Research Article

Why I Cannot Find the Prostate? Behind the Subjectivity of Rectal Exam

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Background. Most physicians use digital rectal examination (DRE) to help detect prostate cancer and to estimate the prostates’ size. The accuracy of DRE is known to be limited. We evaluate the ability of doctors to palpate the whole prostate with DRE. Methods. At time of transrectal ultrasound (TRUS) the distances from the anus to the apex and base of prostates were measured. The TRUS’s distances were compared to the mean index finger length of our clinic doctors. Results. The ability of the urologist to reach and examine the apex, half, three quarters and the whole prostate was in 93.7%, 66.3%, 23.2% and 3.2% of cases respectively. Conclusions. In most cases it was impossible to palpate the whole prostate. Anatomical location and volume of the examined prostate, as well as the length of his own index finger limit DRE and allow the examination of only a small portion of the prostate.

1. Introduction

Digital rectal examination (DRE) is widely used in medicine. A large number of physicians perform it to estimate the prostates’ size and/or for early detection of prostate cancer.

It is well accepted that DRE is a subjective measure and has a high interobserver variability when estimating prostate size. It has been reported that DRE poorly predicted actual prostate size compared to transrectal ultrasound (TRUS) [1]. Others compared radical retropubic prostatectomy specimen weights with prostate weight estimates using TRUS and DRE and found that DRE correlated poorly with prostate weight and that TRUS was superior to DRE [2].

In the prostate specific antigen (PSA) era the role of DRE in early detection of prostate cancer is not clear. This role was evaluated in a number of large trials [3, 4]. The cancer detection rate for DRE was 3.2% and positive predictive value was 21%, PSA or a combination of DRE and PSA were superior to DRE alone for the diagnosis prostate cancer [3]. The positive predictive value and sensitivity of DRE were strongly dependent on PSA level and DRE predicted cancer poorly in patients with low PSA values [4]. Some have suggested that DRE may be unnecessary in patients with PSA values of 3.0 or less [5, 6], since in these patients one would need to perform 289 DREs to find one case of clinically significant prostate cancer.

We believe that in addition to DRE’s subjectivity, there are other inherent limitations to DRE. When performing DRE, prior to reaching any conclusions, the examiner should be able to feel the whole posterior surface of the prostate. However, in clinical practice this may not always be possible. Patient morphometric variables as well as the length of the examiners index finger may impact on the accuracy of DRE. There is no data concerning these factors. One study examined the adequacy of prostate palpation when performing DRE during colonoscopies and concluded that patient positioning and obesity were affecting factors [7]. These observations prompted us to carry out our study.

In this study we tried to explore the limitations of DRE from an anatomical point of view. For this, the distances from anus to the prostate were measured in patients undergoing TRUS. These were compared to the length of index finger of urologists in our clinic. In other words, we examine the ability of urologist to palpate the whole of the prostates posterior surfaces at the time of DRE.
2. Materials and Methods

In a prospective fashion, we examined patients who were referred for prostatic biopsy at our outpatient clinic.

At time of TRUS the distances from the anus to the apex and base of prostates were measured using an 8-MHz biplane probe (B-K medical, 8808 probe). For this purpose the probe was marked with 0.5 cm gradients from the US transverse view crystal out to handle. All TRUS were performed with the patients in the left decubitus position with the knees pulled up to the chest. When the apex of the prostate was viewed in the transverse plain, its depth from the anal verge was noted (anal-apex distance). Then the probe was moved in the cephalic direction until the base of the prostate was visualized and a second measurement was noted (anal-base distance). Index fingers (from the top of the finger to the beginning third interphalangeal joint) of our clinic urologists were measured with a centimeter ruler.

All biopsies were done under local anesthesia using periprostatic block with 20 cc of lidocaine 1%. Systematic transrectal biopsies were obtained using a spring-loaded biopsy gun and 18 G biopsy needle. Patients with BMI 30 or more were excluded from the study.

To analyze the data we divided the prostatic surface length into 4 zones. First is the distal or apical zone that included the distal 25% of the prostate. The second is half prostate, the third zone included 75% of surface, and fourth zone was the whole prostate. The ability of the urologist to palpate all prostatic zones, with DRE was examined comparing the distances measured at the time of TRUS to the mean urologists’ index finger length.

Commercial software (GraphPad Prism) was used for statistical analysis. The results were expressed as mean ± SEM or as median with range. All relationships were assessed by Pearson correlation analysis. Contingency table with Fisher's exact test was used to assess the accuracy of DRE. A level of significance (P value) < 0.05 was considered statistically significant.

