INTENDED USE
The P450 Activity kit is designed to quantitatively measure the enzymatic activity of formaldehyde-producing enzymes such as Cytochrome P450s.
BACKGROUND
The cytochromes P450 (P450s) are a superfamily of heme containing enzymes that display tremendous diversity with regard to substrate specificity and catalytic activity\(^1\),\(^2\). P450s use a plethora of both exogenous and endogenous compounds as substrates in enzymatic reactions. Usually they form part of multicomponent electron transfer reactions (see figure).

Catalysis by the eukaryotic P450 enzymes involves a multistep reaction cycle that includes two steps in which electron transfer is accomplished from a redox partner. The diflavin protein, NADPH cytochrome P450 reductase (reductase) contains both FAD and FMN and can transfer both electrons needed for the catalytic cycle\(^3\). In some P450 reactions, the second electron of the reaction cycle also can be delivered by cytochrome b5\(^4\). The P450 enzymes and cofactors of the mammalian drug-metabolizing system are embedded in the membrane of the endoplasmic reticulum\(^5\). The P450s play a crucial role in the development of new drug entities as drug-drug interactions commonly arise from the inhibition of cytochrome P450 activities.

Lipid plays an important role in the reconstitution of P450-dependent activities after protein purification\(^6\). Most in vitro studies for the reconstitution of P450 activities use dilaurylphosphatidylcholine (DLPC) as the lipid component. The reconstitution of enzymatic activity involves a concentrated incubation of P450, its redox partners (NADPH and reductase), and lipid followed by dilution into the final assay components. The reported preincubation conditions vary significantly\(^7\).

ASSAY PRINCIPLE
The P450 Activity kit is designed to quantitatively measure the enzymatic activity of formaldehyde-producing enzymes such as Cytochrome P450s. The kit is unique in that the fluorescent substrate is not involved in the multicomponent P450 reaction, but measures the product of the demethylation, formaldehyde. No separation or washing is required. The kit has been validated for several P450 systems and should work with any biological system that is producing formaldehyde as a product of demethylation.

The kit provides an optimized buffer for P450, lyophilized vials of the cofactor, NADPH, for the reaction, a stable formaldehyde standard, the Formaldehyde Detection Reagent (FDR) and two 96 well plates for detecting the generated fluorescent signal. The end user will have to provide the microsomal, baculosome system or the recombinant P450, reductase and cytochrome b5 system and any cofactors, etc. necessary for activity, along with any candidate drugs, inhibitors or activators being tested. The reaction should be carried out in our supplied buffer or a similar PBS based buffer system.

Following the P450 NADPH-induced reaction, the generation of formaldehyde can be stopped by addition of a suitable inhibitor, or the supplied stop solution of acetic acid. The FDR is then added to all the wells. If calibration to formaldehyde is needed (for cross lab comparisons) then a formaldehyde standard curve generated from the supplied standard should be run. After a short incubation at 37°C for 30 minutes, the fluorescent product is read at 510 nm in a fluorescent plate reader with excitation at 450 nm. The P450 activity is determined based upon formaldehyde production. We have provided two 96 well plates for measurement but this assay is adaptable for higher density plate formats. If substituting their own plates, the end user should ensure that their black HTS plate is suitable for use with these reagents prior to running samples.
SUPPLIED COMPONENTS

- **Black Half Area 96 Well Plate Two plates**
- **Assay Buffer 60 mL**: A 100 mM potassium phosphate buffer at pH 7.4 containing 0.005% gentamicin.
- **NADPH lyophilized 2 vials**: Reduced β-nicotinamide adenine dinucleotide 2'-phosphate (NADPH) freeze dried with stabilizers and stored in desiccators.
- **Stop Solution 1 mL**: A 1M solution of Acetic Acid in water. CAUTION: Acid solution.
- **Formaldehyde Standard 0.5 mL**: 2,000 μM formaldehyde solution in deionized water. Outer container has formaldehyde absorbing pad. The standard is stable if kept tightly sealed. KEEP TIGHTLY SEALED
- **Formaldehyde Reagent 5 mL**: Special formulation of reagents to detect formaldehyde in solution. Contains ≤ 0.09% sodium azide as a preservative.

PLATE SEALERS

STORAGE INSTRUCTIONS

- All components of this kit should be stored at 4°C until the expiration date of the kit.

OTHER MATERIALS REQUIRED.

