

# User Manual PathHunter<sup>®</sup> IL6R/IL6ST Bioassay Kit

For Chemiluminescent Detection of IL6R/IL6ST Dimerization

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Please read entire user manual before proceeding with the assay. For additional information or Technical Support see contact information below.

### Overview

PathHunter IL6R/IL6ST Bioassay kits provide a robust, highly sensitive, and easy-to-use cell-based functional assay to study potency of anti-IL-6R or anti-IL-6 antibodies and their neutralizing antibodies. The bioassay kits contain all the reagents needed for a complete assay including cells, detection reagents, cell plating reagent, positive control agonist, and assay plates. The pre-validated, frozen cells have been manufactured for single use and are provided in a ready-to-assay format that saves time and adds convenience.

## **Technology Principle**

These assays utilize Enzyme Fragment Complementation (EFC) technology, where the  $\beta$ -galactosidase ( $\beta$ -gal) enzyme is split into two fragments, ProLink (PK) and Enzyme Acceptor (EA). Independently these fragments have no  $\beta$ -gal activity; however, when forced to complement through protein-protein interactions, they form an active  $\beta$ -gal enzyme.

The PathHunter IL6R/IL6ST Bioassay is an application of the DiscoverX Dimerization Assay platform. The assay is designed to detect the ligand-induced interaction of the interleukin-6 receptor (IL-6R) with the interleukin-6 signal transducer protein (IL-6ST). The cells have been engineered to co-express IL-6ST fused to PK, and IL-6R fused to EA. Binding of an agonist to the IL-6R causes multimerization of the IL-6R and IL-6ST receptors, resulting in activation of downstream signaling events. This brings the two  $\beta$ -gal fragments into close proximity, forcing complementation. Heterodimerization of the two receptor chains results in the formation of a functional  $\beta$ -gal enzyme that hydrolyzes the substrate to generate a chemiluminescent signal. Blocking IL-6 binding with an anti- ligand or anti-receptor antibody can prevent this interaction, resulting in a loss of signal.



# **Materials Provided**

List of Components	93-1045Y3-00043	93-1045Y3-00044
PathHunter U2OS IL6R/IL6ST Bioassay Cells (No. of vials)	2	10
PathHunter Bioassay Detection Kit (No. of datapoints)	200	1,000
Detection Reagent 1 (mL)	2	10
Detection Reagent 2 (mL)	8	40
AssayComplete <sup>™</sup> Cell Plating Reagent 5	1 x 100 mL	3 x 100 mL
Protein Dilution Buffer	1 x 50 mL	2 x 50 mL
Control Agonist, IL-6 (No. of vials)	1	1
96-well Clear-Bottom TC Treated, Sterile Plates w/Lid Plates)	2	10

# **Storage Conditions**

#### PathHunter U2OS IL6R/IL6ST Bioassay Cells

Cells are shipped on dry ice and should arrive in a frozen state. Please contact technical support immediately, if the cells received were already thawed.

- Short-term (less than 24 hours): Store vials at -80°C immediately upon arrival. (DO NOT store at -80°C for longer than 24 hours).
- Long-term (greater than 24 hours): Vials should ONLY be stored in the vapor phase of liquid nitrogen.



Safety Warning: A face shield, gloves, and a lab coat should be worn at all times when handling frozen vials. Use tongs to remove cryovials from liquid nitrogen storage, and place the vials immediately on dry ice in a covered container. Wait at least 1 minute for any liquid nitrogen that may be present inside the vial to evaporate. Do not touch the bottom of the vials at any time to avoid inadvertent thawing of the cells.

#### PathHunter Bioassay Detection Kit

Store at -20°C. Once thawed, the detection reagents can be kept at 4°C for up to 4 days. For long-term storage (up to the expiration date listed in the kit's certificate of analysis), the reagent should be aliquoted and stored at -20°C until needed.

For the ten-plate kit, if all the plates will not be used at the same time, we recommend making five aliquots for each of the two detection reagents. Each aliquot will be adequate for two assay plates. Make five aliquots of 2.3 mL each for Detection Reagent 1, and five aliquots of 9.2 mL each for Detection Reagent 2. Sufficient reagent volumes are provided in the kit to make these aliquots.

