

# **Product Sheet**

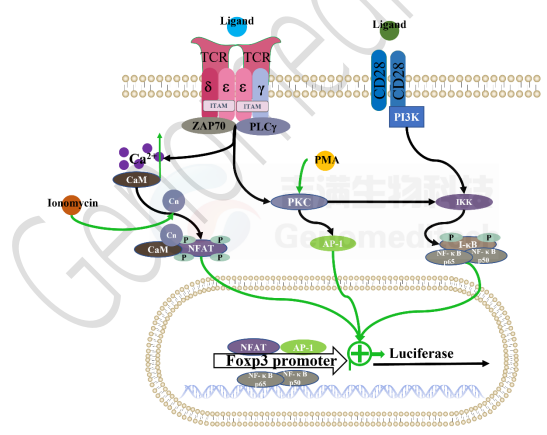
# H\_FOXP3-Promoter Reporter Jurkat Cell Line

Catalog number: GM-C13108

Version 3.3.1.241226

FOXP3 belongs to the forkhead family and is a major transcription factor. Studies have shown that overexpressed FOXP3 in Jurkat cells can compete with NFAT for binding or directly interact with NFAT or NF-κB as a transcriptional repressor of the IL-2 promoter. In humans, the expression of FOXP3 is upregulated in all activated T cells. FOXP3 is essential for the development and function of naturally occurring Treg cells and is involved in the molecular mechanisms that control immune tolerance.

H\_FOXP3-Promoter Reporter Jurkat Cell Line is a clonal stable cell line constructed using lentiviral technology that expresses a FOXP3-Promoter inducible luciferase reporter gene. When PMA and Ionomycin activates downstream signaling pathways, it leads to the expression of luciferase driven by the FOXP3-Promoter. 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 FOXP3-Promoter.





# Specifications

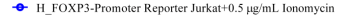
Quantity	5E6 Cells per vial,1 mL		
Product Format	1 vial of frozen cells		
Shipping	Shipped on dry ice		
Storage Conditions	Liquid nitrogen immediately upon receipt		
Recovery Medium	RPMI 1640+10% FBS+1% P.S		
Growth medium	RPMI 1640+10% FBS+1% P.S+0.75 μg/mL Puromycin		
Note	None		
Freezing Medium	90% FBS+10% DMSO		
Growth properties	Suspension		
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.
RPMI 1640	gibco/C11875500BT
Fetal Bovine Serum	Cegrogen biotech/A0500-3010
Pen/Strep	Thermo/15140-122
Puromycin	Genomeditech/GM-040401
РМА/ТРА	Beyotime/S1819
Ionomycin	MCE/HY-13434
GMOne-Step Luciferase Reporter Gene Assay Kit	Genomeditech/GM-040503



### **Figures**



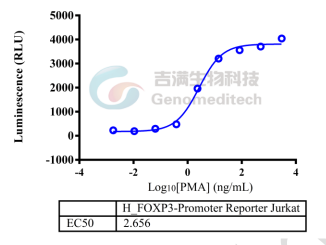


Figure 1 | Response to PMA/TPA+Ionomycin. The H\_FOXP3-Promoter Reporter Jurkat Cell Line (Cat. GM-C13108) at a concentration of 1E5 cells/well (96-well format) was stimulated with serial dilutions of PMA/TPA (Beyotime/S1819) and 0.5 μg/mL Ionomycin (MCE/HY-13434) in assay buffer (RPMI 1640 + 1% FBS + 1% P.S) for 7 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 [16.9]. Data are shown by drug mass concentration.

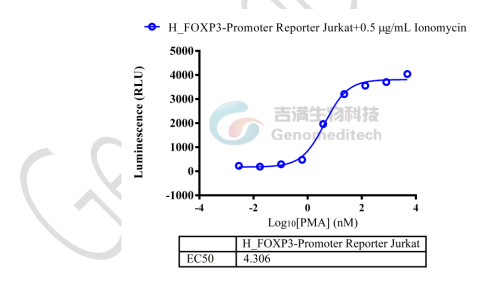


Figure 2 | Response to PMA/TPA+Ionomycin. The H\_FOXP3-Promoter Reporter Jurkat Cell Line (Cat. GM-C13108) at a concentration of 1E5 cells/well (96-well format) was stimulated with serial dilutions of PMA/TPA (Beyotime/S1819) and 0.5 µg/mL Ionomycin (MCE/HY-13434) in assay buffer (RPMI 1640 + 1% FBS + 1% P.S) for 7 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 [16.9]. Data are shown by drug molar concentration.

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## **Cell Recovery**

Recovery Medium: RPMI 1640+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  $-70^{\circ}$ C. Storage at  $-70^{\circ}$ C will result in loss of viability.

- a) 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 complete medium. And dispense the suspension into 1 2 T-25 culture flasks.
- e) 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

- a) Centrifuge at 176 x g for 3 minutes to collect cells.
- b) Resuspend the cells in pre-cooled freezing medium and adjust the cell density to 5E6 cells/mL.
- c) Aliquot 1 mL into each vial.
- d) Place the vial in a controlled-rate freezing container and store at -80°C for at least 1 day, then transfer to liquid nitrogen as soon as possible.

# Cell passage

Growth medium: RPMI 1640+10% FBS+1% P.S+0.75 µg/mL Puromycin

Approximately 48-72 hours after the initial thawing, the cells can be passaged for the first time. After this initial passage, the culture medium can be adjusted to growth medium supplemented with antibiotics. If cells are not passaged within 48 hours, it is recommended to add some fresh recovery medium and place the flask horizontally.

- a) When the cell density reaches 1.5 2E6 cells/mL, subculture the cells. Do not allow the cell density to exceed 2E6 cells/mL.
- b) It is recommended to use T-25 flasks for subculturing.
- c) These cells are suspension cells, and it is recommended to use the "half-medium change" method to maintain optimal cell conditions during passaging.
- d) During passaging, you can directly add fresh growth medium to the culture flask, gently pipette to resuspend the cells, and then transfer the cell suspension to a new T-25 flask for continued culture.

#### Subcultivation Ratio: Maintain cultures at a cell concentraion between 3E5 and 1E6 viable cells/mL.

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#### Medium Renewal: Every 2 to 3 days

#### Notes

- a) These cells are sensitive to density, so please ensure that the cell density is maintained within an appropriate range during culture and subculturing.
- b) During the first passage, pay attention to the nutrient supply; if not subculturing, make sure to add fresh recovery medium every other day as needed.

#### **Related Products**

TCR			
H_IL2-Promoter Reporter Jurkat Cell Line	NFAT-Luc Reporter Jurkat Cell Line		
TCR Knockout Reporter Cell Line(CD4+)	OKT3(CD3 ScFv) CHO-K1 Cell Line		
Anti-CD3-CD19 Bispecific Antibody(Blinatumomab)			

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