Technical Note

Image contrast assessment of metal-based nanoparticles as applications for image-guided radiation therapy

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ABSTRACT

Metal-based nanoparticles (NPs) have been extensively studied for dose enhancement applications in radiation therapy. This study investigated the utility of such NPs for image-guided radiation therapy (IGRT). Phantom images of gold NPs (AuNPs) and titanium peroxide NPs (TiOxNPs) with different concentrations were acquired using IGRT modalities, including cone-beam computed tomography (CBCT). AuNPs induced strong contrast enhancement in kV energy CBCT images, whereas TiOxNPs at high concentrations showed weak but detectable changes. The results indicated that these NPs can be used to enhance IGRT images as well as dose enhancement for treatment purposes.

1. Introduction

Nanoparticles (NPs) have been extensively studied for their use as radiosensitising agents in radiation therapy against cancer [1]. In the reaction with X-rays, metal NPs can generate secondary electrons and consequently reactive oxygen species (ROS) through radiolysis of water molecules, which can damage cancer cells and enhance the X-ray effects [2]. Along with the dose enhancement effects in radiation therapy, their radiosensitising properties can act as potential agents for enhancing the image contrast in computed tomography (CT) because of the X-ray absorption with metal atoms [3]. Thus, by combining these therapeutic and diagnostic enhancements, NPs have been explored for possible theranostic applications.

In current radiation therapy clinical practice, imaging techniques are increasingly used for patient set-up corrections as part of image-guided radiation therapy (IGRT) [4]. Cone-beam CT (CBCT) is among the most common IGRT techniques that allow precise patient set-up and minimisation of interfractional organ motions before the delivery of treatment beams. While standard diagnostic CT uses a fan beam with a single rotation of the kV X-ray source and detector, mounted on linear accelerator (linac) units. Moreover, megavoltage CT (MVCT) using high-energy beams from the linac are also available depending on treatment devices. As IGRT is performed during radiation therapy sessions, metal NPs, which are loaded to enhance the radiation dose in the tumour, can enhance the image contrast in IGRT modalities as well and assist the image registration process by visualising the clear tumour location in soft tissues.

The radiosensitising effects of gold NPs (AuNPs) have been demonstrated in many studies [5-7]. In addition to their therapeutic dose enhancement in radiation therapy, diagnostic enhancement in CT images has also been reported in several studies [8-10]. Although previous studies have used standard diagnostic CT or small animal CT scanners to assess the image contrast ability of AuNPs, none of them have investigated the contrast enhancement in IGRT imaging modalities using AuNPs and other NPs. The aim of this study was therefore to evaluate the contrast ability of metal NPs on IGRT modalities including CBCT and MVCT. Two different types of NPs were employed: AuNPs as high atomic number NPs and titanium peroxide nanoparticles (TiOxNPs) as low atomic number NPs. TiOxNPs were originally synthesised from titanium dioxide nanoparticles (TiO2NPs) with hydrogen peroxide (H2O2) as novel agents for enhancing radiation effects [11]. Owing to their unique...
ability to generate or release \( \text{H}_2\text{O}_2 \), they have shown promising dose enhancement under X-ray irradiation in vitro and in vivo studies [11–13]. Thus, in this study, TiOxNPs were considered a suitable specimen to investigate their image contrast properties as well as AuNPs.

2. Material and methods

2.1. Preparation of NPs

AuNPs were purchased from Nanoprobe (AuroVisiTM, lot number 06S700, Yaphank, NY, USA) and diluted to obtain a concentration of 40 mg/ml using phosphate-buffered saline. According to manufacturer’s instructions, a water-soluble organic ligand shell is coordinated on the gold core. TiOxNPs were synthesised based on a method described in our previous research [11]. The surface of TiOxNPs was functionalised using polyacrylic acid, in accordance with our previous studies [14,15]. The sizes of AuNPs and TiOxNPs were approximately 15 and 50 nm, respectively (supplementary Fig. 1). Final concentrations of NPs used in this phantom study were 0, 1, 5, and 10 mg/ml for each NPs sample, diluted using purified water to achieve the required concentrations.

