Prostaglandin E Metabolite ELISA Kit

Item No. 514531

www.caymanchem.com
Customer Service 800.364.9897
Technical Support 888.526.5351
1180 E. Ellsworth Rd · Ann Arbor, MI · USA
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GENERAL INFORMATION

Materials Supplied

<table>
<thead>
<tr>
<th>Item Number</th>
<th>Item</th>
<th>Quantity/Size</th>
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<tbody>
<tr>
<td>414532</td>
<td>Prostaglandin E Metabolite ELISA Antiserum</td>
<td>1 vial/100 dtn</td>
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<tr>
<td>414530</td>
<td>Prostaglandin E Metabolite AChE Tracer</td>
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<td>414534</td>
<td>Prostaglandin E Metabolite ELISA Standard</td>
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<tr>
<td>400060</td>
<td>ELISA Buffer Concentrate (10X)</td>
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<tr>
<td>400062</td>
<td>Wash Buffer Concentrate (400X)</td>
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<td>400035</td>
<td>Polysorbate 20</td>
<td>1 vial/3 ml</td>
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<td>400005/400007</td>
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<td>96-Well Cover Sheet</td>
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<td>400050</td>
<td>Ellman’s Reagent</td>
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<tr>
<td>400032</td>
<td>Phosphate Buffer</td>
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<td>Carbonate Buffer</td>
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<td>ELISA Tracer Dye</td>
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<tr>
<td>400042</td>
<td>ELISA Antiserum Dye</td>
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</tbody>
</table>

If any of the items listed above are damaged or missing, please contact our Customer Service department at (800) 364-9897 or (734) 971-3335. We cannot accept any returns without prior authorization.
WARNING: THIS PRODUCT IS FOR RESEARCH ONLY - NOT FOR HUMAN OR VETERINARY DIAGNOSTIC OR THERAPEUTIC USE.

Safety Data

This material should be considered hazardous until further information becomes available. Do not ingest, inhale, get in eyes, on skin, or on clothing. Wash thoroughly after handling. Before use, the user must review the complete Safety Data Sheet, which has been sent via email to your institution.

Precautions

Please read these instructions carefully before beginning this assay.

The reagents in this kit have been tested and formulated to work exclusively with Cayman Chemical’s AChE ELISA Kits. This kit may not perform as described if any reagent or procedure is replaced or modified.

When compared to quantification by LC/MS or GC/MS, it is not uncommon for immunoassays to report higher analyte concentrations. While LC/MS or GC/MS analyses typically measure only a single compound, antibodies used in immunoassays sometimes recognize not only the target molecule, but also structurally related molecules, including biologically relevant metabolites. In many cases, measurement of both the parent molecule and metabolites is more representative of the overall biological response than is the measurement of a short-lived parent molecule. It is the responsibility of the researcher to understand the limits of both assay systems and to interpret their data accordingly.

If You Have Problems

Technical Service Contact Information

Phone: 888-526-5351 (USA and Canada only) or 734-975-3888
Fax: 734-971-3641
Email: techserv@caymanchem.com
Hours: M-F 8:00 AM to 5:30 PM EST

In order for our staff to assist you quickly and efficiently, please be ready to supply the lot number of the kit (found on the outside of the box).

Storage and Stability

This kit will perform as specified if stored as directed at -20°C and used before the expiration date indicated on the outside of the box.

Materials Needed But Not Supplied

1. A plate reader capable of measuring absorbance between 405-420 nm.
2. Adjustable pipettes and a repeating pipettor.
3. A source of 'UltraPure' water. Water used to prepare all ELISA reagents and buffers must be deionized and free of trace organic contaminants ('UltraPure'). Use activated carbon filter cartridges or other organic scavengers. Glass distilled water (even if double distilled), HPLC-grade water, and sterile water (for injections) are not adequate for ELISA. NOTE: UltraPure water is available for purchase from Cayman (Item No. 400000).
4. Materials used for Sample Preparation (see page 14).
Background

Prostaglandin $E_2$ (PGE$_2$) is produced by a variety of cell types which, in general, do not contain the enzymes required for metabolism of PGE$_2$. Thus, cultured endothelial cells or osteoblasts will release PGE$_2$ into the culture medium where it will accumulate without appreciable metabolism. The direct assay of PGE$_2$ from the medium is a good way to measure PGE$_2$ production from these cells. PGE$_2$ is rapidly converted in vivo to its 13,14-dihydro-15-keto metabolite (see Figure 1, on page 7), with more than 90% of circulating PGE$_2$ cleared by a single passage through the lungs. Unfortunately, this metabolite is not chemically stable and undergoes a variable amount of degradation to PGA products. For this reason, blood, urine, or other samples from whole animals or humans often contain very little intact PGE$_2$, and measurement of the metabolites is necessary to provide a reliable estimate of actual PGE$_2$ production.