- Incubator capable of accurately maintaining 37°C.
- P450 systems. Microsome, Cerosome, baculosome or supersome P450 systems, or recombinant P450, NADPH/P450 oxidoreductase and cytochrome b5 and Dilaurylphosphatidylcholine (DLPC) as the lipid used for reconstitution.
- Repeater pipet with disposable tips capable of dispensing 25 μL.
- Fluorescence 96 well plate reader capable of reading fluorescent emission at 510 nm, with excitation at 450 nm. Set plate parameters for a 96-well Corning Costar 3694 plate.
- The sensitivity of fluorescent assays is dependant on the capabilities of the plate reader. If your plate reader has adjustable gain you can modify the signals obtained from the assay by increasing or decreasing the gain settings, by changing the aperture settings for monochromator based readers, or by changing the band pass width of the emission and/or excitation filters on some readers. Please review the plate reader manual for details.
- Signals expressed in this insert are Relative Fluorescent Units (RFU) and were obtained on our plate readers. The RFU numbers you obtain may be different from these, but the assay results should be similar.
- Software for converting raw relative fluorescent unit (FLU) readings from the plate reader and carrying out four parameter logistic curve (4PLC) fitting. Contact your plate reader manufacturer for details.

PRECAUTIONS

www.innov-research.com
Ph: 248.896.0145 | Fx: 248.896.0149
As with all such products, this kit should only be used by qualified personnel who have had laboratory safety instruction. The complete insert should be read and understood before attempting to use the product.

Some of the components of this kit contain sodium azide as a preservative, which may react with lead or copper plumbing to form potentially explosive complexes. When disposing of reagents always flush with large volumes of water to prevent azide build-up.

**SAMPLE TYPES AND PREPARATION**

P450 enzyme systems diluted in the supplied Assay Buffer provided or a typical 0.1M phosphate buffer at pH 7.4 are compatible with this assay.

**P450 Demethylating Reaction Conditions**

We have ensured the P450 Activity Assay detects the activity of the 2B4, 2D6 and 3A4 P450 systems. Below we have listed the conditions we used in validating this fluorescent detection system and the ability to quantitate the formaldehyde produced by the Cyp 2B4 P450 enzymatic reaction.

**Typical Cyp 2B4 Enzyme Reaction**

To duplicate wells add 15 μL of P450 enzyme system (equivalent molar ratios of 2B4 P450, Cytochrome P450 Oxidoreductase, and Cytochrome b5 in a pre-sonicated 0.66 mg/mL DLPC solution), followed by 75 μL of the supplied Assay Buffer and 5 μL of P450 substrate. Seal the plate and incubate for 5 minutes at 37°C prior to addition of 5 μL of the reconstituted supplied NADPH activator. Seal the plate again and incubate for 15 minutes at 37°C. Add 5 μL of the supplied Stop Solution followed by the addition of 25μL of the FDR to each well. Reseal the plate and incubate at 37°C for 30 minutes. For calibration purposes to formaldehyde, the 15 μL of P450 enzyme solution is replaced with standards made from the supplied Formaldehyde stock. For calibration purposes to formaldehyde, the P450 enzyme solution is replaced with standards made from the supplied Formaldehyde stock.

**Reaction Overview**

**P450 Reaction**

1. Carry out demethylating enzyme reaction.
2. Stop the reaction (optimal), add FDR.

**Formaldehyde Detection**

3. Incubate at 37°C for 30 minutes, read signal.
4. Calibrate to Formaldehyde generated.

**REAGENT PREPARATION**

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Allow the kit reagents to come to room temperature for 30 minutes. We recommend that all standards and samples be run in duplicate to allow the end user to accurately determine P450 activity. Ensure that all samples have reached optimal temperature for the P450 reaction and have been diluted as appropriate prior to running them in the kit.

**NADPH Preparation**
Remove a vial of NADPH from the desiccator and add 600 μL of the Assay Buffer to the vial and vortex thoroughly. Store any unused reconstituted NADPH at ≤ -20°C for no more than 2 weeks.

**Formaldehyde Standard Preparation**
Label six glass test tubes as #1 through #6. Pipet 400 μL of Assay Buffer into tube #1 and 250 μL into tubes #2-#6. Add 100 μL of the Formaldehyde stock solution to tube #1 and vortex completely. Add 250 μL of tube #1 to tube #2 and vortex completely. Repeat the serial dilutions for tubes #3 through #6. The concentration of formaldehyde in tubes 1 through 6 will be 400, 200, 100, 50, 25, and 12.5 μM.

Use all Standards within 2 hour of preparation.

