Thromboxane $B_2$ ELISA Kit Instructions

Please read all instructions carefully before beginning this assay

PRODUCT #405110
For research use only.

Storage Conditions:
Do not freeze kit components
All other kit components: 4°C

DESCRIPTION
Thromboxane $B_2$ (TXB$_2$) is a stable hydrolyzed product of unstable TXA$_2$ which is derived from PGH$_2$. PGH$_2$ is synthesized from arachidonic acid through the cyclooxygenase pathway. It is a major product following platelet aggregation induced by a variety of agents such as thrombin and collagen. It is produced not only in platelets, but also in other cell types such as fibroblasts and macrophages. Quantitation of thromboxane formation can be made by determining the level of TXB$_2$.

PRINCIPLE OF ASSAY
This is an ELISA (Enzyme-Linked Immunosorbent Assay) for the quantitative analysis of Thromboxane $B_2$ levels in biological fluid. This test kit operates on the basis of competition between the enzyme conjugate and the TXB$_2$ in the sample for a limited number of binding sites on the antibody coated plate.

The sample or standard solution is first added to the microplate. Next, the diluted enzyme conjugate is added and the mixture is shaken and incubated at room temperature for one hour. During the incubation, competition for binding sites is taking place. The plate is then washed removing all the unbound material. The bound enzyme conjugate is detected by the addition of substrate which generates an optimal color after 30 minutes. Quantitative test results may be obtained by measuring and comparing the absorbance reading of the wells of the samples against the standards with a microplate reader at 650 nm. The extent of color development is inversely proportional to the amount of TXB$_2$ in the sample or standard. For example, the absence of TXB$_2$ in the sample will result in a bright blue color, whereas the presence of TXB$_2$ will result in decreased or no color development.
**MATERIALS PROVIDED**

1. **EIA BUFFER:** 30 mL. Provided to dilute enzyme conjugate and TXB₂ standards.
2. **WASH BUFFER (10X):** 20 mL. To be diluted 10-fold with deionized water. This is used to wash all unbound enzyme conjugate, samples and standards from the plate after the one hour incubation.
3. **K-BLUE SUBSTRATE:** 20 mL. Stabilized 3,3’, 5,5’ Tetramethylbenzidine (TMB) plus Hydrogen Peroxide (H₂O₂) in a single bottle. It is used to develop the color in the wells after they have been washed. **LIGHT SENSITIVE.** Keep substrate refrigerated.
4. **EXTRACTION BUFFER (5X):** 30 mL. To be diluted 5-fold with deionized water. This is used for diluting extracted and non-extracted samples.
5. **THROMBOXANE B₂ ENZYME CONJUGATE:** 150 µL. TXB₂ horseradish peroxidase concentrate. Blue capped vial.
6. **THROMBOXANE B₂ STANDARD:** 100 µL. TXB₂ standard at the concentration of 1 µg/mL. Green capped vial.
7. **THROMBOXANE B₂ ANTIBODY-COATED MICROPLATE:** A 96 well MaxiSorp™ Nunc microplate with anti-TXB₂ rabbit antibody precoated on each well. The plate is ready for use as is. **DO NOT WASH!**

**MATERIALS NEEDED BUT NOT PROVIDED**

1. 300 mL deionized water for diluting wash buffer and extraction buffer.
2. Precision pipettes that range from 10 µL-1000 µL and disposable tips.

**NOTE:** If all or several strips are to be used at one time, it is suggested that a multichannel pipette be used.

3. Clean test tubes used to dilute the standards and conjugate.
4. Graduated cylinders to dilute and mix wash buffer and extraction buffer.
5. Microplate reader with 650 nm filter.
6. Plate cover or plastic film to cover plate during incubation.
OPTIONAL MATERIALS:

7. 1 N HCl or Neogen’s Red Stop Solution.
8. Microplate shaker.

If performing an extraction on samples, the following will be required:

9. Methanol
10. Methyl formate
11. 0.1 M Sodium Phosphate buffer, pH 7.5
12. C_{18} Sep-Pak® column (Waters® Corporation)
13. Petroleum ether
14. Nitrogen gas
15. Vortex
16. Centrifuge

WARNINGS AND PRECAUTIONS

1. **DO NOT** use components beyond expiration date.
2. **DO NOT** mix any reagents or components of this kit with any reagents or components of any other kit. This kit is designed to work properly as provided.
3. **DO NOT** pipette reagents by mouth.
4. Always pour substrate out of the bottle into a clean test tube - **DO NOT** pipette out of the bottle. If the pipette tip is unclean, this may result in contamination of the substrate.
5. All specimens should be considered potentially infectious. Exercise proper handling precautions.
6. **DO NOT** smoke, eat or drink in areas where specimens or reagents are being handled.
7. Use aseptic technique when opening and removing reagents from vials and bottles.
8. Keep plate covered except when adding reagents, washing or reading.
9. Kit components should be refrigerated at all times when not in use.

