mice were observed for 4 weeks following inoculation. For evaluating the effect of anatase or rutile TDNPs, different sizes, i.e. 200 μg in 200 μl of phosphate-buffered saline (PBS), were injected into the CT26 subcutaneous tumor. For the NP aspiration model, TDNPs suspended in 200 μl of PBS were aspirated into the nose of Fischer 344 rats under anesthesia. For the NP oral model, TDNPs suspended in 200 μl PBS were administrated to BALB/c mice intragastrically. The toxicity of the different forms of TDNPs (200 μg) to BALB/c mice was determined following subcutaneous or intraperitoneal injection.

**Titanium Gel Plug**
Rutile TDNPs (200 μg) were mixed with melted 1.5% agar-PBS at 37 °C. The suspension was cooled in the lumen of an 18-gauge needle. The needle containing the titanium gel plug was inserted into the tumor and the plug was then expelled. For evaluation of toxicity of the titanium gel plug, one was inserted into the subcutaneous tissue of BALB/c mice without tumors.

**Tissue Titanium**
The subcutaneous tumor (1 g) was digested with proteinase K for 2 h at 45 °C, followed by treatment with 65% nitric acid overnight at 80 °C. The suspension was analyzed using a flameless atomic absorption spectrometer (AAS) to measure the titanium concentration (λ = 226.8 nm), using an AAS titanium standard (Sigma-Aldrich).

**Reverse Transcription-Polymerase Chain Reaction**
Total RNA (1 μg) was synthesized using the ReverTra Ace quantitative PCR (qPCR) RT kit (Toyobo, Osaka, Japan). The primer sets for amplification were listed below. Mouse Tnfa (accession No. D84199.2): upper 5′-AAG ATG GAG GAA GGG CAG TT-3′, lower 5′-GAT CCT GGA GGG GAA GAC AC-3′; mouse Il-1β (accession No. NM_008361.3): upper 5′-GCC CAT CCT GTA CTC AT-3′, lower 5′-AGG CCA CAG GTA TTT TGT CG-3′; mouse Cxcl2 (accession No. NM_009140.2): upper 5′-AGG TTG GGT ACC CTG AA-3′, lower 5′-AGG GAC CAT ATG AGC TAC GAT CAC TT-3′; mouse Hmgb1 (accession No. NM_010439.3): upper 5′-ACA GAG CGG AGA GAG TGA GG-3′, lower 5′-GGG TGC TTC TGT GAC CG-3′; mouse Actb (accession No. NM_007393.4): upper 5′-AGC CAT GTA CGT AGC CAT CC-3′, lower 5′-CTC TCA GCT GTG GAT GA AA-3′. The primers were synthesized by Sigma Genosys (Ishikari, Japan). The PCR conditions were set according to the provider’s instructions. PCR products were electrophoresed on a 2% agarose gel and visualized using ethidium bromide.

**Enzyme-Linked Immunosorbent Assay**
ELISA kits were used to measure concentrations of 4-hydroxy-2-nonenal (4-HNE; Cusabio Biotech Co. Ltd., Wilmington, Del., USA), 8-hydroxy-2′-deoxyguanosine (8-OHdG; Japan Institute for the Control of Aging, Shizuoka, Japan) and high mobility group box-1 (HMGB1; Shino-test Co., Tokyo, Japan), according to the manufacturers’ instructions.
Assessment of Cell Proliferation and Apoptosis

Using 4-μm-thin slice specimens of mouse tumor tissues, Ki-67 immunostaining and terminal deoxynucleotidyl transferase (TdT)-mediated dUTP-biotin nick-end labeling (TUNEL) assay. Immunohistochemistry was performed according to the method previously reported [9]. TUNEL assay was performed using the in situ cell death detection kit, POD (Roche Diagnostics, Indianapolis, Ind., USA). The percent frequency of Ki-67- or TUNEL-positive cells was calculated from the ratio of positive nuclei to 500 examined nuclei.
Statistical Analysis
Statistical significance was calculated by using the two-tailed Fisher exact test, the $\chi^2$ test and the unpaired Mann-Whitney U test with InStat software (Graphpad, Los Angeles, Calif., USA). Statistical significance was defined as a two-sided p value of <0.05.

