Objective. To determine the presence of a possible correlation between prostate specific antigen (PSA) and the findings from digital rectal examination (DRE) in patients with prostate cancer or benign prostatic hyperplasia.

Design. Retrospective, longitudinal, and observational study of diagnostic tests.

Setting. Gregorio Marañón Hospital, Madrid, Spain.

Participants. It included 706 patients with a PSA in the range 4.1-20 ng/mL, studied owing to suspected prostate cancer localised using DRE and transrectal ultrasound, in whom randomised prostate biopsies were performed.

Main measurements. Total PSA and free/total PSA ratio and DRE normal or suspicious were studied as main variables. The outcome variable was the diagnosis of prostatic cancer by biopsy.

Results. With a detection of cancer of 28.2%, there were no statistically significant differences in the PSA or free/total PSA ratio mean values between patients with or without suspicious DRE. The analysis using ROC curves (with a 95% confidence interval) between both groups of patients found the same sensitivity of 95% with a similar specificity of 6% and 10%, respectively, for a PSA of 4.8 ng/mL.

Conclusions. In the PSA range of 4.1-20 ng/mL, the findings of DRE appeared as a variable unrelated to the increase in PSA or the free/total PSA ratio and, therefore are not indicative of a lesser or greater volume of a tumour producing PSA. The performing of this examination could be considered as optional.

Key words: Cancer. Prostate. Diagnosis tests.

Validez diagnóstica del tacto rectal en la era del antígeno específico de la próstata

Objetivo. Determinar la posible correlación entre los valores del antígeno específico de la próstata (PSA) y los hallazgos del tacto rectal en pacientes con cáncer prostático y enfermedad prostática benigna.

Diseño. Estudio de evaluación de pruebas diagnósticas de tipo retrospectivo, longitudinal, con carácter observacional.

Emplazamiento. Hospital Gregorio Marañón, Madrid.

Participantes. Se incluyó a 706 pacientes con PSA en el rango de 4,1-20 ng/ml estudiados por sospecha de cáncer prostático localizado mediante tacto rectal y ecografía transrectal, en los que se realizaron biopsias prostáticas aleatorizadas.

Mediciones principales. Se estudiaron como variables predictoras el PSA total, el cociente PSA libre/total y el tacto rectal categorizado como normal o sospechoso de cáncer. La variable desenlace fue el diagnóstico de cáncer prostático obtenido mediante biopsia.

Resultados. Con una detección de cáncer del 28,2%, no se encontraron diferencias estadísticamente significativas en los valores medios de PSA o PSA libre/total entre los pacientes sin/con tacto rectal sospechoso. El análisis mediante curvas ROC (con un intervalo de confianza del 95%) entre ambos grupos de pacientes encontró para el PSA en el valor de 4,8 ng/ml una misma sensibilidad del 95%, con una especificidad similar, del 6 y el 10%, respectivamente.

Conclusiones. En el rango de PSA 4,1-20 ng/ml, los hallazgos del tacto rectal aparecieron como una variable sin relación con la elevación del PSA ni con el cociente PSA libre/total y, por tanto, no son indicativos de un menor o mayor volumen tumoral productor de PSA. La realización de esta exploración podría ser considerada como opcional.

Palabras clave: Cáncer. Próstata. Pruebas diagnósticas.
Introduction

Some authors recognise that prostate specific antigen (PSA) is the most important tumour marker in the wide field of the study of medicine. In fact, the measurement of PSA to detect the presence of prostate cancer is used universally. However, the clinical value of PSA continues to be controversial since, currently, in asymptomatic patients the majority of scientific organisations (including semFYC [Spanish Society of Family and Community Medicine]) through its PAPPS (Programme of Preventive Activities and Health Promotion) group do not recommend prostate cancer screening. The inability of this test on its own to distinguish between benign diseases of the prostate and localised tumours have given rise to unnecessary prostate biopsies in a large number of men and, in many cases, even repeating these biopsies an indeterminate number of times, with all the costs, both financially and emotionally, which this involves. For this reason, although values of PSA between 4 ng/mL and 20 ng/mL are associated with the possibility of having a cancer confined to one organ and, therefore, potentially curable, it must not be forgotten that only a quarter of these cases will have a cancer and, on the other hand, values <4 ng/mL do not necessarily exclude the possibility of having this neoplasia, therefore this potential for false negatives also has to be taken into account.

