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Optimizing Safety and Accuracy of Prostate Biopsy

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Abstract

Purpose of review—The objective of this article is to examine the safety of prostate biopsy and discuss the emerging role of MRI - ultrasound fusion technology in improving diagnostic accuracy.

Recent findings—Men undergoing prostate biopsy frequently experience minor complications, including hematospermia, hematuria, and infection. Quinolone-resistant bacteria are a growing concern; thus transperineal access or modification of antibiotic prophylaxis based on local antibiograms is now used to avoid infectious complications.

Multiparametric MRI allows visualization of many prostate cancers, and by fusing MRI with realtime ultrasound, a biopsy needle can be directed by a urologist into suspicious regions of interest. Using this new method, detection of clinically significant prostate cancer (csCaP) has increased and the incidence of falsely negative biopsies has decreased.

Summary—Prostate biopsy is generally a safe procedure, and with attention to local patterns of antibiotic resistance, infectious complications can be minimized. MRI - ultrasound fusion has significantly improved the accuracy of prostate biopsy, allowing tracking and targeting not previously possible.

Keywords

MRI - ultrasound fusion; clinically significant prostate cancer (csCaP); prostate biopsy

Introduction

Prostate cancer diagnosis is based on histologic examination of tissue routinely obtained via needle biopsy. (1) An estimated one million prostate biopsies are performed annually in the United States. (2) While typically considered a safe procedure, prostate biopsy infrequently results in severe complications. (3) In recent years, the incidence of infectious complications has increased significantly, which reflects the high prevalence of quinolone-resistant strains of *Escherichia coli*. (2–4) Identifying high-risk men prior to biopsy, and subsequently

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tailoring the selection of antibiotic prophylaxis has been shown to decrease the rate of infectious complications. (3,5)

Transrectal ultrasound (TRUS) - guided prostate biopsy has for three decades been the standard for obtaining tissue for histological diagnosis, but several limitations of this method are now apparent. (6,7) These limitations include frequent diagnosis of clinically insignificant prostate cancer and imprecise sampling, which misclassifies up to 50% of cases compared to radical prostatectomy specimen, and high-grade tumors missed in as many as 30% of cases. (6–8)

Over the past 6-8 years, multiparametric Magnetic Resonance Imaging (mpMRI) has gained widespread use to detect prostate cancer and guide prostate biopsy. (9) This new method provides a major improvement in detection of clinically significant prostate cancer (csCaP), and decreased detection of clinically insignificant prostate cancer. (7,10–16) The objective of the present review is to explore the safety of prostate biopsy, examine current methods for prevention of complications, and discuss the role of MRI – guided prostate biopsy in improving diagnostic accuracy.

Discussion

Safety of prostate biopsy

Prostate biopsy is generally safe with few major, but frequent minor complications. (17,18) Common complications include hematuria, infection, rectal bleeding, and hematospermia. (17,18) In a European study of more than 7,000 prostate biopsies, the most frequent complication was hematospermia, with an incidence of 53.8%. (19) Hematuria (24.3%), significant pain (4.8%), fever (4.1%), and hospital admission (0.7%) were less frequent. (19)

In recent years, there has been a significant increase in the number of infectious complications requiring hospital admission after prostate biopsy. (2) The significant increase in incidence of infectious complications has been attributed to an emergence of quinolone-resistant bacteria. (20) Rates of *E. coli* resistance to quinolones have been reported to be 12%, thus necessitating appropriate antibiotic selection in men undergoing prostate biopsy, especially in those at higher risk for infection. (2,21)

Use of antibiograms to select antibiotic prophylaxis—Local antibiograms should be considered when selecting the appropriate prophylaxis because of the prevalent regional variation seen with antibiotic resistance profiles. (22) In our experience at the University of California at Los Angeles (UCLA), men with no significant risk factors for infection receive antibiotic prophylaxis with ciprofloxacin and ceftriaxone. Our current practice is based on the American Urological Association (AUA) best practice policy statement, which recommends fluoroquinolone prophylaxis in men undergoing prostate biopsy. (23) In men identified as high-risk (e.g. immunosuppression, recent antibiotic exposure, diabetes mellitus, or hospitalization), we administer both ciprofloxacin and ertapenem as antibiotic prophylaxis.

