

#### **PRODUCT DATASHEET**

## Ready-to-Assay<sup>™</sup> ChemiBrite<sup>™</sup> EP<sub>2</sub> Prostanoid Receptor Frozen Cells

#### CATALOG NUMBER: HTS185LRTA

**CONTENTS**: Pack contains 2 vials of mycoplasma-free cells, 1 ml per vial. **STORAGE**: Vials are to be stored in liquid N<sub>2</sub>. Media Component at 4°C (-20°C for prolonged storage).

#### BACKGROUND

Ready-to-Assay<sup>™</sup> GPCR frozen cells are designed for simple, rapid calcium assays with no requirement for intensive cell culturing. Eurofins Discovery Services has optimized the freezing conditions to provide cells with high viability and functionality post-thaw. The user simply thaws the cells and resuspends them in media, dispenses cell suspension into assay plates and, following overnight recovery, assays for calcium response.

ChemiBrite<sup>™</sup> cells express a novel variant of clytin, a calcium-activated photoprotein, to enable sensitive luminescent detection of ligand-induced calcium flux. The ChemiBrite<sup>™</sup> version of clytin contains a mutation that increases its affinity for calcium to a level that permits detection of cytosolic calcium in many cells with greater sensitivity than other photoproteins targeted to the mitochondria. Luminescent calcium assays offer several advantages over fluorescent calcium assays including increased sensitivity and lack of interference from fluorescent compounds.

Prostanoids bind to a family of 8 GPCRs to exert their biological effects (Narumiya and FitzGerald, 2001). The prostanoid PGE2 causes pain, vasodilation, immunosuppression of T cells, bone resorption and promotion of carcinogenesis. Four related GPCRs, EP1, EP2, EP3 and EP4, each bind to PGE2, but the different G protein coupling status of each receptor leads to distinct biological effects. EP2 couples primarily to Gs to increase intracellular cAMP levels. Mice deficient in EP2 receptor showed impaired ovulation and fertilization, salt-sensitive hypertension (Kennedy et al., 1999). It has been shown that EP2 receptors are also involved in cancer associated immunodeficiency. Thus, genetic knockout of the EP2 receptor reduced tumor growth and prolonged survival in mice that had undergone isograft injection of MC26 or Lewis lung carcinoma cells (Yang et al., 2003). The cloned human EP2 receptor-expressing ChemiBrite™ cells were made by stable transfection of HEK293 cells with ChemiBrite™ clytin, the EP2 receptor and a promiscuous G protein to couple the receptor to the calcium signaling pathway. These stability tested cells are ready for luminescent analysis of agonists, antagonists and modulators at the EP2 receptor.

#### **USE RESTRICTIONS**

Please see User Agreement (Label License) for further details. One such restriction is that the contents of the supplied vial(s) are limited to a single use and shall not be propagated and/or re-frozen by licensee.

#### WARNINGS

For Research Use Only; Not for Use in Diagnostic Procedures Not for Animal or Human Consumption

#### GMO

This product contains genetically modified organisms. Este producto contiene organismos genéticamente modificados. Questo prodotto contiene degli organismi geneticamente modificati. Dieses Produkt enthält genetisch modifizierte Organismen. Ce produit contient organismes génétiquement des modifiés. Dit product bevat genetisch gewijzigde organismen. Tämä tuote sisältää geneettisesti muutettuja organismeja. Denna produkt innehåller genetiskt ändrade organismer.

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#### **APPLICATIONS**

Calcium Flux Assays: Luminescent Mode and Fluorescent Mode ; cAMP accumulation

#### **APPLICATION DATA**

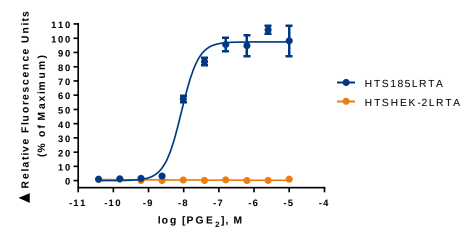


Figure 1. Representative data for activation of  $EP_2$  receptor stably expressed in HEK293 cells induced by Prostaglandin E2 using a fluorescent calcium flux assay.  $EP_2$ -expressing HEK293 cells were seeded at 50,000 cells per well into a 96-well plate, and the following day the cells were loaded with a fluorescent calcium indicator. Calcium flux in response to the indicated ligand with a final concentration of 0.5% DMSO was determined on a Molecular Devices FLIPR<sup>TETRA®</sup> with ICCD camera. Maximal fluorescence signal obtained in this experiment was 12,000 RLU. Similarly parental cells (catalog #: HTSHEK-2L) were tested to determine the specificity of the resulting signal.

