Robot-Guided Open-Loop Insertion of Skew-Line Needle Arrangements for High Dose Rate Brachytherapy

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Abstract—We present a study in human-centered automation that has potential to reduce patient side effects from high dose rate brachytherapy (HDR-BT). To efficiently deliver radiation to the prostate while minimizing trauma to sensitive structures such as the penile bulb, we modified the Acubot-RND 7-axis robot to guide insertion of diamond-tip needles into desired skew-line geometric arrangements. We extend and integrate two algorithms: Needle Planning with Integer Programming (NPIP) and Inverse Planning with Integer Programming (IPIP) to compute skew-line needle and dose plans. We performed three physical experiments with anatomically-correct phantom models to study performance: two with the robot and one control experiment with an expert human physician (co-author Hsu) without the robot. All were able to achieve needle arrangements that meet the RTOG-0321 clinical dose objectives with zero trauma to the penile bulb. We analyze systematic and random errors in needle placement; total RMS error for the robot system operating without feedback ranged from 2.6 mm to 4.3 mm, which is comparable to the RMS error of 2.7 mm obtained in an earlier study for PPI-BT treatment using a robot with 3D ultrasound feedback.

Note to Practitioners Brachytherapy treats cancer by delivering radioactive sources proximal to cancer sites via needles. Current methods use standardized fixed mechanical templates that force needles into parallel arrangements that may prevent needles from reaching prostate volumes blocked by the pubic arch and often require needles to puncture sensitive organs. Skew-line (non-parallel) arrangements of needles can reach targets under the pubic arch and avoid sensitive organs. However these arrangements cannot be achieved with standard templates, motivating the use of automation. We present a human-centered automation system that integrates state-of-the-

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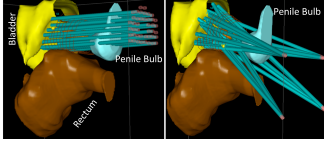


Fig. 1. Left: The current clinical approach to prostate high dose rate brachytherapy (HDR-BT) uses parallel needles guided by a mechanical template. This approach may prevent needles from reaching prostate volumes blocked by the pubic arch and often require needles to puncture sensitive organs (which can produce long-term side-effects). Right: Skew-line needle arrangements facilitated by robot guidance can avoid puncture by reaching under the pubic arch and can minimize trauma to sensitive organs such as the penile bulb which can produce side effects such as incontinence and impotence.

art needle and dose planning algorithms with a modified needle insertion robot. Results suggest that the robot can achieve precision and accuracy comparable to that of expert human physician. This approach has applications to brachytherapy treatment for other organs and to other needle procedures such as biopsy and anesthetic injection.

Index Terms—Brachytherapy, prostate cancer, needle insertion, robot, robot assisted surgery, health care, radiation, steerable needles

I. INTRODUCTION

Each year, over 500,000 cancer patients worldwide are treated with brachytherapy [1], where radioactive sources are placed inside the body close to cancerous tumors ("brachys": Greek for "proximal"). Brachytherapy is an effective treatment for cancers in the prostate, cervix, breast, and other anatomical sites [2]. We focus on prostate treatment, where current approaches often result in side-effects such as incontinence and impotence [3]–[5]. Most side-effects result from needle penetration through sensitive structures (urethra, bladder, rectum, penile bulb, cavernous veins, and neurovascular bundles) [5]–[9].

There are two forms of brachytherapy: prostate permanentseed implant (PPI) and high dose rate (HDR). In PPI-BT, needles implant radioactive seeds with a relatively short halflife (weeks) which are left in the patient after the procedure. In HDR-BT, multiple needles are inserted into the patient. After scanning and planning, a highly radioactive source is automatically moved through each needle using a remote afterloader. The dose distribution is controlled by source dwell times at pre-specified positions along the needles; the source is removed after treatment. This study focuses on HDR-BT.

