A new two-step accurate CT-MRI fusion technique for post-implant prostate cancer

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Abstract

Purpose: To develop an accurate method of fusing computed tomography (CT) with magnetic resonance imaging (MRI) for post-implant dosimetry after prostate seed implant brachytherapy.

Material and methods: Prostate cancer patients were scheduled to undergo CT and MRI after brachytherapy. We obtained the three MRI sequences on fat-suppressed T1-weighted imaging (FST1-WI), T2-weighted imaging (T2-WI), and T2*-weighted imaging (T2*-WI) in each patient. We compared the lengths and widths of 450 seed source images in the 10 study patients on CT, FST1-WI, T2-WI, and T2*-WI. After CT-MRI fusion using source positions by the least-squares method, we decided the center of each seed source and measured the distance of these centers between CT and MRI to estimate the fusion accuracy.

Results: The measured length and width of the seeds were 6.1 ± 0.5 mm (mean ± standard deviation) and 3.2 ± 0.2 mm on CT, 5.9 ± 0.4 mm, and 2.4 ± 0.2 mm on FST1-WI, 5.5 ± 0.5 mm and 1.8 ± 0.2 mm on T2-WI, and 7.8 ± 1.0 mm and 4.1 ± 0.7 mm on T2*-WI, respectively. The measured source location shifts on CT/FST1-WI and CT/T2-WI after image fusion in the 10 study patients were 0.9 ± 0.4 mm and 1.4 ± 0.2 mm, respectively. The shift on CT/FST1-WI was less than on CT/T2-WI (p = 0.005).

Conclusions: For post-implant dosimetry after prostate seed implant brachytherapy, more accurate fusion of CT and T2-WI is achieved if CT and FST1-WI are fused first using the least-squares method and the center position of each source, followed by fusion of the FST1-WI and T2-WI images. This method is more accurate than direct image fusion.

Key words: brachytherapy, fusion, magnetic resonance imaging, post-implant, prostate cancer.
Neoadjuvant hormonal therapy was performed to adjust the prostate volume from 15 to 30 ml.

**Post-implant image acquisition**

Four weeks after prostate brachytherapy, CT and MRI data were acquired for dosimetric analysis according to the recommended schedule [14]. Three imaging sequences were obtained: fat-suppressed T1-weighted imaging (FST1-WI), T2-weighted imaging (T2-WI), and T2*-weighted imaging (T2*-WI) using a Siemens Avanto 1.5T MRI scanner (Siemens, Munich, Germany) with 3 mm-thick-slices. The technical parameters of FST1-WI were as follows: repetition time (TR)/echo time (TE) in ms, 955/150; in-plane resolution, 1.0 × 0.7 mm². The technical parameters of T2-WI were: TR/TE, 5000/100; in-plane resolution, 1.0 × 0.7 mm²; Turbo Spin Echo factor, 15. The technical parameters of T2*-WI were: TR/TE, 600/19; in-plane resolution, 1.1 × 0.7 mm². FST1-WI, T2-WI, and T2*-WI were obtained successively. During PID, the prostate is contoured using T2-WI to estimate prostate volume.

Within 1 h of MRI, a post-implant CT was performed using a GE spiral CT (Hi-Speed Dxi; GE Healthcare, Buckinghamshire, UK). During the CT scan, a urinary catheter (8 Fr) was inserted. Sequences of 2 mm slices were acquired using a 50 mm field of view (FOV), and a pitch of 2 mm/2 mm = 1 (defined as the table feed/total detector width of the collimated beam). No intravenous contrast material was used during the CT or MRI scans.

**Comparison of source images in CT, FST1-WI, T2-WI, and T2*-WI**

We measured the lengths and widths of the source images on CT, FST1-WI, T2-WI, and T2*-WI, and compared the data among the modalities (Fig. 1). We determined the degrees of signal intensity loss on MRI to be those of the seeds. Overlapping source images and extra-prostatic areas that might have increased the inaccuracy of size measurements were excluded from the analysis. Seeds with distinct outlines in an intra-prostatic area were selected for this study. Of the 636 seed sources, 186 were found to be overlapped and located in an extra-prostatic area; these 186 sources were excluded from the study. Consequently, 450 seed sources were measured.