About This Assay

Cayman’s Prostaglandin E Metabolite (PGEM) assay is a competitive assay that converts 13,14-dihydro-15-keto PGA$_2$ and 13,14-dihydro-15-keto PGE$_2$ to a single, stable derivative that can be easily quantified. This assay is, therefore, the method of choice if the samples in question have undergone extensive metabolism prior to collection. The assay has a range from 0.39-50 pg/ml and a sensitivity (80% B/B$_0$) of approximately 2 pg/ml.

Figure 1. Metabolism of PGE$_2$
**Biochemistry of Acetylcholinesterase**

The electric organ of the electric eel, *E. electricus*, contains an avid AChE capable of massive catalytic turnover during the generation of its electrochemical discharges. The electric eel AChE has a clover leaf-shaped tertiary structure consisting of a triad of tetramers attached to a collagen-like structural fibril. This stable enzyme is capable of high turnover (64,000 s⁻¹) for the hydrolysis of acetylthiocholine.

A molecule of the analyte covalently attached to a molecule of AChE serves as the tracer in AChE enzyme immunoassays. Quantification of the tracer is achieved by measuring its AChE activity with Ellman’s Reagent. This reagent consists of acetylthiocholine and 5,5'-dithio-(2-nitrobenzoic acid). Hydrolysis of acetylthiocholine by AChE produces thiocholine (see Figure 3, on page 10). The non-enzymatic reaction of thiocholine with 5,5'-dithio-(2-nitrobenzoic acid) produces 5-thio-2-nitrobenzoic acid, which has a strong absorbance at 412 nm (ε = 13,600).

AChE has several advantages over other enzymes commonly used for enzyme immunoassays. Unlike horseradish peroxidase, AChE does not self-inactivate during turnover. This property of AChE also allows redevelopment of the assay if it is accidentally splashed or spilled. In addition, the enzyme is highly stable under the assay conditions, has a wide pH range (pH 5-10), and is not inhibited by common buffer salts or preservatives. Since AChE is stable during the development step, it is unnecessary to use a ‘stop’ reagent, and the plate may be read whenever it is convenient.

**Description of AChE Competitive ELISAs¹,²**

This assay is based on the competition between Prostaglandin E Metabolite (PGEM) and a PGEM-acetylcholinesterase (AChE) conjugate (PGEM Tracer) for a limited number of PGEM-specific rabbit antiserum binding sites. Because the concentration of the PGEM Tracer is held constant while the concentration of PGEM varies, the amount of PGEM Tracer that is able to bind to the rabbit antiserum will be inversely proportional to the concentration of PGEM in the well. This rabbit antiserum-PGEM (either free or tracer) complex binds to the mouse monoclonal anti-rabbit IgG that has been previously attached to the well. The plate is washed to remove any unbound reagents, and then Ellman’s Reagent (which contains the substrate to AChE) is added to the well. The product of this enzymatic reaction has a distinct yellow color and absorbs strongly at 412 nm. The intensity of this color, determined spectrophotometrically, is proportional to the amount of PGEM Tracer bound to the well, which is inversely proportional to the amount of free PGEM present in the well during the incubation; or

\[
\text{Absorbance} \propto \frac{[\text{Bound PGEM Tracer}]}{[\text{PGEM}]}.
\]

A schematic of this process is shown in Figure 2, below.

---

**Figure 2. Schematic of the AChE ELISA**
Definition of Key Terms

**Blank:** background absorbance caused by Ellman’s Reagent. The blank absorbance should be subtracted from the absorbance readings of all the other wells, including NSB wells.

**Total Activity:** total enzymatic activity of the AChE-linked tracer. This is analogous to the specific activity of a radioactive tracer.