2.2. Imaging modalities

Four different types of medical imaging modalities were applied in this study: standard diagnostic CT, linac-integrated kilovoltage cone-beam CT (CBCT), megavoltage CT (MVCT) in helical tomotherapy, and standard diagnostic magnetic resonance imaging (MRI). Although MRI-linac is on the way to becoming one of the common IGRT modalities [16,17], it was not available in any of our centres, thus, a standard MRI system was used for this study instead. Two milliliters of NP solution for each concentration was placed in holes of a water-equivalent cylinder phantom with a diameter of 15 cm. Thereafter, the phantom was scanned using the imaging modalities. Diagnostic CT images were obtained using an Optima CT580 scanner (GE Healthcare, Waukesha, WI, USA) with a tube potential of 120 kVp and two different tube currents of 100 and 400 mA (for head and body scan protocols, respectively). CBCT images were acquired using two types of Varian linac: Clinac iX and TrueBeam (Varian Medical Systems, Palo Alto, CA, USA). Furthermore, half- and full-rotation CBCT scans were performed with standard acquisition protocols. MVCT images were obtained using Tomotherapy (Accuray, Sunnyvale, CA, USA), wherein 3.5 MV X-ray beams were available for the image guidance [18]. The obtained CT images were imported into the Eclipse software (version 15.6, Varian Medical Systems, Palo Alto, CA, USA) and the Hounsfield units (HUs) for each sample were measured using five different axial slices. The data were represented as mean ± standard deviation. In addition to CT images, T1- and T2-weighted images were acquired using a MAGNETOM Avanto 1.5T MRI scanner (Siemens AG, Munich, Germany) with a turbo spin-echo sequence. Scan parameters for each imaging modality were based on routine clinical settings. The details are summarised in Supplementary Table 1.

2.3. Contrast visibility test

The visibility of AuNPs and TiOxNPs on CT images was assessed to determine whether they were feasible as image contrast agents for clinical use. We prepared a CT image, wherein two samples of NPs and eight samples of water were randomly arranged. Subsequently, four certified medical physicists independently selected two out of the ten samples, thought to be NPs. This test was replicated for different concentrations of the NPs and different sets of CT window width and level (WW/WL). The visibility of the NPs was calculated as a percentage of correct answers. The contrast-to-noise ratio (CNR) was also calculated for quantitative measures [19].

3. Results

For both AuNPs and TiOxNPs the contrast levels in CBCT images were the same as those for standard diagnostic CT regardless of scan parameters (Fig. 1A). Large fluctuations observed on CT attenuation at 1 mg/ml for both NPs were due to nonuniformity and noise of CBCT images. The absolute values of the HUs increased linearly, depending on NP concentrations in both AuNPs and TiOxNPs (supplementary Fig. 2A-F). The overall mean slopes were 26.4 ± 1.3 and 2.7 ± 0.7 HU/(mg/ml) for AuNPs and TiOxNPs, respectively. Evidently, the enhancement effect of AuNPs was much higher than that of TiOxNPs. The HUs of these NPs on MVCT images were much lower than those on the kV CT images, and no visible differences were observed. While the HUs increased linearly with NP concentration on MVCT images (supplementary Fig. 2G), the slopes of linear fits were 1.0 and 0.3 HU/(mg/ml) for AuNPs and TiOxNPs, respectively. Typical NPs images of CT, CBCT, and MVCT are represented in Fig. 1B. Moreover, in T2-weighted MR image, TiOxNPs were negatively enhanced, whereas there were no differences in AuNPs (Fig. 1B and 1C).

The visibility test result showed that images of AuNPs were correctly identified from control images at all concentrations. The contrast of TiOxNPs could be almost visible at the concentrations of 5 and 10 mg/ml on images with a liver window setting; however, that of 1 mg/ml had a poor correct rate with a very low CNR, indicating no visible contrast at this concentration. The visibility and CNR for both NPs are summarised in Table 1.

4. Discussion

This study showed the potential ability of two different types of metal NPs as contrast agents in IGRT modalities. As expected, AuNPs had visible contrast in CBCT images regardless of exposure settings as well as in diagnostic CT images. The results were similar to a previous
caused no significant changes in X-ray attenuation using three different nm AuNPs [8]. They also demonstrated that different sizes of AuNPs were similar to those of previous studies as TiOxNPs caused visible changes in standard CT images [25] whereas there was a dependence on the NP concentration. Furthermore, the findings indicate that metal NPs can be possibly utilised during image registration process in IGRT practice, and can sequentially enhance the irradiated dose for treatment.

