Relationship between two year PSA nadir and biochemical recurrence in prostate cancer patients treated with iodine-125 brachytherapy*

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Abstract Objective: To evaluate the relationship between two year PSA nadir (PSAn) after brachytherapy and biochemical recurrence rates in prostate cancer patients.

Materials and Methods: In the period from January 1998 to August 2007, 120 patients were treated with iodine-125 brachytherapy alone. The results analysis was based on the definition of biochemical recurrence according to the Phoenix Consensus.

Results: Biochemical control was observed in 86 patients (71.7%), and biochemical recurrence, in 34 (28.3%). Mean PSAn was 0.53 ng/ml. The mean follow-up was 98 months. The patients were divided into two groups: group 1, with two year PSAn ≥ 0.5 ng/ml after brachytherapy (74 patients; 61.7%), and group 2, with two year PSAn ≥ 0.5 ng/ml after brachytherapy (46 patients; 38.3%). Group 1 presented biochemical recurrence in 15 patients (20.3%), and group 2, in 19 patients (43.2%) (p < 0.02). The analysis of biochemical disease-free survival at seven years, stratified by the two groups, showed values of 80% and 64% (p < 0.02), respectively.

Conclusion: Levels of two year PSAn ≥ 0.5 ng/ml after brachytherapy are strongly correlated with a poor prognosis. This fact may help to identify patients at risk for disease recurrence.

Keywords: PSA nadir; Brachytherapy; Iodine radioisotopes; Prostatic neoplasms.

Resumo Objetivo: Avaliar os valores de PSA nadir (PSAn) dois anos após braquiterapia e sua relação com a taxa de recidiva bioquímica.

Materiais e Métodos: Cento e vinte pacientes foram tratados no período de janeiro de 1998 a agosto de 2007. Todos os pacientes foram submetidos a braquiterapia isolada. Os resultados foram analisados com base na definição de recidiva bioquímica do Consenso Phoenix.

Resultados: Oitenta e seis pacientes (71,7%) tiveram controle bioquímico e 34 (28,3%) apresentaram recidiva bioquímica. O PSAn médio foi 0,53 ng/ml. O seguimento médio foi 98 meses. Os pacientes foram divididos em dois grupos: grupo 1, com PSAn ≥ 0,5 ng/ml dois anos após a braquiterapia (74 pacientes; 61,7%), e grupo 2, com PSAn ≥ 0,5 ng/ml dois anos após a braquiterapia (46 pacientes; 38,3%). O grupo 1 teve recidiva bioquímica em 15 pacientes (20,3%) e o grupo 2 em 19 pacientes (43,2%) (p < 0,02). A análise da sobrevida livre de doença bioquímica de sete anos, estratificada pelos dois grupos, apresentou valores de 80% e 64% (p < 0,02), respectivamente.

Conclusão: Os níveis de PSAn ≥ 0,5 ng/ml dois anos após a braquiterapia estão fortemente correlacionados com mau prognóstico. Este fato pode ajudar a identificar pacientes com risco de recidiva da doença.

Unitermos: PSA nadir; Braquiterapia; Radioisótopos de iodo; Neoplasia de próstata.

INTRODUCTION

According to the World Health Organization, prostate cancer is the second type of cancer in incidence in men, and also the sixth most frequent cause of death in the male population worldwide¹. In Brazil, 68,880 new cases of prostate cancer were estimated for 2014. Such number of cases corresponds to an estimated risk of 70.42 new cases per 100,000 men².

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Radiotherapy is utilized in the treatment of prostate cancer either as exclusive or adjuvant therapy, or after biochemical recurrence, therefore it represents an important oncologic treatment technique(3-5). Prostatic specific antigen (PSA) test is the main method for detection of biochemical recurrence of prostate cancer. Finding a prognosis marker to identify men at high risk for treatment failure and subsequent death could be useful to select those patients who might benefit from early rescue therapy. Several PSA variables are tested in order to predict biochemical control, including initial PSA (PSAi) and PSA nadir (PSAn)(6).

After radiotherapy, PSA levels gradually decline after a period of 18–24 months. Fluctuations of PSA are not infrequent, with transient increases and subsequent drops called bounces(7). The PSAn is defined as the lowest PSA value measured after utilization of any treatment method. The time required to reach PSAn is different for radical prostatectomy and for external radiotherapy. After surgery, PSA levels decline is immediate, dropping to undetectable levels soon after the intervention, while after radiotherapy the decrease is slow, taking up to 27 months to reach PSAn. Patients who reach PSAn within the first 12 months after radiotherapy present with worse prognosis, with a relative risk for disease recurrence 18 times higher than that observed in those patients whose PSAn occurs two years after such a treatment(8).

