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# **Product Sheet**

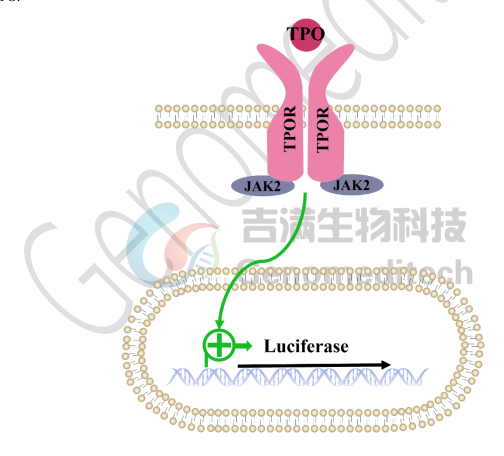
## **H\_TPO Reporter 293 Cell Line**

Catalog number: GM-C30062

Version 3.3.1.241205

TPO (Thyroid Peroxidase) is an important cytokine composed of multiple structural domains, including a catalytic active site and several glycosylation sites. It is primarily synthesized in the liver and is responsible for regulating the production and function of megakaryocytes. Upon binding to its receptor (TPO-R; also known as c-Mpl, CD110), it initiates downstream signaling. TPO also promotes the proliferation and differentiation of megakaryocytes and their precursor cells, directly influencing platelet production.

H\_TPO Reporter 293 Cell Line is a clonal stable cell line constructed using lentiviral technology, constitutive expression of the TPOR gene, along with signal-dependent expression of a luciferase reporter gene. When TPO binds to TPOR, it activates downstream signaling pathways, leading to the expression of luciferase. The luciferase activity measurement indicates the activation level of the signaling pathway and can thus be used to evaluate the in vitro effects of drugs related to TPO.





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#### **Specifications**

**Quantity** 5E6 Cells per vial,1 mL

**Product Format** 1 vial of frozen cells

**Shipping** Shipped on dry ice

Storage Conditions Liquid nitrogen immediately upon receip

**Recovery Medium** DMEM+10% FBS+1% P.S

Growth medium DMEM+10% FBS+1% P.S+4  $\mu$ g/mL Blasticidin+125  $\mu$ g/mL Hygromycin+0.75  $\mu$ g/mL

Puromycin

Note None

Freezing Medium 90% FBS+10% DMSO

**Growth properties** Adherent

**Growth Conditions** 37°C, 5% CO<sub>2</sub>

**Mycoplasma Testing** The cell line has been screened to confirm the absence of Mycoplasma species.

**Safety considerations** Biosafety Level 2

Note It is recommended to expand the cell culture and store a minimum of 10 vials at an early

passage for potential future use.

#### **Materials**

Reagent	Manufacturer/Catalogue No.
DMEM	Gibco/C11995500BT
Fetal Bovine Serum	Cegrogen biotech/A0500-3010
Pen/Strep	Thermo/15140-122
Blasticidin	Genomeditech/GM-040404
Hygromycin	Genomeditech/GM-040403
Puromycin	Genomeditech/GM-040401
Human TPO Protein; His Tag	Genomeditech/GM-87861RP
GMOne-Step Luciferase Reporter Gene Assay Kit	Genomeditech/GM-040503

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### **Figures**

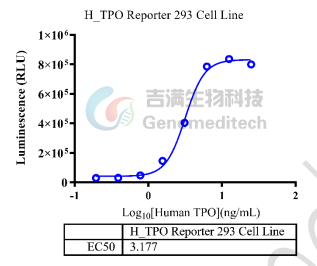


Figure 1 | Response to Recombinant Human TPO; His Tag. The H\_TPO Reporter 293 Cell Line (Cat. GM-C30062) at a concentration of 1.5E4 cells/well (96-well format) was stimulated with serial dilutions of Recombinant Human TPO (Cat. GM-87861RP) in assay buffer (DMEM+1% FBS+1% P.S) for 6 hours. The firefly luciferase activity was measured using the GMOne-Step Luciferase Reporter Gene Assay Kit (Cat. GM-040503). The maximum induction fold was approximately [30.2]. Data are shown by drug mass concentration.