Results

Cytotoxicity of Anatase TDNPs
The cytotoxicity of anatase TDNPs was assessed with or without UVB irradiation (1.5 J/cm²) in CT26 cells (fig. 1a, b). Both treatments showed a TiO₂ concentration-dependent cytotoxicity. However, UVB treatment increased cytotoxicity 2-fold. The cytotoxic effect of anatase TDNPs was confirmed in LL2 cells (fig. 1c) and was found to involve the induction of apoptosis in both LL2 and CT26 cells (fig. 1d).

Induction of Oxidative Stress by Anatase TDNPs
The oxidative stress generated by anatase TDNPs was assessed by measuring 4-HNE and 8-OHdG (fig. 1e, g). 4-HNE levels in CT26 cells treated with various concentrations of anatase TDNPs without UVB were about 80% of those in cells treated with TDNPs with UVB. 8-OHdG levels in CT26 cells treated with anatase TDNPs without UVB were about 60% of those in cells treated with anatase TDNPs and UVB. Various physiological conditions were examined for their effect on the oxidative stress induced by treatment with anatase TDNPs (fig. 1f, h). Acidic condition (pH 6.5), hyperglycemia (400 mg/dl), hyperthermia (38 °C), hypoxia (5% O₂) and advanced glycation end products (AGE, 100 μg/ml glycated bovine serum albumin) increased the levels of both 4-HNE and 8-OHdG in CT26 cells treated with anatase TDNPs.

Induction of Cytokines by TDNPs in Cancer Cells
The expression of TNFα, IL-1β, CXCL2 and HMGB1 mRNA was examined by RT-PCR in TDNP-treated CT26 and LL2 cells (fig. 2a). Expression of these cytokines was upregulated by TDNP treatment in both cancer cell lines.

Effect of TDNPs on Tissue Reactions in the Lung and Colon
To examine the effect of TDNPs on tissues, the lungs and colon of rodents were exposed according to protocol (fig. 2b, c). In rat lungs and mouse colon, granulomas were observed in a dose-dependent manner (fig. 2d–h). In these tissues, oxidative stress, detected by 8-OHdG and the macrophage-activating cytokines MIP-1α and HMGB1, were markedly increased (fig. 2i–k).

Induction of Oxidative Stress by Other NPs
To examine the hypothesis that the induction of oxidative stress is a common feature of NPs, rutile TDNPs, gold NPs and carbon black NPs were tested. The cytotoxicity of all 3 types of NPs was similar in CT26 cells (fig. 3a). The levels of 4-HNE and 8-OHdG were increased in cells treated with the 3 types of NPs (fig. 3b, c) with or without hyperglycemic or hypoxic conditions (fig. 3d, e). The mRNA expression of the macrophage-activating cytokines HMGB1 and MIP-1α was also upregulated by these NPs (fig. 3f).

Effect of Particle Size on the Activities of NPs
The effect of rutile TDNP particle size on the induction of oxidative stress, the expression of cytokines and tissue damage was determined in CT26 cells. The smaller the NPs, the higher were the levels of 4-HNE and 8-OHdG (fig. 3g, h). Using the protocol shown in figure 2b, 5 different sizes of rutile TDNPs were administered into the lungs of rats. Lung weight was increased and the number of granulomas was more pronounced in rats treated with smaller NPs (fig. 3i–l). Smaller NPs also more actively induced HMGB1 expression and the formation of 8-OHdG (fig. 3m, n).

TDNP as a Local Tumor Treatment
The anti-tumor effects of rutile TDNPs in vitro and of a gel plug containing TDNPs that was embedded into a tumor in vivo, were examined. As shown in figure 4a–c, smaller TDNPs showed a more pronounced inhibition of CT26 cell growth as well as relatively more oxidative stress. Six- and 10-nm NPs inhibited cell growth by 60% and caused high levels of 4-HNE and 8-OHdG.