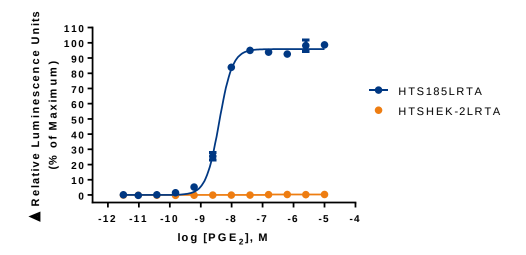


Figure 2. Representative data for activation of EP<sub>2</sub> receptor expressed in HEK293 cells induced by Prostaglandin E2 using a luminescent calcium flux assay. EP<sub>2</sub>–expressing HEK293 cells were loaded with 10µM coelenterazine for 3h at room temperature and calcium flux in response to the indicated ligand was determined on a Molecular Devices FLIPR<sup>TETRA</sup>® with ICCD camera in 96-well format. Luminescence signal obtained in this experiment was 400,000 RLU (Relative Light Units) as measured by area-under-curve for 80s post agonist addition using the provided protocol.

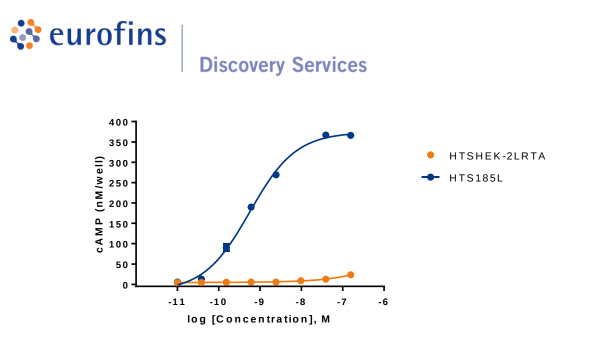


Figure 3. Representative data for activation of EP<sub>2</sub> receptor stably expressed in HEK293 cells induced by PGE2 using a cAMP accumulation assay. EP<sub>2</sub>–expressing HEK293 cells were seeded at 25,000 cells per well into a 96-well plate, and the following day the cells were treated with PGE2 for 10 minutes in the presence of 100  $\mu$ M IBMX and 0.5% DMSO to determine receptor-mediated cAMP generation using a time-resolved fluorescence resonance energy transfer (TR-FRET) assay measured on the BioTek Synergy. Maximal fluorescence signal obtained in this experiment was 400 nmoles cAMP/well. Similarly parental cells (catalog #: HTSHEK-2L) were tested to determine the specificity of the resulting signal.

Table 1. EC<sub>50</sub> values of EP<sub>2</sub>-expressing HEK293 cells

LIGAND	ASSAY	POTENCY EC <sub>50</sub> (nM)	REFERENCE
Prostaglandin E2	Calcium Flux - Fluorescence	8.0	Eurofins Internal Data
Prostaglandin E2	Calcium Flux - Luminescence	4.0	Eurofins Internal Data
Prostaglandin E2	cAMP Accumulation	6.0	Eurofins Internal Data

# **ASSAY SETUP**

#### Luminescence

Table 2. Settings for FLIPR<sup>TETRA</sup>® with ICCD camera option

Option	Setting
Read Mode	Luminescence
Ex/Em	None/None
Camera Gain	280,000
Gate Open	100 %
Exposure Time	0.9 sec
Read Interval	1 sec.
Dispense Volume	50 µl (25 µl for 384-well)
Dispense Height	95 µl (50 µl for 384-well)
Dispense Speed	50 µl/sec
Expel Volume	ΟμΙ
Analysis	Subtract Bias Sample 1



Table 3. Luminescence Assay Materials (Not provided)