Brachytherapy has increasingly been utilized, because of the technological developments in transrectal ultrasonography, availability of radioactive iodine sources adapted for implantation in the prostate gland, and the development of computer-aided planning systems for such therapeutic method(9,10). It is an outpatient procedure, with low morbidity rates, good conformity to the target lesion, excellent dose distribution and better quality of life for the patient(9,10). In the USA, 36% of radiotherapy treatments for prostate cancer consist of brachytherapy(11). The initial success of such a treatment has led to a dramatic increase in its utilization in patients with focal prostate cancer worldwide(12). The advantages of brachytherapy include its conformal property; concentration of high radiation doses within the target volume; and rapid dose decline in adjacent structures such as the bladder and rectum(10,13).

Several studies have demonstrated the relationship between PSAn after definitive radiotherapy and biochemical disease-free survival (BDFS)(14-16). Although most patients experience a decrease in PSA levels after definitive radiotherapy, only few of them reach undetectable PSA levels(16). Several authors have proposed a cut-off point for various PSAn levels, ranging from 0.2 to 4.0 ng/ml, as a predictive factor for biochemical recurrence in patients submitted to radiotherapy, brachytherapy or a combination of both treatments, either with or without androgen suppression(6,15,17,18).

The present study was aimed at evaluating the relationship between PSAn values two years after iodine-125 brachytherapy and chances of biochemical prostate cancer recurrence.

MATERIALS AND METHODS

The study was duly approved by the Committee for Ethics in Research of Hospital Universitário Clementino Fraga Filho – Universidade Federal do Rio de Janeiro, Brazil.

From January 1998 to August 2007, 120 consecutive prostate cancer patients (T1-T2) referred by different urologists were submitted to isolated brachytherapy. All the patients underwent follow-up from January 2001 to December 2012. The patients were classified into three categories of recurrence risk groups, also utilized by D’Amico(19), based on prognostic factors such as PSAi, Gleason score and clinical stage, which evaluate the risk for biochemical recurrence after any type of treatment:

- Low risk: PSAi < 10.0 ng/ml, Gleason ≤ 6 and clinical stage = T1b-T2a.
- Intermediate risk: PSAi between 10.1 and 19.9 ng/ml or Gleason = 7 or clinical stage = T2b.
- High risk: PSAi ≥ 20.0 ng/ml or Gleason ≥ 8 or clinical stage ≥ T2c.

PSAi is defined as the last recorded PSA value before the beginning of any treatment. Those patients with a probable life expectancy of less than five years; those with previous transurethral resection causing major deformity of the prostatic urethra; and those with high surgical risk and distant metastases were excluded from the study as per contraindications proposed by the American Brachytherapy Society(20).

All the patients were treated with isolated brachytherapy. Neoadjuvant hormone therapy with one or more luteinizing hormone-release agonists and oral anti-androgens were prescribed by the assisting urologists, for two to four months, whenever the initial prostate volume was above 50 g, with the purpose of reducing the prostate volume, and were interrupted before brachytherapy planning. None of the patients was submitted to external radiotherapy.

The brachytherapy technique proposed by the authors is described in detail in another article(21). Post-implant dosimetry evaluates the technical quality of the implant and the quantified radiation dose delivered to the bladder and to the adjacent rectum, based on pelvic computed tomography performed 30 days after brachytherapy. Criteria for appropriate post-implant dosimetry included V100 > 80%, D90 > 90%, and V150 < 60% for iodine-125 seeds(22). In Brazil, only iodine-125 seeds are commercially available for low dose rate brachytherapy.

The patients were followed-up every four or six months by means of total PSA level testing and clinical and radiological evaluation, as necessary, in the event of PSA levels increase.

The parameter utilized to evaluate the results was total PSA. The Phoenix Consensus(23) is adopted to define biochemical recurrence, established as PSAn + 2 ng/ml or in cases where any rescue treatment is initiated. PSAn is the lowest total PSA value recorded after any treatment.
Several authors have proposed a cut-off value for different PSAn levels, ranging from 0.2 to 4.0 ng/ml, as predictive factor for biochemical recurrence. In the present study, the groups were divided according to PSAn < 0.5 ng/ml or ≥ 0.5 ng/ml as the cut-off value.