In vivo treatment with a TDNP gel plug induced tumor cell death in the tumor (fig. 4d). In degraded tumor
cells, TDNP aggregations were found (fig. 4d inset). Titanium contents in the tumors were higher in 10-nm TDNP-treated tumors than those in 70-nm TDNP-treated tumors (fig. 4e). In 10-nm TDNP-treated tumors, TUNEL-positive apoptotic cells were increased and Ki-67-positive proliferating cells were decreased in comparison to those in tumors treated with gel alone (fig. 4f). Intratumoral insertion of the TDNP gel plug inhibited tumor growth in an NP size-, dose- and time-dependent manner (fig. 4g, h).

The local effect of TDNPs in the nontumoral tissues was evaluated by insertion of a TDNP gel plug into the subcutaneous tissue of the mouse with no tumor (fig. 4i). Thick fibrosis occurred in the insertion site in the subcutaneous tissue; however, no alteration was observed at the fascia and skeletal muscle. The systemic effect of TDNPs was evaluated by measuring urinary excretion of 8-OHdG as an index of general oxidative toxicity. Mice without a tumor were treated with rutile TDNPs or TDNP gel (10 nm, 100 μg). Subcutaneous or intraperitoneal administration of TDNPs suspended in PBS resulted in high concentrations of 8-OHdG in the urine. TDNP gel, administered intraperitoneally, melted and yielded the same levels of urine 8-OHdG (fig. 4j). In contrast, TDNP gel inserted into the subcutaneous tissue was thick with fibrosis and provided lower levels of urinary 8-OHdG than in the other groups (fig. 4j). These findings suggest that intratumoral administration of a TDNP gel was effective for inhibiting the growth of cancer cells with low toxicity to the host.

**Fig. 3.** Effect of NP composition and size on indices of toxicity and oxidative stress. a–c Changes in cell number, 4-HNE and 8-OHdG in CT26 cells treated with 400 μg/ml rutile TDNPs (10 nm), gold (10 nm) or carbon black (CB, 15 nm). d, e Effect of high glucose concentration (Glc, 400 mg/ml) and hypoxia (5% O2) on 4-HNE and 8-OHdG in CT26 cells treated with 200 μg/ml of TDNPs (10 nm), gold (10 nm) or CB (15 nm). f Expression of MIP-1α and HMGB1 mRNA in CT26 cells treated with 200 μg/ml TiO2 (10 nm), gold (10 nm) or CB (15 nm). g, h Effect of treating CT26 cells with different sizes of rutile TDNPs (200 μg/ml) on 4-HNE and 8-OHdG levels. i Effect of different sizes of rutile TDNPs on lung damage in the rat aspiration model (n = 5 for each group). j, k Rat alveoli 4 weeks after treatment with TDNPs. HE. j In rats treated with 70-nm TDNPs, minimal granulomatous changes are evident. k In rats treated with 10-nm TDNPs, marked granulomatous changes and thickening of the alveolar wall are evident. Number of granulomatous lesions (l), concentrations of HMGB1 (m) and concentrations of 8-OHdG (n) in the rat lungs following treatment with different sizes of TDNPs. Data are expressed as means ± SD from 5 animals or 3 independent trials. * p < 0.05.

**Discussion**

In this study, the toxicity of anatase and rutile TDNPs was determined with and without concomitant UV irradiation. TDNPs caused oxidative stress and induced the production of inflammatory cytokines in vivo. These effects were enhanced by UV irradiation. Smaller NPs generated more pronounced levels of oxidative stress and cytokines than did larger NPs. NPs caused granulomatous inflammation in the lung and intestine. Importantly, TDNPs damaged tumor tissue following intratumoral administration. TDNP administered into tumors formed micrometer-level aggregation, as reported previously [10], and stayed in degraded tumor tissue.

We found that treating CT26 cells with TDNPs produced oxidative stress without UV. This observation was consistent with other reports [11–13]. We also confirmed that gold and carbon black NPs induced oxidative stress. These results indicate that oxidative stress is a common feature of treating cells with NPs.

The mechanism by which NPs generate oxidants in an acellular system is reported to be based upon pro-oxidative functional groups at the NP surface and redox cycling of transition metals [14, 15]. We used NPs devoid of transition metals to evaluate acellular oxidant production. With such NPs, surface and size might be pivotal factors for generating oxidants. A size effect was reported for carbon black NPs [12]. In addition, the surface area of carbon black NPs correlated with the generation of oxidants.