In the current approach to prostate HDR-BT, hollow needles are inserted into the prostate through the perineum. The insertion is performed manually by the physician using real-time imaging using a trans-rectal ultrasound probe and a rigid template with parallel holes. As illustrated in Figure 1(left), the rigid template requires that all needles be parallel. This restriction often results in puncture of healthy organs such as the penile bulb and related vasculature, and can prevent access to some sections of the prostate due to pubic arch interference. Alternatively, skew-line (nonparallel, non-intersecting) needle arrangements as shown in Figure 1(right), can avoid puncturing delicate structures and be angled to reach under the pubic arch. Recently a "freehand" approach that does not require the template was proposed by physicians to allow skew-line needle arrangements [10]. However the freehand approach requires a high degree of skill and clinical proficiency. This paper explores the use of a robot to guide skew-line needle arrangements in HDR-BT.

In previous work, we developed the IPIP algorithm to compute HDR-BT dose plans [11] and the NPIP algorithm for computing skew-line needle arrangements [12]. In simulation we have shown that these algorithms can generate patient-specific skew-line needle arrangements that avoid sensitive organs and meet treatment dose objectives. This study integrates these planning algorithms with the Acubot-RND needle guiding robot [13] illustrated in Figure 2(left). Experiments suggest that a human-centered automation system can successfully implant skew-line needle arrangements that avoid puncturing non-prostate structures, meet clinical radiation dose objectives, with mean RMS error between planned and actual dwell points between 2-4 mm.

II. BACKGROUND AND RELATED WORK

Automation can benefit a variety of medical applications: surgical robotics [14], remote diagnosis [15], radiation biodosimetry [16],health analytics [17] and monitored anesthesia control [18]. Okamura et al [19] provides a detailed description of recent advances in medical and healthcare robotics.

The clinical HDR-BT workflow has six main steps: preimplant patient scanning, needle planning, needle insertion, post-implant patient scanning, dose planning, and dose delivery. Existing research has explored planning systems for computing optimal dose distributions for both PPI- and HDR-BT [11], [20]–[24]. Since the set of possible dose distributions depends on the implanted needle arrangement, planning systems like Prostate Implant Planning Engine for Radiotherapy (PIPER) [20] and Hybrid Inverse Planning and Optimization (HIPO) [24] incorporate the positioning of needles into their dose planning model. However, these approaches were developed for the standard parallel needle template, which has a smaller search space: fewer than 100

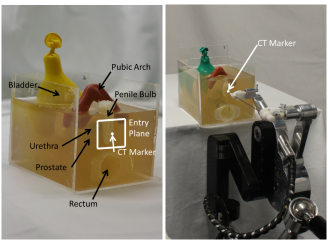


Fig. 3. Prostate phantom (5in x 5in x 8in) (left) and insertion setup (right). The anatomy modeled in the phantom includes: prostate; urethra; bladder; penile bulb; pubic arch; and rectum. A CT marker is centered on the square entry zone for calibration. As shown in the right image, the Acubot-RND is registered to the CT-marker.

candidate parallel needles in contrast to 200-300 candidates for skew-line needles.

In contrast to active needle steering using bevel-tips or cannuli [25]–[31], this study explores how a symmetric (diamond-tip) needle can be steered to a desired configuration within tissue by precisely positioning and orienting its primary axis outside the body.

Prior research in automated needle insertion has explored devices that address the clinical challenges of space constraints and safety requirements for needle insertion robots specially designed for prostate brachytherapy with transrectal ultrasound guidance [32]–[35]. Several of these devices can potentially insert skew-line needles, but they focus on PPI-BT and are not fully integrated with needle planners [36]–[40] The Acubot-RND was designed for PPI-BT and is operated by a manual joystick [13], [41], [42]. In this study, we modify the Acubot-RND with an interface to our needle planning software.

A recent study by Long et al [43], used the PROSPER image-guided robotic brachytherapy system [35] to perform multiple needle insertions into a gelatin phantom using intra-operative feedback from a 3-D ultrasound system. As noted in the Discussion section, we obtain similar error values without using ultrasound feedback.

The present study focuses on HDR-BT and integrates automated needle planning system with open-loop robot guided insertion using the Acubot-RND. The needle and dose planning systems are discussed in Section IV and the modifications to the Acubot-RND are discussed in Section V. This is a revised and expanded version of a paper presented at the IEEE International Conference on Automation Science and Engineering (CASE) [44]. This paper is rewritten throughout, with an expanded related work section and detailed analysis of random vs. systematic error.