Different studies have been carried out to determine if the combination of digital rectal examination (DRE) along with PSA measurement is a more useful way of increasing the detection of prostate cancer. In primary care, the reliability of this examination is hampered by the fact that the mean number of prostate consultations is only 3.92 per week, according to a questionnaire sent to 15 000 primary care doctors by the Spanish Prostatic Health Council, with hardly any data on the inter-observer or intra-observer reproducibility of this examination. With the sole use of DRE as an investigational method of prostate cancer, 67% to 88% of these types of neoplasias show up as apparently localised cancer at the time of diagnosis, but in reality this figure is much lower in the final histopathology analysis of operated patients. In the study by the American College of Surgeons in 1999, almost 60% of the males who took part had an apparently localised cancer in a clinical stage at the time of diagnosis. In another study, it has been reported that only 33% of the males studied in the era before PSA with a cancer detected using DRE had a histologically localised disease in the prostate gland. The measurement of PSA as a tumour marker has given rise to a spectacular change in the detection of prostate cancer and has established a reasonable doubt on continuing to systematically use DRE in clinical practice, and which is the objective of this study.

Methods

The study is designed as a retrospective evaluation of diagnostic tests: data was collected from 706 patients, suspected of having prostate cancer with a PSA ranging from 4.1 ng/mL to 20.0 ng/mL, sent from the primary care clinic. These cases were attended to by a reference specialist during the period November 2002-February 2004 in the Gregorio Marañón Hospital, and an ultrasound was performed with random sextant prostate biopsies. The inclusion criteria were: >40 years and a biopsy with a conclusive result. The exclusion criteria were all the situations capable of interfering with the baseline PSA value and previous prostate surgery. DRE, performed in all the cases, was categorised as normal, or suspicious of cancer by the same urologist who carried out the ultrasound. The prostate glands which were only found to be enlarged were not classified as abnormal, in accordance with the aims of this study.

The measurement of PSA in the serum of the patients was carried out, before the biopsy, using an equimolar immunochemiluminescent method (Immulite 2000 PSA, Los Angeles, USA). The free/total PSA fraction was analysed using the solid phase, 2-site sequential chemiluminescent immunomasssay. Additionally, the PSAD (prostate specific antigen density) was measured in all patients by measuring the prostatic volume in all cases before performing the biopsy. The number of biopsies was 6 in each case, independent of the prostate volume present, with the aim of being able to uniformly evaluate the results obtained.

Descriptive statistical analysis was performed on the measured variables, for which the mean, median, and standard deviation was used for the quantitative variables and the absolute and percentage
frequencies for the categorical or qualitative variables. The discriminatory power of PSA for the results benignity or malignancy of the biopsy according to the data from the DRE were evaluated using ROC (receiver operating characteristic) curves, which allowed cut-off points capable of reaching a sensitivity of 95% to be set, and its corresponding specificity was also evaluated. The statistical comparisons between cancer and benign prostatic hyperplasia were performed using the Student t test and the Mann-Whitney test, considering values of \( P < .05 \) as statistically significant.

**Results**

From the total patients studied (\( n=706 \)), the presence of cancer was detected in 199 cases (28.2%). The distribution of total PSA values and the findings of DRE are shown in Table 1. 61.0% of the sample (431/706) had a PSA value between 4 ng/mL and 10 ng/mL. The DRE was suggestive of being benign in 80% of cases (565/706), and a DRE with suspicion of malignancy was obtained in 20% (141/706). These percentages were almost identical (81% and 19%) in the subgroup with PSA values of 4 ng/mL to 10 ng/mL in which the overlap between benign prostatic hyperplasia (BHP) and cancer is normally more frequent.

On analysing the total number of diagnosed cancers (199 adenocarcinomas), the digital examination with suspicion of malignancy found in 141 patients, confirmed this diagnosis in 86 cases (61%) (positive predictive value of the test), but cancer will only be detected regardless of the suspicion in rectal examination in 55 cases (39%). The DRE, apparently normal in 565 patients, did not detect the cancer actually present in 113 cases (20%) and corresponds to an absence of cancer in 452 cases (80%) (negative predictive value). This difference between the results of the DRE and the final diagnosis was significantly different (\( P < .0001 \)), with a sensitivity of 43.2% and a specificity of 89.2% (Figure 1). In the PSA range of 4.1 ng/mL to 20.0 ng/mL studied, the PSA acquired a better specificity (56%) at a cut-off value of 9 ng/mL, but with a sensitivity of only 52%. The analysis of free/total PSA showed an apparent optimal cut-off point at 25%, with a sensitivity of 93%, losing only 7% of the cancers, but with a specificity of only 6%. The PSAD in the standardised cut-off point of 0.15 ng/mL obtained a sensitivity of 82% and a specificity of 45%.