The antibiogram at UCLA reports bacteria susceptibilities from urine isolates within the local patient population. The five most common urine isolates are shown in the 2015 antibiogram (Table 1). E.coli, which is only 78% susceptible to ciprofloxacin, is 93% susceptible to ceftriaxone and 99% susceptible to ertapenem. Following the dictates of the most current antibiogram allows us to modify antibiotic prophylaxis for high-risk patients on a contemporaneous basis.

Antibiotic prophylaxis for prostate biopsy—Liss et al. recently described an increased risk of infection (OR 3.98, $p<0.001$) and hospital admission (OR 4.77, $p<0.001$) after transrectal biopsy, in men with fluoroquinolone-resistant positive rectal culture. (24) Targeted antibiotic prophylaxis, based on rectal swab culture results, has been shown to decrease the rate of infection after biopsy. (5) Taylor et al. reported their experience with 22 men found to have positive rectal swab cultures prior to biopsy. (25) All men received targeted antibiotic prophylaxis; there were no infectious complications after biopsy. (25) At UCLA, we augment antibiotic prophylaxis in every patient considered high-risk, based on the above guidelines, thus allowing us to avoid the extra measure of rectal swab cultures.

Transperineal biopsy—Transperineal biopsy is an alternative to the transrectal route, especially because of fewer infections. (26,27) In a recent systematic review of 165 articles on the subject, transrectal biopsy was associated with higher rates of hospitalization (1.1% vs. 0.9%) and sepsis (0.8% vs. 0.1%) compared to transperineal biopsy, while urinary retention was more common using the transperineal approach (4.2% vs 0.9%). (28) Transperineal biopsies are poorly tolerated in a clinic setting, thus general anesthesia is customarily used, which limits the widespread use of this approach.

Repeat biopsies—Bokhorst et al. recently evaluated the risk of complications from serial prostate biopsies in men on active surveillance. (29) The number of previous biopsies did not significantly predict the risk of infection (OR 1.04). (29) The type of antibiotic prophylaxis used was the only significant predictor of infection after biopsy. (29)

Transrectal ultrasound – guided prostate biopsy

Transrectal ultrasound (TRUS) – guided prostate biopsy has for more than three decades been the method for diagnosing prostate cancer. (6) Hodge et al. originally compared the use of TRUS - guided biopsy in a directed versus random systematic manner, ultimately leading to the widespread use of this technique. (30) Several limitations of TRUS - guided biopsy have since been described, including underestimation of tumor grade, inadequate sampling of tumor (e.g. anterior zone), and over-detection of low-risk prostate cancer. (6,7,31)

When whole organs are studied, Gleason score upgrading beyond biopsy findings is common. (8,32) Cohen et al. evaluated radical prostatectomy specimens of 2,890 men diagnosed with prostate cancer via TRUS-guided biopsy. (33) Of those men, 36% of radical prostatectomy specimens revealed a higher grade than previously diagnosed on initial biopsy. (33)

Use of MRI to detect prostate cancer

Since the 1980's, magnetic resonance imaging (MRI) has been used to image the prostate gland. (34) Advances in technology have led to the development of multiparametric MRI (mpMRI), thus allowing enhanced detection of clinically significant prostate cancer (csCaP). (34,35) Multiparametric MRI combines anatomic T2-weighted images with functional and physiological assessments, including diffusion weighted imaging (DWI) and dynamic contrast-enhanced (DCE) MRI. (34,36,37)

In 2012, the Prostate Imaging – Reporting and Data System (PI-RADS) was introduced to standardize the reporting of mpMRI results. (37) PI-RADS utilizes a five-grade scoring system based on the likelihood that mpMRI findings correlate with csCaP within the prostate. (34,37) Grade 1 lesions are “highly unlikely” to contain csCaP, whereas grade 5 lesions are considered “highly likely” to represent csCaP. (34,37)