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<th>Std 1</th>
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<th>Std 4</th>
<th>Std 5</th>
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<td><strong>Addition</strong></td>
<td>Stock</td>
<td>Std 1</td>
<td>Std 2</td>
<td>Std 3</td>
<td>Std 4</td>
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<td><strong>Volume of Addition (μL)</strong></td>
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<td><strong>Final Conc (μM)</strong></td>
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<td>200</td>
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ASSAY PROTOCOL

1. P450 reaction volume should be no more than 100 μL in each well including all cofactors, inhibitors and activators so that 25 μL of FDR can be added to each well for detection.
2. Use the plate layout sheet on the back page of the insert to aid in proper sample and standard identification.

P450 Reaction

3. Pipet 95 μL of Assay Buffer as a Zero standard, standards or samples including all cofactors, substrates and/or inhibitors into the duplicate wells in the black plate. Seal with the plate sealer and incubate for 15 minutes at 37°C.
4. Add 5 μL of the reconstituted NADPH to each well, seal the plate and incubate at 37°C for 15-60 minutes (incubation time varies and is based upon the system and microsomes used - see pages 7 and 13).
5. Add 5μL of Stop Solution to each well.

Formaldehyde Detection

6. Add 25 μL of the Formaldehyde Detection Reagent to each well using a repeater pipet.
7. Gently tap the sides of the plate to ensure adequate mixing of the reagents.
8. Incubate at 37°C for 30 minutes. Room temperature incubation will yield approximately 75% of the fluorescent signal generated with 37°C incubation.
9. Set plate parameters for a 96-well Corning Costar 3694 plate. Read the fluorescent signal from each well in a plate reader capable of reading the fluorescent signal at 510 nm with excitation at 450 nm. Please contact your plate reader manufacturer for suitable filter sets. This assay requires a plate reader with efficient fluorescence optics. Please refer to page 6 for details on increasing sensitivity.
CALCULATION OF RESULTS
Average the duplicate FLU readings for each standard and sample. Create a standard curve by reducing the data using the 4PLC fitting routine on the plate reader, after subtracting the mean FLUs for the zero standard. The sample activity obtained should be multiplied by the dilution factor to obtain neat sample values.

Typical Data
Cyp 2B4 P450 Demethylation of Benzphetamine
Low Sensitivity (Gain Setting at Low)

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<th>pmoles/100μL of P450 System</th>
<th>Mean FLU</th>
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Typical Standard Curve
Typical 2B4 P450 Standard Curve
Low Sensitivity - (Plate Reader Gain Set on Low)
INTERFERENCE STUDIES
The following additives were added to the 2B4 P450 enzyme reaction with its substrate to test for interference with the signal generation.

Organic Solvents
Additions of 1% and 5% of the organic solvents ethanol, methanol, N,N-dimethylformamide and dimethylsulfoxide were tested in the 2B4 P450 assay. DMF and DMSO at 5% and ethanol at 1% showed small negative effects on the signal (< 9% decrease). Methanol at 1% gave a 15% drop in signal and at 5% a 20% decrease in P450 generated signal.

Preservatives
0.09% sodium azide or 0.005% gentamicin in the assay buffer increased the P450 generated signal by 20%. 0.09% Kathon inhibited the signal by 83.2%.
Solvent studies described here were carried out with a 62.5 mM potassium phosphate buffer at pH 7.4 containing 0.005% gentamicin. Preservative studies were carried out in the same buffer with various added preservative concentrations.
Phosphate buffers with molarities up to 0.5M with the 2B4 P450 system are compatible with the assay.
CEROSOME/MICROSOME EXPERIMENTS
We tested the assay with a number of microsomal preparations using both rat and human liver native microsomes and with P450 co-expressed in Saccharomyces cerevisiae. In each case the microsomal preparation was incubated with substrate (1 mM erythromycin for 3A4, 20 mM dextromethorphan for 2D6 and 20 mM benzphetamine for 2B4) and NADPH at 37°C for 60 minutes and stopped using the provided Stop Solution. Production of formaldehyde from the individual P450 systems was then detected by addition of FDR and incubation at 37°C for 30 minutes. Fluorescence intensity was then measured.
P450 INHIBITION EXPERIMENTS

Inhibition of 2B4 P450 Activity
The 2B4 P450 inhibitor, 1-chlorophenyl imidazole (CPI), was added to P450 enzyme system with its substrate at concentrations ranging from 0 to >200 µM in the microtiter plate well. The following graph shows the effect on 2B4 P450 signal.
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