The descriptive statistics values for PSA, the free/total PSA ratio and the PSAD compared to DRE are shown in Table 2. When the total sample of all the included subjects is evaluated, there was no statistically significant difference between the mean value for PSA between patients with a digital examination negative for cancer (9.9 ng/mL) and those in whom cancer was suspected (10.2 ng/mL), and the mean value obtained for the free/total PSA ratio was similar between normal digital examination (15.5%) and those who had a digital examination with a suspicion of cancer (15.0%).

The ROC curves of the PSA in patients with digital examination indicative of benignity and in the cases of digital examination with a suspicion of neoplasia were analysed. The areas under the curve obtained in patients with normal or abnormal DRE were 0.516 and 0.608, respectively, with overlapping 95% confidence interval values. The cut-off point required to obtain 95% sensitivity was the same in both groups (4.8 ng/mL) with a specificity of 6% and 10%, which meant that the information contributed by DRE did not have a significant impact on the results obtained over the detection of cancer using PSA (Figure 2).
The debate on whether biopsy has to be recommended in males with a DRE with no suspicion of cancer and a PSA between 4 ng/mL and 10 ng/mL, the common range for BPH and localised cancer, has been the subject of controversy for a long time. For a decade it was accepted, with an almost total consensus, that a prostate biopsy had to be indicated when the DRE was suspicious or the PSA was >10 ng/mL. However, since the publication of the multicentre study led when the DRE was suspicious or the PSA was >10 ng/mL. Thus, in a large multicentre study carried out on 6630 males, in 18% of those who had a biopsy performed, this idea had to change, since in those cases with PSA values of 4 ng/mL and 10 ng/mL with a digital rectal examination not indicative of malignancy 21% were found with cancer, a similar percentage to that found in the cases with a suspicious digital examination (21%).

During all this process it is evident that the primary care doctor plays a fundamental role. Some publications show that, in certain health environments outside our country, the family doctor systematically carries out the DRE in 84% of patients >50 years who consult with micturation symptoms of the lower urinary tract, and the PSA is also frequently measured in these patients.

Thus, in a large multicentre study carried out in our country with 587 biopsied patients, the total PSA was shown to be a more sensitive technique for the detection of cancer than a DRE only, since of the 131 tumours detected according to the PSA value, 41 (31.2%) would not have been detected if DRE had been the only criteria to indicate biopsy. However, the positive predictor value of digital examination was 43.7%, higher than the PSA (24% for a cut-off of 4 ng/mL). Only a PSA value >10 ng/mL in the aforementioned study had a positive predictive value similar to digital examination (46.2%). This increased positive predictive value for digital examination contrasts with data published by other authors, who obtained a positive predictive value of 22%–39%. Thus, Benson et al mention 46.3% and Babaian et al, 51%, and a better positive predictive value with the combination of the 2 parameters being quoted in the majority of series reviewed. In a study of prostate biopsies it was found that when the palpable nodule in the DRE was unilateral, the probability of finding a positive biopsy was similar, in the side supposedly affected as well as in the other side, concluding that the detection of these cancers could be due to chance. As regards studies carried out on large population groups with PSA>4 ng/mL, in the Prostate Cancer Prevention Trial, which studied 2950 patients with a negative DRE and a biopsy performed on all of them (with a PSA in that range it is considered “normal”), as well as in the study carried out by Carvalhal et al in 2703 males with a DRE with a suspicion of cancer and a PSA<4 ng/mL, they found fairly similar rates of cancer detection (15.2% and 13.9%, respectively).

As regards the PSA value in the 4-10 ng/mL range to detect a suspicious digital examination (AUC=0.704).

**Discussion**

The debate on whether biopsy has to be recommended in males with a DRE with no suspicion of cancer and a PSA between 4 ng/mL and 10 ng/mL, the common range for BPH and localised cancer, has been the subject of controversy for a long time. For a decade it was accepted, with an almost total consensus, that a prostate biopsy had to be indicated when the DRE was suspicious or the PSA was >10 ng/mL.
On comparing population studies with PSA, the positive predictive value of DRE, although the findings of DRE did not correlate with the prostate weight. By omitting the DRE there is a low risk of missing potentially curable cancers.