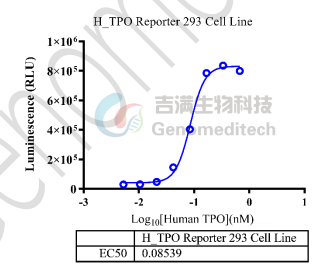


Figure 2 | Response to Recombinant Human TPO; His Tag. The H\_TPO Reporter 293 Cell Line (Cat. GM-C30062) at a concentration of 1.5E4 cells/well (96-well format) was stimulated with serial dilutions of Recombinant Human TPO (Cat. GM-87861RP) in assay buffer (DMEM+1% FBS+1% P.S) for 6 hours. The firefly luciferase activity was measured using the GMOne-Step Luciferase Reporter Gene Assay Kit (Cat. GM-040503). The maximum induction fold was approximately [30.2]. Data are shown by drug molar concentration.

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**Cell Recovery** 

Recovery Medium: DMEM+10% FBS+1% P.S

To insure the highest level of viability, thaw the vial and initiate the culture as soon as possible upon receipt. If upon arrival, continued storage of the frozen culture is necessary, it should be stored in liquid nitrogen vapor phase and not at -70°C.

Storage at -70°C will result in loss of viability.

Thaw the vial by gentle agitation in a 37°C water bath. To reduce the possibility of contamination, keep the O-ring

and cap out of the water. Thawing should be rapid (approximately 2 - 3 minutes).

b) Remove the vial from the water bath as soon as the contents are thawed, and decontaminate by dipping in or spraying

with 70% ethanol. All of the operations from this point on should be carried out under strict aseptic conditions.

c) Transfer the vial contents to a centrifuge tube containing 5.0 mL complete culture medium and spin at approximately

176 x g for 5 minutes. Discard supernatant.

d) Resuspend cell pellet with the recommended recovery medium. And dispense into appropriate culture dishes.

Incubate the culture at 37°C in a suitable incubator. A 5% CO<sub>2</sub> in air atmosphere is recommended if using the medium

described on this product sheet.

Cell Freezing

Freezing Medium: 90% FBS+10% DMSO

Centrifuge at 176 x g for 3 minutes to collect cells.

Resuspend the cells in pre-cooled freezing medium and adjust the cell density to 5E6 cells/mL. b)

Aliquot 1 mL into each vial. c)

Place the vial in a controlled-rate freezing container and store at -80°C for at least 1 day, then transfer to liquid d)

nitrogen as soon as possible.

Cell passage

Growth medium: DMEM+10% FBS+1% P.S+4 µg/mL Blasticidin+125 µg/mL Hygromycin+0.75 µg/mL Puromycin

For the first 1 to 2 passages post-resuscitation, use the recovery medium. Once the cells have stabilized, switch to a growth

medium.

Subculturing is necessary when the cell density reaches 80%. It is recommended to perform subculturing at a ratio of

1:3 to 1:4 every 2-3 days. Ensure that the density does not exceed 80%, as overcrowding can lead to reduced viability

due to compression.

Remove and discard culture medium. b)

Briefly rinse the cell layer with PBS to remove all traces of serum that contains trypsin inhibitor. c)

d) Add 1.0 mL of 0.25% (w/v) Trypsin-EDTA solution to dish and observe cells under an inverted microscope until cell

layer is dispersed (usually within 30 to 60 seconds at 37°C).

Note: To avoid clumping do not agitate the cells by hitting or shaking the flask while waiting for the cells to detach.

Cells that are difficult to detach may be placed at 37°C to facilitate dispersal.

f) Add 2.0 mL of growth medium to mix well and aspirate cells by gently pipetting.



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g) After centrifugation, resuspend the pellet and add appropriate aliquots of the cell suspension to new culture vessels.

h) Incubate cultures at 37°C.

Subcultivation Ratio: A subcultivation ratio of 1:3 - 1:4 is recommended

Medium Renewal: Every 2 to 3 days

**Notes** 

a) Upon initial thawing, a higher number of dead cells is observed, which is a normal phenomenon. Significant

improvement is seen after adaptation. Once the cells reach a stable state, the number of dead cells decreases after

subculturing and the cell growth rate becomes stable.

b) Ensure that the cell density does not exceed 80%, as overcrowding may lead to reduced viability due to compression.

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