A simple technique for the generation of institution-specific nomograms for permanent prostate cancer brachytherapy

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Abstract

Purpose: Nomograms once had a vital role in prostate brachytherapy practice. Although some of their functions have been assumed by computerized dosimetry, many programs still find them useful to determine the number and strength of seeds to be ordered in advance of the implant. As it has been demonstrated that brachytherapists differ in their implant practices and preferences (in regard to seed distribution and total implanted activity), we propose a simple technique for generating institution-specific nomograms.

Material and methods: Using the data generated by previous implants, we demonstrate a simple technique, utilizing ubiquitous software, for generating nomograms to predict seed number, strength, and total implant activity.

Results: Despite wide variations in the degree of post-implant swelling (+12% to +56%), nine of the first ten implants using the generated nomogram have met all target dose parameters.

Conclusions: It is appropriate for each institution to develop nomograms specific to their prostate brachytherapy technique and constraints. Nomograms can estimate the number and strength of seeds to be ordered, and may partially compensate for the effects of implant-induced swelling.

Key words: brachytherapy, medical dosimetry, nomograms, prostate cancer.

Purpose

Mathematical models (nomograms [1-6], nomographs [7], algorithms [8], reference [9] or look-up tables [10,11], and tie-line charts [12]) have long been used as aids for prostate brachytherapy. They are formulations to ‘pre-determine the total required isotope activity for performing transperineal interstitial permanent prostate brachytherapy’ from the pre-implant prostate volume [1]. They can be derived from statistical analysis of clinical data [3-5,10], or based upon a theoretical model of the activity needed to subsume a spheroidal volume with a prescription dose cloud [1,6-8,11,12]. Although their role has diminished in the era of computerized dosimetry [13,14], nomograms are still used as an independent check of pre-plans, and some brachytherapists utilizing intra-operative planning rely on them to determine the number and strength of seeds in their pre-implant seed order.

As implant preferences (prescription dose, seed distribution pattern, extra-capsular seed placement, organ constraints) differ among institutions, it has been recommended that each institution derive nomograms reflective of their program’s implant philosophy [3,15]. We relate a simple method, utilizing ubiquitous spreadsheet software, to generate institution-specific nomograms for the calculation of seed number and strength for permanent prostate brachytherapy.

Material and methods

The process presented here relies on the analysis of previous implants. For purposes of illustration, we present the process we used to generate a nomogram based on the first 40 consecutive Cs-131 (131Cs) implants performed when our program transitioned to caesium sources (this study, including review of patient records, has been approved by our institution’s review board). The implants had been pre-planned, without intentional placement of extracapsular seeds. Post-implant dosimetry was based upon CT scans performed two weeks after implantation.

To generate a nomogram by this process, you will need:

1. Spreadsheet software; we used Excel 2011 for Mac, 14.1 (Microsoft Corporation, Redmond, WA, USA).
2. Post-implant dosimetry for 30 or more prostate implants, that had been analyzed by prostate brachytherapy dosimetry.
Table 1. The post-implant dosimetric outcomes

|        | Range       | Median | Standard deviation |
|--------|-------------|--------|--------------------|
| D90    | 101-124%    | 112%   | 7.4                |
| V150   | 42-70%      | 52%    | 9.0                |
| R100   | 0.00-0.78 mL| 0.05 mL| 0.29 mL            |

Table 1. The post-implant dosimetric outcomes

The process.

1. Generate a spreadsheet that contains data from patient records and post-implant dosimetry. Column headings are:
   - a) case identifier;
   - b) pre-implant prostate volume;
   - c) number of seeds implanted;
   - d) activity per seed;
   - e) ‘actual’ total implant activity (the product of columns ‘c’ and ‘d’);
   - f) ‘optimal’ total implant activity (see step 2).

2. Define prescription dose and constraints. Ours were:
   - prescription dose: 110 Gy for monotherapy, 70 Gy for partial implants;
   - constraints: a) D90 range: 100-120% of prescription dose; b) optimal D90: midpoint of the D90 range (in our case, 110% of prescription dose; for monotherapy, 121 Gy); c) V150 constraint: ≤ 60% of prostate; d) V100rectum constraint: < 0.75 mL.

3. Using the spreadsheet software, generate Cartesian (scatter) plots with ‘optimal’ total implant activity (Y axis) as a function of ‘prostate volume’ (X axis). Fit to a linear or power regression. Display the equation and R2.

4. Using the spreadsheet software, generate Cartesian (scatter) plots with ‘optimal’ total activity (Y axis) as a function of ‘prostate volume’ (X axis). Fit to a linear or power regression. Display the equation and R2.

5. You now have equations that can predict the appropriate number of seeds and total implant activity for an implant of a prostate of a given volume. Or, you can generate a ‘look-up table’. With the equations generated in steps 3 and 4, create a look-up table for the range of volumes likely to be implanted (say, 20-60 mL).

The columns being:

- a) prostate volume (every integer in the range of volumes);
- b) number of seeds (solve for Y, using the equation generated in step 2, where X is the prostate volume);
- c) ‘optimal’ total implant activity (solve for Y, using the equation generated in step 3, where X is the prostate volume);
- d) activity per seed (divide column c by column b).