Description	Supplier and Product Number
HBSS	Invitrogen: 14025
HEPES 1M Stock	Millipore Sigma: H3537
Quest Fluo-8 <sup>™</sup> , AM	AAT Bioquest: 21080
Prostaglandin E2 ligand	Cayman: 14010
Non-Binding 96/384 well Plates (for ligand prep)	Corning: 3605/ 3574
Black (clear Bottom) cell assay plates	Corning: 3904/ 3712
Coelenterazine-h (250µg). Prepare to 10mM	Promega: S2011

#### **Fluorescence**

Table 4. Settings for FLIPR<sup>TETRA®</sup> with ICCD camera option

Option	Setting
Read Mode	Fluorescence
Ex/Em	Ex470_495 / Em515_575
Camera Gain	2000
Gate Open	6 %
Exposure Time	0.53
Read Interval	1s
Dispense Volume	50 µl (25 µl for 384-well)
Dispense Height	95 μl (50 μl for 384-well)
Dispense Speed	50 µl/sec
Expel Volume	0 µl
Analysis	Subtract Bias Sample 1

Table 5. Fluorescence Assay Materials (Not provided)

Description	Supplier and Product Number
HBSS	Invitrogen: 14025
HEPES 1M Stock	Millipore Sigma: H3537
Probenicid	Sigma: P8761
Quest Fluo-8 <sup>™</sup> , AM	AAT Bioquest: 21080
Prostaglandin E2 ligand	Cayman: 14010
Non-Binding 96/384 well Plates (for ligand prep)	Corning: 3605/ 3574
Black (clear Bottom) cell assay plates	Corning: 3904/ 3712

#### **cAMP**

Table 6. Settings for Plate Reader

Option	Setting
Excitation	300 nm
Emission	665/620 nm

Table 7. cAMP Assay Materials (Not provided)

Description	Supplier and Product Number
HEPES 1M Stock	Millipore Sigma: H3537
IBMX Buffer	Sigma #I5879
96-Well Flat Bottom Microtiter Plates	Costar #3917
Non-Binding 96 well Plates (for ligand prep)	Costar: #3789
Prostaglandin E2 ligand	Cayman: 14010
cAMP Hi Range Kit	CisBio # 62AM6PEC



### Assay Protocol – Luminescence

- 1. Immediately upon receipt, thaw cells or place cells in liquid nitrogen.
- 2. Thaw cells rapidly by removing from liquid nitrogen and immediately immersing in a 37°C water bath. Immediately after ice has thawed, sterilize the exterior of the vial with 70% ethanol.
- 3. Add 1mL of pre-warmed Media Component to each vial of cells. Place contents from two vials into a 15 mL conical tube and bring the volume to 10 mL of Media Component.
- 4. Centrifuge the cell suspension at 190 x g for four minutes
- 5. Remove supernatant and add 10.5 mL of pre-warmed Media Component to resuspend the cell pellet.
- 6. Seed cell suspension into appropriate assay microplate (100 μL/well for 96-well plate, 25 μL/well for 384-well plate).
- 7. When seeding is complete, place the assay plate at room temperature for 30 minutes.
- 8. Move assay plate to a humidified  $37^{\circ}C 5\% CO_2$  incubator for 18-24 h.
- 9. Next day, prepare Assay buffer (HBSS, 20mM HEPES, pH 7.4) and Loading buffer (Assay buffer with 10μM coelenterazine). Note: Please prepare coelenterazine stock according to Manufacturer's Recommendations at 10mM to allow for 1:1000 dilution into Loading buffer (10uM final concentration). It is critical that coelenterazine solution is prepared at room temperature and is protected from light.
- 10. Remove plate from incubator and quickly invert plate on an absorbent pad and blot to remove all Media Component.
- Add Loading buffer to assay plate (100 μL/well for 96-well plate). Incubate plate for 1.5 h at room temperature, protected from light.
- 12. Prepare ligands in assay buffer at 3x final concentration in non-binding plates. Use Buffer Only Control Wells for Background Subtraction.
- 13. Create protocol for ligand addition. Please refer to FLIPR<sup>TETRA®</sup> settings provided in Table 2. Set time course for 180 s, with ligand addition at 10 s.
- 14. After the run is complete, apply subtract bias on sample 1. Export data to analyze using the area under the curve statistic.