PROCEDURAL NOTES

1. It is not necessary to allow reagents to warm to room temperature before use.
2. Desiccant bag must remain in foil pouch with unused strips. Keep zip-lock pouch sealed when not in use to maintain a dry environment.
3. Always use different pipette tips for the buffer, enzyme conjugate, standards and samples.
4. Before pipetting a reagent, rinse the pipette tip three times with that reagent (i.e. fill the tip with the desired amount of reagent and dispense back into the same vial - repeat 2 times). Now the tip is properly rinsed and ready to dispense the reagent into your well or test tube.
5. When pipetting into the wells, **DO NOT** allow the pipette tip to touch the inside of the well, or any of the reagents already in the well. This can result in cross contamination.
6. Standards and samples should be assayed in duplicate.
7. To quantitate, always run samples alongside a standard curve. If testing a sample that is not extracted, standards should be diluted in the same type of medium being tested. This medium should be known to be negative.
8. Gently mix specimens and reagents before use. Avoid vigorous agitation.
9. When using only partial amounts of a kit, it is recommended to transfer the appropriate volume of each reagent to a clean vessel for repeated dispensing. This will reduce reagent contamination by repeated sampling from the original container.
10. The enzyme conjugate is most stable in its concentrated form. Dilute only the volume necessary for the amount of strips currently being used.
11. Before taking an absorbance reading, wipe the outside bottom of the wells with a lint-free wiper to remove dust and fingerprints.
12. Before opening the enzyme conjugate and standard vial, tap vial in an upright position to remove any liquid in the cap.
SAMPLE PREPARATION

This assay is non-species specific. Usually, urine and tissue culture supernatant can be assayed directly by diluting them with the diluted extraction buffer. Plasma and most other mediums will need to be extracted.

EXTRACTION OF TXB$_2$

1. Add 0.2 mL of methanol to 1 mL of biological fluid and vortex.
2. For tissue, homogenize it in 15% methanol in 0.1 M sodium phosphate buffer, pH 7.5 (100 mg in 1 mL methanol - buffer). Centrifuge the homogenate for five (5) minutes. Collect the supernatant in a clean tube.
3. Precondition the C$_{18}$ Sep-Pak® column (Waters® Corporation) by washing the column with 2 mL of methanol followed by 2 mL of water.
4. Apply the above sample into the column and adjust the flow rate to 1 mL per minute.
5. Wash the column with 2 mL of 15% methanol in water followed by 2 mL of petroleum ether.
6. The TXB$_2$ is eluted by 2 mL of methyl formate.
7. Evaporate methyl formate eluate with a stream of nitrogen gas.
8. Dissolve the residue with 1 mL of diluted extraction buffer and assay 50 µL in duplicates.
9. If the concentration is higher than the high range of the standard curve, the samples in # 8 need to be further diluted and assayed.

NOTE: Extraction buffer must be diluted 5 fold with deionized water before use. Any precipitant present must be brought into solution before dilution.

TEST PROCEDURES

1. Prepare standards as follows:

<table>
<thead>
<tr>
<th>Standard</th>
<th>Preparation</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>stock solution 1 µg/mL (provided in green capped vial)</td>
</tr>
<tr>
<td>B</td>
<td>take 20 µL of A, add to 980 µL of EIA buffer and mix=20 ng/mL</td>
</tr>
<tr>
<td>C</td>
<td>take 200 µL of B, add to 1.8 mL of EIA buffer and mix=2 ng/mL</td>
</tr>
<tr>
<td>D</td>
<td>take 200 µL of C, add to 1.8 mL of EIA buffer and mix=0.2 ng/mL</td>
</tr>
</tbody>
</table>

Continue standard preparation following Scheme I.