Mean and median values were calculated for results analysis. The following statistical methods were utilized: chi-square test for categorical variables and Student’s t test for continuous variables. The adopted significance level was 5% probability (p ≤ 0.05), with a confidence interval of 95% (CI95%). The BDFS was analyzed by means of the Kaplan-Meier curve and the log-rank test. The statistical analysis was carried out by means of the software SPSS for windows, release 13 (SPSS Inc.; Chicago, USA).

RESULTS

In total, 120 patients were analyzed: 34 patients (28.3%) with biochemical recurrence, and 86 (71.7%) presented with biochemical control. Mean PSAn was 0.53 ng/ml (0.01–2.00; CI95%; 0.45–0.61) and the mean follow-up period was 98 months (29–144; CI95%; 91–104).

In those patients with biochemical control, the mean PSA value was 0.27 ng/ml (0.01–0.98; CI95%; 0.23–0.31), and in those patients with biochemical recurrence, it was 0.96 ng/ml (0.50–2.00; CI95%; 0.83–1.08), with p < 0.035. The mean time for onset of biochemical recurrence was 36 months (5–93; CI95%; 26–45).

The patients were divided into two groups, as follows: group 1, with 74 patients (61.7%) presenting with PSAn < 0.5 ng/ml two years after brachytherapy; and group 2, with 46 patients (38.3%) presenting with PSAn ≥ 0.5 ng/ml two years after brachytherapy. The clinical characteristics of the patients are described in Table 1.

In group 1, 15 patients (20.3%) with biochemical recurrence, while in group 2, 19 patients (43.2%) had biochemical recurrence with p < 0.05. The analysis of risk factors such as age (p = 0.13), PSAi (p = 0.4), prostate volume (p = 0.85), dosimetric parameters (p > 0.08), risk groups (p = 0.3), CS (p = 0.06) and Gleason score (p = 0.25) did not demonstrate any significant association between the two groups.

The analysis of BDFS after seven years of follow-up in groups 1 and 2 was 80% and 64% respectively (Figure 1), demonstrating statistical significance of PSAn as a predictive factor of biochemical recurrence (p < 0.02).

DISCUSSION

Miller et al. presented questions on the predictive value of PSAn and whether or not it could be utilized to predict subsequent failure after prostate cancer radiotherapy. Investigators are looking for an early predictor of failure in order to allow for an earlier institution of rescue treatment. However, PSAn might generally take several years to occur, with its lower plateau not being reached even after 8–10 years, continuing to decrease slowly in some cases. As a result, PSAn would be of little practical clinical value. A much more reasonable approach would be to utilize the lowest PSA value evaluated during a definite and relatively short time interval (for example, 12 months) after completion of radiotherapy. Some authors report that PSAn is significantly related to clinical failure. The present study includes the PSAn values two years after iodine-125 brachytherapy and the chances of biochemical recurrence in the treatment of prostate cancer.

Kapadia et al. have reviewed several studies with different absolute post-treatment PSAn levels indicative of bio-

| Table 1—Groups’ characteristics. |
|----------------------------------|
|                                | Group 1           | Group 2           | p    |
| Mean (CI95%)                   | Mean (CI95%)      | p                |
| Age (years)                    | 68 (67–69)        | 66 (65–67)        | 0.13 |
| PSAi (ng/ml)                   | 10.7 (9.7–11.8)   | 11.2 (10–12.4)    | 0.4  |
| Volume (g)                     | 39.6 (37.9–41.2)  | 35 (33.4–36.6)    | 0.85 |
| V100 (%)                       | 83.6 (81.7–85.6)  | 87.3 (84.8–89.8)  | 0.25 |
| D0 (%)                         | 97.6 (95.3–100)   | 101 (97.8–104.3)  | 0.08 |
| V150 (%)                       | 49.7 (47.3–52.1)  | 56.7 (54–59.3)    | 0.1  |
| Risk groups                    | Patients (%)      | Patients (%)      | p    |
| Low                             | 11 (12.2%)        | 5 (16.7%)         | 0.3  |
| Intermediate                   | 52 (57.8%)        | 15 (50%)          |      |
| High                            | 27 (30%)          | 10 (33.3%)        |      |
| PSAi < 0.5 ng/ml               | 80 (88.8%)        | 25 (83.3%)        | 0.25 |
| 10.1–20 ng/ml                  | 5 (5.6%)          | 3 (10%)           |      |
| > 20 ng/ml                     | 5 (5.6%)          | 2 (6.7%)          |      |
| Clinical stage                  |                  |                  |      |
| T1c–T2a                        | 16 (17.8%)        | 8 (26.7%)         | 0.06 |
| T2b–T2c                        | 74 (82.2%)        | 22 (73.3%)        |      |
| Gleason score                   |                  |                  |      |
| 2–6                            | 80 (88.8%)        | 25 (83.3%)        |      |
| 7                              | 5 (5.6%)          | 3 (10%)           |      |
| 8–9                            | 5 (5.6%)          | 2 (6.7%)          |      |
| NHT                            | Yes              | Yes              | 0.52 |
|                                | 46 (51%)          | 16 (53%)          |      |
|                                | No               | 44 (49%)          |      |