III. PROBLEM STATEMENT

The RTOG-0321 clinical protocol [45] established recommendations for a set of dosimetric indices that are correlated

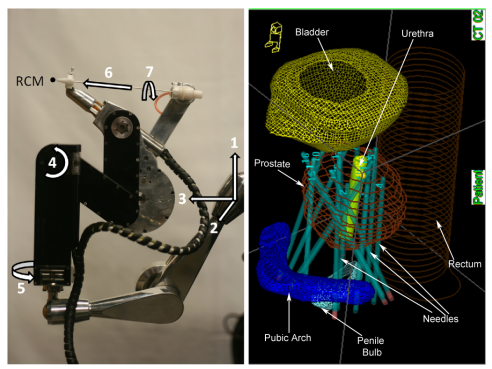


Fig. 2. The left figure shows the 7-DoF Acubot-RND robot used for this study. It has a 3-DoF Cartesian stage (1,2 and 3), a 2 DoF rotating center of motion (4 and 5), needle insertion (6) and needle rotation (7). The right figure shows a skew-line needle arrangement implanted by the robot system into a phantom as viewed after CT-Scan

with positive patient outcomes. In these indices, V_d^s , is the volume of structure s that receives at least d percent (eg., 75%, 100%, 150%) of a specified reference radiation dose (typically 950 cGy).

For the prostate, the value of $V_d^{\rm Prostate}$ is specified as a percentage of the total prostate volume, thus $V_{100}^{\rm Prostate}>=90\%$ specifies that at least 90% of the prostate volume should receive at least 100% of the specified reference radiation dose. For other structures such as the bladder, penile bulb, rectum, and urethra, $V_d^{\rm s}$ is specified in cubic centimeters, thus $V_{125}^{\rm Urethra}<=0.1$ cc specifies that no more than 1 cc of the urethra should receive more than 125% of the reference dose. The RTOG-0321 recommendations are summarized in the second column of Table I. Note that $V_{200}^{\rm Body}=0$ cc specifies that no non-organ volume of the body should receive 200% of reference radiation dose.

The treatment requires a sequence of steps: A 3D model of patient anatomy is obtained from a CT scan and manually segmented into organs. We then (1) plan a needle arrangement, if such exists, that lies within the workspace of the robot, avoids non-prostate organs/structures, and meets RTOG-0321 dose requirements, (2) transform this plan into a set of corresponding robot set-points so that each needle starting position and orientation guides a human novice who inserts needles to the indicated depth. (3) perform a second CT scan, compute a dose plan for the actual needle arrangement and report RTOG-0321 dose indices.

To quantify the damage to sensitive organs and structures, we propose a trauma metric equal to the total intersection volume: The trauma metric for structure s is:

$$T^s = \sum_k A_k L_k^s,$$

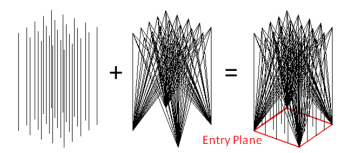


Fig. 4. The candidate needle set is the set of needles that are available during needle planning. As shown in the figure, the candidate needle set for this study consisted of: parallel lines, and skew-lines. The entry plane, which represents the bounded region on the perineum within which needles can enter the phantom is also depicted.

where A_k is the cross sectional area of needle k and L_k^s is the length of needle k puncturing structure s. The needles have a circular cross sections, hence, $A_k = \pi d^2/4$ in mm², where d is needle diameter.

IV. PLANNING SKEW-LINE NEEDLE ARRANGEMENTS AND DOSE DISTRIBUTIONS

To plan skew-line needle arrangements and dose plans, we modified the NPIP needle planning algorithm [12] to use a more comprehensive sample set of candidate needles and we incorporated it with the IPIP dose planning algorithm [11]. These references include details on these planners with experiments and sensitivity analysis.

NPIP accepts as input patient anatomy, the prostate target, obstacles such as the pubic arch and penile bulb, and the defined needle entry zone to search for an arrangement of skew-line needles that: (1) includes approximately 16 needles (the standard at the UCSF clinic), (2) avoids the pubic

arch bone and other sensitive organs, (3) offers dwell points that can deliver a dose plan that meets RTOG-0321 dose objectives, (4) minimizes for the trauma metric.

