

## **Total Prostate Specific Antigen** (tPSA)

Product Code: 2125-300

Intended Use: The Quantitative Determination of Total Prostrate Specific Antigen (tPSA) Concentration in Human Serum by a Microplate Immunoenzymometric assay

## **SUMMARY AND EXPLANATION OF THE TEST**

Prostate Specific antigen (PSA) is a serine protease with chymotrypsin-like activity (1,2). The protein is a single chain glycoprotein with a molecular weight of 28.4 kDA (3). PSA derives its name from the observation that it is a normal antigen of the prostrate but is not found in any other normal or malignant tissue

PSA is found in benign, malignant and metastatic prostrate cancer. Since prostate cancer is the second most prevalent form of male malignancy, the detection of elevated PSA levels plays an important role in the early diagnosis. Serum PSA levels have been found too more useful than prostatic acid phosphatase (PAP) in the diagnosis and management of patients due to increased sensitivity (4).

In this method, PSA calibrator, patient specimen or control is first added to a streptavidin coated well. Biotinylated monoclonal and enzyme labeled antibodies (directed against distinct and different epitopes of PSA) are added and the reactants mixed. Reaction between the various PSA antibodies and native PSA forms a sandwich complex that binds with the streptavidin coated to the well.

After the completion of the required incubation period, the enzyme-PSA antibody bound conjugate is separated from the unbound enzyme-PSA conjugate by aspiration or decantation. The activity of the enzyme present on the surface of the well is quantitated by reaction with a suitable substrate to produce

The employment of several serum references of known prostate specific antigen (PSA) levels permits the construction of a dose response curve of activity and concentration. From comparison to the dose response curve, an unknown specimen's activity can be correlated with PSA concentration.

### PRINCIPLE

## Immunoenzymometric assay (TYPE 3):

The essential reagents required for an immunoenzymometric assay include high affinity and specificity antibodies (enzyme and immobilized), with different and distinct epitope recognition, in excess, and native antigen. In this procedure, the immobilization takes place during the assay at the surface of a microplate well through the interaction of streptavidin coated on the well and exogenously added biotinylated monoclonal anti-PSA antibody.

Upon mixing monoclonal biotinylated antibody, the enzymelabeled antibody and a serum containing the native antigen, reaction results between the native antigen and the antibodies, without competition or steric hindrance, to form a soluble sandwich complex. The interaction is illustrated by the following

Btn Ab<sub>(m)</sub> =Biotinylated Antibody (Excess Quantity)

Ag<sub>PSA</sub> = Native Antigen (Variable Quantity)

EnzAb<sub>(p)</sub> = Enzyme labeled Antibody (Excess Quantity)

 $Enz_{Ab_{(p)}}Ag_{PSA}$ - $Btn_{Ab_{(m)}}$ =Antigen-Antibodies Complex

k<sub>2</sub> = Rate Constant of Association

k<sub>-a</sub> = Rate Constant of Dissociation

Simultaneously, the complex is deposited to the well through the high affinity reaction of streptavidin and biotinylated antibody. This interaction is illustrated below:

Enz Ab<sub>(D)</sub>-Ag<sub>PSA</sub>-Btn Ab<sub>(m)</sub>+Streptavidin<sub>C.W.</sub>⇒Immobilized complex

Streptavidin<sub>C.W.</sub> = Streptavidin immobilized on well

Immobilized complex = complex bound to the solid surface

After equilibrium is attained, the antibody-bound fraction is separated from unbound antigen by decantation or aspiration. The enzyme activity in the antibody-bound fraction is directly proportional to the native antigen concentration. By utilizing several different serum references of known antigen values, a dose response curve can be generated from which the antigen concentration of an unknown can be ascertained.

### REAGENTS

#### Materials Provided:

A. Prostrate Specific antigen (PSA) 1ml/vial - Icons A-F Six (6) vials of references PSA Antigen at levels of O(A), 5(B), 10(C), 25(D), 50(E) and 100(F) ng/ml. Store at 2-8°C. A preservative has been added.

> Note: The calibrators, human serum based, were calibrated using a reference preparation, which was assayed against the 1st IS 96/670.

B. PSA Enzyme Reagent —13ml/vial - Icon One (1) vial containing enzyme labeled antibody,

biotinylated monoclonal mouse IgG in buffer, dye, and preservative. Store at 2-8°C.

## C. Streptavidin Coated Plate -- 96 wells - Icon

One 96-well microplate coated with streptavidin and packaged in an aluminum bag with a drying agent. Store

## D. Wash Solution - 20 ml - Icon

One (1) vial containing a surfactant in buffered saline. A preservative has been added. Store at 2-30°C.

