Association of the Serum Vascular Endothelial Growth Factor Levels With Benign Prostate Hyperplasia and Prostate Malignancies

Mohammad Reza Sharif 1; Amirreza Shaabani 2; Hossein Mahmoudi 1; Hassan Nikoueinejad 3; Hossein Akbari 4; Behzad Einollahi 3

1Department of Pediatrics, Kashan University of Medical Sciences, Kashan, IR Iran
2Department of Urology, Kashan University of Medical Sciences, Kashan, IR Iran
3Department of Nephrology and Urology Research Center, Baqiyatallah University of Medical Sciences, Tehran, IR Iran
4Department of Biostatistics, Kashan University of Medical Sciences, Kashan, IR Iran

*Corresponding author: Hassan Nikoueinejad, Department of Nephrology and Urology Research Center, Baqiyatallah University of Medical Sciences, Baqiyatallah Hospital, Mollasadra Ave., Vanak Sq., P.D. Box: 19395-5487, Tehran, IR Iran. Tel: +98-2181262073, Fax: +98-3615551112, E-mail: hnikoueinejad@yahoo.com

Received: September 11, 2013; Revised: November 11, 2013; Accepted: March 15, 2014

1. Background

Prostate cancer is the fourth cause of death in men worldwide (9.2%) being surpassed only by lung cancer (18%). Due to the magnitude of the impact over public health, strong efforts have been made aiming the prevention, early diagnosis, and treatment of this disease (1).

As we enter into the postgenomics era, novel biomarkers of prostate cancer will invariably emerge (2). Angiogenesis is a critical component of prostate cancer development and progression (2, 3) and some studies have shown a close association between increased microvascular density and Gleason score (an evaluating system for prostate cancer based on microscopic features of prostate cells) in advanced prostate cancer (2).

Vascular endothelial growth factor (VEGF), which is expressed at elevated levels in several tumor types, is an important hypoxia-inducible proangiogenic protein and a potent inducer of endothelial cell growth (4, 5). The clinical importance of VEGF in the concept of tumor growth is supported by the fact that most tumors produce VEGF and inhibition of VEGF-induced angiogenesis significantly reduces tumor growth in vivo (6). Although the serum levels of VEGF in many types of cancer have been correlated with stage of the disease (7), the validity of the VEGF as a prognostic marker of prostate cancer is controversial. Some studies stated that the plasma (8) as well as urinary (9) levels of this factor had a prognostic value in both localized (8, 10, 11) and metastatic (8, 9) prostate cancer, especially in early stages of the tumor (12, 13). In addition, few studies have introduced the value of VEGF as a screening test for prostate cancer, as a tool for its staging, as a target for therapeutic strategies (14), and as the only significant prognostic factor of disease-specific survival (5).

Implication for health policy/practice/research/medical education:

We aimed to compare serum level of VEGF in patients with prostate cancer with the levels in patients with benign prostate hyperplasia and healthy people to determine if it would be a prognostic marker as well as differentiating factor in such patients. This study was performed for the first time in Iranian population.
On the other hand, some studies revealed that the expression of VEGF was not correlated with any clinicopathological parameters among prostate cancer specimens (15). It was stated that the plasma levels of VEGF did not have any association with the clinical staging (3, 7), the form (benign vs. malignant) (16) and the progression (localized vs. metastatic) (17) of the prostate cancer. These controversies are also seen in benign prostate hyperplasia (BPH). Some researchers came to the conclusion that serum level of VEGF as well as its expression is increased in the benign patterns like BPH (4) but others believed that its expression would decrease along with progression of the disease (15).

2. Objectives

For the first time in Iranian population, we aimed to compare the serum levels of VEGF in patients with prostate cancer with the levels in patients with BPH as well as healthy people and to determine whether it would be a prognostic marker as well as differentiating factor in such patients.