It is advisable to order a few extra seeds to compensate for seed loss or the discovery of a larger than anticipated prostate volume at time of implantation. The nomogram can assist in making appropriate adjustments in total implant activity in the operating theatre if the prostate volume is determined to be different than measured preoperatively (e.g., due to the action of hormonal therapy).

Results

Our first 10 131Cs implants guided by the nomogram generated by this method were analyzed. The post-implant dosimetric outcomes are summarized in Table 1.

Not surprisingly, the degree of post-implant swelling, expressed as the ratio of post-implant prostate volume (on CT performed 2 weeks after implantation) to pre-implant prostate volume (measured by ultrasound) greatly influenced post-implant dosimetry metrics. The median ratio was 1.28, but the range was 1.12-1.56. The implant with the greatest degree of swelling (ratio 1.56) had the ‘coolest’ D90 (101%) and V150 (42%). The implant with the least swelling (ratio 1.12) had the ‘warmest’ D90 (124%) and V150 (70%); the latter was the only implant in which constraints were not met.

The volume of tissue encompassed by the prescription dose on the post-implant CT scans ranged from 1.46 to 1.69 times the pre-implant prostate volume (mean: 1.55, standard deviation: 0.08).

Discussion

When the clinicians and physicists at New York’s Mt. Sinai Hospital began their prostate implant program (1990), they adopted the Memorial Hospital nomogram to determine the total activity to implant for a given prostate volume. It became apparent, however, that the nomogram developed for Memorial’s homogenous seed distribution was inappropriate for Sinai’s peripherally-loaded implants. In a series of 5 iterations over 5 years, the Sinai program developed an institution-specific nomogram to satisfy their needs [16]. When our institution switched from 125I to 131Cs seeds, we relied on guidelines that had been developed elsewhere. We, too, were disappointed with our dosimetric outcomes and endeavored to develop a nomogram that reflected our implant pattern and dosimetric constraints.

Nomograms had once been used to determine total implant activity, seed number, and spacing [7]. We believe that computer planning has reduced, but not eliminated the utility of nomograms. Nomograms can be used for quality control and to maintain consistency among implant plans. The VariSeed software utilizes a ‘Nomogram Planning Module’ to generate a complete treatment plan (including total activity, needle, and source positions), but relies on a nomogram entered by the operator. It may be
unsatisfactory to adopt a ‘generic’ nomogram obtained from medical literature or provided by a seed distributor, as it has been demonstrated that experienced brachytherapists espousing the same implant philosophy and prescribing to the same dose will implant the same volume with a different number of seeds and total activity [17]. Indeed, the variation in total implant activity can exceed 40% [15]. It has therefore been recommended that a nomogram be generated to reflect an institution’s own practices [3,15].

It has long been appreciated that the edema engendered by prostate implantation can degrade implant quality [18-20]. If the magnitude of swelling could be predicted, it would be a simple matter to compensate by a reciprocal increase in implant strength; but the magnitude is highly variable and unpredictable [18-20]. Two strategies have been commonly used to reduce the impact of swelling on implant quality:

1. Devising a treatment plan in which the target volume encompasses the prostate with a several millimeter margin [21]. Use of our nomogram has resulted in the delivery of prescription dose to a volume 47-69% greater than the pre-implant prostate volume (coinciding with previously observed day-1 swelling), without exceeding normal tissue constraints or placement of extra-capsular seeds.

2. Utilizing intra-operative treatment planning with computer optimization [13]. Unless the brachytherapist orders seeds in bulk (impractical for small programs and those utilizing short half-life radionuclides), a method is required for predicting the number and strength seeds that will be used.

It has been proposed that nomograms can be used to at least partially compensate for implant-induced edema [22]; our experience suggests this to be so. The technique we have proposed for nomogram generation utilizes ubiquitous software; it has been utilized by one of the authors (JNA) for over a decade. His nomogram has been revised whenever his technique was modified or a new radionuclide was adopted (the process facilitates changing of radionuclides, as the post-implant CT scans performed after implantation with the ‘old’ radionuclide can be used to generate a nomogram for the ‘new’ one).

Nomograms can be based on either linear or power equations. Historically, power equations have been most often used, but it had previously been demonstrated that, for the range of volumes typically implanted, a power curve is almost ‘linear’ [23]; our current experience confirms this (Fig. 1). Indeed, the calculated difference between our linear and power equations was less than one seed for prostate volumes between 25 and 45 mL (data not shown). We suggest that practitioners generate both power and linear equations, and use the equation that has the better fit (higher \( R^2 \)). We wish to stress that nomograms may have a supplementary role in treatment planning, and do not diminish the centrality of computer dosimetry in brachytherapy planning.

Conclusions

Computer dosimetry has diminished, but not eliminated the role of nomograms in prostate brachytherapy. They remain useful for estimating the total activity and number of sources to be implanted. Our experience suggests that they can partially compensate for the unpredictable degree of post-implant edema. Brachytherapists should use nomograms that reflect their own implant practices, tissue constraints, and philosophy. We have demonstrated a simple method for the generation of institution-specific nomograms for permanent prostate brachytherapy.

Disclosure

Authors report no conflict of interest.

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