If reagents will be used for a single plate, then the remaining Detection Reagents can be frozen. The detection reagents can be thawed and frozen for a total of three times without loss in performance.

#### AssayComplete Cell Plating Reagent 5

Once thawed, the Cell Plating Reagent can be stored at 4°C for up to 4 weeks. For longer storage (up to the expiration date listed in the kit's Certificate of Analysis), the reagent should be aliquoted and stored at -20°C until needed. Avoid multiple freeze-thaw cycles.

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To make aliquots suitable for testing one assay plate each, 30 mL of reagent per aliquot can be dispensed and frozen down.

#### **Protein Dilution Buffer**

Once thawed, the Protein Dilution Buffer can be stored at 4°C for up to 4 weeks. For longer storage (up to the expiration date listed in the kit certificate of analysis), the reagent should be aliquoted and stored at -20°C until needed. Avoid multiple freeze-thaw cycles.

To make aliquots suitable for testing one assay plate each, 10 mL of reagent per aliquot should be dispensed and frozen down. This amount may vary depending on your stock sample concentrations and should be adjusted accordingly.

#### Recombinant Human IL-6 Control Agonist

Store at -20°C until ready to use (up to the expiration date listed in the kit's Certificate of Analysis). Centrifuge the vial prior to opening to maximize recovery. Reconstitute to a concentration of 100  $\mu$ g/mL by adding 200  $\mu$ L of Protein Dilution Buffer and gently shake (do not vortex) for ten minutes to increase solubility. Reconstituted ligand is stable for 12 months at -20 to -80°C, or for 1 week at 2-8 °C.

#### 96-Well Tissue Culture Treated Plates

Store at room temperature.

### **Additional Materials Required**

The following equipment and additional materials are required to perform these assays:

Material	Ordering Information	
V-Bottom 96-well ligand dilution plates	92-0011	
Multimode or luminescence plate reader	Refer to Instrument Compatibility Chart at discoverx.com/instrument-compatibility	
Disposable reagent reservoir	Thermo Fisher Scientific, Cat. No. 8094 or similar	
Single and multichannel micro-pipettors and pipette tips		

## **Protocol Schematic**

Quick-Start Procedure: In a 96-well tissue culture treated plate, perform the following steps:



## **Detailed Protocols**

#### Day 1: PathHunter Bioassay Cell Preparation\_

The following protocol is for thawing and plating frozen PathHunter U2OS IL6R/IL6ST Bioassay cells from cryovials.

- 1. Before thawing the cells, ensure that all the necessary materials are set up in the tissue culture hood. This includes:
  - a. One 25 mL reagent reservoir
  - b. One 15 mL conical tube
  - c. A micropipettor (P1000) set to dispense 1 mL
  - d. A multichannel pipette and tips set to dispense 80 µL
  - e. A bottle of Cell Plating Reagent 5 (CP5, pre-warmed in a 37°C water bath for 15 minutes)
  - f. A white-walled, opaque-bottom 96-well assay plate
- 2. Dispense 9.6 mL of CP5 into the 15 mL conical tube
- 3. Remove the cryovial from liquid nitrogen and immediately place in dry ice.



**DO NOT** use heated water bath to thaw the vial. Wipe down the cryovial quickly with 70% EtOH, and immediately bring it into the tissue culture hood. Hold the cryovials at the cap, **DO NOT** touch the sides or bottom of the vial to avoid thawing of the cell pellet.

- 4. Add 1 mL of pre-warmed CP5 from the 15 mL conical tube to the cryovial to thaw the cell pellet. The reagent should be added slowly along the wall of the vial. Mix the cells gently by pipetting up and down several times to break up any clumps. Transfer the cell suspension to the conical tube containing the remaining 8.6 mL of CP5. Remove all medium/suspension in the tube to ensure maximum recovery of all the cells from the vial.
- 5. Mix the tube by gently inverting the tube several times to ensure that the cells are properly mixed in the medium, without creating any froth in the suspension, and immediately pour the suspension into the 25 mL reagent reservoir.
- 6. Add 80 μL of cells to each well of the 96-well assay plate using the multichannel pipette. Replace lid and leave the plate at room temperature for 15 minutes to allow the cells to settle uniformly, and to minimize potential edge effects. Gently place the assay plate in a tissue culture incubator set to 37°C and 5% CO<sub>2</sub> for 24 hours before proceeding with the assay.