CI95%, confidence interval at 95%; NHT, neoadjuvant hormone therapy.

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after the seeds implant followed by subsequent decrease, generally 12 to 24 months after treatment, being described as an increase of PSA must be taken into consideration as it occurs in 35% of the cases. In their analysis, Zelefsky et al. have indicated that the reference time span for metastases and mortality. If an increase in PSA levels was observed during follow-up, PSA tests were performed at shorter intervals (30 days), until the cause of the biochemical failure could be found, in order to avoid the PSA bounce effect. Kaminski et al. have suggested that post-radiotherapy PSA is more significant in patients at a high risk for biochemical recurrence, such as in the case of patients at advanced clinical stages of disease, those presenting with high Gleason score or high PSA. In the present study, patients from all risk classes were included, and none of them demonstrated statistically significant differences.

Neoadjuvant hormone therapy has been utilized by some patients and that might affect the PSA levels due to the changes in the PSA levels. However, in the present study, PSA was not significantly influenced by the utilization of neoadjuvant hormone therapy. Kaminski et al. have demonstrated that the prostate volume is predictive for PSA, as larger prostates might lead to higher PSA values. In the present study, the evaluation of prostate volume and PSA did not demonstrate any statistical difference.

Zelefsky et al. have found out that the PSA level was predictive of biochemical control at long term, since relatively low PSA levels seem to reflect high intraprostatic doses delivered by interstitial brachytherapy as a result from the ablution of malignant and normal cells on the production of PSA in the prostate epithelium, as the prostatic epithelial cells are responsible for the PSA serum levels, and higher radiation doses in the prostate volume are, probably, directly related to the decrease in such an antigen levels. Thus, one might conclude that higher \( D_{90} \) and \( V_{100} \) would be related to lower PSA values, demonstrating the importance of the radiation dose as higher doses are associated with lower PSA values and better biochemical control. However, the present study did not demonstrate any influence of \( D_{90} \) and \( V_{100} \) on the PSA values.

Pinkawa et al. report that PSA is highly relevant for the prognosis, since all their patients with PSA < 0.1 ng/ml remained free of recurrence (100%) while those with PSA < 0.5 ng/ml remained free of recurrence in 87% of the cases. In the present study, the BDFS rate for PSA < 0.5 ng/ml was 80%, very close to that in the mentioned study.

The rate of five-year BDFS found by Nickers et al. and Cury et al. was 95% for PSA < 1.0 ng/ml, and 57% for PSA ≥ 1.0 ng/ml. Lee et al. recorded rates of 99% and 49%, respectively, for PSA < 1.0 ng/ml and PSA ≥ 1.0 ng/ml. Such data are consistent with those from the present study. Critz et al. have reported that all patients with PSA > 1.0 ng/ml after five years have greater probability of biochemical recurrence, in spite of the utilization of brachytherapy in association with radiotherapy. Other authors have found a five-year BDFS of 32% for patients with PSA ≥ 1.0 ng/ml. In the present study, with seven years of actuarial analysis of BDFS of patients with PSA ≥ 0.5 ng/ml, the BDFS was 64% two years after isolated brachytherapy.

Recently, Chen et al. concluded that PSA < 0.49 ng/ml two years after radiotherapy translated into better long term biochemical control, as does the present study.

The present study had the following limitations: retrospective analysis, patients referred by different assisting urologists, staging performed by means of transrectal ultrasonography and difficulty in comparing data with those reported by other authors because of samples heterogeneity both in Brazil and abroad.