3. Results and Discussion

Between March and June 2010, ninety-five men were included in the study. The median age was 64 (range 49–82). Median PSA was 6.94 ng/mL (range 1.56–347). In Thirty-six (38%) men this was not the first biopsy session. The median number of biopsies was 12 (range 8–16). In 15 men (16%) the DRE was suspicious for prostate cancer. Median TRUS prostate volume was 53 mL (range 13.7–301) and the median DRE estimated prostate volume was 40 mL (range 10–80). The correlation between TRUS measured volumes and DRE estimated volumes was not high (Pearson $r = 0.42$, $P < 0.0006$) (Figure 1).

The median anal-apex distance was 5 cm (range 3–7.5), and anal-base distance was 10.3 cm (range 7.3–15.7). The median length of our urologists index fingers was 8.25 cm (range 7–9, $N = 7$). Thus in most cases it was impossible to palpate the whole posterior surface of the prostates. In fact, the ability of the urologist to reach and examine the apex, half prostate, three quarters, and the whole prostate was in 93.7%, 66.3%, 23.2%, and 3.2% cases, respectively. There was a good correlation between anal-base distance and the TRUS volume of prostate (Pearson $r = 0.72$, $P < 0.001$) (Figure 2).

Twenty-nine (30.5%) cases of prostate cancer were diagnosed. The sensitivity and specificity of DRE for the diagnosis of prostate cancer were 21% and 86%, respectively. The positive predictive value of DRE was 40%. According to the Fisher’s exact test the DRE was an inaccurate exam ($P = 0.38$).

This study demonstrated that in most cases it was impossible to palpate the whole posterior surface of the prostate by DRE. This is first time that distance from anal verge to prostate was recorded. Our findings may explain why DRE is a poor predictor of prostate volume and has low sensitivity for the detection prostate cancer.

3.1. DRE and Prostate Volume. The ability to estimate prostate volume is very important before surgical intervention, brachytherapy, benign prostatic hyperplasia management,
and calculating PSA density [8–10]. According to the available literature, DRE is an inaccurate test. DRE underestimates prostate size, particularly if prostate volume is greater than 30 mL [1]. In our study a strong correlation between the anal-base distance and prostate volume was demonstrated. It follows, then, that in larger prostates even more of the prostatic surface would be beyond the reach of the palpating finger making it even harder to estimate its volume. Therefore DRE volume estimations may correlate well with TRUS measured volumes in small glands. In our study we found that DRE estimated volume did not correlate well with TRUS measured volume (Pearson $r = 0.42$). Even worse correlation was reported by Loeb et al. in their large study [2]. Although, in their study they compared DRE estimated prostatic volume with the actual weight of radical prostatectomy specimens. Based on our, and others' findings, DRE is not a good predictor of actual prostatic volume though it may help distinguish small prostates from large ones and may estimate precisely prostatic volumes in patients with small glands.

3.2. DRE and Prostate Cancer Screening. The debate concerning prostate cancer screening is still underway. Even after the publication of the results of large trials from USA [11] and Europe [12] the question “to screen or not to screen” remains unanswered.

In the US, the Prostate, Lung, Colorectal, and Ovarian (PLCO) Cancer Screening Trial, Andriole et al. [11] reported no mortality benefit from combined screening with PSA testing and DRE over a median followup of 11 years. In the European Randomized Study of Screening for Prostate Cancer (ERSPC) trial, Schröder et al. [12] reported that PSA screening without DRE was associated with a 20% relative reduction in the death rate from prostate cancer at a median followup of 9 years, with an absolute reduction of about 7 prostate cancer deaths per 10,000 men screened. In the PLCO study DRE was used as a screening tool whereas in the ERSPC trial DRE was used only in a small portion of the patients. One may conclude then that DRE is not helpful in detection prostate cancer. On the other hand, DRE is an integral part of the neurological function of the lower urinary tract. However, the examiner has to consider factors that limit the accuracy of DRE such as the anatomical location and volume of the examined prostate, as well as the length of his own index finger. One must remember that in many patients these factors limit DRE and allow the examination of only a small portion of the prostate.

4. Conclusions

DRE is still an important and inexpensive tool for the physician. One may gain information about the prostate size and consistency as well as the neurological function of the lower urinary tract. However, the examiner has to consider factors that limit the accuracy of DRE such as the anatomical location and volume of the examined prostate, as well as the length of his own index finger. One must remember that in many patients these factors limit DRE and allow the examination of only a small portion of the prostate.

References