Table 1
Visibility of AuNPs and TiOxNPs in a CT phantom image.

| NPs    | Concentration | CNR | WW/WL | Visibility (%) |
|--------|---------------|-----|-------|----------------|
| AuNPs  | 1 mg/ml       | 7.4 | Abdomen 100 | 90 |
|        | 5 mg/ml       | 35.8| Abdomen 100 | 90 |
|        | 10 mg/ml      | 72.9| Abdomen 100 | 90 |
| TiOxNPs| 1 mg/ml       | 0.6 | Abdomen 12.5 | 80 |
|        | 5 mg/ml       | 2.9 | Abdomen 12.5 | 80 |
|        | 10 mg/ml      | 5.9 | Abdomen 50 | 70 |

CNR, contrast-to-noise ratio; WW, window width; WL, window level. WW/WL = 350/50, 150/50, 400/40 for abdomen, liver, and pelvis settings, respectively.

study by Dong et al., wherein enhanced image contrast was observed with the attenuation rate of 23.7 HU/(mg/ml) in diagnostic CT using 50 nm AuNPs [8]. They also demonstrated that different sizes of AuNPs caused no significant changes in X-ray attenuation using three different clinical CT systems with different tube potentials, possibly suggesting no size effects on the contrast enhancement in CBCT images, even though single-sized AuNPs were employed in our study. Despite the enhanced contrast of AuNPs in kV CT images, a low contrast enhancement could be observed in MVCT images even at a 10 mg/ml concentration of AuNPs. Theoretically, the Compton scattering becomes a dominant process for MV X-rays in the interaction with materials having a high atomic number, such as gold, and the X-ray cross-section is smaller than that in the interaction with kV energy photons, in which the photoelectric effect is dominant. In contrast to low X-ray attenuation of AuNPs in MVCT images, there are several in vitro studies that report significant radiosensitising effects of AuNPs in response to MV X-ray irradiation [20,21]. The discrepancy between therapy and image contrast enhancements is possibly due to biological factors, including intracellular ROS and oxidative stress induced by AuNPs [22].

The contrast enhancement caused by TiOxNPs in CT images were much weaker than those by AuNPs. This was because titanium has lower oxidative stress induced by AuNPs [22] and mass attenuation coefficient (\(\mu/\rho = 0.2721\) cm\(^2\)/g) than those of gold (79, 5.158 cm\(^2\)/g, respectively, at 100 keV photons) [23], resulting in less interaction with X-rays than gold. However, our qualitative results of the visibility test indicated that TiOxNPs with a concentration greater than 5 mg/ml could still be identified in certain soft tissues similar to water HU. The CNR of 3 is possibly a threshold for NPs contrast to be observable, which agrees with the Rose criterion [24]. A few studies have investigated the CT image contrast enhancement of TiOxNPs which are foundational materials of TiOxNPs. One previous study evaluated it with up to 1 mg/ml TiOxNPs and found no visible changes in standard CT images [25], whereas another study showed that TiOxNPs with a concentration of 15 mg/ml had 26.6 ± 4 HU in CT images indicating that they were detectable on a CT scanner even at low concentrations [26]. Our results for TiOxNPs were similar to those of previous studies as TiOxNPs caused visible changes in CT images at high concentrations. However, TiOxNPs induced negative contrast in T2-weighted MRI, whereas AuNPs did not. Our previous study using TiOxNPs also demonstrated the change in T2 relaxation time with TiOxNPs [25]. The negative enhancement of NPs may show darkened tumour regions in T2-weighted MRI, when they are used as radiosensitisers for tumours, which may be, however, unfavourable in IGRT using MRI-linac.

There are several limitations in this study. The contrast enhancements were assessed in a phantom with uniform contrast around NPs samples; however, visual appearances may be changed in a human body where surrounding tissues have different contrast. Similar to other in vivo studies that demonstrated the contrast enhancement of AuNPs in CT images using mice [3,27], future studies should be conducted to determine the contrast enhancement ability of NPs with subsequent image registration in clinical situations using IGRT modalities. Another limitation is cellular cytotoxicity of NPs at high concentrations, which is not considered in this study as phantom-based experiments. While a higher concentration of NPs may cause higher contrast enhancement in images, it can also increase cytotoxicity in cells. Therefore, in vitro and in vivo studies are warranted to find optimal concentrations of NPs as a clinical theranostic agent in future.

In conclusion, both AuNPs and TiOxNPs had contrast enhancement in kV energy CBCT images as well as in standard diagnostic CT images, whereas there was a dependence on the NP concentration. Furthermore, the findings indicate that metal NPs can be possibly utilised during image registration process in IGRT practice, and can sequentially enhance the irradiated dose for treatment.

Conflicts of interest
None.

Declaration of Competing Interest
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data
Supplementary data to this article can be found online at https://doi.org/10.1016/j.jphro.2021.11.003.

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