3. Patients and Methods

This study was performed under KAUMS (Kashan University of Medical Sciences) Ethics Committee-approved protocol and written informed consent was obtained from all participants. The study protocol conformed to the ethical guidelines of the 1975 Helsinki Declaration. Exclusion criteria consisted of prior surgery or any other treatment of prostate cancer or BPH as well as any active infection or inflammatory process. From May 2011, serum samples were collected from 158 men. Forty-four of them were new cases of BPH based on clinical and sonographic criteria (diffused enlargement of transitional zone of prostate nearby the base of bladder with nodular and heterogenic echo). We recruited 57 age-assimilated healthy control men without any complaint or sign of inflammatory/noninflammatory diseases. In those with an abnormal rectal exam and a prostate-specific antigen (PSA) > 4, we proved the diagnosis of cancer by biopsy. We considered just new cases without any surgical procedures to assess TMN; therefore, it was impossible to evaluate the staging of the patients. Five patients with BPH attended with a mildly elevated PSA levels; therefore, we confirmed no evidence of malignancy in them with histological findings on the biopsy specimens.

All patients and controls had a blood sample drawn into a tube with no anticoagulant. In BPH, patients’ blood samples were taken after the diagnosis and before any treatment, and in cancer suspects (according to rectal exam findings and PSA) before prostatic biopsy. Blood samples of biopsy-confirmed cancerous patients were enrolled in the study. Therefore, nobody had any surgical and medical treatment at the time of sampling. The serum stored at −20°C until analysis. Concentration of serum VEGF was measured using a quantitative immunosassay technique, namely enzyme-linked immunosorbent assay (ELISA), by a commercially available ELISA kit (eBioscience, USA), according to the manufacturer’s instruction. Samples were analyzed twice and mean VEGF levels reported in picogram per milliliter in each group. All data were analyzed by SPSS v.17 (SPSS Inc, Chicago, Illinois, USA) using descriptive statistics, namely chi-square and fissure exact tests. Analytical data were analyzed via ANOVA and student t-test. We used a logistic regression model to assess the effective factors on prostate cancer risk. Then the impact of all factors on VEGF was studied simultaneously by covariance analytic model. Using receiver operating characteristic (ROC) curve and area under curve (AUC), we tried to determine the sensitivity and specificity of VEGF as a diagnostic marker of prostate cancer. Moreover, using analysis of the principle components, we calculated linear combination of VEGF and PSA variables with equal co-efficiencies. Finally, we calculated the specificity, sensitivity and AUC of combined variable to predict prostate cancer.

4. Results

Table 1 shows the mean ages, serum levels of VEGF as well as PSA, and prostate volume in each group. The serum level of VEGF and PSA were significantly higher in the cancer group in comparison to the BPH and healthy groups (P < 0.001).

Table 2 shows the correlation coefficient of VEGF with Gleason score, PSA, and prostate volume. There was a meaningful correlation between VEGF and PSA in prostate cancer group (r = 0.425, P = 0.004). Using a logistic regression model, we determined that the age (P < 0.001) and VEGF (P < 0.001) as risk factors of prostate cancer. Adjusting the age effect, VEGF had an association with prostate cancer risk (P < 0.001).

Using ROC curve and area under curve (AUC) (Figure 1), we determined the sensitivity and specificity of VEGF as a diagnostic marker of prostate cancer. In the best cutoff point of VEGF = 188.2, sensitivity and specificity were 86.4% and 70.2%, respectively, and AUC was determined as 0.876. Odds ratio for such diagnosis was calculated as 14.9 with the 95% confidence interval of 3.89-57.1. Moreover, calculating the sensitivity and specificity of the variable taken from the combination of VEGF and PSA as a diagnostic marker of prostate cancer (Figure 1), we determined a sensitivity of 91% and a specificity of 78% at the cutoff point of 102 and AUC was determined as 0.876. Odds ratio for such diagnosis was calculated as 35 with a 95% confidence interval of 6.3-194.2.
Table 1. Study Groups, Mean Ages, Vascular Endothelial Growth Factor, Prostate-Specific Antigen, and Prostate Volume Values a, b