**NSB (Non-Specific Binding):** non-immunological binding of the tracer to the well. Even in the absence of specific antibody a very small amount of tracer still binds to the well; the NSB is a measure of this low binding. Do not forget to subtract the Blank absorbance values.

**B₀ (Maximum Binding):** maximum amount of the tracer that the antibody can bind in the absence of free analyte.

**%B/B₀ (%Bound/Maximum Bound):** ratio of the absorbance of a particular sample or standard well to that of the maximum binding (B₀) well.

**Standard Curve:** a plot of the %B/B₀ values versus concentration of a series of wells containing various known amounts of analyte.

**Dtn:** determination, where one dtn is the amount of reagent used per well.

**Cross Reactivity:** numerical representation of the relative reactivity of this assay towards structurally related molecules as compared to the primary analyte of interest. Biomolecules that possess similar epitopes to the analyte can compete with the assay tracer for binding to the primary antibody. Substances that are superior to the analyte in displacing the tracer result in a cross reactivity that is greater than 100%. Substances that are inferior to the primary analyte in displacing the tracer result in a cross reactivity that is less than 100%. Cross reactivity is calculated by comparing the mid-point (50% B/B₀) value of the tested molecule to the mid-point (50% B/B₀) value of the primary analyte when each is measured in assay buffer using the following formula:

\[
\text{% Cross Reactivity} = \left( \frac{\text{50% B/B₀ value for the primary analyte}}{\text{50% B/B₀ value for the potential cross reactant}} \right) \times 100\%
\]
NOTE: Water used to prepare all ELISA reagents and buffers must be deionized and free of trace organic contaminants (‘UltraPure’). Use activated carbon filter cartridges or other organic scavengers. Glass distilled water (even if double distilled), HPLC-grade water, and sterile water (for injections) are not adequate for ELISA. UltraPure water may be purchased from Cayman (Item No. 400000).

Buffer Preparation

Store all diluted buffers at 4°C; they will be stable for about two months.

1. **ELISA Buffer Preparation**

   Dilute the contents of one vial of ELISA Buffer Concentrate (10X) (Item No. 400060) with 90 ml of UltraPure water. Be certain to rinse the vial to remove any salts that may have precipitated. **NOTE: It is normal for the concentrated buffer to contain crystalline salts after thawing. These will completely dissolve upon dilution with water.**

2. **Wash Buffer Preparation**

   5 ml vial Wash Buffer Concentrate (400X) (96-well kit; Item No. 400062): Dilute to a total volume of 2 liters with UltraPure water and add 1 ml of Polysorbate 20 (Item No. 400035).

   **OR**

   12.5 ml vial Wash Buffer Concentrate (400X) (480-well kit; Item No. 400062): Dilute to a total volume of 5 liters with UltraPure water and add 2.5 ml of Polysorbate 20 (Item No. 400035).

   Smaller volumes of Wash Buffer can be prepared by diluting the Wash Buffer Concentrate 1:400 and adding Polysorbate 20 (0.5 ml/liter of Wash Buffer).

   **NOTE:** Polysorbate 20 is a viscous liquid and cannot be measured by a regular pipette. A positive displacement pipette or a syringe should be used to deliver small quantities accurately.

3. **Phosphate Buffer**

   Prepare a 1 M Phosphate Buffer solution by dissolving the contents of the 100 dtm vial of Phosphate Buffer (Item No. 400032) in 30 ml UltraPure water, or dissolve the contents of one of the 250 dtm vials of Phosphate Buffer (Item No. 400032) in 75 ml UltraPure water.

4. **Carbonate Buffer**

   Prepare a 1 M Carbonate Buffer solution by dissolving the contents of the 100 dtm vial of Carbonate Buffer (Item No. 400027) in 25 ml UltraPure water, or dissolve the contents of the 500 dtm vial of Carbonate Buffer (Item No. 400027) in 125 ml UltraPure water.

5. **PGEM Assay Buffer**

   Prepare 20 ml of PGEM Assay Buffer by combining 13 ml ELISA Buffer, 3 ml Carbonate Buffer, and 4 ml Phosphate Buffer. This quantity of buffer should be more than sufficient to complete one 96-well plate.
**Sample Preparation**

This assay has been validated for a wide range of samples including urine, and plasma. Proper sample storage and preparation are essential for consistent and accurate results. Please read this section carefully before beginning the assay.