SCHEME I

<table>
<thead>
<tr>
<th>Standards</th>
<th>ng/mL</th>
<th>EIA buffer (µL added)</th>
<th>C standard µL</th>
<th>D standard µL</th>
</tr>
</thead>
<tbody>
<tr>
<td>$S_0$</td>
<td>0</td>
<td>as is</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>$S_1$</td>
<td>0.004</td>
<td>980</td>
<td>-</td>
<td>20</td>
</tr>
<tr>
<td>$S_2$</td>
<td>0.01</td>
<td>950</td>
<td>-</td>
<td>50</td>
</tr>
<tr>
<td>$S_3$</td>
<td>0.02</td>
<td>900</td>
<td>-</td>
<td>100</td>
</tr>
<tr>
<td>$S_4$</td>
<td>0.04</td>
<td>800</td>
<td>-</td>
<td>200</td>
</tr>
<tr>
<td>$S_5$</td>
<td>0.1</td>
<td>500</td>
<td>-</td>
<td>500</td>
</tr>
<tr>
<td>$S_6$</td>
<td>0.2</td>
<td>-</td>
<td>-</td>
<td>as is</td>
</tr>
<tr>
<td>$S_7$</td>
<td>0.4</td>
<td>800</td>
<td>200</td>
<td>-</td>
</tr>
</tbody>
</table>
2. Determine the number of wells to be used.
3. Dilute the TXB$_2$ enzyme conjugate. Add 1 µL of enzyme conjugate into 50 µL total volume of EIA buffer for each well assayed. For the whole plate, add 110 µL of the enzyme conjugate into 5.5 mL total volume of EIA buffer. Mix the solution thoroughly.
4. Add 50 µL of standards (S) or unknown (U) (some samples may require diluting) to the appropriate wells in duplicate.

**See Scheme II for suggested template design.**

5. Add 50 µL of the diluted enzyme conjugate to each well. Use 8-channel pipette or 12-channel pipette for rapid addition.
6. Mix by shaking plate gently. A microplate shaker may be used.
7. Cover plate with plastic film or plate cover and incubate at room temperature for one hour.

**NOTE:** Keep plate away from drafts and temperature fluctuations.

8. Dilute concentrated wash buffer with deionized water (i.e. 20 mL of wash buffer plus 180 mL of deionized water). Mix thoroughly.
9. After incubation, dump out the contents of the plate. Tap out contents thoroughly on a clean lint-free towel.
10. Wash each well with 300 µL of the diluted wash buffer. Repeat for a total of three washings. An automated plate washer can be used, however, increase wash cycles from three to five.
11. Add 150 µL of substrate to each well. Use multichannel pipette for best results. Mix by shaking plate gently.
12. Incubate at room temperature for 30 minutes.
13. Gently shake plate before taking a reading to ensure uniform color throughout each well.
14. Plate is read in a microplate reader at 650 nm. If a dual wavelength is used, set W$_1$ at 650 nm and W$_2$ at 490 nm.
15. If accounting for substrate background, use 2 to 8 wells as blanks with only substrate in the wells (150 µL/well). Subtract the average of these absorbance values from the absorbance values of the wells being assayed.

**NOTE:** Some microplate readers can be programmed to do these subtractions automatically when reading the plate. Consult your instrument manual.

**OPTIONAL TEST PROCEDURES**

16. Add 50-100 µL of 1 N HCl or Neogen’s Red Stop Solution to each well to stop enzyme reaction.
17. Read plate at 450 nm, if 1N HCl solution was used. Read plate at 650 nm, if Neogen’s Red Stop Solution was used.
18. Plot the standard curve and estimate the concentrations of the samples from the curve. See "CALCULATIONS."

**NOTE:** Absorbance readings will approximately double when stopped with acid. If absorbance readings are too high for measuring with your microplate reader, decrease the substrate incubation approximately 10 minutes but no more than 15 minutes.

**SCHEME II**

<table>
<thead>
<tr>
<th></th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
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<th>6</th>
<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>$^0$</td>
<td>$^0$</td>
<td>$^1$</td>
<td>$^1$</td>
<td>$^9$</td>
<td>$^9$</td>
<td>$^{17}$</td>
<td>$^{17}$</td>
<td>$^{25}$</td>
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<td>$^{33}$</td>
</tr>
<tr>
<td>B</td>
<td>$^1$</td>
<td>$^1$</td>
<td>$^2$</td>
<td>$^2$</td>
<td>$^{10}$</td>
<td>$^{10}$</td>
<td>$^{18}$</td>
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<td>$^{26}$</td>
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<tr>
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<td>$^7$</td>
<td>$^7$</td>
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<td>$^{39}$</td>
</tr>
<tr>
<td>H</td>
<td>$^7$</td>
<td>$^7$</td>
<td>$^8$</td>
<td>$^8$</td>
<td>$^{16}$</td>
<td>$^{16}$</td>
<td>$^{24}$</td>
<td>$^{24}$</td>
<td>$^{32}$</td>
<td>$^{32}$</td>
<td>$^{40}$</td>
<td>$^{40}$</td>
</tr>
</tbody>
</table>
CALCULATIONS