Using a similar scoring system, the likelihood of detecting csCaP in 825 men with grade 3 regions of interest (ROIs) at UCLA is shown in Figure 1. (10) Men with grade 3 ROIs on mpMRI carry a 24% chance of csCaP on prostate biopsy, compared to an 80% chance of csCaP in those men with grade 5 ROIs. (10)

Recently, Cash et al. reported cancer detection rates (CDRs) and their correlations with PI-RADS score in 408 patients undergoing prostate biopsy. (38) Men with PI-RADS grade 3, 4, and 5 lesions on mpMRI were reported as having clinically significant CDRs of 66% (19/29), 74% (70/94), and 95% (94/99), respectively. (38) Comparable results were reported by Radtke et al., with a positive predictive value of 68% and 81% for PI-RADS 4 and 5 lesions to harbor csCaP, respectively. (39)

The accuracy of PI-RADS version 1.0 has been validated in a meta-analysis with a combined area under the curve (AUC) of 0.82 in receiver operating characteristics curve analysis. (40) However, a learning curve for the PI-RADS scoring system occurs among radiologists for both PI-RADS version 1.0 and the recently updated version 2.0. (41,42) In addition, PI-RADS grading is not as accurate in the transition zone as it is in the peripheral zone. (42)

Applying targeted biopsy cores in MRI suspicious lesions can be prone to a variety of targeting errors. In a cohort of 120 men who underwent radical prostatectomy after MRI – ultrasound fusion biopsy, 8% of the targeted biopsies missed the MRI suspicious lesion. (14) Using rigid co-registration, Hadaschik and Simpfendörfer described a mean targeting error of 2-3mm. (13,43)

MRI – guided prostate biopsy

Suspicious lesions identified on mpMRI can be targeted during fusion biopsy. (44) Three methods of MRI targeted biopsy exist: cognitive fusion, in-bore, and MRI - ultrasound fusion. (44) Cognitive fusion biopsy requires the operator to infer the location of suspicious lesions on MRI using TRUS images. (45) In-bore biopsy is usually performed by a radiologist and involves targeting a suspicious lesion within the MRI tube. (46) Landmarks and targets for biopsy are defined using the preceding diagnostic mpMRI. (47) MRI -

ultrasound fusion biopsy utilizes registration or fusion software to identify and target a lesion on MRI during TRUS - guided biopsy. (46)

Increased detection of significant prostate cancer—Use of MRI – ultrasound fusion to guide biopsy results in increased detection of clinically significant prostate cancer (csCaP). (7,15,48,49) In the largest series reported to date, Filson et al. recently evaluated the combined use of targeted and systematic biopsy in detecting csCaP in 1042 men. (10) Combining both techniques diagnosed more csCaP (n=289) than targeted biopsy (n=229) or systematic biopsy (n=199) alone. (10) ROI grade was the strongest predictor of csCaP on fusion biopsy. (10) When compared to men with a grade 3 ROI, those with a grade 5 ROI were found to have 9 times the odds of csCaP (OR, 9.05). (10) Table 2 shows the likelihood of cancer detection in biopsy naïve patients, depending on degree of suspicion on mpMRI. (10)

Sonn et al. investigated the ability of MRI - ultrasound fusion biopsy to detect prostate cancer in 105 men with prior negative biopsy and persistently elevated prostate-specific antigen (PSA). (11) Targeted and systematic biopsies identified prostate cancer in 36 men (34%) altogether. (11) Using Gleason 3+4 or maximal core length (MCL) 4mm as the definition for clinical significance, 21 of the 23 men (91%) diagnosed with cancer on targeted biopsy had csCaP. (11) By contrast, 15 of the 28 men (54%) had csCaP on conventional systematic biopsy. (11) The combination provided greater sensitivity in detection of csCaP than either alone.

Figure 2 graphically displays current UCLA targeted and systematic biopsy data in 320 men with a prior negative biopsy and persistently elevated PSA undergoing initial MRI - ultrasound fusion biopsy. Targeted biopsy diagnosed csCaP in 73 men, compared to only 53 men diagnosed with csCaP using systematic biopsy alone. Combining the two methods diagnosed csCaP in 86 men.