### General Precautions

- All samples must be free of organic solvents prior to assay.
- Samples should be assayed immediately after collection; samples that cannot be assayed immediately should be stored at -80°C.
- Samples of rabbit origin may contain antibodies which interfere with the assay by binding to the mouse anti-rabbit plate. We recommend that all rabbit samples be purified prior to use in this assay.
- AEBSF (Pefabloc SC®) and PMSF inhibit AChE. Samples containing these protease inhibitors should not be used in this assay.

### Testing for Interference

In general, urine samples may be derivatized, diluted with PGE Metabolite Buffer and added directly to the assay well. Plasma, serum, and other heterogeneous mixtures such as lavage fluids and aspirates often contain contaminants which can interfere in the assay. It is best to check for interference before embarking on a large number of sample measurements. To test for interference, derivatize one or two test samples, then dilute with PGE Metabolite Buffer to obtain at least two different dilutions of each sample between ~2 and 50 pg/ml (i.e., between 20-80% B/B₀). If the two different dilutions of the sample show good correlation (differ by 20% or less) in the final calculated PGEM concentrations, purification is not required. If you do not see good correlation of the different dilutions, purification is advised. The protocol described on page 16 is one such method.

### Plasma

Analysis of plasma samples without purification may lead to inconsistent results. We recommend using the purification protocol described below for all plasma samples.

Plasma samples should be collected in vacutainers containing sodium citrate, heparin, or EDTA. Vacutainers can also be supplemented with indomethacin to give a final concentration of at least 10 µM. Indomethacin will prevent ex vivo formation of prostaglandins, which have the potential to interfere with this assay (although most prostaglandins do not appear to exhibit any cross reactivity (see page 31)). Samples that cannot be assayed immediately should be stored at -80°C.
Sample Purification

Acetone Precipitation

Acetone precipitation is recommended for removal of proteins from heterogeneous samples. For some samples, acetone precipitation may be the only purification that is required.

1. Aliquot samples into clean tubes.
2. Add four volumes of ice-cold acetone.
3. Incubate at -20°C for a minimum of thirty minutes.
4. Pellet proteins by centrifugation in a tabletop centrifuge at a minimum of 400 x g for five minutes.
5. Transfer supernatants to clean tubes.
6. Dry under a gentle stream of nitrogen.
7. Resuspend samples in the original volume of ELISA Buffer.
8. The samples are now ready for derivatization.

ASSAY PROTOCOL

Derivatization of Standards and Samples to PGEM

Derivatization Hints
- Allow the derivatization to proceed overnight to ensure that all the PGE₂ metabolites derivatize completely.
- Derivatize all standards and samples for the same amount of time.

Derivatization of the PGEM ELISA Standard

Equilibrate a pipette tip in ethanol by repeatedly filling and expelling the tip with ethanol several times. Using the equilibrated pipette tip, transfer 100 µl of the PGEM ELISA Standard (Item No. 414534) into a clean test tube, then dilute with 900 µl UltraPure water. The concentration of this solution (the bulk standard) will be 40 ng/ml.

Aliquot 50 µl of this solution into a clean tube and dilute to a total volume of 1 ml with ELISA Buffer (i.e., add 950 µl). Add 300 µl of Carbonate Buffer and incubate at 37°C overnight. Then add 400 µl Phosphate Buffer and 300 µl ELISA Buffer. This solution is 1,000 pg/ml.

Derivatization of the Samples

Aliquot 500 µl of each sample into a clean test tube. Add 150 µl of Carbonate Buffer and incubate overnight at 37°C. Then add 200 µl of Phosphate Buffer and 150 µl of ELISA Buffer. Depending on your sample type, the samples may now be ready to use directly in the assay.

NOTE: Because of the high salt concentration in the PGEM Standards, it is crucial that PGEM Assay Buffer is used rather than ELISA Buffer to dilute samples if further dilution is necessary.