Oxidative stress produced in response to TDNP administration was enhanced by high glucose concentration, acidic pH, hypoxia, a high temperature and the presence of AGE. This enhancement might be associated with the direct effects of NPs on cells to produce free radicals and other oxidants. In addition, NP-derived oxidative stress in vivo may involve mitochondria or NAD(P)H oxidase [13, 14]. The conditions that enhance TDNP-mediated oxidative stress may affect the redox status of mitochondrial oxidative phosphorylation. Such conditions are present in patients with diabetes and ischemia and NPs may be more toxic to such individuals.

Paradoxically, NPs might play a role in scavenging oxidants generated by hyperglycemia in an acellular system [16, 17]. In hypoxia, NPs derived from fullerene showed antioxidant effects in the wound-healing process [18]. These findings suggest that the particle surface might provide NPs with antioxidant properties, although our current data suggest that NPs are more likely to adversely affect these disease-associated conditions. It is noteworthy that cancer tissues are commonly hypoxic and...
Fig. 4. Effects of rutile TDNPs on cancer cells in vivo and in vitro. 

- a–c: Effects of treating CT26 cells for 48 h with different sizes of rutile TDNPs (200 μg/ml) on cell number, 4-HNE and 8-OHdG.
- d–f: Effect of TDNP gel plug on tumor. 
  - d: Massive cell death (asterisk) in the CT26 tumor treated with 10-nm TDNPs. Bar: 100 μm. Inset: Aggregation of TDNPs in degraded tumor cells. 
  - e: Titanium contents in the tumors were measured by AAS. 
  - f: TUNEL-positive apoptotic cells and Ki-67-positive proliferating cells in the CT26 tumor treated with 10-nm TDNP or gel alone. Inset: Representative in the tumor treated with 10-nm TDNP. 

- g: Effect of different sizes of rutile TDNPs in a gel on the in vivo growth of CT26 subcutaneous tumors (n = 5 mice in each group). 
- h: Dose-dependent effects of 10- and 70-nm rutile TDNP gel on the in vivo growth of CT26 subcutaneous tumors (n = 5 mice in each group). 
- i: Effect of the TDNP gel plug on nontumoral tissue: it was inserted into the subcutaneous tissue of tumor-negative mice. Subcutaneous fibrosis (asterisk).
- j: Effect of intraperitoneal and subcutaneous rutile TDNP gel on urinary 8-OHdG after 1 week. Data are expressed as means ± SD of 4 mice per group. SD from 5 animals or 3 independent trials.
Acidic [19]. This may enhance the oxidative stress produced by NPs, and suggests that NPs may be an effective anticancer treatment.

In our in vitro and in vivo experiments, inflammatory cytokines, such as MIP-1α and HMGB1, were produced by cancer cells and noncancerous tissues. These cytokines might enhance oxidative stress and activate granulomatous macrophages [20, 21]. This is recognized as the third mechanism of NP-associated oxidative stress [14]. HMGB1, a late-phase inflammatory cytokine, is secreted from macrophages activated with IL-1β or TNFα and promotes their secretion [21]. This vicious cycle between inflammatory cytokines and oxidative stress plays an important role in the cellular and tissue damage caused by NPs.

NPs generally induce oxidative stress, activate caspase-3 and induce apoptosis [22]. Thus, nonselective diffusion of NPs has the potential to evoke damage in noncancerous tissues. To avoid this potential toxicity, rutile TDNPs were intratumorally administered in a gel plug. Our data show that TDNPs administered in this manner induced apoptosis, suppressing the growth of remnant cancer cells otherwise stimulated by HMGB1 released from dead cells [23].

Overall, our data suggest that the local use of TDNPs is an effective cancer treatment. A single, localized dose of TDNPs might produce oxidative stress and inhibit cancer growth for a prolonged period of time with minimal side effects. In further study, examination of the antitumor effect and the side effects of long-term administration of TDNPs are needed.

Acknowledgments

This work was supported in part by grants-in-aid for Scientific Research from the Japan Society for the Promotion of Science (11017371) and the Ministry of Health, Labor and Welfare, Japan (11103430).

References


12 Koike E, Kobayashi T: Chemical and biological oxidative effects of carbon black nanoparticles. Cemosphere 2006;65:946–951.