Siddiqui et al. recently compared the diagnostic accuracy of conventional TRUS - guided biopsy, with MRI - ultrasound fusion prostate biopsy. (7) In their study, targeted biopsy diagnosed 30% more high-risk prostate cancers (173 vs. 122 cases, $p < .001$) and 17% fewer low-risk prostate cancers (213 vs. 258 cases, $p = .002$). (7) Thus, use of targeted biopsy clearly yields an improved detection of clinically significant prostate cancer when compared to conventional systematic biopsy. (7,11)

Screening for active surveillance—Active surveillance is a favorable management strategy for many men with low-risk prostate cancer. (50) In 1994, the Epstein histological criteria were introduced to define clinically insignificant prostate cancer and determine eligibility for active surveillance. (51) Based on their results, clinically insignificant prostate cancer had no Gleason 4 disease, no more than 2 cores involved, and no core with >50% involvement. (51,52) The Epstein histological criteria were determined using biopsy specimens obtained on conventional TRUS biopsy. (51)

Hu et al. recently applied MRI – guided biopsy to men fulfilling the Epstein criteria on conventional biopsy. (52) In this study, 113 men enrolled in the UCLA active surveillance

program who met Epstein criteria at initial diagnosis, subsequently underwent confirmatory biopsy using MRI - ultrasound fusion. (52) Targeted biopsy resulted in reclassification in 41 men (36%) beyond the Epstein criteria. (52) On further analysis of reclassified men, 26 (23%) were reclassified due to Gleason grade 7 and 15 (13.3%) were reclassified due to higher volume Gleason 6 disease. (52) Men with mpMRI ROI grade 4 or 5 were reclassified more often than those men with mpMRI ROI grade 2 or 3 (OR 3.2, $p=0.006$). (52) Figure 3 details these results graphically. (52)

Using the Prostate Cancer Research International Active Surveillance (PRIAS) protocol, data from the Heidelberg group report similar results. (53) Men whose active surveillance was based on an initial TRUS – guided biopsy had a significantly higher probability of upgrading due to pathological progression on MRI – ultrasound fusion confirmatory biopsy, compared to men whose active surveillance was based on initial MRI – ultrasound fusion biopsy. (53) Targeting suspicious lesions on mpMRI may decrease sampling error seen with conventional biopsy, thus allowing accurate detection of clinically insignificant prostate cancer in men on active surveillance. (52)

Tracking of tumors—Sonn et al. investigated the use of an MRI - ultrasound fusion device (Artemis) to electronically monitor specific sites of prostate cancer in men on active surveillance. (54) In this study, 53 men enrolled in an active surveillance program underwent an initial MRI - ultrasound fusion biopsy, followed by a repeat tracking biopsy of prostate cancer specific sites. (54) On initial biopsy, all positive biopsy sites were mapped and tracked to enable specific resampling on subsequent biopsy. (54) Figure 4 illustrates the method of resampling prior positive sites using the MRI - ultrasound fusion device.

Repeat tracking biopsy revealed prostate cancer in 29 sites (39%), of which 14 (49%) had csCaP. (54) Cancer detection rate was directly related to cancer core length (CCL) and presence of tumor within an mpMRI ROI at initial biopsy. (54) On repeat biopsy, there was a 71% CDR when CCL was ≥ 4 mm, compared to a 14% CDR when CCL was < 1 mm. (54) When prostate cancer specific sites were located within an mpMRI ROI and the initial CCL was > 4 mm, 5 of 6 (83%) revealed cancer on repeat biopsy. MRI ROI grade was the strongest predictor of prostate cancer on repeat biopsy (OR 1.48). (54) In men on active surveillance, electronic tracking of specific tumor sites may improve monitoring.

