Investigation of Fiducial Marker Migration during the Treatment of Bladder Cancer with Image-Guided Radiation Therapy

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Abstract

Surgically implanted fiducial markers are widely used in the treatment of prostate, lung, and breast cancers with Image-Guided Radiation Therapy (IGRT). Fiducial markers are typically non-migrating gold seeds with various dimensions for different site applications. To prevent markers from migrating in implanted soft tissue, their surfaces are cross-cut using a special knurling process. Implanted fiducial markers are felt to constitute a more reliable reference coordinate system than surrounding bony landmarks for a moving target and are used for point-based image registration. Studies have shown that fiducial markers are safe, stable, and can provide superior tumor tracking and targeting capability during both treatment planning and treatment delivery. At our institution, implanted fiducial markers are now being used for target volume localization in bladder cancer patients treated with IGRT. In this paper, we present our initial experience with this novel treatment procedure in terms of fiducial marker migration.

Keywords: Bladder cancer; IGRT; IMRT; OBI; Fiducial marker

Introduction

In 2017, approximately 79,030 Americans were diagnosed with bladder cancer and 16,870 died of this disease [1]. The most common type of bladder cancer is urothelial carcinoma, which accounts for approximately 90% cases in the United States. Based on a 7-year study, the overall 5-year relative survival for all bladder cancer patients was 77.4%. For patients with muscle invasive bladder cancer, excellent local control rates can be achieved with either primary cystectomy or radiation therapy in combination with chemotherapy. However, the majority of the patients with bladder cancer are over 65 years of age, with the median age of 73 at the time of diagnosis [1,2]. For many of these patients, Chemo Radiation Therapy (CRT) is the only safe and curative option. In addition, many younger patients who have had a Complete Response (CR) to induction chemotherapy will prefer bladder preservation therapy. Thus, consolidative CRT is also an attractive treatment option for this group of patients [3,4]. Regardless of the treatment algorithm, delivering high radiation dose safely and accurately to the target in the bladder is a complex process. Due to bladder’s anatomical structure and central location in the pelvis, there are unique technical challenges involved in treating bladder cancer using conventional radiation therapy. Firstly, the bladder is a hollow organ with elastic muscular wall. Under normal physiological conditions, its volume undergoes periodic and dramatic variation, from empty to full expansion. The bladder volume change can be as large as 600 ml. This imposes a huge technical challenge on target localization for a safe and accurate radiation delivery. Even though patients are instructed to drink the same amount of water daily prior to each radiation treatment, many have difficulty maintaining a full bladder secondary to irritation by tumor, inflammation related to radiation treatment or due to compression by the immobilization device or mold. Secondly, the position of the bladder relative to the adjacent pelvic bone can be displaced by patient position on the treatment couch. This positioning displacement or organ shift is particularly apparent when the bladder is full. To compensate for these uncertainties in target localization, the most common approach is to expand the Clinical Target Volume (CTV) three-dimensionally by a large margin at least equal to the distance change in bladder’s radial direction. As a result, the planning target volume (PTV) will include a sizable shell of normal tissue, resulting in unnecessary radiation-induced side effects and damage. Furthermore, the bladder is surrounded by the large and small intestines which are very sensitive to radiation. This is particularly the case for the
small intestines, which have a lower dose tolerance. Thus, radiation dose escalation for bladder cancer is challenging without an accurate and reproducible system for target localization in the bladder. The latest generation of medical Linear Accelerators (LINAC) is equipped with an On-Board Imaging System (OBI) in the Kilo-Volts (KV) x-ray range. The modern OBI is capable of producing KV images with a quality close to that of diagnostic x-ray images. However, the low electron density of soft tissue, including bladder tissue, does not exhibit sufficient intrinsic contrast on KV images, which results in poor image contrast. Consequently, target localization based on inherent soft tissue contrast cannot be reliably performed with KV imaging. Although image registration using bony landmarks can be performed using KV imaging, this technique simply cannot track bladder displacement and bladder volume change. An alternative approach to KV imaging would be to use an imaging contrast agent through a catheter during radiation treatment. However, this approach is also impractical because it results in patient discomfort, increases setup time, and poses an infection risk to the patient. In addition, the contrast agent can significantly alter the electron density of the bladder and impact the radiation attenuation coefficient, which can, in turn, affect the delivered radiation dose distribution. Without a reliable image-guidance, radiation therapy in this setting can result in overdosing adjacent critical organs and/or under dosing the target. Over the past few years, surgically implanted fiducial markers have been increasingly used in the treatment of prostate, lung, and breast cancers with Image-Guided Radiation Therapy (IGRT) [5,6]. At our institution, implanted fiducial markers are now being used for target volume localization in bladder cancer patients treated with IGRT. However, unlike other soft tissue sites, such as the prostate and the breast, implanting fiducial markers safely into the thin bladder wall is a more complicated surgical procedure. Our initial assessment of the use of bladder fiducial marker placement is that it significantly improves the accuracy of bladder volume localization and greatly reduces the treatment margin. More importantly, it provides an opportunity for further radiation dose escalation, which may lead to improved tumor control and overall survival. In this paper, we present our initial experience with this novel technique for the treatment of muscle invasive bladder cancer and, in particular, our preliminary assessment of its target localization accuracy and reliability.