The planner uses integer programming: it is not complete (guaranteed to find such an arrangement if one exists) nor does it always produce an optimal solution. NPIP was modified to use non-uniform sampling to generate the candidate needle set and an additional constraint: all needles in the solution must have mutual clearance of $\gamma.$ The parameter γ specifies the distance between the medial axes of a pair of needles. For a non intersecting needle pair, $\gamma \geq d,$ where d is the needle diameter. We chose a conservative value of $\gamma=2d$ to allow for deviations during insertion.

The prostate volume is discretized into a rectangular grid of sample points, with spacing of 4 mm in the x- and ydirections and 3 mm in the z-direction (the inter-plane CT sample distance). This produced approximately 1000 points for each case. NPIP takes as input this set of sample points and a user-specified parameter, δ . NPIP generates a candidate needle set (line segments) and searches for a subset of these candidate needles where every point within the prostate is within δ of at least one needle. A high value of δ allows needles to cover more volume, producing needle arrangements with fewer needles. To normalize across prostate volume, we set $\delta = 33\%$ of the radius of a sphere with equivalent volume to the prostate and iteratively increased or decreased it to obtain a solution with 16 needles. NPIP uses heuristics to solve an integer program so there are no time or performance guarantees, but for the cases we considered, NPIP computes solutions within 120 seconds (see Experiments section).

The needle arrangements computed by NPIP are given as input to the Inverse Planning by Integer Program (IPIP) dose planning algorithm [11]. Given the set of needles, IPIP computes a set of dwell times (spaced 5 mm apart within each needle) for the radioactive source that maximize $V_{100}^{\rm Prostate}$ subject to the RTOG-0321 dose requirements. For the three phantom cases we studied, IPIP found solutions within 10 seconds with values as reported in Table I.

V. THE ACUBOT-RND ROBOT

The Acubot-RND robot system was designed and constructed at at Johns Hopkins University to guide needle insertion for permanent-seed (PPI-BT) treatment [13]. Hardware specifications for the Acubot-RND, including spatial resolutions and maximum ranges for each degree of freedom are reported in Fichtinger et al [34].

A. Robot guided needle insertion

As shown in Figure 2 (left), the Acubot-RND is a 7-DoF robot with three stages: The first is the 3-DoF Cartesian Positioning Stage (CPS), The second is the 2-DoF Rotating Center of Motion (RCM) that sets needle angle keeping the needle tip position fixed, and the third is the 2-DoF Rotating Needle Driver Module (RND) that can rotate and insert needles automatically.

The phantom is draped during the experiments. For this study we position the first stage manually during calibration

and we send computed commands to the second stage to orient the needle prior to insertion. We then send a command to the third stage to insert the needle to a pre-specified end point without feedback. At this point a human novice (co-author Garg, an IEOR graduate student with no clinical experience) manually retracts each needle leaving behind a stylet in tissue.

B. Digital Interface

The needle entry plane with CT marker defines the coordinate frame. We modified the Acubot-RND, augmenting the manual joystick operation with a digital interface that allows commanding specific offsets in tip position from the center of the entry zone, and specific pairs of angular offsets from the normal to the plane of needle entry zone.

A needle plan defines a set of i needles, each specified with two points: \underline{p}_0^i in the entry plane, and \underline{p}_1^i at the desired distal tip of the inserted needle, where x and y components of \underline{p}^i span the entry plane in horizontal and vertical directions; and the z component points into the phantom volume. The insertion depth for needle i is d_i , the Euclidean distance between the points. The angles for angle needle i, defined as rotations in the associated planes, are:

$$\theta_{xz} = atan2 \ (x_1 - x_0, z_1 - z_0)$$

$$\theta_{yz} = atan2 \ (y_1 - y_0, z_1 - z_0)$$

These angles are specified as joint angles for the RCM.