Improved reflection of final pathology—Gleason score concordance from conventional systematic biopsy to radical prostatectomy has been described as weak. (33) Shaw et al. reported a misclassification rate of up to 50% on conventional TRUS – guided prostate biopsy compared to radical prostatectomy specimen. (8) More recently, Le et al. investigated the use of MRI - ultrasound fusion biopsy as a predictor of final pathology on radical prostatectomy specimen in 54 men. (55)

Figure 5 depicts a case scenario of a patient enrolled in the study. MRI - ultrasound fusion included systematic mapping biopsy and targeted biopsy. (55) Each method alone revealed a concordance rate of 54%. (55) In contrast, both modalities combined resulted in a concordance rate of 81%. (55) Combining both modalities significantly improved the predictive accuracy of prostate biopsy. (55)

Radtke et al. recently combined a transperineal saturation biopsy with MRI – ultrasound fusion biopsy. (14) Compared to radical prostatectomy specimen, the combined approach detected 96% of index lesions and 97% of significant multifocal lesions. (14)

Conclusion

Biopsy currently remains the only method to confirm a diagnosis of prostate cancer. (1,56) Common complications of prostate biopsy include hematuria, rectal bleeding, hematospermia, and infection, but these are generally self-limiting. (9,18) In recent years, infectious complications have increased significantly, largely due to an emergence of quinolone-resistant bacteria. (2,20) Judicious selection of antibiotic prophylaxis, based on local antibiograms will decrease the risk of infectious complications. (22,23)

Diagnostic accuracy of prostate biopsy has improved with the use of MRI - ultrasound fusion. (7,10,11,38) When combined with conventional biopsy, the detection of csCaP is significantly enhanced. (7,10) Targeted biopsy also enables more accurate sampling of the highest grade tumor component, which may provide an improvement in selection of candidates for active surveillance. (55) MRI - ultrasound fusion biopsy improves accuracy of prostate cancer detection in several specific applications; however, further research is necessary to define its overall advantage, cost-effectiveness, and decrease target errors. (7,10,11,14,15,38,57)

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Key points

1. Prostate biopsy safely diagnoses prostate cancer with few major, but frequent minor complications.
2. Local antibiograms should be considered when selecting antibiotic prophylaxis.
3. Use of targeted prostate biopsy results in detection of more clinically significant prostate cancer (csCaP) than conventional biopsy.
4. Combining targeted and systematic biopsy via MRI- ultrasound fusion results in greater sensitivity for detection of csCaP than either alone.

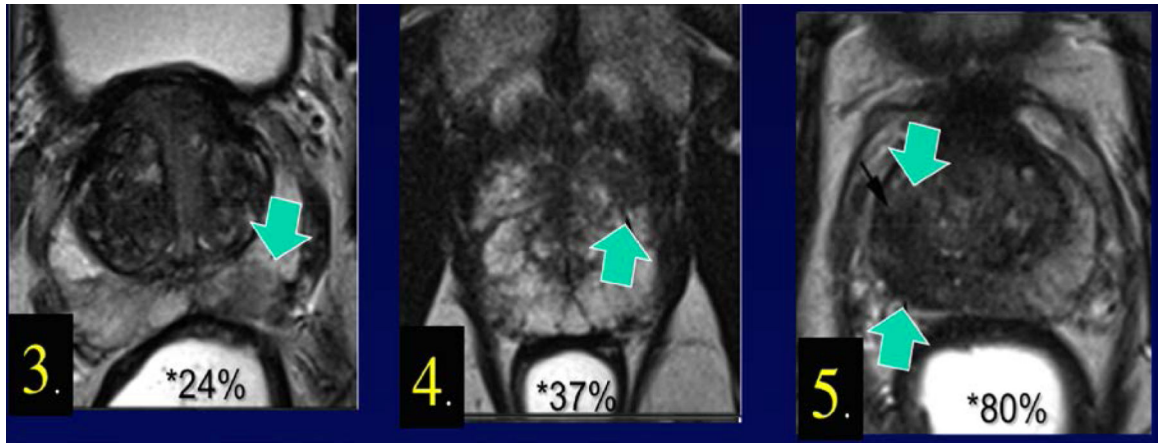


Figure 1. Percent likelihood of clinically significant prostate cancer (csCaP) based on region of interest (ROI) grade, (N=825). (10) Region of interest grade is shown at lower left corner of each section. Chance of csCaP is directly related to ROI grade.