Thus, although it is admitted that prostate screening which includes/without DRE suspicion of cancer. The positive predictive value of DRE, although being high (61%), cannot identify 20% of the cancers present. The findings of DRE did not correlate with the PSA values and they are not indicative of the tumour volume in localised cancer.

What This Study Contributes

- In the 4-20 ng/mL PSA range, the PSA mean appears to be the same between patients with/without DRE suspicion of cancer.
- The positive predictive value of DRE, although being high (61%), cannot identify 20% of the cancers present.
- The findings of DRE did not correlate with the PSA values and they are not indicative of the tumour volume in localised cancer.

What Is Known About the Subject

- PSA is a more sensitive technique than digital rectal examination for the detection of cancer.
- On comparing population studies with PSA <4 ng/mL, the detection of cancer is similar with/without suspicious DRE.
- By omitting the DRE there is a low risk of missing potentially curable cancers.

Key points

- PSA is a more sensitive technique than digital rectal examination for the detection of cancer.
- On comparing population studies with PSA <4 ng/mL, the detection of cancer is similar with/without suspicious DRE.
- By omitting the DRE there is a low risk of missing potentially curable cancers.

References

Prostate specific antigen (PSA) is a protein synthesised in the prostate tissue. Its main function is to liquefy the seminal coagulate and facilitate the transport of spermatozoa through the vas deferens to the urethra. Its application in clinical practice is based on its use as an immunological marker of prostate cancer, both in screening and in the follow up of this neoplasia. However, on being an organ specific protein and not a tumour specific one, it can be elevated without there being prostate cancer. Thus, for certain values of PSA it does not have an increased specificity and does not allow a clear distinction between benign prostatic hyperplasia (BPH) and localised prostate neoplasia.1

With the aim of increasing the specificity of this screening test, that is, to diagnose the same number of neoplasias but reducing the number of negative biopsies and the consequent anxiety which arises from this suspected diagnosis in patients, different ways of quantifying serum PSA values have been proposed. These different ways of evaluating PSA2,3 have a special use in the so-called “grey area” (or maximum overlap between the possibility that the increased PSA may be the consequence of benign or malignant prostatic disease), which is situated in the values of PSA between 4 ng/mL and 10 ng/mL.

Although some scientific societies recommend the use of PSA for the screening of prostate cancer, other societies do not recommend its use, because they consider that, currently, there is not sufficient proof to demonstrate that a screening programme might have an impact on the morbidity and mortality due to prostate cancer. In our country the Programme of Preventive Activities and Health Promotion (PAPPS) and

Key Points

- PSA is an organ-specific protein and is not tumour-specific.
- The Programme of Preventive Activities and Health Promotion (PAPPS) and the Spanish Society of Family and Community Medicine do not recommend the use of PSA as a screen for prostate cancer.
- Digital rectal examination is a very useful examination, by which we obtain information which we can only obtain with this examination.
the Spanish Society of Family and Community Medicine do not recommend it.\(^4\)

A recent study published in JAMA points out that no PSA value offered sufficient reliability for screening and demonstrates that at no cut-off point can a specificity and sensitivity be found at least reasonably close.\(^5\)

On the other hand, what effect the fact of adding digital rectal examination to the PSA has on the positive predictive value, the sensitivity and specificity can be seen (Table).\(^6\) Thus, if the patient has opted for screening for prostate cancer, this must be optimised by looking to increase the sensitivity by using the PSA together with digital rectal examination. Digital rectal examination is a very useful examination, easy to perform, by which we obtain information which we can only obtain by carrying out this examination, and will be important when deciding subsequent action (diagnosis and treatment). It is a straightforward examination within the capabilities of all doctors. The argument of inexperience is not valid, given that it is an examination which should be performed frequently, and it will be this frequency which will enable us to have and maintain the necessary skill. Digital rectal examination is an essential element within the basic package of physical examinations carried out by any primary care doctor.

With the performing of digital rectal examination we will obtain data which will enable us to evaluate the morphology, the size, the consistency, the mobility, the regularity of its limits, the presence of nodules and the sensitivity of the prostate.\(^7\)

### References