Expression of ERG Protein and TMRPSS2-ERG Fusion in Prostatic Carcinoma in Egyptian Patients

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Abstract

AIM: Prostate cancer (PCa) is the second most common cancers in men worldwide. Its incidence can be influenced by several risk factors including genetic susceptibility. Therefore the search for the expression of a certain gene (ERG) and its rearrangement could give us clues for proper identification of PCas. And the study of ERG expression and its comparison to FISH in Egyptian patients can show whether ERG immunophenotype could be used instead of FISH, as it is cheaper.

MATERIALS AND METHODS: This study was performed on 85 cases of PCa, showing 30 cases with HGPIN and 30 cases of prostatic hyperplasia. All were immunohistochemistry stained using ERG monoclonal rabbit antihuman antibody was used (clone: EP111). FISH analysis was performed in 38 biopsies of PCa cases to detect TMRPSS2-ERG rearrangement using the FISH ZytoLight TriCheck Probe (SPEC TMRPSS2-ERG).

RESULTS: ERG expression was found in 26% of PCa cases and 20% of HGPIN cases. FISH analysis showed fusion of 21 cases of PCa (out of 22 cases showing ERG immunoexpression).

CONCLUSION: Our findings emphasise that only malignant and pre-malignant cells and not benign cells from the prostate stain positive. ERG expression may offer a simpler, accurate and less costly alternative for evaluation of ERG fusion status in PCa.

Introduction

Prostatic carcinoma (PCa) is a heterogeneous disease process with a varied spectrum of light microscopic morphologic as well as biological features [1-2]. Identification of prostate malignant acini can sometimes present a diagnostic challenge for pathologists since PCas can mimic benign prostatic glands [3] and the architectural or cytologic clues for the diagnosis of carcinoma may not always be seen in small foci of suspicious glands. Histopathological diagnosis of PCa can be established by transrectal ultrasound-guided (TRUS) biopsy [4] after an abnormal finding on digital rectal examination or finding an augmentation in prostate-specific antigen (PSA) level [5].

Prostatic intraepithelial neoplasia (PIN) consists of pre-existing prostatic ducts and acini lined by cytotologically atypical cells. HGPIN is considered the precancerous lesion of PCa [6, 7]. There is a strong association between HGPIN and PCa. The incidence of HGPIN in radical prostatectomy specimens for PCa is remarkably high, approximately 85–100% of specimens [6].

About 4% of contemporary prostate needle biopsies contain collections of small acini that are suspicious for cancer and are reported as atypical small acinar proliferation (ASAP) suspicious for but not diagnostic of malignancy. Prostate cancer has been identified in subsequent biopsies in the majority
of cases of ASAP, indicating that this finding is a significant predictor of cancer [8]. Histopathological diagnosis of small focus carcinomas in prostatic needle biopsies is often assisted by IHC [9].

One of the earliest methods of detection of this cancer is PSA, which has been used to screen since the late 1980s [10]. The early disease diagnosis in the Western world is due to the improvement in and widespread availability of the measurement of PSA, including PSA density, PSA volume and adjusted age-specific PSA ranges [5]. The free to total (f/t) PSA ratio was shown, and this ratio is usually lower in patients with PCa than in those with BPH [11]. PCa patients with higher Gleason score correlated with increased total PSA and decreased free PSA [12].

ERG (v-ets erythroblastosis virus E26 oncogene homolog [avian], chromosome 21q22.3) is only seen in prostate cancer. It was first reported as an overexpression of the oncogene ERG at the transcript level in 50% of clinically localised and metastatic PCa samples [13]. Numerous studies showed that the basis for this overexpression was a result of ERG fusion with the promoter region of the androgen-induced TMRPSS2-ERG gene, the most common variant [14-17] HGPIN showing TMRPSS2-ERG fusion is associated with co-existing TMRPSS2-ERG PCa. The detection of isolated TMRPSS2-ERG fusion in HGPIN would improve the positive prognostic value of finding TMRPSS2-ERG fusion PCa in subsequent biopsies [18].