### Assay Protocol – Fluorescence

- 1. Immediately upon receipt, thaw cells or place cells in liquid nitrogen.
- 2. Thaw cells rapidly by removing from liquid nitrogen and immediately immersing in a 37°C water bath. Immediately after ice has thawed, sterilize the exterior of the vial with 70% ethanol.
- Add 1mL of pre-warmed Media Component to each vial of cells. Place contents from two vials into a 15 mL conical tube and bring the volume to 10 mL of Media Component.
- 4. Centrifuge the cell suspension at 190 x g for four minutes
- 5. Remove supernatant and add 10.5 mL of pre-warmed Media Component to resuspend the cell pellet.
- 6. Seed cell suspension into appropriate assay microplate (100 μL/well for 96-well plate, 25 μL/well for 384-well plate).
- 7. Move assay plate to a humidified  $37^{\circ}C 5\% CO_2$  incubator for 18-24 h.
- 8. Next day, prepare Assay buffer (HBSS, 20mM HEPES, 2.5 mM Probenicid, pH 7.4) and Loading buffer (Assay buffer with 5 mM Fluo8 Dye). *Note: Please prepare Fluo8 stock according to Manufacturer's Recommendations*
- 9. Remove plate from incubator and quickly invert plate on an absorbent pad and blot to remove all Media Component.
- 10. Add Loading buffer to assay plate (100 μL/well for 96-well plate). Incubate plate for 1.5 h at room temperature, protected from light.
- 11. Prepare ligands in assay buffer at 3x final concentration in non-binding plates. Use Buffer Only Control Wells for Background Subtraction.
- 12. Create protocol for ligand addition. Please refer to FLIPR<sup>1ETRA</sup>® settings provided in Table 2. Set time course for 180 s, with ligand addition at 10 s.
- 13. After the run is complete, apply subtract bias on sample 1. We recommend using Negative Control Correction with Buffer Only Wells. Export data to according to research needs. For most Calcium Flux analysis using Export of Max Signal to end of run is sufficient.



#### Assay Protocol – cAMP

- 1. Immediately upon receipt, thaw cells or place cells in liquid nitrogen.
- 2. Thaw cells rapidly by removing from liquid nitrogen and immediately immersing in a 37°C water bath. Immediately after ice has thawed, sterilize the exterior of the vial with 70% ethanol.
- 3. Add 1mL of pre-warmed Media Component to each vial of cells. Place contents from two vials into a 15 mL conical tube and bring the volume to 10 mL of Media Component.
- 4. Centrifuge the cell suspension at 190 x g for four minutes
- 5. Remove supernatant and add 20 mL of pre-warmed Media Component to resuspend the cell pellet.
- 6. Seed cell suspension into appropriate assay microplate (100 μL/well for 96-well plate, 25 μL/well for 384-well plate).
- 7. When seeding is complete, place the assay plate at room temperature for 30 minutes.
- 8. Move assay plate to a humidified 37°C 5% CO<sub>2</sub> incubator for 18-24 h.
- 9. Prepare Assay Buffer (HBSS, 20mM HEPES, pH 7.4)
- 10. Prepare anti-cAMP-Cryptate and cAMP-D2 conjugate stocks. Please prepare stocks according to the Manufacturers Recommendations for reconstitution volume to be used according to Appendix-2 of the insert (Two-step protocol). Mix gently. Store at 10 to 35 °C after reconstitution. Reagent can only go through one cycle of freeze and thaw.
- 11. Prepare 1M IBMX Buffer: add 450 µL of DMSO to 1 vial of 100 mg IBMX powder.
- 12. Next day, prepare 2mM IBMX Solution: Place the 1M IBMX Buffer and the Assay Buffer in 50-60°C water bath for 10 min to 15 min. Add 10 μL of 1M IBMX Buffer to 4.990 mL of Assay Buffer. Mix by vortex. Place the 2 mM of IBMX Buffer into 34-40°C Incubator for at least 10 minutes or until ready to add to cells.
- 13. Prepare 25uL/well of compounds for testing.
- 14. Remove the cell assay plate from previous day from cell culture incubator. Quickly invert plate on an absorbent pad and blot to remove all Media Component. Add 25 µL of the 2 mM IBMX Buffer to the plate with the seeded GLP-1/HEK cells. Tap plate gently 3-4 times. Cover plate and incubate inside 34-40°C incubator, static until ready for compound addition.
- 15. Add 25 μL compounds, internal control and test sample dilutions to cell assay plate. Cover plate and incubate for 15 to 25 min at 20-25°C
- 16. Prepare fresh working dilutions of 1:24 of anti-cAMP-Cryptate and cAMP-D2 conjugate in Lysis Buffer. Protect from light. Do not vortex.
- 17. Add 125 μL of cAMP-D2 solution into 2,875 μL of Lysis Buffer for total volume of 3 mL.
- 18. Add 125 μL of anti-cAMP-Cryptate solution into 2,875 μL of Lysis Buffer for total volume of 3 mL.
- It is imperative that detection reagents are added to plate in the following order: Add 25 μL cAMP-D2 conjugate/lysis buffer solution to each well of cell assay plate. Add 25 μL anti-cAMP-Cryptate/lysis buffer solution to each well of cell assay plate.
- 20. Cover with aluminum plate sealer and incubate the cell assay plate 20-25°C, for 60 to 65 min (If available,
- use gentle plate shaker). Protect from light.
- 21. Read plate on a plate reader with 330 nm (excitation) and 665/620 nm (emission).
- 22. Calculate Ratio Emission 665/620 nm.