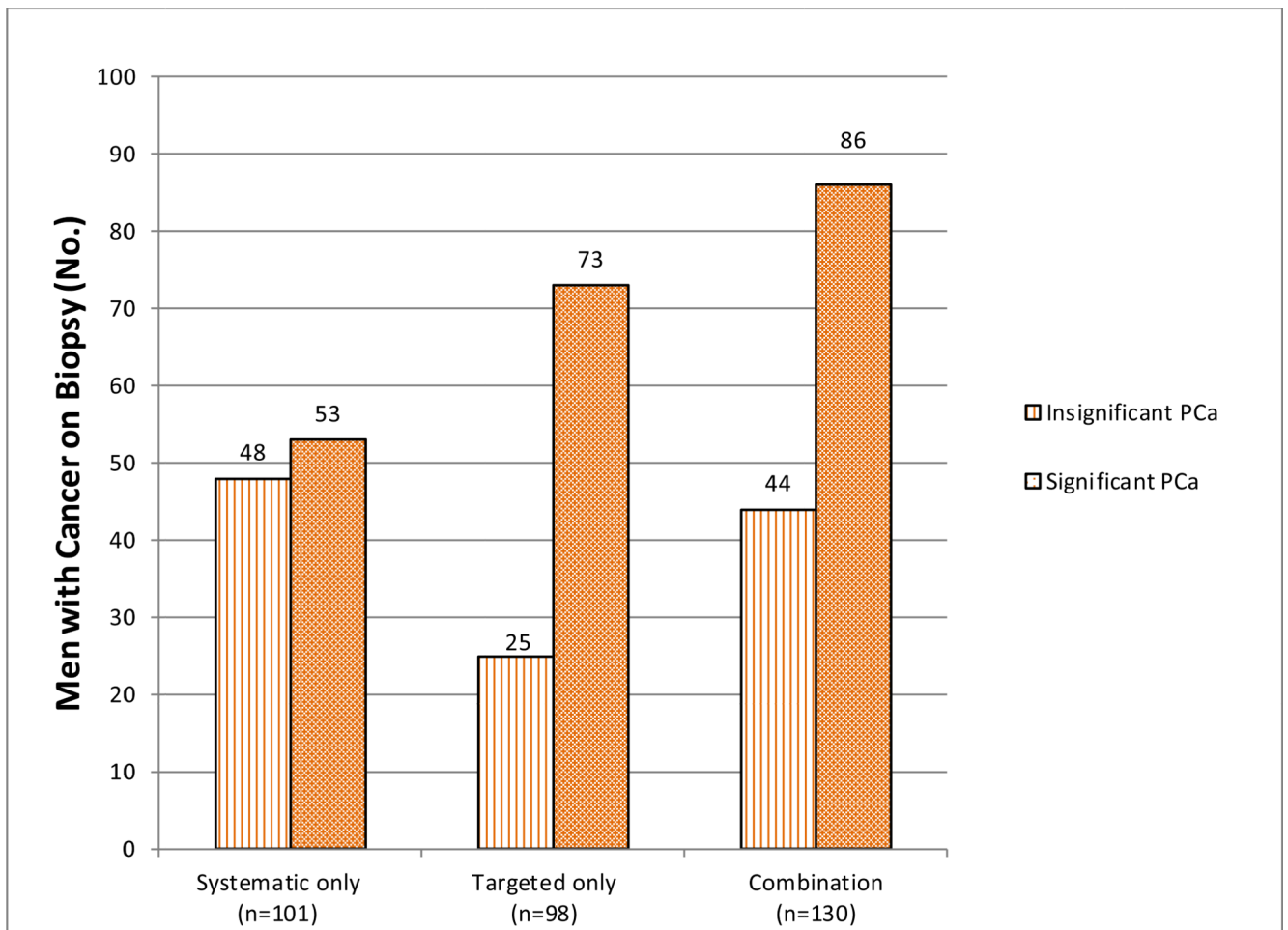


Figure 2.

This graph shows the number of subjects with a prior negative biopsy and persistently elevated PSA diagnosed with significant and insignificant cancers depending on biopsy method. The combination of systematic and targeted biopsy results in detection of more csCaP than either alone.