## E. Substrate A --7ml/vial - Icon SA

One (1) bottle containing tetramethylbenzidine (TMB) in buffer. Store at 2-8°C.

## F. Substrate B -- 7ml/vial - Icon SB

One (1) bottle containing hydrogen peroxide (H2O2) in buffer. Store at 2-8°C.

# G. Stop Solution -- 8ml/vial - Icon



One (1) bottle containing a strong acid (1N HCI). Store at

## H. Product Instructions.

Note 1: Do not use reagents beyond the kit expiration date. Note 2: Opened reagents are stable for sixty (60) days when stored at 2-8°C.

Note 3: Above reagents are for a single 96-well microplate

### Required But Not Provided:

- 1. Pipette(s) capable of delivering 25 & 50µl volumes with a precision of better than 1.5%.
- 2. Dispenser(s) for repetitive deliveries of 0.100ml and 0.300ml volumes with a precision of better than 1.5%.
- 3. Microplate washers or a squeeze bottle (optional).
- 4. Microplate Reader with 450nm and 620nm wavelength absorbance capability.
- Absorbent Paper for blotting the microplate wells.
- 6. Plastic wrap or microplate cover for incubation steps.
- Vacuum aspirator (optional) for wash steps.
- 9. Quality control materials

### **PRECAUTIONS**

#### For In Vitro Diagnostic Use Not for Internal or External Use in Humans or Animals

All products that contain human serum have been found to be non-reactive for Hepatitis B Surface Antigen, HIV 1&2 and HCV Antibodies by FDA licensed reagents. Since no known test can offer complete assurance that infectious agents are absent, all human serum products should be handled as potentially hazardous and capable of transmitting disease. laboratory procedures for handling blood products can be found in the Center for Disease Control / National Institute of Health. "Biosafety in Microbiological and Biomedical Laboratories," 2nd Edition, 1988, HHS Publication No. (CDC) 88-8395.

## SPECIMEN COLLECTION AND PREPARATION

The specimens shall be blood, serum in type and the usual precautions in the collection of venipuncture samples should be observed. For accurate comparison to established normal values, a fasting morning serum sample should be obtained. The blood should be collected in a plain redtop venipuncture tube without additives or anti-coagulants. Allow the blood to clot. Centrifuge the specimen to separate the serum from the cells.

Samples may be refrigerated at 2-8°C for a maximum period of five (5) days. If the specimen(s) cannot be assayed within this time, the sample(s) may be stored at temperatures of -20°C for up to 30 days. Avoid repetitive freezing and thawing. When assayed in duplicate, 0.050ml of the specimen is required.

## **QUALITY CONTROL**

Each laboratory should assay controls at levels in the low, normal and elevated range for monitoring assay performance. These controls should be treated as unknowns and values determined in every test procedure performed. Quality control charts should be maintained to follow the performance of the supplied reagents. Pertinent statistical methods should be employed to ascertain trends. Significant deviation from established performance can indicate unnoticed change in experimental conditions or degradation of kit reagents. Fresh reagents should be used to determine the reason for the variations

## REAGENT PREPARATION:

## 1 Wash Buffer

Dilute contents of wash solution to 1000ml with distilled or deionized water in a suitable storage container. Store at room temperature 20-27°C for up to 60 days.

## 2. Working Substrate Solution

Pour the contents of the amber vial labeled Solution 'A' into the clear vial labeled Solution 'B'. Place the yellow cap on the clear vial for easy identification. Mix and label accordingly. Store at 2 - 8°C.

Note: Do not use the working substrate if it looks blue.

## **TEST PROCEDURE**

Before proceeding with the assay, bring all reagents, serum references and controls to room temperature (20 - 27°C).

- 1. Format the microplates' wells for each serum reference. control and patient specimen to be assayed in duplicate. Replace any unused microwell strips back into the aluminum bag, seal and store at 2-8°C.
- 2. Pipette 0.025 ml (25µl) of the appropriate serum reference, control or specimen into the assigned well.
- 3. Add 0.100 ml (100ul) of the PSA Enzyme Reagent to each well. It is very important to dispense all reagents close to the bottom of the coated well.
- 4. Swirl the microplate gently for 20-30 seconds to mix and
- 5. Incubate 30 minutes at room temperature.
- 6. Discard the contents of the microplate by decantation or aspiration. If decanting, tap and blot the plate dry with
- 7. Add 300µl of wash buffer (see Reagent Preparation Section), decant (tap and blot) or aspirate. Repeat two (2) additional times for a total of three (3) washes. An automatic or manual plate washer can be used. Follow the manufacturer's instruction for proper usage. If a squeeze bottle is employed, fill each well by depressing the container (avoiding air bubbles) to dispense the wash. Decant the wash and repeat two (2) additional
- 8. Add 0.100 ml (100µl) of working substrate solution to all wells (see Reagent Preparation Section). Always add reagents in the same order to minimize reaction time differences between wells.