# HOST CELL

HEK293

## **EXONGENOUS GENE EXPRESSION**

Human PTGER2 cDNA (Accession Number: NM\_000956; see CODING SEQUENCE below) and a proprietary mutant clytin photoprotein expressed in a bicistronic vector

#### **CODING SEQUENCE**

ATG	GGC	AAT	GCC	TCC	AAT	GAC	TCC	CAG	TCT	GAG	GAC	TGC	GAG	ACG	CGA	CAG	TGG	CTT	CCC	CCA	GGC	GAA	
М	G	Ν	A	S	Ν	D	S	Q	S	Ε	D	С	E	Т	R	Q	W	L	P	Ρ	G	E	
AGC	CCA	GCC	ATC	AGC	TCC	GTC	ATG	TTC	TCG	GCC	GGG	GTG	CTG	GGG	AAC	CTC	ATA	GCA	CTG	GCG	CTG	CTG	
S	Р	А	I	S	S	V	М	F	S	A	G	V	L	G	Ν	L	I	А	L	А	L	L	
GCG	CGC	CGC	TGG	CGG	GGG	GAC	GTG	GGG	TGC	AGC	GCC	GGC	CGC	AGG	AGC	TCC	СТС	TCC	TTG	TTC	CAC	GTG	
GCG	CGC	CGC	TGG	CGG	GGG	GAC	GIG	GGG	TGC	AGC	GCC	GGC	CGC	AGG	AGC	TCC	CIC	TCC	TIG	TIC		CAC	CAC GIG



A R R W R G D V G C S A G R R S S L S L F H V CTG GTG ACC GAG CTG GTG TTC ACC GAC CTG CTC GGG ACC TGC CTC ATC AGC CCA GTG GTA CTG GCT TCG E L V F T D L L G T C L I S P V L A S TAC GCG CGG AAC CAG ACC CTG GTG GCA CTG GCG CCC GAG AGC CGC GCG TGC ACC TAC TTC GCT TTC GCC Y A R N Q T L V A L A P E S R A C T Y F A F А ATG ACC TTC TTC AGC CTG GCC ACG ATG CTC ATG CTC TTC GCC ATG GCC CTG GAG CGC TAC CTC TCG ATC M T F F S L A T M L M L F A M A L E R Y L S I Н Ρ Y F Y Q R R V S R S G G L A V L Ρ V A V S L L F C S L P L L D Y G O Y V O Y C P G ACC TGG TGC TTC ATC CGG CAC GGG CGG ACC GCT TAC CTG CAG CTG TAC GCC ACC CTG CTG CTG CTT CTC F I R H G R T A Y L Q L Y A T L L L L ATT GTC TCG GTG CTC GCC TGC AAC TTC AGT GTC ATT CTC AAC CTC ATC CGC ATG CAC CGC CGA AGC CGG I V S V L A C N F S V I L N L I R M H R R S R AGA AGC CGC TGC GGA CCT TCC CTG GGC AGT GGC CGG GGC GGC CCC GGG GCC CGC AGG AGA GGG GAA AGG G P S L G S G R G G P G A R R R R С G E GTG TCC ATG GCG GAG GAG ACG GAC CAC CTC ATT CTC CTG GCT ATC ATG ACC ATC ACC TTC GCC GTC TGC V S M A E E T D H L I L A I M T I T F A V C TCC TTG CCT TTC ACG ATT TTT GCA TAT ATG AAT GAA ACC TCT TCC CGA AAG GAA AAA TGG GAC CTC CAA S L P F T I F A Y M N E T S S R K E K W D L 0 GCT CTT AGG TTT TTA TCA ATT AAT TCA ATA ATT GAC CCT TGG GTC TTT GCC ATC CTT AGG CCT CCT GTT A L R F L S I N S I I D P W V F A I L R P P V CTG AGA CTA ATG CGT TCA GTC CTC TGT TGT CGG ATT TCA TTA AGA ACA CAA GAT GCA ACG CAA ACT TCC V L C C R I S L R T Q D A T-T Q T S M R S L R L TGT CCT ACA CAG TCA GAT GCC AGT AAA CAG GCT GAC CTT TGA C P T Q S D A S K Q A D L Stp