For plasma samples, the following ethyl acetate extraction protocol is suggested after derivatization.
Acidification and Ethyl Acetate Extraction
We suggest the following procedure for samples requiring additional purification following derivatization:
1. Acidify the sample to ~pH 4 by the addition of 1M acetate buffer. (To avoid having to measure the pH of each individual sample, adjust the pH of an equivalent volume of sample matrix to pH 4.0 using the 1M acetate buffer. NOTE: For samples of different volumes, the amount of acid should be adjusted to maintain this ratio of acid to sample.)
2. Add two volumes of ethyl acetate to each derivatized sample.
3. Vortex thoroughly to mix.
4. Centrifuge samples at a minimum of 200 x g for five minutes.
5. Transfer the ethyl acetate (upper layer) to a clean glass tube.
6. Repeat this extraction three more times, combining all of the ethyl acetate layers from each individual sample.
7. Dry samples under a gentle stream of nitrogen.
8. Resuspend samples in PGEM Assay Buffer.
9. The samples are now ready to use in the assay

NOTE: Because of the high salt concentration in the PGEM Standards, it is crucial that PGEM Assay Buffer is used rather than ELISA Buffer to dilute samples if further dilution is necessary.

Preparing the Standard Curve
NOTE: Because of the high salt concentration in the 1,000 pg/ml solution, all the points of the standard curve must contain the same salt concentration. Thus, when performing the serial dilution, use the PGEM Assay Buffer.

To prepare the standard for use in ELISA: Obtain eight clean test tubes and number them #1 through #8. Aliquot 950 µl PGEM Assay Buffer to tube #1 and 500 µl PGEM Assay Buffer to tubes #2-8. Transfer 50 µl of the derivatized standard (1,000 pg/ml) to tube #1 and mix thoroughly. Serially dilute the standard by removing 500 µl from tube #1 and placing it in tube #2; mix thoroughly. Next, remove 500 µl from tube #2 and place it into tube #3; mix thoroughly. Repeat this process for tubes #4-8. These diluted standards should not be stored for more than 24 hours.

Figure 4. Preparation of the PGEM standards
Preparation of Assay-Specific Reagents

PGEM AChE Tracer
Reconstitute the PGEM AChE Tracer as follows:

100 dtn PGEM AChE Tracer (96-well kit; Item No. 414530): Reconstitute with 6 ml ELISA Buffer.

OR

500 dtn PGEM AChE Tracer (480-well kit; Item No. 414530): Reconstitute with 30 ml ELISA Buffer.

Store the reconstituted PGEM AChE Tracer at 4°C (do not freeze!) and use within four weeks. A 20% surplus of tracer has been included to account for any incidental losses.

Tracer Dye Instructions (optional)
This dye may be added to the tracer, if desired, to aid in visualization of tracer-containing wells. Add the dye to the reconstituted tracer at a final dilution of 1:100 (add 60 µl of dye to 6 ml tracer or add 300 µl of dye to 30 ml of tracer).

PGEM ELISA Antiserum
Reconstitute the PGEM ELISA Antiserum as follows:

100 dtn PGEM ELISA Antiserum (96-well kit; Item No. 414532): Reconstitute with 6 ml ELISA Buffer.

OR

500 dtn PGEM ELISA Antiserum (480-well kit; Item No. 414532): Reconstitute with 30 ml ELISA Buffer.

Store the reconstituted PGEM ELISA Antiserum at 4°C. It will be stable for at least four weeks. A 20% surplus of antiserum has been included to account for any incidental losses.

Antiserum Dye Instructions (optional)
This dye may be added to the antiserum, if desired, to aid in visualization of antiserum-containing wells. Add the dye to the reconstituted antiserum at a final dilution of 1:100 (add 60 µl of dye to 6 ml antiserum or add 300 µl of dye to 30 ml of antiserum).
Plate Set Up

The 96-well plate(s) included with this kit is supplied ready to use. It is not necessary to rinse the plate(s) prior to adding the reagents. NOTE: If you do not need to use all the strips at once, place the unused strips back in the plate packet and store at 4°C. Be sure the packet is sealed with the desiccant inside.

Each plate or set of strips must contain a minimum of two blanks (Blk), two non-specific binding wells (NSB), two maximum binding wells (B0), and an eight point standard curve run in duplicate. NOTE: Each assay must contain this minimum configuration in order to ensure accurate and reproducible results. Each sample should be assayed at two dilutions and each dilution should be assayed in duplicate. For statistical purposes, we recommend assaying samples in triplicate. A suggested plate format is shown in Figure 5, below. The user may vary the location and type of wells present as necessary for each particular experiment. The plate format provided below has been designed to allow for easy data analysis using a convenient spreadsheet offered by Cayman (see page 26, for more details). We suggest you record the contents of each well on the template sheet provided (see page 34).