VI. PHYSICAL EXPERIMENTS

To evaluate the performance of the NPIP and IPIP algorithms and robot hardware, we constructed three nearly identical physical phantoms in the clinic at UCSF: Ph1, Ph2 and Ph3. Each includes anatomically-correct organ structures of similar density as human tissue and suspended in a translucent gelatin medium. Harder bone structures like that of the pubic arch is constructed from modeling clay. The organ structures include urethra, prostate, bladder, penile bulb, pubic arch and rectum as shown in figure 3. The square entry zone has dimension 45 mm, consistent with clinical practice as shown in Figure 3, relative to an example candidate needle set in Figure 4.

We performed end-to-end needle insertion procedures with 16 needles on each phantom using the robot for the first two(Ph1 and Ph2) and an expert human physician for the third phantom (Ph3).

Each experiment includes these steps (with step 2 omitted for the expert human physician who used his clinical intuition to determine a needle plan):

- 1) Perform first CT-Scan and 3D segmentation of organs.
- 2) Plan desired Needle configuration using NPIP and calculate dose plan IPIP.
- 3) Implant Needles with robot or with expert human.
- 4) Perform second CT-scan of phantom with needles.
- 5) Perform dose planning using IPIP.

A. Robot Experiments

A side view of an implanted phantom Ph1 is shown with needle configuration A1 in Figure 6. Robot-assisted implant of needles was performed on two phantoms, Ph1 and Ph2. The needle entry zone is a square on the surface of the phantom centered on the CT marker. As in typical clinical cases, the entry zone is $45~\text{mm} \times 45~\text{mm}$ as shown in figures 3 and 4. We place a radio-opaque CT-marker at the center of each entry zone to register the coordinate system of the planning algorithm with the robot.

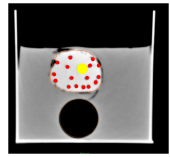
1) Pre-Implant Scanning and Planning: CT scans of tissue phantoms, before and after all 16 needles are inserted, were taken in 3 mm thick slices. The contoured prostate volumes for the three phantoms were 39 cc, 32 cc, and 37 cc. The total phantom volume was 750 cc.

The organs of the phantom and the CT marker were contoured in 3D using the Nucletron Oncentra® Dynamic Planning Environment. Using Oncentra, we added a 2 mm margin to the outer contour of the penile bulb. These 3D organ models were exported to NPIP and IPIP. A reference dose of 950 cGy is commonly prescribed for prostate HDR-BT; we used this level as reference in all cases.

For Ph1 and Ph2, there were 287 and 229 candidate needles respectively. NPIP used a δ value of 6.5 mm for Ph1 and 6.0 mm for Ph2 to produce solutions with 16 needles. γ value was chosen to be twice the needle diameter, 4 mm. For Ph1 and Ph2, we define two needle arrangements the planned needle arrangements, P1 and P2, and the actual needle arrangements, A1 and A2.

All computation was performed using Matlab R2011a on a Lenovo ThinkPad with an Intel i5-2410M processor and 4GB of RAM. The integer program optimization was done using the Matlab interface for the Mosek Optimization Toolbox v.6. The complete run for planning using NPIP less than 70 seconds for both Ph1 and Ph1; and IPIP runs took 10 seconds for both Ph1 and Ph2.

2) Robot Experiments on Ph1 and Ph2: After the initial CT scan, the robot and phantom are clamped to a worktable, leveled, and manually calibrated as follows (1) the robot is manually moved to an initial state with first needle tip at the registration mark and aligned normal to the entry plane by moving to specified x and y offsets and confirming that it just touches the surface at each point. Figure 3 shows the Acubot-RND and phantom in such an initial state. We used a standard 18-gauge diamond-tip brachytherapy needle (COOK Biotech) of length 15 cm and 2 mm diameter hollow sheath that housed a rigid stylet. To implant needle arrangements in Ph1 and Ph2, the Acubot-RND was brought into each specified position and orientation where a needle was inserted by the robot until the pre-specified depth in phantom tissue. The insertion depth was marked on a stylet and it was manually pushed through the hollow needle in the phantom by the novice operator, and needle is retracted to leave the stylet in the phantom. The stylets were used as a proxy for needles in phantom to minimize interference to robot during subsequent needle insertions.