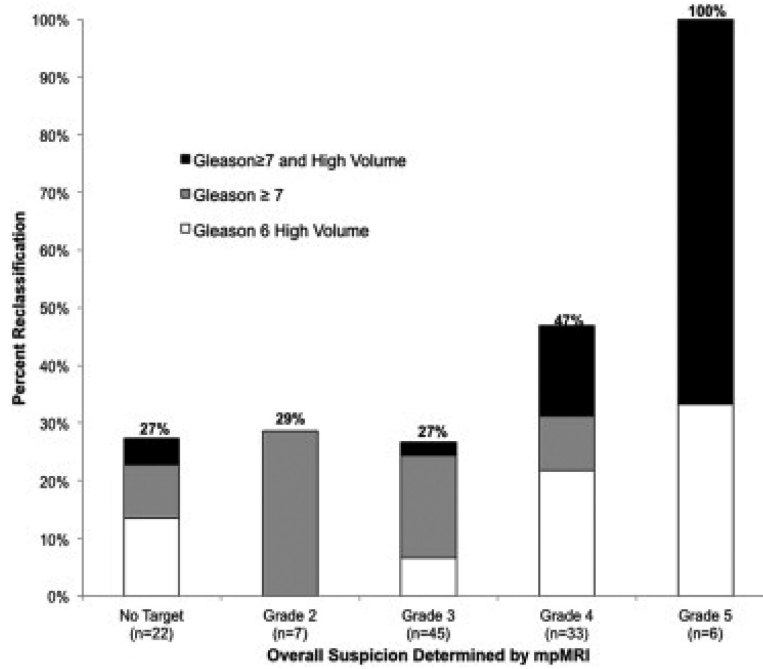


Figure 3. Effect of MRI grade on reclassification beyond Epstein criteria using mpMRI-US biopsy. (52) The higher the suspicion grade (UCLA scoring system), the greater the chance of reclassification beyond traditional Epstein criteria.

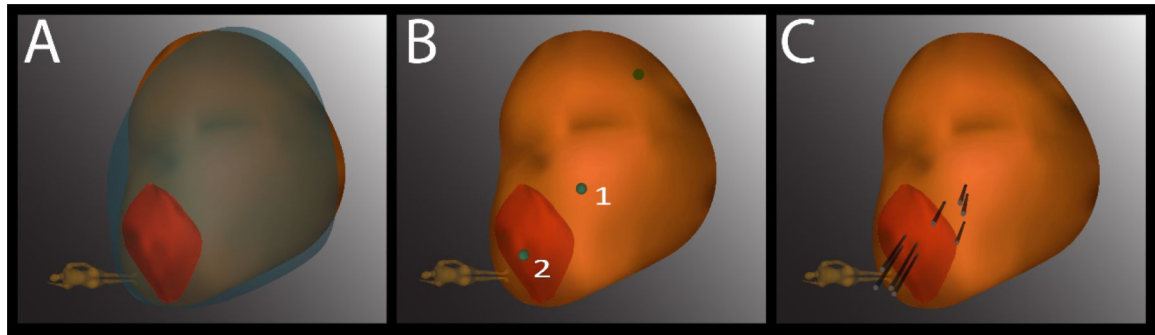


Figure 4.

Example of resampling of prior positive biopsy sites using the MRI - ultrasound fusion (Artemis) device. (A) A 3-dimensional model of the prostate from a second biopsy (brown) is superimposed on the model from a first biopsy (blue), revealing a close match in shape and size. The superimposed model is created in real time at second biopsy. An MRI target (red) is displayed in the model. (B) The location of prior positive sites (1 and 2) is mapped by the Artemis device. Site 1 is a systematic site and site 2 is from the MRI-targeted core. (C) A total of 4 cores (black cylinders) are taken from each site.