Materials and Methods

Fiducial Marker Implantation

Our bladder IGRT treatment protocol uses Visicoil gold fiducial markers (IBA Dosimetry, Bartlett, TN) as the reference points for the point-based image registration. Visicoil fiducial markers are a helical, flexible, and thin gold coil. They come with various diameters and lengths. Usually, smaller diameter markers are optimized for KV and Cone Beam CT (CBCT) visualization and the larger ones are designed for Mega-Volts (MV) Electronic Portal Imaging Device (EPID) visualization. Visicoil fiducial markers are small enough so that they will not introduce metal streak artifacts on a CT scan and alter the CT number distribution in the soft tissue near the implantation site [7]. Thus, they are ideal for soft tissue application and IGRT treatment. Visicoil fiducial markers are pre-loaded on a needle-carrier delivery
device and can be implanted to the patient quickly. For our routine IGRT treatment, 3 to 4 Visicoil fiducial markers are implanted into the detrusor muscle wall of the bladder near the site of tumor resection [8]. Fiducial marker placement is typically performed at the time of Trans Urethral Resection (TUR), with a needle applicator passing through a standard adult rigid cystoscope.

IGRT Treatment Planning and Delivery

The patient was simulated and scanned by a Brilliance Big Bore CT scanner (Phillips Medical Systems, Cleveland, OH, USA) in the supine position one week after the placement of fiducial markers. The patient was immobilized in a customized thermoplastic mold to minimize patient movement during the procedure. The patient drank GoLytely (Braintree Laboratories, Braintree, MA) to empty the bowel the evening before CT scan. A rectal catheter was placed to localize the rectum during the scanning procedure. A Foley catheter was not used for the localization of the urethra. CT images were acquired (3-mm slice thickness) of the pelvis. An Intensity-Modulated Radiation Therapy Plan (IMRT) was computed using an in-house treatment planning system (Top Module). The CTV, urethra, rectum, bowel, and bladder were delineated on the treatment planning CT scan by a radiation oncologist. The PTV was created by adding a 1-cm margin around the CTV, except at the interface of the prostate and the rectal wall, where a 6-mm margin was used. The planner delineated the femoral heads to include them in the final dose calculation and statistics. In addition, a series of PTV-concentric tuning structures were also created to improve the target dose conformity and homogeneity and confine the dose hot spots to the PTV. Using Boolean operations, the overlapped structures were optimized independently so that the optimizer could steer hot spots away from the critical structures. Most treatment plans consisted of 5 coplanar beams at 225º, 285º, 0º, 75º, and 135º in Varian IEC scale. Given a set of dose limits and dose–volume constraints, the plan was optimized by minimizing a quadratic objective function using an iterative gradient search algorithm. The quadratic objective function was constructed as the sum of squares of differences between the desired and actual doses. The algorithm computed the optimal intensity map for each beam such that the dose distribution resulting from all beams met the dose constraints specified by the planner. If the criteria for plan acceptance were not met, a trade-off between the target dose coverage and the Organ-At-Risk (OAR) constraints would have to be made. Once optimal intensity maps were obtained, Multiple-Leaf Collimator (MLC) sequences were subsequently generated using the Dynamic MLC (DMLC) technique. Based on the leaf sequences, the final dose distribution was then computed using a pencil-beam algorithm. During patient treatment, the radiation plan was delivered on a Varian Trilogy linear accelerator (Varian Medical Systems, Palo Alto, CA).