ERG's oncogenic potential is known because of its involvement in Ewing's sarcoma and leukaemia [17]. Additionally, ERG is a highly specific endothelial marker. It is expressed in small-caliber vessels adjacent to PCa using IHC. Hence, it can be used as an internal staining control, regardless of ERG rearrangement status in cancer [20].

The aim of the study was to determine the frequency of ERG expression in PCa of Egyptian patients and in HGPIN lesions juxtaposed to positively expressed PCa cases as well as to assess the concordance between ERG immunoexpression and TMRPSS2-ERG fusion status.

Patients and Methods

The present study was conducted on 115 prostatic cases including 85 patients with PCa, 30 of them containing PIN in the adjacent area of the tumour and 30 patients diagnosed with prostatic hyperplasia. Ten biopsies from patients with hemangioma served as a control group: they were five males and five females. According to the procedure, 1) there were 28 cases of TRUS cores, each case consisting of 6 tissue core samples, in total = 28 cases x 6 cores = 168 cores.

We chose the cases of most presentable areas and cores were representing a composite score, 2) 24 TURP, eight blocks from each case or 24 x 8 = 192 blocks. We chose blocks containing tumours, 3) 33 prostatectomies: we processed 20 blocks from both lobes = 33x20 = 660 blocks, and we chose blocks showing most tumour areas representing a composite score. Total blocks=162+192+660=1114 blocks, also 30 TRUS cases with benign prostatic hyperplasia, 30x6 cores=180 block. We reconstructed 15 tissue microarrays.

Histopathological study

Sections were cut from the paraffin block on the microtome, 4-micron thickness, and were stained with Haematoxylin and Eosin for histopathological diagnosis. Scoring was assessed for pathological evaluation of cases using Gleason scoring system. The PCa cases were sorted into three groups according to the current grading criteria of the ISUP 2014 modified GS (ISUP-GS) [23]: Group 1: GS≤ 6 (including Grade group 1) included 34 cases, Group 2: GS= 7 (including Grade Group 2 & 3) included 25 cases, and Group 3: GS ≥ 8 (including Grade group 4&5) included 26 cases.

Immunohistochemical (IHC) technique

ERG expression was evaluated using a commercial rabbit anti-ERG monoclonal antibody (cloneEP111; Dako, Denmark A/S). Deparaffinization, hydration of the slides, and blocking with pre-antibody solution (20 min) where performed in Dako PT Link (Code PT100/PT101). Then, a protocol template was created. The staining steps and incubation times were pre-programmed into the Autostainer Link software (Dako Autostainer). These were diluting anti-ERG primary antibody (1:50 for 20 min at room temp); applying Poly-HRP anti-rabbit IgG (20 min); applying DAB (20 min, Sigma Fast DAB tablets, Sigma-Aldrich, St. Louis MO); counter staining with EnVision FLEX hematoxylin (5 min); and dehydration, clearing, mounting, and covering.

Interpretation of immunostaining

ERG expression is normally observed in lymphocytes and endothelium. These cells were used as positive internal controls for the ERG staining but were not included in the evaluation of ERG fusion status, and only nuclear staining was considered to be positive.

The prostatic sections were examined using a Zeiss light microscope (Oberkochen, Germany). The number of positively stained cells with the highest
expression recorded within ten successive fields (x400) was counted per section.

Each case was independently assessed by two pathologists (HO and KH) using an H-score system obtained by multiplying the intensity of the stain (0: no staining; 1: weak staining; 2: moderate staining; 3: intense staining) by the percentage (0–100) of the cell showing that staining intensity (H-score range 0–300). Any nuclear staining positivity (H-score >0) was considered as indicative of ERG expression [22].

**Fluorescence in situ hybridization (FISH) technique**

The formation of TMRPSS2-ERG fusions leads to disruption of the ERG gene locus. Of the 115 prostatic tissue specimens, 38 specimens were tested with the FISH approach including different Gleason scores using a previously described triple-color FISH assay (Zytolight SPEC TMRPSS2/ERG Tricheck probe), that detects the hybridization signals of the labelled ERG gene (21q22.2) which appear green and orange, whereas the hybridization signals of the labelled TMRPSS2 gene (21q22.3) appear blue [23].