|                  | Control | BPH      | Cancer   | P Value |
|------------------|---------|----------|----------|---------|
| Age, y           | 67.2 ± 7.2 | 68.7 ± 8.3 | 66.5 ± 9.1 | 0.38    |
| 40-59            | 7 (12.3) | 6 (10.5) | 9 (20.5) |         |
| 60-69            | 26 (45.6) | 25 (43.9) | 24 (54.5) |         |
| ≥ 70             | 24 (42.1) | 26 (45.6) | 11 (25)  |         |
| VEGF, pg/mL      | 124.9 ± 50.8 | 147.5 ± 80 | 356.3 ± 181.1 | < 0.001 |
| PSA, ng/mL       | 1.94 ± 1.94 | 3.86 ± 2.3 | 38.7 ± 17.5 | < 0.001 |
| Prostate volume  | 37.4 ± 15.5 | 71.7 ± 25.5 | < 0.001 |         |

a Abbreviations: BPH, benign prostate hyperplasia; PSA, prostate-specific antigen; VEGF, vascular endothelial growth factor.

b Data are presented as mean ± SD or No. (%).

Table 2. Correlation Coefficient of Vascular Endothelial Growth Factor With Gleason Score, Prostate-Specific Antigen, and Prostate Volume a

|                  | Gleason Score | PSA            | Prostate Volume |
|------------------|---------------|----------------|-----------------|
| Control          | -             | r = -0.01, P = 0.962 | r = -0.176, P = 0.381 |
| BPH              | -             | r = 0.07, P = 0.61 | r = -0.059, P = 0.665 |
| Prostate cancer  | r = -0.089, P = 0.566 | r = 0.425, P = 0.004 | -   |

a Abbreviations: BPH, benign prostate hyperplasia; PSA, prostate-specific antigen

5. Discussion

Morphology-based approaches, especially Gleason scoring, combined to clinical parameters of PSA and T stage have provided clinicians some important prognostic information about prostate cancer. Recent successes have served to cultivate the growing interest in discovering more molecular-based prognostic factors (2). Such biomarker should be quickly quantifiable in accessible biological fluids without any overlapping to untreated and healthy people; moreover, they should be consistent, cost-effective, readily interpretable by clinicians, prostate-specific, able to differentiate cancerous prostate and its stages from prostatic hyperplasia, evaluate the survival of the patients as well as response to treatment (18). VEGF, which induces vascular permeability and stimulates endothelial cell growth, is now recognized as a key factor required for growth of the tumors (6) and might be a prognostic factor in several tumors such as prostate cancer (19). It seems that VEGF has some of such above-mentioned advantages. According to our study, it may differentiate cancerous prostate from benign hyperplasia and according to Duque et al. study (7), it can discriminate metastatic disease from its localized form. In this regards, VEGF may resolve the drawback of PSA, which is tissue-specific rather than prostate cancer-specific and its serum concentration might be affected by several benign conditions (1, 18). Such false-positive results of PSA may substantially necessitate further diagnostic evaluation, impose extra expenses, and lead to the employing more invasive procedures. It has been revealed that for every patient who benefits from PSA diagnosis-initiated treatment, 47 patients undergo unnecessary biopsy and other treatments because of false-positive PSA test results. Conversely, efforts to prevent such over diagnosis may lead to some delays in the treatment of aggressive
and potentially life-threatening cancers (18). Some drawbacks of PSA are also related to its false-negative results. It has been demonstrated that prostate cancer may be detected in about 15% of men with normal or very low levels of total PSA, thereby making it difficult to reliably rule out the possibility of cancer at any PSA level (20). Considering that there are not reliable biomarkers as diagnostic as well as prognostic factors for prostate cancer (21), and in line with some other studies on tissue (10, 22) and serum (3), our study demonstrated that VEGF might be a potential biomarker for prostate cancer. According to some previous studies, however, there is not any consensus on the VEGF level expression as well as its prognostic and predictive value in prostate cancer and results are so controversial (22, 23). Such conflicting results could be due to different factors such as using different sample sizes, different stages of the cancer, and sensitivity of the used assays (22). Therefore, considering VEGF expression alone has a limited value in prostate cancer and according to our results, it seems that considering a combination of PSA and a proper cut-off point for VEGF might result in overcoming such drawbacks.