**Fusion method**

We accomplished the registration using 12 geometric centers of the seed distribution by a least-squares method [15] between CT and each MRI sequence (FST1-WI and T2-WI) using the Variseed 8.0 software. The two distinct outline seeds were selected on either side (right and left lobe) at the base, middle, and apex of the prostate (12 seed centers total), and the 12 source centers were used for seed match registration. Extra-prostatic sources were not used. The width of the T2*-WI source images appeared nearly twice as long as the seed clarifying sequences (FST1-WI). Because of the considerable loss of T2*-WI signal intensity, many of the sources overlapped. Detection of the seed centers on T2*-WI was difficult enough that an estimation...
of the fusion accuracy between CT images and T2*-WI was not performed.

**Estimation of the fusion accuracy**

In each patient, the source location shifts on CT/FST1-WI and CT/T2-WI were calculated from source location data using the Variseed Source Location Export function in Variseed 8.0. The mean shifts for all sources were calculated and estimated for each patient.

**Statistics**

To assess the fusion accuracy between CT and each MRI sequence, each data point was compared using the paired-sample t-test. Analyses were performed using SPSS, version 18 (SPSS Inc., Chicago, IL, USA). P values < 0.05 were considered statistically significant.

**Results**

The estimated mean prostate volume was 21.6 ml (range 13.1-32.8 ml). The median number of intra-prostatic seeds per patient was 45 (range 29-59, with a total of 450 sources). The measured lengths and widths of the seeds were 6.1 ± 0.5 mm [mean ± standard deviation], and 3.2 ± 0.2 mm on CT, 5.9 ± 0.4 mm and 2.4 ± 0.2 mm on FST1-WI, 5.5 ± 0.5 mm and 1.8 ± 0.2 mm on T2-WI, and 7.8 ± 1.0 mm and 4.1 ± 0.7 mm on T2*-WI, respectively (Fig. 2). The measured source location shifts on CT/FST1-WI and CT/T2-WI after image fusion in the 10 study patients were 0.9 ± 0.4 mm and 1.4 ± 0.2 mm, respectively. The shift on CT/FST1-WI was less than on CT/T2-WI (p = 0.005).

**Discussion**

This study compared the accuracy of seed match registration using the least squares method with different MRI sequences and concluded that the sequences that more precisely define the seeds resulted in a better match. Additional MRI sequences were then matched to this registration, under the assumption that they would match since these images were obtained at the same time from the same position. The present study determined that FST1-WI was the most suitable MRI sequence for CT/MRI fusion. The differences in the size of the source images between CT and FST1-WI were the smallest; thus, we accomplished the registration easily and accurately by using the geometric centers of the source images on each image by the least-squares method. We invented a new CT-MRI fusion technique using FST1-WI-namely, a two-step image-fusion method using two sets of images (CT and FST1-WI, and FST1-WI and T2-WI). In the first step, fusion of CT images and FST1-WI was performed by matching selected seed positions on each image. The least squares method was used during the first step, because McLaughlin et al. previously reported that registration could be accurately accomplished using seed positions by this method [15]. In the next step, both CT and FST1-WI obtained from the first step could be accurately matched to T2-WI, because FST1-WI and T2-WI were co-registered. This two-step image fusion technique provides high-quality CT-MRI fusion images. Also, in a clinical setting, the accuracy of the image fusion obtained from the two-step fusion technique can be evaluated by eye as the shift in seed between the fused CT images and FST1-WI. This is a critically important method for evaluating the fusion accuracy.

Computed tomography and T2-WI were fused more accurately using the two-step image-fusion technique than using direct image fusion. We compared the two-step image-fusion and direct image-fusion measuring source location shifts on CT/FST1-WI and CT/T2-WI after the seed match registration using 12 geometric seed centers by the least-squares method. The shift was significantly less on CT/FST1-WI than on CT/T2-WI (p = 0.005).

The two-step fusion method is an approximation, and a residual mismatch may require an anatomic match and correction. We acknowledge that errors may occur with the least squares method and that one sequence may merely lead to a better match than another. At the mid-prostate, the contour of the prostate is relatively clear on CT. After seed to seed registration, CT images of the mid-prostate may reveal that seed to seed registration by the least squares method is in fact only an approximation and not always adequate. This overlay of mid-prostate
images will enable identification of residual mismatches that require an anatomic match and correction.

Katayama et al. demonstrated that T2*-WI is useful for the detection of seeds without intravenous contrast media [16]. However, metallic implants such as seed sources cause considerable loss of signal intensity in T2*-WI (Fig. 1) [17]. Detection of the center of implanted seeds by T2*-WI is difficult.