Sections were heated for 10 min at 70°C, pretreated by dewaxing and proteolysis. Preparation of slides was carried out according to manual. TMRPSS2/ERG probe was added, and slides were denatured for 10 min at 75°C. Then slides were transferred to a humidity chamber and hybridised at 37°C overnight. Cell nuclei were stained with DAPI/Dura Tech-Solution.

**Interpretation of FISH**

Evaluation of the ERG rearrangement is carried out by fluorescence microscopy (Olympus X51) [using four filters red (FITC), green, blue and DAPI] using a × 100 oil immersion objective lens. The microscope is attached to high-resolution video camera (Jale) and monitor. We captured and interpreted photos using hardware (Cytovision 2.3, USA). In nuclei without TMRPSS2-ERG rearrangements, two green/orange fusion signals and two blue signals appeared. An ERG translocation without the involvement of TMRPSS2 was indicated by the split of one green/orange fusion signal, resulting in a separated green signal, near one blue signal, and a separated orange signal. On 21q22.13-q22.3 locus affected by a 21q22.2 deletion resulting in the TMRPSS2-ERG, fusion is indicated by one separate orange signal co-localizing with one blue signal and the loss of one green signal.

**Statistical analyses**

SPSS software version 18 was used for data management and analysis. We did median and interquartile range and compared the medians using Mann-Whitney U-test. Qualitative data were presented as frequencies and percentages. Tests were considered statistically significant when P< 0.05.

**Results**

**ERG protein expression was analysed by IHC** in 115 patients who underwent TRUS, TURP, RP, open prostatectomy and radical cystoprostatectomy at Theodor Bilharz Research Institute in 2014 and 2015. 85 cases were histopathologically diagnosed as PCa, 30 cases of HGPIN and ERG positivity was correlated to age, PSA, and histological parameters. Patients' mean age was 71 years (range 50–88), GS in PCa cases was Group 1< 7 in 34 (40.1%) including Grade group 1, Group 2 = 7 in 25 (29.4%) including Grade Group 2 and 3, and Group 3> 7 in 26 (30.5%) cases. It was expanded by additional FISH analyses of 40 cases; 38 diagnosed as PCa and two benign cases.

**ERG expression by IHC**

The mean age of ERG positive cases is 67 years, which is slightly lower than the ERG negative cases that are 73 years old with no statistical significance. Median total PSA (tPSA) shows a higher level in Group 2 & 3 (PCa with GS>7) than Group 1 (PCa with GS<7) with no statistical significance (Table 1).

| Group | GS range | No. of cases | Total PSA Median | Interquartile Range Median |
|-------|----------|--------------|------------------|----------------------------|
| Group 1 | GS ≤ 6   | 34           | 16.75 (1–64)     | 14 (0.8–800)              |
| Group 2 | GS = 7   | 25           | 31.03 (0.8–500)  | 12.3 (4.5–5000)           |
| Group 3 | GS ≥8    | 26           | 216.75 (4–2700)  | 0.24 (0.4–0.95)           |

Group 1: GS 6 (including Grade group 1) included 34 cases; Group 2: GS = 7 (including Grade Group 2 &3) included 25 cases; Group 3: GS ≥8 (including Grade group 4 &5) included 26 cases; Using Mann-Whitney U-test, we found no statistical difference between different groups.

**Table 1: PSA in correlation to different Grade Groups**

| Group | No. of cases | Total PSA Median | Interquartile Range Median |
|-------|--------------|------------------|----------------------------|
| Group 1 | 34           | 16.75 (1–64)     | 14 (0.8–800)              |
| Group 2 | 25           | 31.03 (0.8–500)  | 12.3 (4.5–5000)           |
| Group 3 | 26           | 216.75 (4–2700)  | 0.24 (0.4–0.95)           |

**Interquartile Range/Median tPSA in ERG positive cases was 126.95 (16-300), and ERG negative cases were 00.00 (Table 2).**

**Table 2: Total PSA in correlation to ERG expression by H Score**

| ERG expression | No. of cases | Interquartile Range Median |
|----------------|--------------|----------------------------|
| ERG + ≤ 10     | 2            | 126.95 (16-300)            |
| ERG + >10      | 6            | 0.00                       |
| ERG - ≤ 10     | 55           | 0.00                       |
| ERG - >10      |              | 0.00                       |
ERG overexpression was found in 22 (26.0%) out of the 85 PCa cases; 2 cases show mild intensity, 6 cases show moderate intensity, and 14 cases show severe intensity. GS groups were sorted into low (GS<7), intermediate (GS =7) and high (GS>7), revealing a higher prevalence with the higher intensity of GS = 7 tumours (50%) in the ERG + group. ERG was positive in 6 of 34 specimens (17.6%) with GS<7, in 11 of 25 specimens (44%) with GS=7, and in 5 of 26 specimens (19.2%) with GS>7.