Results

In order to investigate the magnitude of migration of the fiducial markers in the bladder during the course of treatment, we studied an 84-year old male with bladder cancer recently treated at our department. This patient was felt to be a good candidate for tracking the pattern of the fiducial marker migration and assessing the effect of bladder volume on bladder target localization. The patient had 3 Visicoil fiducial markers implanted in the bladder wall and was treated with a two-phase IGRT plan: 1) initial phase using 25 fractions [180 cGy/fx] with an empty bladder, 2) the second phase using 11 fractions [180 cGy/fx] with a full bladder. The full bladder volume at the time of phase 2 simulation was 262.6 cc. The Visicoil fiducial markers implanted into the bladder wall had a diameter 0.35 mm. The initial distances among those fiducial markers were 3.48, 3.10, and 2.52 cm, respectively, as measured on the treatment planning CT scan. Prior to each daily treatment, the patient was setup in reference to a pair of orthogonal Digitally Reconstructed Radiographs (DRR) at the gantry angles of 45º and 315º, respectively (Figure 1). The KV OBI was set at 50 cm away from the MV radiation isocenter. The accuracy of the patient setup was evaluated by registering the 3 fiducial markers on the orthogonal pair of radiographic images acquired by the KV OBI with those on the DRRs. The image registration was performed using a rigid body model, which yielded three translational shifts and one rotational shift (yaw). The patient shifts were corrected by automatically moving the treatment couch to the target position. Following the shift correction, a new pair of KV images was then acquired to confirm the patient setup to be within our institutional tolerance (±1.0 mm). Following the shift correction, a new pair of KV images was then acquired to confirm the patient setup to be within our institutional tolerance (±1.0 mm). After a satisfactory patient setup had been achieved, the radiation treatment was delivered. Post-treatment, the daily KV images were analyzed off-line using the Varian image registration software, Offline Review. To facilitate our analysis, we defined the three fiducial markers as A, B, and C, respectively, and measured the baseline distance between each pair of the fiducial markers on the DRs obtained on the simulation date. The baseline distances were d AB=2.34cm, d BC=2.73cm, and d CA=2.22cm at the gantry angle of 315º and d AB=1.2cm, d BC=1.37cm, and d CA=2.33cm at the gantry angle of 45º. These were the distances projected onto the isocenter plane as viewed by the MV radiation source and, therefore, were gantry angle-dependent. To assess the fiducial marker migration as a function of time, we also measured the distance between each pair of the fiducial markers on the daily KV images for each treatment fraction. (Figure 2) shows the distances as measured on daily KV images at the gantry angle of 315º as a function of treatment fraction number. (Figure 3) shows the distances as a function of treatment fraction number at the gantry angle of 45º. Solid markers on the plots indicate treatment fractions with empty bladder (phase 1) and hollow markers indicate those treated with full bladder (phase 2). As seen in the plots, despite the significant bladder volume changes, the distances between fiducial markers were not drastically changed. When the plots were fitted to a linear trend line, we found that the distances decreased at a very small rate of approximately 0.010 cm/fx. In specific, the respective rates were r AB=0.008 ± 0.001cm/fx, r BC=-0.011 ± 0.002cm/fx, and r CA=-0.009 ± 0.001cm/fx at the gantry angle of 315º. The corresponding values at the gantry angle of 45º were r AB=0.004 ± 0.001cm/fx, r BC=-0.008 ± 0.001cm/fx, and r CA=-0.010 ± 0.001cm/fx. When examining the full and empty bladder data together, we found that the mean distances were d AB=2.16 ± 0.12cm, d BC=2.39 ± 0.15cm, and d CA=2.23 ± 0.13cm at the gantry angle of 315º and d AB=1.2cm, d BC=1.37cm, and d CA=2.33cm at the gantry angle of 45º.
d_{BC}=2.70 \pm 0.11\text{cm}, \text{and } d_{CA}=2.17 \pm 0.25\text{cm} \text{ at the gantry angle of 315º}