1. After the substrate background has been subtracted from all absorbance values, average all of your duplicate well absorbance values.
2. The average of your two $S_0$ values is now your $B_0$ value. ($S_i$ now becomes $B_i$, etc.)
3. Next, find the percent of maximal binding (%B/$B_0$ value). To do this, divide the averages of each standard absorbance value (now known as $B_1$ through $B_7$) by the $B_0$ absorbance value and multiply by 100 to achieve percentages.
4. Graph your standard curve by plotting the %B/$B_0$ for each standard concentration on the ordinate (y) axis against concentration on the abscissa (x) axis. Draw a curve by using a curve-fitting routine (i.e. 4-parameter or linear regression).
5. Divide the averages of each sample absorbance value by the $B_0$ value and multiply by 100 to achieve percentages.
6. Using the standard curve, the concentration of each sample can be determined by comparing the %B/$B_0$ of each sample to the corresponding concentration of TBX$_2$ standard.
7. If the samples were diluted, the concentration determined from the standard curve must be multiplied by the dilution factor.

TYPICAL STANDARD CURVE

[Graph showing a typical standard curve with concentration on the x-axis and %B/$B_0$ on the y-axis.]

TYPICAL DATA

**NOTE:** "Typical data" is a representation. Variances in data will occur. Optical density readings may fluctuate during the shelf-life of the kit, but the %B/$B_0$ should remain comparable. Measuring wavelength: 650 nm

<table>
<thead>
<tr>
<th>Standard</th>
<th>Standard Concentration (ng/mL)</th>
<th>Optical Density (Absorbance Value)</th>
<th>%B/$B_0$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$S_0$ ($B_0$)</td>
<td>0</td>
<td>1.277</td>
<td>100</td>
</tr>
<tr>
<td>$S_1$ ($B_1$)</td>
<td>0.004</td>
<td>1.147</td>
<td>90</td>
</tr>
<tr>
<td>$S_2$ ($B_2$)</td>
<td>0.01</td>
<td>0.996</td>
<td>78</td>
</tr>
<tr>
<td>$S_3$ ($B_3$)</td>
<td>0.02</td>
<td>0.820</td>
<td>64</td>
</tr>
<tr>
<td>$S_4$ ($B_4$)</td>
<td>0.04</td>
<td>0.619</td>
<td>48</td>
</tr>
<tr>
<td>$S_5$ ($B_5$)</td>
<td>0.1</td>
<td>0.361</td>
<td>28</td>
</tr>
<tr>
<td>$S_6$ ($B_6$)</td>
<td>0.2</td>
<td>0.221</td>
<td>17</td>
</tr>
<tr>
<td>$S_7$ ($B_7$)</td>
<td>0.4</td>
<td>0.137</td>
<td>11</td>
</tr>
</tbody>
</table>
### CROSS REACTIVITY

<table>
<thead>
<tr>
<th>Compounds</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thromboxane B₂</td>
<td>100.0%</td>
</tr>
<tr>
<td>2,3-Dinor-thromboxane B₂</td>
<td>30.0%</td>
</tr>
<tr>
<td>Prostaglandin D₂</td>
<td>1.21%</td>
</tr>
<tr>
<td>Prostaglandin E₂</td>
<td>0.08%</td>
</tr>
<tr>
<td>11-Dehydro-thromboxane B₂</td>
<td>0.07%</td>
</tr>
<tr>
<td>Prostaglandin F₂α</td>
<td>0.06%</td>
</tr>
<tr>
<td>6-Keto-prostaglandin F₁a</td>
<td>0.05%</td>
</tr>
<tr>
<td>Prostaglandin F₁a</td>
<td>0.02%</td>
</tr>
<tr>
<td>Arachidonic acid</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Leukotriene B₄</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Prostaglandin A₂</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Prostaglandin B₂</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>13,14-Dihydro-15-keto-prostaglandin F₂α</td>
<td>&lt;0.01%</td>
</tr>
</tbody>
</table>

### REFERENCES


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Technical assistance is available Monday-Friday between 8:00 a.m. and 6:00 p.m. EST.