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ORIGINAL PAPER

Multiparametric MRI/TRUS fusion prostate biopsy: The transperineal approach

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SUMMARY

Objectives: To evaluate the detection rate for clinically significant prostate cancer (csPCa) of transperineal multiparametric MRI/TRUS (magnetic resonance imaging/transrectal ultrasound) fusion targeted biopsy (TPBx).

Materials and Methods. We previously reported our experience in the detection for csPCa (Gleason score > 6 and/or more than two positive cores and or GPC > 50%) performing TPBx; from 1032 men (median age 63 years) with negative digital rectal examination and previous negative biopsy underwent repeat transperineal saturation biopsy (SPBx; reference test) for the suspicion of cancer. All the patients underwent pelvic mpMRI and in the lesions characterized by a PI-RADS score > 3 (524 cases) a TPBx (four cores) was added to SPBx.

Results: None had significant complications (Clavien-Dindo grade I) from prostate biopsy that needed hospital admission; a T1c PCa was found in 372/1032 (36%) patients and 272 (73.1%) of them were classified as csPCa. The detection rate for overall cancers vs csPCa using SPBx vs TPBx PI-RADS > 3 vs TPBx PI-RADS > 4 was equal to 90.3 and 95.6% vs 72% and 83.8% vs 48.4 and 60.3%, respectively; SPBx missed 12 (4.5%) suggestive as being csPCa vs 44 (16.2%) and 108 (39.7%) missed by TPBx PI-RADS > 3 and TPBx PI-RADS > 4, respectively. In detail, 40/228 (17.5%) csPCa were diagnosed in the anterior zone of the gland by TPBx.

Conclusions: Although mpMRI is mandatory to improve the accuracy of prostate biopsy in the diagnosis of csPCa, still today, the systematic biopsy should be performed, anyway, in addition to targeted cores (PI-RADS score > 3) or alone (negative mpMRI) in the presence of clinical parameters suspicious for cancer; in this respect, the transperineal approach reset the risk of sepsis and improve the diagnosis of csPCa located in the anterior zone of the gland.

Key words: Prostate cancer; multiparametric MRI; transrectal fusion biopsy; transperineal fusion biopsy; saturation prostate biopsy.

INTRODUCTION

Prostate cancer (PCa) is the most frequent tumor diagnosed in older men with more than 1 million biopsies a year performed in the United States (1). The transrectal prostate biopsy approach is associated with an increased risk of infection with an estimated hospital admission and sepsis equal to 2.5 (3) and 3.5% (4), respectively; in addition, the rate of overdiagnosis in men enrolled in screening protocols is equal to 50% of the cases (2). Therefore, the main goal is to reduce the number of unnecessary biopsies and diagnose only clinically significant PCa. In this respect, multiparametric magnetic resonance imaging (mpMRI) combined with TRUS (transrectal ultrasound) fusion targeted biopsy has improved the accuracy of standard biopsy schemes in detecting clinically significant prostate cancer (csPCa), especially, in case of a repeat biopsy (5-7) and in the reevaluation of men enrolled in active surveillance (AS) programs (8-10). Although the accuracy of mpMRI/fusion targeted biopsy has been evaluated in a lot of series, very few papers have compared the detection rate for PCa or/and complications of the different MRI/TRUS fusion platforms and/or approaches in the same population (11-14); in this respect, the standard transperineal biopsy in comparison with the transrectal procedure has demonstrated a higher accuracy in diagnosing PCa located in the anterior zone of the gland (15) resetting the risk of sepsis (16).

In this report, the advantages of the transperineal fusion targeted biopsy (TPBx) in the diagnosis of csPCa (17) have been evaluated.

MATERIALS AND METHODS

We previously reported our experience in the detection for csPCa performing TPBx (18); from January 2011 to February 2018 1032 men (median age 63 years) with negative digital rectal examination and previous negative biopsy underwent repeat transperineal saturation biopsy (SPBx; reference test) for the suspicion of cancer (increasing or persistently elevated PSA values). All the patients underwent pelvic mpMRI; SPBx (median of 30 cores; range: 28-34 cores) was performed transperineally using a GE Logiq P6 ecograph (General Electric; Milwaukee, WI, USA) supplied with a bi-planar trans-rectal probe (5-7.5
MHz) using a tru-cut 18-gauge needle (Bard; Covington, GA, USA) under sedation and antibiotic prophylaxis (one tablet daily of levofloxacin 500 mg for 3 days) (18). All mpMRI examinations were performed using a 3.0 Tesla scanner. (ACHIEVA 3T; Philips Healthcare Best, the Netherlands) equipped with surface 16 channels phased-array coil placed around the pelvic area with the patient in the supine position; multi-planar turbo spin-echo T2-weighted, axial diffusion weighted imaging, axial dynamic contrast enhanced MRI were performed for each patient. The mpMRI lesions characterized by a PI-RADS score > 3 were considered suspicious for cancer; moreover, two radiologists blinded to pre-imaging clinical parameters evaluated the mpMRI data separately and independently. In the presence of mpMRI lesions suggestive of cancer (524 cases) a fusion targeted biopsy (four cores) was added to SPBxc cognitive transperineal fusion biopsy in 229 cases, mpMRI TRUS fusion biopsy (GE Logiq E9, General Electric; Milwaukee, WI, USA) in 169 cases and transperineal fusion guided-biopsies (Hitachi 70 Arietta ecograph, Chiba, Japan) in 126 cases, respectively (7); the targeted biopsies of the anterior zone lesions were always performed transperineally. The GE Logiq E9 and Hitachi Arietta 70 platforms allowed processing of a software based rigid registration of pelvic mpMRI and TRUS (end-fire probe and biplanar probe, respectively) by the use of a fusion device; moreover, an electromagnetic tracking system showed the needle localization. The data were collected following the START criteria (13).