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</table>

Figure 5. Sample plate format

Performing the Assay

Pipetting Hints

- Use different tips to pipette each reagent.
- Before pipetting each reagent, equilibrate the pipette tip in that reagent (i.e., slowly fill the tip and gently expel the contents, repeat several times).
- Do not expose the pipette tip to the reagent(s) already in the well.

Addition of the Reagents

1. **PGEM Buffer**
   Add 50 µl ELISA Buffer and 50 µl of PGEM Buffer to NSB wells. Add 50 µl PGEM Buffer to B0 wells.

2. **PGEM ELISA Standard**
   Add 50 µl from tube #8 to both of the lowest standard wells (S8). Add 50 µl from tube #7 to each of the next two standard wells (S7). Continue with this procedure until all the standards are aliquoted. The same pipette tip should be used to aliquot all the standards. Before pipetting each standard, be sure to equilibrate the pipette tip in that standard.

3. **Samples**
   Add 50 µl of sample per well. Each sample should be assayed at a minimum of two dilutions. Each dilution should be assayed in duplicate (triplicate recommended).

4. **PGEM AChE Tracer**
   Add 50 µl to each well except the TA and the Blk wells.

5. **PGEM ELISA Antiserum**
   Add 50 µl to each well except the TA, the NSB, and the Blk wells.
Table 1. Pipetting summary

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<th>ELISA Buffer</th>
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<th>Tracer</th>
<th>Antiserum</th>
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<td>-</td>
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<td>Std/Sample</td>
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Incubation of the Plate
Cover each plate with plastic film (Item No. 400012) and incubate for 18 hours at room temperature.

Development of the Plate
1. Reconstitute Ellman’s Reagent immediately before use (20 ml of reagent is sufficient to develop 100 wells):

   **100 dtn vial Ellman’s Reagent (96-well kit; Item No. 400050):** Reconstitute with 20 ml of UltraPure water.

   **OR**

   **250 dtn vial Ellman’s Reagent (480-well kit; Item No. 400050):** Reconstitute with 50 ml of UltraPure water.

   **NOTE:** Reconstituted Ellman’s Reagent is unstable and should be used the same day it is prepared; protect the Ellman’s Reagent from light when not in use. Extra vials of the reagent have been provided should a plate need to be re-developed or multiple assays be run on different days.

2. Empty the wells and rinse five times with Wash Buffer.
3. Add 200 µl of Ellman’s Reagent to each well
4. Add 5 µl of tracer to the TA wells.
5. Cover the plate with plastic film. Optimum development is obtained by using an **orbital shaker** equipped with a large, flat cover to allow the plate(s) to develop in the dark. This assay typically develops (i.e., B₀ wells ≥0.3 A.U. (blank subtracted)) in **60-90 minutes**.

Reading the Plate
1. Wipe the bottom of the plate with a clean tissue to remove fingerprints, dirt, etc.
2. Remove the plate cover being careful to keep Ellman’s Reagent from splashing on the cover. **NOTE:** Any loss of Ellman’s Reagent will affect the absorbance readings. If Ellman’s Reagent is present on the cover, use a pipette to transfer the Ellman’s Reagent into the well. If too much Ellman’s Reagent has splashed on the cover to easily redistribute back into the wells, wash the plate three times with Wash Buffer and repeat the development with fresh Ellman’s Reagent.
3. Read the plate at a wavelength between 405 and 420 nm. The absorbance may be checked periodically until the B₀ wells have reached a minimum of 0.3 A.U. (blank subtracted). The plate should be read when the absorbance of the B₀ wells in the range of 0.3-1.5 A.U. (blank subtracted). If the absorbance of the wells exceeds 2.0, wash the plate, add fresh Ellman’s Reagent and let it develop again.
Many plate readers come with data reduction software that plots data automatically. Alternatively a spreadsheet program can be used. The data should be plotted as either %B/B_0 versus log concentration using a four-parameter logistic fit or as logit B/B_0 versus log concentration using a linear fit. NOTE: Cayman has a computer spreadsheet available for data analysis. Please contact Technical Service or visit our website (www.caymanchem.com/analysis/elisa) to obtain a free copy of this convenient data analysis tool.