#### Day 2: Sample Preparation

The following protocol is designed for testing purified biologics. The PathHunter assays can also be run in the presence of high levels of serum or plasma, without adversely impacting assay performance. In some assays, standard curves of the control can be prepared in neat serum or plasma and added directly to cells without further dilution. For best results, the optimized minimum required dilution of crude samples should be empirically determined.

A 1:3 serial dilution for the Agonist/Control Ligand, IL-6 has been used in this protocol. The volumes listed below are designed for running samples from one dose-response curve in triplicate on the assay plate (Refer to the Representative Plate Map for Agonist/Inhibitor Dose-Response Curve).



Control Ligand Serial Dilutions: Make eleven 3-fold serial dilutions of the antibody in a dilution plate or vials.

- Prepare the reference agonist (IL-6) dose-response curve, which will serve as a positive control in this assay. Agonist is prepared at 5X the desired final concentration as it will be diluted by adding 20 μL to the 80 μL of medium present in the assay plate.
  - a. Add 80 μL of Protein Dilution Buffer (PDB) to wells A2 to A12 of a dilution plate (e.g. a V-bottom polypropylene 96-well dilution plate, DiscoverX, Cat. No. 92-0011 or similar).
  - b. Add 200 μL of Reconstitution Buffer to the IL-6 vial containing 20 μg of lyophilized powder to make a 100 μg/mL stock solution. Gently shake (do not vortex) for ten minutes to completely dissolve the powder.
  - c. Make a 1:50 dilution by adding 98 μL of PDB to a non-binding microcentrifuge tube. Add 2 μL of the 100 μg/mL IL-6 stock to this tube. Mix thoroughly by pipetting up and down several times with the pipettor set to at least 100 μL.
  - d. Add 475 μL of PDB to another non-binding microcentrifuge tube, and then add 25 μL of the IL-6 1:50 dilution that was just prepared earlier in Step 1c. This results in a 100 ng/mL solution (5X the final 20 ng/mL curve top).
  - e. Transfer 120 µL to well No. A1 of the dilution plate.
  - f. Using a fresh pipet tip, transfer 40 μL from well No. A1 into well No. A2 and mix by pipetting up and down several times. Replace the pipette tip, and transfer 40 μL from well No. A2 into well No. A3. Mix by pipetting up and down several times. Repeat this process until well No. A11 is reached, resulting in an eleven point, 1:3 dilution series. No ligand is transferred to any wells in column 12 as these will serve as the negative control wells.
- 2. Prepare the reference IL-6R antagonist antibody curve at 6X the desired final concentration, with 11-point dilution series. The dilution scheme for each antibody will need to be empirically determined. Note: No antibody is added to any wells in column 12, as these will be the negative control wells.