**RESULTS**

The median total PSA and prostate weight was 8.6 ng/ml (range: 3.5-46 ng/ml) and 57 grams (range: 20-135 grams), respectively; a mpMRI PI-RADS score > 3 vs > 4 was found in 524/1032 (50.7%) vs 272/1032 (26.3%) cases, respectively (18). None had significant complications (Clavien-Dindo grade I) from prostate biopsy that needed hospital admission; moreover, the mpMRI procedure was well tolerated and successfully performed in all cases (men with claustrophobia, cardiac pacemaker and hip replacement were not included in the study). A total of 272 patients was classified as csPCa: a Gleason score 3 + 4 vs 4 + 3 vs 4 + 4 vs 4 + 5 was found in 112 (41.1%) vs 60 (22.1%) vs 50 (18.4%) vs 50 (18.4%) cases. The Gleason score was directly correlated with PI-RADS score; moreover, the median prostate weight and PSA density were equal to 45 grams and 0.21 vs. 63 grams and 0.15 in men with PCa vs. normal parenchyma, respectively. A normal parenchyma was diagnosed in the remaining 660/1032 (64%) men. The detection rate for overall cancers vs csPCa using SPBxc vs. TPBx PI-RADS > 3 vs. TPBx PI-RADS > 4 was equal to 90.3 and 95.6% vs 72% and 83.8% vs 48.4 and 60.3%, respectively; in detail, SPBxc missed 12 (4.5%) suggestive as being csPCa vs 44 (16.2%) and 108 (39.7%) missed by TPBx PI-RADS > 3 and TPBx PI-RADS > 4, respectively (18). The detection rate for csPCa performing cognitive transperineal fusion biopsy vs transrectal vs transperineal MRI/TRUS fusion biopsy was superimposable and equal to 35.8% (82/229) vs 35.5% (60/172) vs 36.5% (46/126), respectively (18); in detail, 40/228 (17.5%) csPCa were diagnosed in the anterior zone of the gland by transperineal targeted approach.

In definitive, in our series (18), mpMRI significantly reduced the number of unnecessary repeat prostate biopsies (about 50% of the cases using a PI-RADS score > 3); on the other hand, our results suggested that the patients should be informed of the significantly false-negative rate for csPCa of TPBx in the presence of PI-RADS > 3 (16.2%) or > 4 (39.7%).

**DISCUSSION**

The improvement of diagnostic imaging by mpMRI has allowed to perform targeted biopsies of suspicious area increasing the accuracy in the diagnosis of csPCa (19) resulting predictive of definitive Gleason score with a higher detection rate of cancer for each core in comparison with standard prostate biopsy schemes. The detection rate for PCa of mpMRI is between 39% and 59% (20) with an incidence of cancer located only in the anterior zone
equal to 20% (21, 22). Although mpMRI is strongly recommended in men candidate to repeat biopsy or enrolled in AS protocols, still today, extended or SPBx should be always combined with mpMRI/TRUS fusion biopsy because the false negative rate of mpMRI (15-20% of PCa with low volume and Gleason score > 7) and the variable diagnostic accuracy reported using the different mpMRI/TRUS fusion biopsy platforms (18). The targeted biopsy of mpMRI suspicious areas could be performed using “in-bore” mpMRI-guidance, real-time mpMRI/TRUS imaging fusion or performing cognitive mpMRI/TRUS biopsies; although the detection rate for csPCA is superimposable performing cognitive vs fusion targeted biopsy (23), many papers have demonstrated a higher accuracy in favour of the fusion technique (24); in fact, the detection rate for csPCA is directly correlated with the expertise of the surgeon and the accuracy of the MRI/TRUS fusion platforms.

In addition, few data have been reported regarding the accuracy of transrectal vs transperineal mpMRI/TRUS fusion approach in diagnosing clinically significant PCa (12). In this respect, the two approaches are provided of a superimposable detection rate for PCa, but, at the same time, the transperineal approach allows to easily and better reach the anterior zone of the gland (16, 25) resetting the risk of sepsis (15, 26, 27). The estimated targeting error of transperineal stereotactic biopsies is below 1 millimeter (28); therefore, the higher accuracy in the diagnosis of csPCA using the template combined with stereotactic fusion platforms (Figures 1, 2) improve the accuracy in the reevaluation of men enrolled in Active Surveillance (29) protocols and/or the treatment and reevaluation of men enrolled in clinical trials and candidate to focal therapy (30).

Conclusions

Although mpMRI is mandatory in order to improve the accuracy of prostate biopsy in the diagnosis of csPCA, still today, the systematic biopsy should be performed, anyway, in addition to targeted cores (PIRADS score > 3) or alone (negative mpMRI) in the presence of clinical parameters suspicious for cancer; in this respect, the transperineal approach reset the risk of sepsis and improve the diagnosis of csPCA located in the anterior zone of the gland.

References