## DO NOT SHAKE THE PLATE AFTER SUBSTRATE ADDITION

- 9. Incubate at room temperature for fifteen (15) minutes.
- 10. Add 0.050ml (50µl) of stop solution to each well and mix gently for 15-20 seconds. Always add reagents in the same order to minimize reaction time differences
- 11. Read the absorbance in each well at 450nm (using a reference wavelength of 620-630nm to minimize well imperfections) in a microplate reader. The results should be read within thirty (30) minutes of adding the stop solution.

## **CALCULATION OF RESULTS**

A dose response curve is used to ascertain the concentration of PSA in unknown specimens

- 1. Record the absorbance obtained from the printout of the microplate reader as outlined in Example 1.
- 2. Plot the absorbance for each duplicate serum reference versus the corresponding PSA concentration in ng/ml on linear graph paper (do not average the duplicates of the serum references before plotting).
- 3. Draw the best-fit curve through the plotted points.
- 4. To determine the concentration of PSA for an unknown, locate the average absorbance of the duplicates for each unknown on the vertical axis of the graph, find the intersecting point on the curve, and read the concentration (in ng/ml) from the horizontal axis of the graph (the duplicates of the unknown may be averaged as indicated). In the following example, the average absorbance (1.142) intersects the dose response curve at (23.6 ng/ml) PSA concentration (See Figure 1).

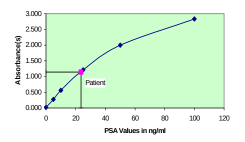
Note: Computer data reduction software designed for ELISA assays may also be used for the data reduction.

### **EXAMPLE 1**

| Sample<br>I.D.  | Well<br>Number | Abs (A) | Mean<br>Abs (B) | Value<br>(ng/ml) |
|-----------------|----------------|---------|-----------------|------------------|
| Cal A           | A1             | 0.019   | 0.019           | 0                |
|                 | B1             | 0.019   |                 |                  |
| Cal B           | C1             | 0.279   | 0.276           | 5                |
|                 | D1             | 0.273   |                 |                  |
| Cal C           | E1             | 0.567   | 0.563           | 10               |
|                 | F1             | 0.559   |                 |                  |
| Cal D           | G1             | 1.248   | 1.213           | 25               |
| <b>*</b> **** - | H1             | 1.179   |                 |                  |
| Cal E           | A2             | 2.051   | 1.999           | 50               |
| -               | B2             | 1.947   |                 |                  |
| Cal F           | C2             | 2.892   | 2.833           | 100              |
|                 | D2             | 2.775   |                 | . 30             |
| Patient         | E3             | 1.186   | 1.142           | 23.6             |
|                 | F3             | 1.099   |                 | _3.0             |

\*The data presented in Example 1 and Figure 1 is for illustration only and **should not** be used in lieu of a dose response curve prepared with each assay.

Figure 1



## Q.C. PARAMETERS

In order for the assay results to be considered valid the following criteria should be met:

- 1. The absorbance (OD) of calibrator F should be  $\geq$  1.3.
- Four out of six quality control pools should be within the established ranges.

### LIMITATIONS OF PROCEDURE

### A. Assay Performance

- It is important that the time of reaction in each well is held constant for reproducible results. Pipetting of samples should not extend beyond ten (10) minutes to avoid assay drift. If more than one (1) plate is used, it is recommended to repeat the dose response curve.
- Addition of the substrate solution initiates a kinetic reaction, which is terminated by the addition of the stop solution. Therefore, the addition of the substrate and the stopping solution should be added in the same sequence to eliminate any time deviation during reaction.
- 3. Plate readers measure vertically. Do not touch the bottom of
- Failure to remove adhering solution adequately in the aspiration or decantation wash step(s) may result in poor replication and spurious results.
- Sample(s), which are contaminated microbiologically, should not be used in the assay. Highly lipemeic or hemolysed specimen(s) should similarly not be used.
- Patient specimens with PSA concentrations above 100 ng/ml may be diluted (for example 1/10 or higher) with normal female serum (PSA = 0 ng/ml) and re-assayed. The sample's concentration is obtained by multiplying the result by the dilution factor (10).