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- 3. Remove the assay plate from the 37°C and 5% CO<sub>2</sub> incubator and bring into the tissue culture hood.
- 4. Transfer 20 μL of the anti-IL6R antibody from the dilution plate to the appropriate wells of the assay plate (Refer to the Representative Plate Map for Agonist/Inhibitor Dose-Response Curve).
- 5. Transfer 20 µL from the agonist reference curve in the dilution plate as prepared in Step 1, to the appropriate wells of the assay plate.
- 6. Incubate for 3 hours at 37°C and 5% CO<sub>2</sub>.
- 7. Agonist challenge for an anti-IL-6R antibody curve: The EC<sub>80</sub> of the DiscoverX IL-6 was determined to be approximately 2 ng/mL. If using IL-6 from a different vendor, the EC<sub>80</sub> should be determined empirically, prior to testing samples. Prepare the agonist challenge at 6X the desired final concentration. The following steps will provide enough agonist challenge for a single antagonist curve run in triplicate:
  - a. Dilute 2 μL of the 100 μg/mL stock with 98 μL of PDB in a non-binding Eppendorf tube and mix thoroughly by pipetting up and down several times to make a 1:50 dilution.
  - b. Dilute 18 µL of this 1:50 dilution mixture from Step 2a with 3 mL of PDB in a fresh 15 mL conical tube and mix thoroughly by pipetting up and down several times.
  - c. Transfer the solution prepared in step 7 into a reagent reservoir, and add 20 µL/well to the appropriate wells of the assay plate containing the anti-IL6 antibody.
- Add 20 μL from each well of the agonist challenge prepared in Step 7 to the appropriate wells of the assay plate containing the anti-IL-6 antibody.
- 9. Incubate the assay plate at 37°C and 5% CO<sub>2</sub> for 3 hours, before continuing on to the detection steps.

#### Day 2: Detection\_

- Add 10 µL of Detection Reagent 1 to each well of the assay plate. Place the plate onto an orbital shaker at 350 rpm for 1 minute to cause even mixing.
  PathHunter Bioassay Detection
- 2. Incubate the plate at room temperature for 15 minutes in the dark.
- 3. Add 40  $\mu L$  of Detection Reagent 2 to each well of the assay plate.
- 4. Incubate the plate at room temperature for one hour in the dark.
- 5. Read sample on a standard luminescence plate reader at 0.1 to 1 second/well for photomultiplier tube (PMT) readers or 5-10 seconds for imager.

Note: For crude biologic samples, gently removing the liquid from all wells and replacing with 100 µL of Cell Plating Reagent before the addition of the detection reagents can result in higher signal. Additional Cell Plating Reagent is necessary for this method.



PathHunter Bioassay Detection Reagents are light sensitive, thus incubation in the dark is necessary

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#### Representative Plate Map for Agonist/Inhibitor Dose-Response Curve



**Assay Plate Map:** This plate map shows two interdigitated 11-point dose curves, with 3 replicates per dose point, for a test and reference sample tested with their appropriate dilution schemes.

### **Typical Results**

Refer to the cell line-specific Certificate of Analysis, included the kit, for representative example of the assay performance.

### **Troubleshooting Guide**

Problem	Potential Cause	Proposed Solution	
No response	Improper thawing procedure	Refer to thawing instructions in PathHunter Bioassay Cell Preparation section of of this user manual.	
	Improper ligand used or improper ligand incubation time	See datasheet for recommended ligand and assay conditions.	
	Improper preparation of ligand (agonist or antagonist)	Refer to vendor specific datasheet to ensure proper handling, dilution, and storage of ligand.	
	Improper time course for induction	Optimize time course of induction with agonist and antagonist.	
Low or no signal	Improper preparation of detection re- agents	Detection reagents should ideally be prepared just prior to use and are sensitive to light.	
	Problem with microplate reader	Microplate reader should be in luminescence mode. Read at 0.1-1 second/well.	
Experimental S:B does not match datasheet value	Incorrect incubation temperature	Confirm assay conditions.	
		Check and repeat assay at correct incubation temperature as indicated on the assay datasheet.	
	Improper preparation of ligand (agonist or antagonist)	Some ligands are difficult to handle. Confirm the final concentration of ligands.	
	Suboptimal agonist challenge concen- tration	Perform agonist curve to reassess $EC_{80}$ with the ligand provided in the kit. Perform antibody titrations with $EC_{80}$ and $EC_{90}$ agonist challenge concentrations to re-optimize assay window.	
$EC_{50}$ is right-shifted	Improper ligand handling or storage	Check ligand handling requirements.	
	Difference in agonist binding affinity	Confirm that the ligand used is comparable to the ligand in the Product Certificate of Analysis.	
	Problems with plate type and compound stability	Hydrophobic compounds should be tested for solubility and may may require evaluation in alternative dilution buffers.	
		Non-binding surface plates may be necessary for hydrophobic compounds.	

For additional information or technical support, please contact technical support listed below.

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