#### **RELATED PRODUCTS**

Product Number	Description
HTSHEK-2L	ChemiScreen™ HEK293 Parental Cell Line (control cells)
HTS185M	ChemiScreen <sup>™</sup> EP <sub>2</sub> Prostanoid family receptor membrane prep
HTS081L	ChemiBrite <sup>™</sup> TP Prostanoid family Stable Cell Line
HTS081LRTA	Ready-to-Assay™ ChemiBrite™ TP Prostanoid Receptor Frozen Cells
HTS185L	ChemiBrite <sup>™</sup> HEK stable EP <sub>2</sub> Prostanoid Receptor Cell Line
HTS185LRTA	Ready-to-Assay <sup>™</sup> ChemiBrite <sup>™</sup> EP <sub>2</sub> Prostanoid Receptor Frozen Cells
HTS142L	ChemiBrite <sup>™</sup> HEK stable EP₄ Prostanoid Receptor Cell Line
HTS142LRTA	Ready-to-Assay™ ChemiBrite™ EP₄ Prostanoid Receptor Frozen Cells
HTS091L	ChemiBrite <sup>™</sup> HEK stable DP Prostanoid Receptor Cell Line
HTS091LRTA	Ready-to-Assay™ ChemiBrite™ DP Prostanoid Receptor Frozen Cells
HTS092L	ChemiBrite <sup>™</sup> HEK stable EP <sub>3</sub> Prostanoid Receptor Cell Line
HTS092LRTA	Ready-to-Assay <sup>™</sup> ChemiBrite <sup>™</sup> EP <sub>3</sub> Prostanoid Receptor Frozen Cells
HTS093L	ChemiBrite <sup>™</sup> HEK stable FP Prostanoid Receptor Cell Line
HTS093LRTA	Ready-to-Assay™ ChemiBrite™ FP Prostanoid Receptor Frozen Cells
HTS099L	ChemiBrite <sup>™</sup> HEK stable EP <sub>1</sub> Prostanoid Receptor Cell Line
HTS099LRTA	Ready-to-Assay™ ChemiBrite™ EP <sub>1</sub> Prostanoid Receptor Frozen Cells
HTS131L	ChemiBrite <sup>™</sup> HEK stable IP1 Prostanoid Receptor Cell Line
HTS131LRTA	Ready-to-Assay™ ChemiBrite™ IP1 Prostanoid Receptor Frozen Cells



#### REFERENCES

- 1. Kennedy CR *et al.*(1999) Salt-sensitivity hypertension and reduced fertility in mice lacking the prostaglandin EP<sub>2</sub> receptor. *Nat. Med.* 5:217-220.
- 2. Narumiya S and FitzGerald GA (2001) Genetic and pharmacological analysis of prostanoid receptor function. *J. Clin. Invest.* 108: 25-30.
- 3. Yang N *et al.* (2003) Cancer-associated immunodeficiency and dendritic cell abnormalities mediated by the prostaglandin EP<sub>2</sub> receptor. *J. Clin. Invest.* 111: 727–735.

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