Several other image fusion methods can be used, such as fusion by bony anatomy, mutual information (MI) registration, and visual manual seed match registration, using available software. Fusion by bony anatomy is sub-optimal in terms of accuracy and reproducibility. Roberson et al. showed that MI registration using large-image data sets of the same anatomic volume was more substantial than seed match CT/T2 MR registration, and approximately 30 min was needed for MI registration compared with several hours for seed match registration [18]. However, given the independent motion of the prostate compared with pelvic soft tissue, MI registration using anatomic image data sets will be inaccurate. Our two-step intra-prostatic seed match MR registration can be achieved as quickly as and is more accurate than MI registration for the prostate. Visual manual seed to seed registration on T2-weighted MRI to CT images using available software is commonly performed; however, this method is more time consuming than our two-step method. As the next step of research, it would be interesting to quantitatively compare our method to mutual information or simply landmark registration.

Maletz et al. reported a method [9] requiring the use of specialized MimVista software. Conversely, in the present study, image fusion and evaluation were performed using the more common Variseed software. To evaluate image fusion quality, Maletz et al. compared dose volume parameters for the target volume. However, the parameters they used were dependent upon prostate contouring and the dose distribution after implantation. The evaluation of dose volume parameters is not a substituted method for measuring fusion accuracy. In the present study, we examined the distance of the same seed between fused CT and MR images to evaluate the fusion accuracy. Since it is not dependent on dose volume parameters, the evaluation methodology is preferred.

In the present study, seed match registration was performed using the least squares method with 12 geometric centers between CT images and each MRI sequence (FST1-WI and T2-WI). Registration using 12-seed centers may be a practical first step; without a systematic comparison of seed numbers, we cannot conclude that 12 is an adequate number. Roberson et al. attempted to define all seeds on T2-WI during seed match registration [18]. However, it is difficult to visualize all implanted seed sources, and the inclusion of vague sources may decrease fusion accuracy.

It is possible to estimate PID using MRI alone. However, visualizing all implanted seeds on MRI, even on FST1-WI, is difficult. Detecting seed source positions is easier using CT, particularly using the seed-finder facility in Variseed 8.0. Thus, the CT-MRI fusion method was advocated [4,19].

In this study, the post-implant prostate volume was not large (mean, 21.6 ml; range, 13.1-32.8 ml) because of neoadjuvant hormonal therapy to adjust the prostate volume. In cases involving a large prostate, the activity per seed in accordance with prostate volume will be adequately selected. Our two-step image-fusion technique does not rely heavily on prostate volume; therefore, our results may be extrapolated to larger prostates. In this study, of the 636 seed sources, 186 (29%) were found to be overlapping and located in an extra-prostatic area. In cases that involve larger prostates with stronger activity per seed, fewer overlapping source images may be found than in this study.

One limitation of the present study was the assumption that since the images were acquired at the same time, the optimally matched sequence can be matched to other MRI sequences. Magnetic resonance imaging scans take several minutes to perform, and multiple sequences require at least 15 minutes of acquisition time, during which the bowel and bladder may displace the prostate from the initial scan position.

Another fundamental limitation of the present study was that the seed position was displaced by slice imaging (CT or MRI) in proportion to the slice thickness. In fact, when seed length and width variation were plotted against MRI sequence (Fig. 2), all imaging methods including CT led to an overestimation of the length and width compared with the actual parameters (4.55 × 0.97 mm). Due to the shift of every seed by CT and MRI, the seed center was shifted as well. With seed length consistently longer than the slice thickness, it creates uncertainty regarding the actual seed position. This uncertainty is unavoidable as long as seed position is obtained using slice imaging in proportion to the slice thickness.

Another limitation was the potential variation in the appearance of different source models, and these differences may affect the fusion accuracy of the two-step fusion technique. However, the ability of FST1-WI to more precisely define the sources can result in a better seed match. Additional measurements using different source models will need to be obtained in the future.

Conclusions

Computed tomography and T2-WI were fused more accurately using the two-step image-fusion technique than using direct image fusion. The two-step technique was performed by first fusing CT and FST1-WI by the least-squares method using each source center position, followed by fusion of the first-step image with a T2-WI.

Disclosure

Authors report no conflict of interest.

References