Calculations

Preparation of the Data

The following procedure is recommended for preparation of the data prior to graphical analysis.

NOTE: If the plate reader has not subtracted the absorbance readings of the blank wells from the absorbance readings of the rest of the plate, be sure to do that now.

1. Average the absorbance readings from the NSB wells.
2. Average the absorbance readings from the B_0 wells.
3. Subtract the NSB average from the B_0 average. This is the corrected B_0 or corrected maximum binding.
4. Calculate the B/B_0 (Sample or Standard Bound/Maximum Bound) for the remaining wells. To do this, subtract the average NSB absorbance from the S1 absorbance and divide by the corrected B_0 (from Step 3). Repeat for S2-S8 and all sample wells. (To obtain %B/B_0 for a logistic four-parameter fit, multiply these values by 100.)

NOTE: The TA values are not used in the standard curve calculations. Rather, they are used as a diagnostic tool; the corrected B_0 divided by the actual TA (10X measured absorbance) will give the %Bound. This value should closely approximate the %Bound that can be calculated from the Sample Data (see page 28). Erratic absorbance values and a low (or no) %Bound could indicate the presence of organic solvents in the buffer or other technical problems (see page 32 for Troubleshooting).

Plot the Standard Curve

Plot %B/B_0 for standards S1-S8 versus PGEM concentration using linear (y) and log (x) axes and perform a 4-parameter logistic fit.

Alternative Plot - The data can also be linearized using a logit transformation. The equation for this conversion is shown below. NOTE: Do not use %B/B_0 in this calculation.

\[
\text{logit} \left( \frac{B}{B_0} \right) = \ln \left[ \frac{B / B_0}{1 - B / B_0} \right]
\]

Plot the data as logit (B/B_0) versus log concentrations and perform a linear regression fit.

Determine the Sample Concentration

Calculate the B/B_0 (or %B/B_0) value for each sample. Determine the concentration of each sample using the equation obtained from the standard curve. NOTE: Remember to account for any concentration or dilution of the sample prior to the addition to the well. Samples with %B/B_0 values greater than 80% or less than 20% should be re-assayed as they generally fall out of the linear range of the standard curve. A 20% or greater disparity between the apparent concentration of two different dilutions of the same sample indicates interference which could be eliminated by purification.
Performance Characteristics

Sample Data

The standard curve presented here is an example of the data typically produced with this kit; however, your results will not be identical to these. You must run a new standard curve. Do not use the data below to determine the values of your samples. Your results could differ substantially.

<table>
<thead>
<tr>
<th>Raw Data</th>
<th>Average</th>
<th>Corrected</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Activity</td>
<td>1.752</td>
<td>1.965</td>
</tr>
<tr>
<td>NSB</td>
<td>-0.001</td>
<td>-0.003</td>
</tr>
<tr>
<td>B₀</td>
<td>0.674</td>
<td>0.726</td>
</tr>
</tbody>
</table>

NSB corrected values:

<table>
<thead>
<tr>
<th>Dose (pg/ml)</th>
<th>Raw Data</th>
<th>Corrected</th>
<th>%B₀/B₀</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>0.138</td>
<td>0.140</td>
<td>0.122</td>
</tr>
<tr>
<td>25</td>
<td>0.230</td>
<td>0.232</td>
<td>0.215</td>
</tr>
<tr>
<td>12.5</td>
<td>0.335</td>
<td>0.337</td>
<td>0.309</td>
</tr>
<tr>
<td>6.25</td>
<td>0.441</td>
<td>0.443</td>
<td>0.433</td>
</tr>
<tr>
<td>3.13</td>
<td>0.518</td>
<td>0.520</td>
<td>0.523</td>
</tr>
<tr>
<td>1.56</td>
<td>0.592</td>
<td>0.594</td>
<td>0.584</td>
</tr>
<tr>
<td>0.78</td>
<td>0.640</td>
<td>0.642</td>
<td>0.620</td>
</tr>
<tr>
<td>0.39</td>
<td>0.649</td>
<td>0.651</td>
<td>0.689</td>
</tr>
</tbody>
</table>

Table 2. Typical results

Assay Range = 0.39-50 pg/ml
Sensitivity (defined as 80% B₀/B₀) = 2 pg/ml
Mid-point (defined as 50% B₀/B₀) = 8-14 pg/ml

The sensitivity and mid-point were derived from the standard curve shown above. The standard was diluted with PGEM Buffer.