In line with Souliotis et al. (15), we showed that serum VEGF might not be increased in prostatic hyperplasia. In addition, there were no statistical differences in serum VEGF level between BPH patients and healthy people in our study; a concept which was confirmed by some other studies (7) and is important to discriminate benign patterns from malignant ones. We suggest for the first time that VEGF combined with PSA, may be used as a powerful screening test to evaluate the risk of prostate cancer. We propose complementary researches using larger sample size to certificate the diagnostic validity of serum VEGF at different stages.

Our study had some advantages. First, relatively proper sample size and associated standard deviations yielded a proper power to detect differences in subgroup analyses. Second, all of our patients were newly diagnosed and there were no essential factors affecting the comparability of the groups including using different kinds of medical or surgical treatments. Such uniformity of the patients especially in the cancerous arm was seen less in other studies. In a case-control study, Vancliff et al. (24) showed that some alleles of VEGF gene have significant correlation with prostate cancer risk (P = 0.04). They supported the fact that genetics can affect VEGF expression. This means that VEGF may have different prognostic values in prostate cancer at different populations. Therefore, we should investigate the VEGF gene polymorphism in our population in the future studies.

Acknowledgements

This study was funded and supported by Deputy of Research, Kashan University of Medical Sciences (KAUMS), Grant No. 9275.

Authors’ Contributions

Ali Reza Sharif introduced more patients and revised the final article. Hassan Nikoueinejad developed the original idea and the protocol, performed the tests, edited, and submitted the manuscript. Alireza Shaabani followed the patients, gathered the blood samples, abstracted the data, and prepared the manuscript. Hossein Mahmoudi contributed to the development of the study, visited, and introduced the patients. Hossein Akbari analyzed the data. Behzad Einollahi revised the manuscript for important intellectual content.

Financial Disclosure

There was no conflict of interest.

Funding/Support

The study was funded by Kashan University of Medical Sciences, Kashan, Iran.

References

1. Miotto A Jr, Srougi M, Brito GA, Leite KM, Nesarullah AJ, Ortiz V. Value of various PSA parameters for diagnosing prostate cancer in men with normal digital rectal examination. Int Braz J Urol. 2004;30(2):99-13.
2. Chakravarti A, Zhai GG. Molecular and genetic prognostic factors of prostate cancer. World J Urol. 2003;21(1):265-74.
3. Shariat SF, Anwari VA, Lamb DJ, Shah NW, Wheeler TM, Slawin KM. Association of preoperative plasma levels of vascular endothelial growth factor and soluble vascular cell adhesion molecule-1 with lymph node status and biochemical progression after radical prostatectomy. J Clin Oncol. 2004;22(9):1655-63.
4. Jackson MW, Bentel JM, Tilley WD. Vascular endothelial growth factor (VEGF) expression in prostate cancer and benign prostatic hyperplasia. J Urol. 1997;157(6):2235-38.
5. Green MM, Hilel CT, Shanks JH, Bottomley IC, West CM, Cowan RA, et al. Expression of vascular endothelial growth factor (VEGF) in locally invasive prostate cancer is prognostic for radiotherapy outcome. Int J Radiat Oncol Biol Phys. 2007;67(1):84-90.
6. Translational Research: the Role of VEGF in Tumor Angiogenesis. Proceedings of a symposium. Washington, DC, USA. November 16, 1999. Oncologist. 2000;5 Suppl 1:13-57.
7. Duque JL, Loughlin KR, Adam RM, Kantoff P, Mazzucchi E, Freeman MR. Measurement of plasma levels of vascular endothelial growth factor in prostate cancer patients: relationship with clinical stage, Gleason score, prostate volume, and serum prostate-specific antigen. Clinics (Sao Paulo). 2006;61(5):401-8.
8. Peyromaure M, Badoual C, Camparo P, Grabar S, Goulvestre C, Fulla Y, et al. Plasma levels and expression of vascular endothelial growth factor A in human localized prostate cancer. Oncol Rep. 2007;18(1):345-9.
9. Miyake H, Muramaki M, Kurahashi T, Yamakana K, Hara I. Urinary levels of vascular endothelial growth factor in patients with prostate cancer as a predictor of disease progression. Anticancer Res. 2005;25(5):3645-9.
10. Peyromaure M, Camparo P, Badoual C, Desczaeaud A, Dinh-Xuan AT. The expression of vascular endothelial growth factor is associated with the risk of cancer progression after radical prostatectomy. BJU Int. 2007;99(5):1150-1.
11. Yang J, Wu HF, Qian LX, Zhang W, Hua LX, Yu ML, et al. Increased expressions of vascular endothelial growth factor (VEGF), VEGF-C and VEGF receptor-3 in prostate cancer tissue are associated with tumor progression. Asian J Androl. 2006;8(2):169-75.
12. Latil A, Bieche I, Pesche S, Valeri A, Fournier G, Cussenot O, et al.
13. Kaushal V, Mukunyadzi P, Dennis RA, Siegel ER, Johnson DE, Kohli M. Stage-specific characterization of the vascular endothelial growth factor axis in prostate cancer: expression of lymphangiogenic markers is associated with advanced-stage disease. *Clin Cancer Res.* 2005;11(2 Pt 1):584–93.