CONCLUSIONS

The results from the present study suggest that PSA can be a predictive factor for biochemical recurrence. Levels of PSA > 0.5 ng/ml two years after brachytherapy are strongly associated with a poor prognosis. Such a fact can be useful to identify patients at risk for disease recurrence.

A longer follow-up time and a higher number of patients are necessary to confirm data from the present study.

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REFERENCES

1. Castro HAS, Iared W, Shigueoka DC, et al. Contribuição da densidade do PSA para predizer o câncer da próstata em pacientes com valores de PSA entre 2.6 e 10.0 ng/ml. Radiol Bras. 2011;44:205-9.

2. Brasil. Ministério da Saúde. Instituto Nacional de Câncer. Estima-tiva 2014 – Incidência de câncer no Brasil. Rio de Janeiro: INCA;
Franca CAS et al. / Relationship between two year PSA nadir and biochemical recurrence

2014. [Acessado em 17 de março de 2014]. Disponível em: http://www.inca.gov.br/estimativa/2014/estimativa-24012014.pdf.

3. Rolim AEH, Costa LJ, Ramalho LMP. Percepções da radioterapia na região orofacial e seu tratamento. Radiol Bras. 2011;44:388–95.

4. Silva AV, Galante C, Manzi FR. Efeito da radiação ionizante sobre o paladar em pacientes submetidos a radioterapia para a região da cabeça e pescoço. Radiol Bras. 2011;44:297–300.

5. Trindade BM, Campos TPB. Sistema computacional para dosimetria de nêutrons e fótons baseado em métodos estocásticos aplicado a radioterapia e radiologia. Radiol Bras. 2011;44:109–16.

6. Tseng YD, Chen MH, Beard CJ, et al. Posttreatment prostate specific antigen nadir predicts prostate cancer specific and all cause mortality. J Urol. 2012;187:2068–73.

7. Zelfsmy MJ, Shi W, Yamada Y, et al. Postradiation therapy 2-year prostate specific antigen nadir as a predictor of long-term prostate cancer mortality. Int J Radiat Oncol Biol Phys. 2009;75:1350–6.

8. Franca CAS, Vieira SL, Penna ABCBC. Definição de recidiva bioquímica após tratamento radioterápico do câncer de próstata localizado: revisão de literatura. Rev Bras Cancerol. 2008;54:57–61.

9. Ragde H, Blasko JC, Grimm PD, et al. Interstitial iodine-125 radiation without adjuvant therapy in the treatment of clinically localized prostate carcinoma. Cancer. 1997;80:442–53.

10. Trindade BM, Campos TPR. Sistema computacional para dosimetria de nêutrons e fótons baseado em métodos estocásticos aplicado a radioterapia e radiologia. Radiol Bras. 2011;44:109–16.

11. Nickers P, Albert A, Waltregny D, et al. Prognostic value of PSA kinetics following external-beam radiotherapy and temporary (Ir-192) or permanent (I-125) brachytherapy for prostate cancer. Int J Radiat Oncol Biol Phys. 2007;67:327–33.

12. Franca CAS, Vieira SL, Carvalho ACP, et al. Localized prostate cancer with intermediate- or high-risk features treated with combined external beam radiotherapy and iodine-125 seed brachytherapy. Brachytherapy. 2010;9:307–12.

13. Symon Z, Griffith KA, McLaughlin PW, et al. Dose escalation for localized prostate cancer: substantial benefit observed with 3D conformal therapy. Int J Radiat Oncol Biol Phys. 2003;57:384–90.

14. Kapadia NS, Olson K, Sandler HM, et al. Interval to biochemical failure as a biomarker for cause-specific and overall survival after dose-escalated external beam radiation therapy for prostate cancer. Cancer. 2012;118:2059–68.

15. Hanlon AL, Diratzouin H, Hanks GE. Posttreatment prostate specific antigen nadir highly predictive of distant failure and death from prostate cancer. Int J Radiat Oncol Biol Phys. 2002;53:297–303.

16. Kaminski JM, Hanlon AL, Horwitz EM, et al. Relationship between prostate volume, prostate-specific antigen nadir, and biochemical control. Int J Radiat Oncol Biol Phys. 2002;52:888–92.