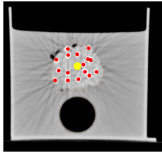


Fig. 5. Cross-sectional view of an actual needle arrangement inserted by an expert human physician without the robot (left) and one inserted by a novice human guided by the robot (right). Both are considered successful as they meet the RTOG dose objectives without penetrating the penile bulb.

- 3) Expert Human Physician Experiment on Ph3: Coauthor Dr. I-Chow Hsu is a certified radiation oncologist at UCSF with a specialization in brachytherapy and over 18 years of clinical experience. He performed insertion on Ph3 for comparison. We performed a CT scan of Ph3 as above. Dr. Hsu used his expert intuition to determine a needle plan. He inserted 16 standard HDR-BT needles into phantom Ph3 under trans-rectal ultrasound (TRUS) guidance using the UCSF-developed "freehand" technique [10]. A HAWK 2102 EXL TRUS system from B-K Medical was used for ultrasound imaging.
- 4) Post-Implant CT Scan: After executing all implants, another CT scan is performed on the phantom. The needles are segmented and organs are contoured to determine the needle configuration actually implanted, A_i .

VII. RESULTS

The RTOG-0321 clinical requirements and results from all three experiments, planned and actual for the robot, and actual for the human, are summarized in Table I. For all 3 cases, clinical requirements were met and performance with the robot was comparable to that of an expert human physician.

The expert human physician experiment was completed in under 15 minutes. Each robot experiment required approximately 45 minutes due to calibration and slow needle insertion speeds by the novice. We also note that the expert human physician had the benefit of ultrasound feedback while the needle insertions for the robot experiments were performed without ultrasound or visual feedback.

Figure 5 shows cross-section of the needle arrangements implanted by the expert (left) and by a novice with the robot guide (right).

Table I lists the clinical dose index and trauma metrics, the The difference between the values obtained from planned vs actual needle arrangements are relatively minor. An exception is the difference in $V_{75}^{\rm Bladder}$ values for P2 and A2 which were 0.3 cc and 0.8 cc, respectively. They are both below the clinically acceptable limit for this criterion: 1 cc. This discrepancy is due to some needles not being inserted far enough into the prostate. This is mainly due to placement error in manual step of the needle insertion. Since no dwell positions are available at the apex of the prostate, IPIP increases the dwell times at the distal ends of the needles

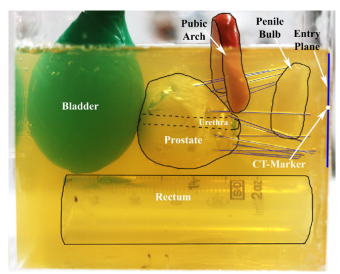


Fig. 6. Anatomically-correct phantom Ph1 with robot-implanted needle configuration A1. The organ boundaries and actual needles positions are highlighted. All sensitive structures were spared needle puncture.

to achieve coverage, but this produces a higher-than-desired dose to the bladder.

Although actual needles could puncture the penile bulb due to placement error, the puncture volume in all planned and actual cases, for the robot and the human, was zero (0 cc). Also, no needles intersected the pubic arch.

Robot Placement Error

We next consider the total, systematic, and random errors between planned and actual needle arrangements in the two robot experiments (there is no planned needle arrangement for the third experiment). We sample needle position at 1 millimeter intervals producing 60 sample points per needle. We use same sampling procedure for planned and actual needle configurations. Hence using all needles in the arrangement we generate two sets of corresponding points: a set of planned points P and set of actual points A.

The total error in mm between any pair of planned and actual points is the distance between them. Table II summarizes mean, min, and max RMS error (RMSE) along each dimension and d, the Euclidean distance. The total placement error is the RMS distance over all planned and implanted needle points. For Ph1 and Ph2, the total RMS errors were $2.6 \, \text{mm}$ and $4.3 \, \text{mm}$ respectively.