Table 3: H score in correlation to different Grade Groups

| Grading Groups | ERG H-SCORE Interquartile Range/Median |
|----------------|----------------------------------------|
| Group 1 (n=34) | 124.4 (16-300)                         |
| Group 2 (n=25) | 138.4 (52-300)                         |
| Group 3 (n=26) | 207.9 (14.4-300)                       |

Group 1: GS≤ 6 (including Grade group 1) included 34 cases; Group 2: GS= 7 (including Grade Group 2&3) included 25 cases; Group 3: GS ≥ 8 (including Grade group 4&5) included 26 cases. Using Mann-Whitney U-test, we found no statistical difference between different groups.

H score was increased in group 2 compared to group 1 with no statistical significance (Table 3).

FISH procedure was conducted on 38 cases diagnosed as prostatic carcinoma. FISH analysis showed TMPRSS2-ERG fusion of 21 cases specimen selected of PCa (out of 22 cases that showed ERG immunoexpression).

The gene fusion occurs as a result of either a chromosomal translocation or an interstitial deletion, where 13 cases show interstitial deletion, and 8 cases show chromosomal translocation. Therefore about 61.9% of cases show deletion and 38.1% show translocation (Table 4; Fig. 3).

Table 4: H score positive IHC cases in correlation to FISH positive cases

| ERG by IHC | Number of negative cases | Number of positive cases | Total |
|------------|--------------------------|--------------------------|-------|
| Negative   | 16                        | 0                        | 16    |
| Positive   | 1                         | 21                       | 22    |
| Total      | 17                        | 21                       | 38    |

ERG overexpression was found in 22 (26.0%) out of 85 PCa cases. TMPRSS2-ERG fusion in the specimen selected, 21 cases of the prostatic carcinoma (out of 38 cases) showed fusion. The gene fusion occurs as a result of either a chromosomal translocation or an interstitial deletion, where 13 cases show interstitial deletion, and 8 cases show chromosomal translocation. Therefore about 61.9% of cases show deletion and 38.1% show translocation.
We found a strong association between ERG immunohistochemical overexpression and TMRPSS2-ERG rearrangement detected by FISH with 95.5% concordance, 100% sensitivity and 94.4% specificity.

![Image](https://example.com/image1.png)

FIGURE 3: (A) case of prostatic adenocarcinoma, Gleason score 3+3, showing malignant cells with normal expression of TMRPSS2-ERG as merged picture or triple Bandpass filter set (orange, green and blue) of SPEC TMRPSS2-ERG TriCheck Probe (FISH, magnification × 1000), (B) case of prostatic adenocarcinoma, Gleason score 4+4, showing rearranged SPEC TMRPSS2-ERG as fusion associated deletion in malignant cells with merged picture or triple Bandpass filter set (fused orange and blue, missing green) (arrows) (FISH, SPEC TMRPSS2-ERG TriCheck Probe, magnification × 1000), (C) case of prostatic adenocarcinoma, Gleason score 5+4, showing rearranged SPEC ERG translocation not affecting TMRPSS2 in malignant cells with merged picture or triple Bandpass filter set (fused orange and blue, presence of green but away) (arrows) (FISH, SPEC TMRPSS2-ERG TriCheck Probe, magnification × 1000)

**Discussion**

Globally, prostate cancer is reported to be the sixth most common type of cancer, and the second most common in men, especially in Europe, North America, and parts of Africa [24-25]. Studies are underway to evaluate new tests for PCa that could distinguish more aggressive cancers from those less likely to be lethal, to identify men at higher risk of developing prostate cancer, and to enable more efficient use of PSA testing [26]. Diagnosis, treatment and recommendations of PCa are based on biopsy assessments. Hence, to affect therapeutic decision making, a biomarker must be proved in biopsies [23].