17. Bowes D, Crook JM, Wallace K, et al. Results of a surgically derived nomogram to predict Gleason score upgrading applied to a cohort of patients with “favorable-risk” prostate cancer treated with permanent seed brachytherapy. Urology. 2012;80:649–55.

18. [No authors listed]. Consensus statement: guidelines for PSA following radiation therapy. American Society for Therapeutic Radiology and Oncology Consensus Panel. Int J Radiat Oncol Biol Phys. 1997;37:1035–41.

19. Sylvester JE, Grimm PD, Blasko JC, et al. 15-Year biochemical free survival in clinical Stage T1-T3 prostate cancer following combined external beam radiotherapy and brachytherapy; Seattle experience. Int J Radiat Oncol Biol Phys. 2007;67:57–64.

20. Nag S, Beyer D, Friedland J, et al. American Brachytherapy Society (ABS) recommendations for transperineal permanent brachytherapy of prostate cancer. Int J Radiat Oncol Biol Phys. 1999;44:789–99.

21. Franca CAS, Vieira SL, Bernabe AJ, et al. The seven-year preliminary results of brachytherapy with iodine-125 seeds for localized prostate cancer treated at a Brazilian single-center. Int Braz J Urol. 2007;33:752–63.

22. Miller N, Smolkin ME, Bissomette E, et al. Undetectable prostate specific antigen at 6-12 months: a new marker for early success in hormonally treated patients after prostate brachytherapy. Cancer. 2005;103:2499–506.

23. Roach M 3rd, Hanks G, Thames H Jr, et al. Defining biochemical failure following radiotherapy with or without hormonal therapy in men with clinically localized prostate cancer: recommendations of the RTOG-ASTRO Phoenix Consensus Conference. Int J Radiat Oncol Biol Phys. 2006;65:965–74.

24. Miller N, Smolkin ME, Bissonette E, et al. Undetectable prostate specific antigen at 6-12 months: a new marker for early success in hormonally treated patients after prostate brachytherapy. Cancer. 2005;103:2499–506.

25. Pollack A, Hanlon AL, Horwitz EM, et al. Prostate cancer radiotherapy dose response: an update of the fox chase experience. J Urol. 2004;171:1132–6.

26. Symon Z, Griffith KA, McLaughlin PW, et al. Dose escalation for localized prostate cancer: substantial benefit observed with 3D conformal therapy. Int J Radiat Oncol Biol Phys. 2003;57:384–90.

27. Abdel-Wahab M, Mahmoud O, Merrick G, et al. ACR Appropriateness Criteria® external-beam radiation therapy treatment planning for clinically localized prostate cancer. J Am Coll Radiol. 2012;9:233–8.

28. Pinkawa M, Piroth MD, Holy R, et al. Prostate-specific antigen kinetics following external-beam radiotherapy and temporary (Ir-192) or permanent (I-125) brachytherapy for prostate cancer. Radiother Oncol. 2010;96:25–9.

29. Nickers P, Albert A, Waltregny D, et al. Prognostic value of PSA nadir < or = 4 ng/mL within 4 months of high-dose radiotherapy for locally advanced prostate cancer. Int J Radiat Oncol Biol Phys. 2006;65:73–7.

30. Cury FL, Hunt D, Roach M 3rd, et al. Prostate-specific antigen kinetics after short-term hormone therapy plus external-beam radiotherapy and outcome in patients treated on Radiation Therapy Oncology Group study 9413. Cancer. 2013;119:1999–2004.

31. Lee WR, Hanlon AL, Hanks GE. Prostate specific antigen nadir following external beam radiation therapy for clinically localized prostate cancer: the relationship between nadir level and disease-free survival. J Urol. 1996;156(2 Pt 1):450–3.

32. Critz FA, Williams WH, Holladay CT, et al. Post-treatment PSA < or = 0.2 ng/mL defines disease freedom after radiotherapy for prostate cancer using modern techniques. Urology. 1999;54:968–71.

33. DeWitt KD, Sandler HM, Weinberg V, et al. What does postradiotherapy PSA nadir tell us about freedom from PSA failure and progression-free survival in patients with low and intermediate-risk localized prostate cancer? Urology. 2003;62:492–6.

34. Chen LN, Suy S, Uhm S, et al. Stereotactic body radiation therapy (SBRT) for clinically localized prostate cancer: the Georgetown University experience. Radiat Oncol. 2013;8:58.

Radiol Bras. 2014 Mar/Abr;47(2):89–93