14. Delongchamps NB, Peyromaure M. The role of vascular endothelial growth factor in kidney and prostate cancer. *Can J Urol.* 2007;14(5):3669–77.

15. Soullietis N, Karyotis I, Delakas D, Spandidos DA. Expression analysis of peptide growth factors VEGF, FGF2, TGFβ1, EGF and IGF1 in prostate cancer and benign prostatic hyperplasia. *Int J Oncol.* 2006;29(2):305–14.

16. Walsh K, Sherwood RA, Dew TK, Mulvin D. Angiogenic peptides in prostatic disease. *BJU Int.* 1999;84(9):1081–3.

17. Mao K, Badoual C, Camparo P, Delongchamps NB, Vieillefond A, Dinh-Xuan AT, et al. The prognostic value of vascular endothelial growth factor (VEGF)A and its receptor in clinically localized prostate cancer: a prospective evaluation in 100 patients undergoing radical prostatectomy. *Can J Urol.* 2008;15(5):4257–62.

18. Madu CO, Lu Y. Novel diagnostic biomarkers for prostate cancer. *J Cancer.* 2010;1:150–77.

19. Jones A, Fujiyama C, Turner K, Fuggle S, Cranston D, Bicknell R, et al. Elevated serum vascular endothelial growth factor in patients with hormone-escaped prostate cancer. *BJU Int.* 2000;85(3):276–80.

20. Thompson IM, Pauler DK, Goodman PJ, Tangen CM, Lucia MS, Parnes HL, et al. Prevalence of prostate cancer among men with a prostate-specific antigen level < or =4.0 ng per milliliter. *N Engl J Med.* 2004;350(22):2239–46.

21. Sutcliffe P, Hummel S, Simpson E, Young T, Rees A, Wilkinson A, et al. Use of classical and novel biomarkers as prognostic risk factors for localised prostate cancer: a systematic review. *Health Technol Assess.* 2009;13(5):i, xi-xiii 1-219.

22. Pan L, Baek S, Edmonds PR, Roach M, 3rd, Wolkov H, Shah S, et al. Vascular endothelial growth factor (VEGF) expression in locally advanced prostate cancer: secondary analysis of radiation therapy oncology group (RTOG) 8610. *Radiat Oncol.* 2010;5:100.

23. Concato J, Jain D, Uchio E, Risch H, Li WW, Wells CK. Molecular markers and death from prostate cancer. *Ann Intern Med.* 2009;150(9):595–601.

24. VanCleave TT, Moore JH, Benford ML, Brock GN, Kalbfleisch T, Baumgartner RN, et al. Interaction among variant vascular endothelial growth factor (VEGF) and its receptor in relation to prostate cancer risk. *Prostate.* 2010;70(4):349–52.