\text{and } d_{AB}=1.17 \pm 0.09\text{cm}, d_{BC}=1.49 \pm 0.11\text{cm}, \text{and } d_{CA}=2.27 \pm 0.13\text{cm}

\text{at the gantry angle of 45º. They did not drift greatly from the baseline values. These findings were consistent with the results reported previously for prostate patients.}

Compared to the magnitude of the fiducial marker migration, the daily couch shifts necessary to correct the random patient setup errors and the bladder displacement were much larger. For this patient, the mean daily couch shifts in the Vertical (VRT), Longitudinal (LNG), and Lateral (LAT) directions were $C_{\text{VRT}}=0.07 \pm 0.35\text{cm/fraction}$, $C_{\text{LNG}}=0.56 \pm 0.30\text{cm/fraction}$, and $C_{\text{LAT}}=-0.39 \pm 0.36\text{cm/fraction}$, respectively, for the empty bladder treatments and $C_{\text{VRT}}=-0.27 \pm 0.28\text{cm/fraction}$, $C_{\text{LNG}}=1.21 \pm 0.41\text{cm/fraction}$, and $C_{\text{LAT}}=-0.20 \pm 0.43\text{cm/fraction}$ for the full bladder treatments (Figure 4). These findings demonstrate that bladder target localization using implanted fiducial markers is reliable because the coordinate system established by the fiducial markers remains relatively fixed in space as compared to the magnitude of the random patient setup errors and the bladder displacement in the pelvis. We also performed statistical analysis of the fiducial marker migration. Figure 5 shows the histogram of the fiducial marker migration $d_{CA}$ as an example. It followed a normal distribution with a mean of 2.20 cm and the standard deviation of 0.14 cm. The mean was effectively equal to the $d_{CA}$ value (2.22 cm) measured at the date of patient simulation. For about 95% of the treatment fractions, $d_{CA}$ fell between 2.1 and 2.4 cm.

Conclusions

Visicoil fiducial markers have been found to be stable in prostate cases. Several studies have shown that the typical marker migration over an entire treatment course is about 1.0 mm. Similar investigations have been performed in lung, breast, and GI cases. However, bladder Visicoil fiducial marker implantation is a relatively new clinical application and the fiducial marker migration in the bladder has not been fully investigated. Our analysis shows that the distances between two neighboring fiducial markers implanted in the bladder shrank at a mean rate of -0.043±0.082mm per fraction and were not correlated with the bladder volume change. The typical numerical order of fiducial marker migration for the entire treatment course was about 1.0 mm, which was much smaller than the daily couch shifts. This provides a concrete evidence for validating this procedure as a viable IGRT technique in bladder cancer treatment. The use of Visicoil as fiducial markers in bladder cancer patients for IGRT treatment is still a new technique. Implantation of Visicoil fiducial markers into a thin and dynamic bladder wall requires significant clinical experience and extensive training. Currently, the number of bladder cancer patients implanted with Visicoil fiducial markers for IGRT treatment is very limited. More patient data using the bladder Visicoil fiducial markers need to be analyzed to ensure a better understanding of the causes and patterns of fiducial marker migration.

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References