We decompose total error into systematic and random components by computing the least-squares rigid transformation between the pairs of point sets [46]. Specifically, we compute the rotation matrix, R, and the translation vector, T, which minimizes the least-squares error over the whole point set,

$$\sum \|P - (RA + T)\|_2,$$

where P is the vector of planned points and A is the vector of actual points. The associated translations and rotation values define the systematic error. The α , β , and γ values are the rotations in the Euler angles reported in degrees. The Euler angles are computed as:

		Phantom 1		Phantom 2		Phantom 3	
Metric	RTOG Req.	P1	A1	P2	A2	A3	
V ₁₀₀ ^{Prostate}	$\geq 90\%$	99.0	97.0	96.0	96.0	98.0	
$V_{150}^{Prostate}$	$\leq 45\%$	39.0	40.0	40.0	37.0	37.0	
$V_{75}^{Bladder}$	$\leq 1 \text{ cc}$	0.00	0.00	0.30	0.80	0.30	
$V_{100}^{\rm Bladder}$	= 0 cc	0.00	0.00	0.00	0.00	0.00	
$ m V^{Bulb}_{75}$	$\leq 1 \text{ cc}$	0.00	0.00	0.00	0.00	0.00	
$ m V_{100}^{Bulb}$	= 0 cc	0.00	0.00	0.00	0.00	0.00	
$V_{75}^{ m Rectum}$	$\leq 1 \text{ cc}$	0.06	0.00	0.00	0.00	0.00	
$V_{100}^{ m Rectum}$	= 0 cc	0.00	0.00	0.00	0.00	0.00	
$V_{125}^{Urethra}$	$\leq 0.1 \text{ cc}$	0.06	0.05	0.04	0.06	0.07	
$V_{150}^{Urethra}$	= 0 cc	0.00	0.00	0.00	0.00	0.00	
V_{200}^{Body}	= 0 cc	0.00	0.00	0.00	0.00	0.00	
$T^{\overline{\mathrm{Bulb}}}$	min	0.00	0.00	0.00	0.00	0.00	

TABLE I

THIS TABLE LISTS THE CLINICAL DOSE INDEX AND TRAUMA METRICS,
THE RTOG-0321 REQUIREMENTS, AND THE VALUES FROM EACH
EXPERIMENT, PH1 AND PH2 USING THE ROBOT, AND PH3 BY AN EXPERT
HUMAN PHYSICIAN. COLUMNS P1 AND A1 ARE THE DOSE VALUES
ACHIEVED BY IPIP FOR THE PLANNED AND ACTUAL NEEDLE
ARRANGEMENTS RESPECTIVELY FOR PH1. THE SAME FOR P2, A2, AND
PH2. A3 FOR THE THIRD PHANTOM PH3 IS BASED ON THE NEEDLES AS
ACTUALLY IMPLANTED BY THE EXPERT HUMAN PHYSICIAN (WHO DID
NOT PLAN A NEEDLE ARRANGEMENT)

$$\alpha = \sin^{-1}(r_{1,3}), \beta = \cos^{-1}\left(\frac{r_{1,1}}{\cos(\alpha)}\right), \gamma = \cos^{-1}\left(\frac{r_{3,3}}{\cos(\alpha)}\right),$$

where $r_{i,j}$ is the element of R in the ith row and the jth column. The errors for Phantom 1 and Phantom 2 are shown in Table II.

The random error is the residual error after the actual points are compensated by the least square transformation. Note that systematic and random components do not sum to the total error due to rotations.

Total random error for Ph1 and Ph2 are 1.4 mm and 2.4 mm respectively. Table II summarizes the results.

The superposition of the planned (blue) and implanted (red) needles is shown in Figure 7, as well as the planned and adjusted needle arrangements (green).

VIII. DISCUSSION AND FUTURE WORK

This paper describes the system architecture, algorithms, hardware, and experiments with a human-centered automation system for inserting skew-line needle arrangements for HDR-BT. We report results with an open-loop robot guide system that uses CT scans before insertion and does not use sensor feedback during insertion, and results from an experiment performed by an expert human physician using ultrasound guidance. These results, in a controlled experimental setup with phantom tissues, suggest that skew-line needle arrangements can be planned and executed with a robot guide to achieve the RTOG-0321 clinical treatment

