# Protocol of RXR-BLA HEK 293T Cell-based Assay for Highthroughput Screening

# DOCUMENT: RXR-BLA\_TOX21\_SLP\_Version1.0 TITLE: Protocol of RXR-BLA HEK 293T Cell-based Assay for High-throughput Screening

# ASSAY RFERENCES:

Assay Target	Cell Lines	Species	Tissue of Origin	Assay Readout	Assay Provider	Toxicity Pathway
Retinoid X Receptor alpha (Recombinant)	НЕК 293Т	Human	Embryonic kidney cells	Beta- lactamase reporter	Invitrogen	NR signaling

# **QUALITY CONTROL PRECAUTIONS:**

1. -Cell culture is maintained by passaging twice a week and should not reach more than 90% confluence

2. -The assay should be performed in black-clear bottom 1536 well plates, so the bottom of the plates should not be touched

#### **MATERIALS and INSTRUMENTS:**

Supplies/Medium/Reagent	Manufacturer	Vender/Catalog Number	
-Phenol red-free DMEM	-Invitrogen	-Invitrogen/21063	
-DMEM	-Invitrogen	-Invitrogen/10569	
-Dialyzed FBS	-Invitrogen	-Invitrogen/26400	
-Charcoal stripped FBS	-Invitrogen -Invitrogen/12676		
-NEAA	-Invitrogen	-Invitrogen/11140	
-Sodium pyruvate	-Invitrogen	-Invitrogen/11360	
-HEPES	-Invitrogen	-Invitrogen/15630	
-Penn-strep	-Invitrogen	-Invitrogen/15140	
-Hygromycin B	-Invitrogen	-Invitrogen/10687	
-Zeocin	-Invitrogen	-Invitrogen/R25001	
-Recovery Cell Culture Freezing Medium	-Invitrogen	-Invitrogen/12648	
-0.05% Trypsin-EDTA	-Invitrogen -Invitrogen/25300		

-Black-clear bottom 1536 well plates	-Greiner	-Greiner/789092F	
-9-cis-Retinoic acid (Agonist control compound)	-Enzo Life Sciences	-Enzo Life Sciences/BML- GR101-0005	
-Multidrop COMBI	-Thermo Electron Corporation -Thermo Electron Corpo		
-BioRAPTR FRD dispenser	-Beckman Coulter	-Beckman Coulter	
-LiveBLAzer B/G FRET substrate	-Invitrogen	-Invitrogen/K1028	
-CellTiter-Glo(R) One Solution Assay	-Promega	-Promega/G8462	
-Envision Plate Reader	-Perkin Elmer	-Perkin Elmer	
-ViewLux Plate Reader	-Perkin Elmer	-Perkin Elmer	

### **PROCEDURE:**

- 1. Cell handling:
  - 1.1. Media Required:

Component	Growth Medium	Assay Medium	Thaw Medium	Freezing Medium
-Phenol red-free DMEM	-	-98%	-	-
-DMEM	-90%	-	-90%	-
-Dialyzed FBS	-10%	-	-10%	-
-Charcoal stripped FBS	-	-2%	-	-
-NEAA	-0.1mM	-0.1mM	-0.1mM	-
-Sodium pyruvate	-1mM	-1mM	-1mM	-
-HEPES	-25mM	-	-25mM	-
-Penn-strep	-100U/ml- 100ug/ml	-100U/ml- 100ug/ml	-100U/ml- 100ug/ml	-
-Hygromycin B	-100ug/ml	-	-	-
-Zeocin	-100ug/ml	-	-	-
-Recovery Cell Culture Freezing Medium	-	-	-	-100%

#### 1.2. Thawing method

1.2.1 -1ml frozen cells of RXRalpha-bla were taken in pre-warmed 10ml of thaw medium for centrifuging

1.2.2 -2-3ml of the thaw medium is taken to resuspend the pellet 1.2.3 -The cells were seeded in T-75 flask at 2 million cells.

#### 1.3. Propagation method

1.3.1 -The cells are detached using 0.05% Trypsin

1.3.2 -The cells are further passaged at a density of 4-5 million cells per T-225 flask

2. Assay Protocol

2.1 -Rinse the cells with DPBS and detach them by using 0.05% Trypsin and centrifuge

2.2 -Resuspend the pellet with assay medium

2.3 -Plate the cells in black-clear bottom 1536 well plate at 2000/well/6uL through 8 tip of a plate dispenser (Multi drop)

2.4 -Incubate at 37C for 5hrs

2.5 -Transfer 23nL of the compounds from the library collection and positive control through Pintool

2.6 -Incubate at 37C for 16hrs

2.7 -Add 1uL of CCF4 dye using a single tip of a plate dispenser (Bioraptr)

2.8 -Incubate at room temperature for 2hrs

2.9 -Read the fluorescence intensity through Envision plate reader

2.10 -Then add 4uL of CellTiter-Glo reagent using a single tip of a plate dispenser (Bioraptr)

2.11 - Incubate at room temperature for 30 min

2.12 -Read the luminescence intensity through ViewLux plate reader

3. Assay Performance

RXRα-bla (9-cis Retinoic acid; Agonist control)	Online Validation Agonist (Mean ± SD)	
EC50	8.40 ± 2.30 nM (n = 27)	
S/B	2.19 ± 0.09	
CV (%)	5.03 ± 0.31* (n = 18)	
Z'	0.49 ± 0.07	

\*CV values shown represent average of all plates excluding top 3 compound concentration plates.