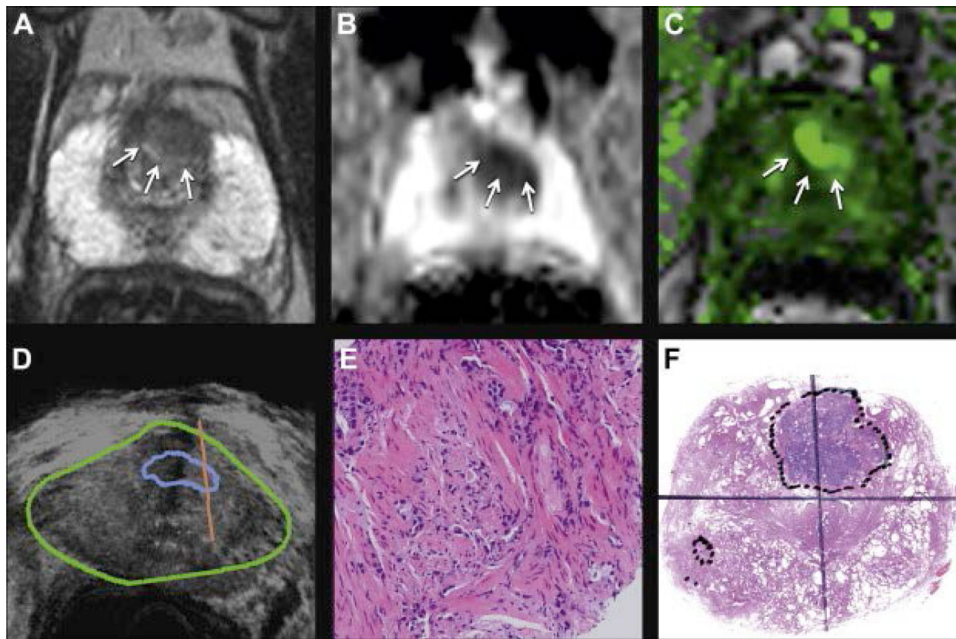


Figure 5. 68 year old man with PSA 8.3 ng/ml underwent mp-MRI, including T2-weighted (A), diffusion weighted (B) and DCE (C) imaging, followed by fusion biopsy (D). Mapping biopsy revealed Gleason 3+3 but targeted biopsy (E) revealed Gleason 4+5 disease (reduced from $\times 20$). GS on whole mount prostatectomy specimen was Gleason 4+3 with tertiary pattern 5 (F, reduced from $\times 1$). Fusion biopsy, which included MRI targeted ROI, predicted highest Gleason grade at final pathology. (55)

Table 1
 Ronald Regan UCLA Hospital Outpatient Antibigram: Adults (>21 y.o.) Gram-negative Bacteria – Urine Isolates, % Susceptible

| Organism | No. Isolates | Ampicillin | Cefazolin | Cefepime | Ceftriaxone | Ertapenem | Imipenem | Meropenem | Gentamicin | Ciprofloxacin | Nitrofurantoin | Trimethoprim-sulfamethoxazole |
|------------------------|--------------|------------|-----------|----------|-------------|-----------|----------|-----------|------------|---------------|----------------|-------------------------------|
| Enterobacter cloacae | 75 | R | R | 96 | - | 89 | 97 | 99 | 97 | 92 | 12 | 85 |
| Escherichia coli | 5379 | 53 | 89 | 96 | 93 | 99 | 99 | 99 | 90 | 78 | 93 | 78 |
| Klebsiella pneumoniae | 523 | R | 91 | 98 | 93 | 99 | 99 | 99 | 97 | 95 | 29 | 86 |
| Proteus mirabilis | 253 | 81 | 94 | 99 | 96 | 98 | 10 | 99 | 92 | 80 | R | 79 |
| Pseudomonas aeruginosa | 118 | R | R | 88 | R | R | 77 | 85 | 96 | 72 | R | R |

Risk of prostate cancer on fusion biopsy based on mpMRI findings in biopsy naïve patients. (10)

Table 2

| Systematic Plus Targeted | | | | | |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|--|
| Maximum ROI grade | Negative | GS 3+3=6 | GS 3+4=7 | GS 4+3=7 | |
| No lesion/grade 1-2 | 38 (68%) | 11 (20%) | 4 (7%) | 3 (5%) | |
| Grade 3 | 60 (46%) | 27 (21%) | 32 (25%) | 10 (8%) | |
| Grade 4 | 36 (33%) | 28 (26%) | 21 (19%) | 24 (22%) | |
| Grade 5 | 1 (3%) | 5 (14%) | 8 (23%) | 21 (60%) | |