In this study, no association between the immunoeexpression of ERG and the clinical parameters including age, and PSA level was found (no statistical significance). This is in agreement with the study by Rubio-Briones et al. which showed no association of ERG status and preoperative PSA [27].

On the other hand, a previous PSA screening study reported a higher frequency of ERG+ cancer in low PSA patients, however, ‘low PSA’ was defined as ≤10 ng /ml[28]. However, two potential limitations in our current study are the small sample size and the inability to separate age and PSA entirely. Age-adjusted PSA cut points ensure that a young man with a lower PSA is more likely to get a biopsy than an older man with the same PSA level.

Data presented in this study demonstrated that ERG was overexpressed in 26% of the PCa group whereas most of the studies that have been done in Western countries demonstrated that ERG gene is overexpressed in approximately 50% in patients with PCa [28-31]. However, our results are higher than those for Asian patients. For example, Chinese prostate cancer patients showed ERG overexpression in 14.9% only [37]. Japan and Korea also show similar low frequency [32-34].

This study showed ERG expression in 20% of the HGPIN cases (HGPIN in association with PCa in the same cases). This is in agreement with other studies demonstrating that ERG was expressed in 27% of HGPIN and that ERG-positive HGPIN is strongly associated with positive ERG expression PCa in the specimen [35-36]; this indicates that ERG fusions appear early in the development of the disease.

Yu and his co-workers found that TMRPSS2-ERG plays a central role as a "malignant regulatory switch" that shuts down androgen signalling inhibiting normal prostate differentiation [37]. Our study showed that ERG is expressed in the PCa cases with Gleason score (GS) equal to 6 or higher. This is in agreement with other studies that support the role for TMRPSS2-ERG fusion in the initiation of carcinogenesis, as it is found in early lesions and typically homogenously maintained within high-grade tumours [38-41]. Previous papers have reported an association with low-grade prostate tumours [42-44]; whereas other studies support an association with high-grade ones [45].

ERG expression showed no correlation to different GS groups with any statistical significance. Our results, in agreement with many studies, did not find any association between the rearrangement and Gleason grade [47-51]. Conversely, previous studies showed that higher GS were associated with an increased frequency of TMRPSS2-ERG fusion event; ERG expression is inversely related to GS [51-53]. A recent study reported that the expression levels of TMRPSS2-ERG and ERG mRNA are related to more aggressive tumours [54]. However, there is controversy about the role of TMRPSS2-ERG in the development and progression of PCa [51-55].

The standard assessment for TMRPSS2-ERG is FISH [23]. In our research study, we have compared the result of ERG immunoeexpression with TMRPSS2-ERG fusion status that revealed high
sensitivity (100%) and high specificity (94.4%) with a concordance of 95.5%. This is in agreement with Berg and his colleagues’ study that showed a sensitivity of 100% and a specificity of 95.5% for an ERG rearrangement of 100%. This is in agreement with Jiang and his colleagues’ study that showed correlation with 100% sensitivity and 88.9% specificity [56]. Furthermore, many former studies have displayed high sensitivities (95.7%–99.1%) and specificities (92.3%–99.4%) [57–58]. Consequently, our research applied ERG protein overexpression to determine the presence or absence of TMPRSS2-ERG rearrangement. We established that detection of ERG expression by IHC technique could be used as a surrogate marker for ERG rearrangement, in agreement with Svensson et al. and Berg and his coworkers [23, 57].

In conclusion, the prevalence of ERG protein expression is 26% in Egyptian patients with PCa. ERG immunoexpression was present in 20% of HGPIN lesions exclusively when juxtaposed to fusion positive PCa, implying that this is an early event in the process of PCa formation. The standard immunohistochemistry has high accuracy for defining ERG fusion status in men with PCa. Consequently, ERG expression may offer a simpler, accurate and less costly alternative for evaluation of ERG fusion status in PCa. The association between ERG expression and prostate cancer based on the Egyptian population should be further investigated.

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