## B. Interpretation

- If computer controlled data reduction is used to interpret the results of the test, it is imperative that the predicted values for the calibrators fall within 10% of the assigned concentrations
- 2. PSA is elevated in benign prostrate hypertrophy (BPH). Clinically an elevated PSA value alone is not of diagnostic value as a specific test for cancer and should only be used in conjunction with other clinical manifestations (observations) and diagnostic procedures (prostate biopsy). Free PSA determinations may be helpful in regard to the discrimination of BPH and prostrate cancer conditions (5).

## **EXPECTED RANGES OF VALUES**

Healthy males are expected to have values below 4 ng/ml (4).

# TABLE I Expected Values for the PSA Elisa Test System

Healthy Males <4 ng/ml

It is important to keep in mind that establishment of a range of values which can be expected to be found by a given method for a population of "normal"-persons is dependent upon a multiplicity of factors: the specificity of the method, the population tested and the precision of the method in the hands of the analyst. For these reasons each laboratory should depend upon the range of expected values established by the Manufacturer only until an in-house range can be determined by the analysts using the method with a population indigenous to the area in which the laboratory is located.

### PERFORMANCE CHARACTERISTICS

#### A. Precision

The within and between assay precisions of the tPSA AccuBindTM ELISA test system were determined by analyses on three different levels of control sera. The number, mean value, standard deviation and coefficient of variation for each of these control sera are presented in Table 2 and Table 3.

TABLE 2 Within Assay Precision (Values in ng/ml)

| Sample  | N  | Х    | S.D. | C.V. |
|---------|----|------|------|------|
| Level 1 | 20 | 0.7  | 0.05 | 7.1% |
| Level 2 | 20 | 4.5  | 0.20 | 4.4% |
| Level 3 | 20 | 28.3 | 1.07 | 3.7% |

TABLE 3
Between Assay Precision\* (Values in ng/ml)

| Sample  | N  | x    | S.D. | C.V.  |
|---------|----|------|------|-------|
| Level 1 | 10 | 0.8  | 0.09 | 11.3% |
| Level 2 | 10 | 4.3  | 0.25 | 5.8%  |
| Level 3 | 10 | 27.5 | 1.42 | 5.2%  |

<sup>\*</sup>As measured in ten experiments in duplicate.

### B. Sensitivity

The tPSA AccuBind™ ELISA test system has a sensitivity of 0.012 ng. This is equivalent to a sample containing 0.5 ng/ml tPSA concentration.

#### C. Accuracy

The tPSA AccuBind™ ELISA method was compared with a reference Elisa method. Biological specimens from low, normal, and elevated concentrations were assayed. The total number of such specimens was 241. The least square regression equation and the correlation coefficient were computed for the tPSA AccuBind™ ELISA test method in comparison with the reference method. The data obtained is displayed in Table 4.

| TABLE 4         |              |                       |             |
|-----------------|--------------|-----------------------|-------------|
|                 | Least Square |                       |             |
|                 |              | Regression            | Correlation |
| Method          | Mean         | Analysis              | Coefficient |
| This Method (X) | 5.62         | y = -0.0598 + 0.98(X) | 0.987       |
| Reference (Y)   | 5.57         |                       |             |

Only slight amounts of bias between the tPSA AccuBind $^{TM}$  ELISA methodand the reference method are indicated by the closeness of the mean values. The least square regression equation and correlation coefficient indicates excellent method agreement.

## E. Specificity:

No interference was detected with the performance of tPSA AccuBind™ ELISA test system upon addition of massive amounts of the following substances to a human serum pool.

| Acetylsalicylic Acid | 100 µg/ml   |
|----------------------|-------------|
| Ascorbic Acid        | 100 µg/ml   |
| Caffeine             | 100 µg/ml   |
| CEA                  | 10 μg/ml    |
| AFP                  | 10 μg/ml    |
| CA-125               | 10,000 U/ml |
| hCG                  | 1000 IU/ml  |
| hLH                  | 10 IU/ml    |
| hTSH                 | 100 mIU/ml  |
| hPRL                 | 100 μg/ml   |

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| Size           |    | 96(A)    | 192(B)      |
|----------------|----|----------|-------------|
| Reagent (fill) | A) | 1ml set  | 1ml<br>set  |
|                | B) | 1 (13ml) | 2<br>(13ml) |
|                | C) | 1 plate  | 2<br>plates |
|                | D) | 1 (20ml) | 1<br>(20ml) |
|                | E) | 1 (7ml)  | 2<br>(7ml)  |
|                | F) | 1 (7ml)  | 2<br>(7ml)  |
|                | G) | 1 (8ml)  | 2<br>(8ml)  |

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