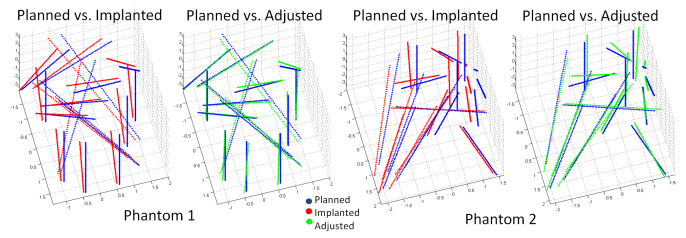


Fig. 7. Superposition of planned (blue) and implanted (red) needle arrangement for Phantom 1 and Phantom 2. Although no sensitive structure was punctured in the implanted needle arrangement and all dose objectives were met, there was non-zero placement error. The placement error was separated into systematic and random error. Upon compensation for the systematic error, the adjusted needle arrangement (green) fits better to the planned configuration.

	Phantom 1			Phantom 1			
Total RMS	Min	Max	Mean	Min	Max	Mean	
X	0.5	2.5	1.4	0.8	2.7	1.9	
У	0.2	2.5	1.6	0.7	3.2	2.3	
z	0.1	3.0	1.5	0.9	5.3	3.1	
d	1.3	4.1	2.6	2.0	6.3	4.3	
Random							
X	0.0	0.8	0.5	0.1	2.3	1.2	
У	0.1	1.1	0.5	0.1	1.8	1.1	
Z	0.1	2.3	1.2	0.0	5.1	1.8	
d	0.2	2.5	1.4	0.8	5.2	2.4	
Systematic							
ϵ_x	1.2			0.9			
ϵ_y	1.4			2.2			
ϵ_z	0.8			2.4			
α	1.8			1.3			
β	-0.9			0.8			
γ	1.6			3.5			

TABLE II

ERROR ANALYSIS: TOTAL ERRORS ARE RMS ERRORS (IN MM) MEASURED IN PHANTOMS POST-IMPLANT. RANDOM ERRORS ARE RMS ERRORS (IN MM) AFTER COMPENSATION FOR SYSTEMATIC ERROR. THE x-,y- and z- rows list RMS errors in each direction. d is the overall RMS error. Systematic errors are obtained by least square point set matching. (ϵ_i in mm and angles in degrees)

objectives while avoiding puncture of sensitive structures such as the penile bulb.

Long et al, 2012 [43] used the PROSPER robot system (developed for PPI-BT), to insert glass bead markers into a gelatin prostate phantom. After an initial insertion, the needle tip and target bead were measured using 3D ultrasound and needle tip was adjusted along the insertion axis until error was minimized. Using such intra-operative feedback, the PROSPER system achieved position errors of 2.7 mm. This error, between needle tips and target points, is relevant for PPI-BT. For HDR-BT, we report RMS error along the entire needle which contains dwell positions. We were able

to achieve RMS errors of 2.6 mm and 4.3 mm, which are comparable to the error achieved in the closed-loop PROSPER system. In future work, we will perform additional experiments with more complex anatomy, for example enlarged prostates where it may be difficult to avoid pubic arch interference and to treat cancers in other organs. We will study how NPIP and IPIP may be enhanced with higher-resolution sampling, where cloud computing may make it feasible to compute plans that are more robust to uncertainty in anatomy and needle motion.

We will also explore how calibration can be enhanced with additional CT markers to reduce systematic error and perform experiments to explore how needle insertion order and needle rotation (rifling) may affect needle insertion accuracy. We will also explore how feedback control can be used during insertion.

Some studies like [47], [48] have explored use of MRI for real-time scanning. Tovar-Arriaga et al [49] and Ji [50] proposed workflows for needle insertion using CT and MRI feedback respectively. [51], [52] have studied accuracy of needle placements in real-time MRI tracking. Real-time feedback from either CT or MRI has to deal with trade-of between spatial resolution and temporal resolution. CT can be used for feedback, but it results in radiation exposure to patient. MRI (magnetic resonance imaging) is relatively slow, requires that all needles and guiding equipment be nonferrous, and has issues with image warping in larger imaging volumes. As Ultrasound is safe and and prvides real-time imaging, we will explore how it can be incorporated for active needle guidance.

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