Figure 6. Typical standard curve
**Precision:**
The intra- and inter-assay CVs have been determined at multiple points on the standard curve. These data are summarized in the graph on page 29 and in the table below.

<table>
<thead>
<tr>
<th>Dose (pg/ml)</th>
<th>%CV* Intra-assay variation</th>
<th>%CV* Inter-assay variation</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>8.1</td>
<td>18.2</td>
</tr>
<tr>
<td>25</td>
<td>5.4</td>
<td>7.2</td>
</tr>
<tr>
<td>12.5</td>
<td>5.9</td>
<td>8.3</td>
</tr>
<tr>
<td>6.25</td>
<td>5.5</td>
<td>11.2</td>
</tr>
<tr>
<td>3.13</td>
<td>12.8</td>
<td>8.0</td>
</tr>
<tr>
<td>1.56</td>
<td>25.1</td>
<td>13.4</td>
</tr>
<tr>
<td>0.78</td>
<td>23.7</td>
<td>39.3</td>
</tr>
<tr>
<td>0.39</td>
<td>N.D.</td>
<td>123</td>
</tr>
</tbody>
</table>

Table 3. Intra- and inter-assay Variation
*%CV represents the variation in concentration (not absorbance) of 40 repetitions of each point on the standard curve as determined using a reference standard curve.

**Cross Reactivity:**

<table>
<thead>
<tr>
<th>Compound</th>
<th>Cross Reactivity</th>
</tr>
</thead>
<tbody>
<tr>
<td>13,14-dihydro-15-keto PGE$_1$ *</td>
<td>100%</td>
</tr>
<tr>
<td>13,14-dihydro-15-keto PGE$_2$ *</td>
<td>100%</td>
</tr>
<tr>
<td>Bicyclo Prostaglandin E$_1$</td>
<td>38%</td>
</tr>
<tr>
<td>13,14-dihydro-15-keto PGD$_2$ *</td>
<td>0.08%</td>
</tr>
<tr>
<td>13,14-dihydro-15-keto PGF$_{2a}$ *</td>
<td>0.02%</td>
</tr>
<tr>
<td>Arachidonic Acid</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Leukotriene B$_4$</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>tetranor-PGEM</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>tetranor-PGFM</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Prostaglandin D$_2$</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Prostaglandin E$_1$</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>6-keto Prostaglandin E$_1$</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Prostaglandin E$_2$</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Prostaglandin F$_{1a}$</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>6-keto Prostaglandin F$_{1a}$</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Prostaglandin F$_{2a}$</td>
<td>&lt;0.01%</td>
</tr>
<tr>
<td>Thromboxane B$_2$</td>
<td>&lt;0.01%</td>
</tr>
</tbody>
</table>

Table 4. Cross Reactivity of the PGEM ELISA
*Derivatized per Assay Protocol
Troubleshooting

<table>
<thead>
<tr>
<th>Problem</th>
<th>Possible Causes</th>
<th>Recommended Solutions</th>
</tr>
</thead>
</table>
| Erratic values; dispersion of duplicates | A. Trace organic contaminants in the water source  
  B. Poor pipetting/technique | A. Replace activated carbon filter or change source of UltraPure water |
| High NSB (>0.100) | A. Poor washing  
  B. Exposure of NSB wells to specific antibody | A. Re-wash plate and redevelop |
| Very low B₀ | A. Trace organic contaminants in the water source  
  B. Plate requires additional development time  
  C. Dilution error in preparing reagents | A. Replace activated carbon filter or change source of UltraPure water  
  B. Return plate to shaker and re-read later |
| Low sensitivity (shift in dose response curve) | Standard is degraded | Replace standard |
| Analyses of two dilutions of a biological sample do not agree (i.e., more than 20% difference) | Interfering substances are present | Purify sample prior to analysis by ELISA³ |
| Only Total Activity (TA) wells develop | Trace organic contaminants in the water source | Replace activated carbon filter or change source of UltraPure water |

Additional Reading

Go to www.caymanchem.com/514531/references for a list of publications citing the use of Cayman’s PGEM Metabolite ELISA Kit.

References

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