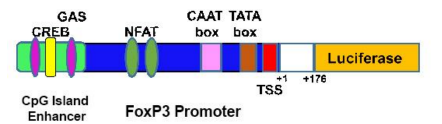


# Data Sheet Foxp3 Reporter (Luc) - Jurkat Recombinant Cell Line Catalog # 60628

# Description

Human Foxp3 luciferase reporter construct is stably integrated into the genome of Jurkat T-cells. The firefly luciferase gene is controlled by a human Foxp3 promoter and an enhancer-like conserved noncoding sequence upstream of the Foxp3 promoter.



**Figure 1.** Illustration of Foxp3 promoter region with representative transcription factor binding sites and enhancer regions.

### Background

Foxp3, belonging to the forkhead family, is a master transcription factor that expresses exclusively in regulatory T cells, a subset of CD4+ T cells. Regulation of Foxp3 is critical for maintaining immunological tolerance. Over-expression of Foxp3 is known to suppress effector T cell activation.

### Host Cell

Jurkat (Human Acute T-Cell Leukemic) Cell Line, Clone E61. Suspension cells.

# Format

Each vial contains  $\sim 2 \times 10^6$  cells in 1 ml of 10% DMSO in FBS.

### Storage

Store in liquid nitrogen immediately upon receipt.

### Culture Media

**Thaw Medium 2 (BPS Cat. #60184):** RPMI 1640 medium (Thermo Fisher, Cat. #A1049101) supplemented with 10% FBS (Thermo Fisher, Cat. #26140079), 1% Penicillin/Streptomycin (Hyclone #SV30010.01).

**Growth Medium 2B (BPS Cat. #79530):** Thaw Medium 2 (BPS Cat. #60184) and 1 mg/ml G418 (Thermo Fisher, Cat. #11811031).



### **Recommended Culture Conditions**

*Frozen Cells*: Prepare a 50 ml conical tube and a T-25 culture flask with 5 ml of pre-warmed Thaw Medium 2 (**no G418**). Quickly thaw cells in a 37°C water bath with constant and slow agitation. Clean the outside of the vial with 70% ethanol and immediately transfer the entire contents to the conical tube with Thaw Medium 2 (**no G418**) and rock the tube the tube gently. Centrifuge the cells at 200 x g for 3 minutes. Re-suspend the cells in 6 ml of pre-warmed Thaw Medium 2 (**no G418**) and transfer the entire content to the T25 culture flask containing Thaw Medium 2 (**no G418**). Avoid pipetting up and down, and gently rock the flask to distribute the cells. Incubate the cells in a humidified 37°C incubator with 5% CO<sub>2</sub>. Forty-eight hours after incubation, centrifuge cells at 250 x g for 5 minutes and re-suspend to fresh Thaw Medium 2 (**no G418**). Continue to monitor growth for 2-3 days and change medium to remove dead debris. If slow cell growth occurs during resuscitation, increase FBS to 15% for the first week of culture. Switch to Growth Medium 2B (**containing G418**) after multiple cell colonies (in clumps) start to appear (indicative of healthy cell division). We recommend passing cells for 3 passages after thawing before using them in the luciferase assay.

*Subculture*: When cells reached 80% confluency, transfer cells to a 50 ml conical tube and centrifuge cells at 200 x g for 5 minutes. Wash cells once with PBS (without Magnesium or Calcium) and re-suspend cells in 10 ml pre-warmed Growth Medium 2B (**contains G418**); gently pipette up and down to dissociate cell clumps. <u>*This cell line is "clumpy" (indicates healthy growth)*. It is highly advised to dissociate the clumps by gentle pipetting during passaging or seeding for assays. Dispense 2-4 ml of the cell suspension into a new T-75 flask containing pre-warmed 15 ml Growth Medium 2B. Incubate cells in a humidified 37°C incubator with 5% CO<sub>2</sub>. Freeze cells in freezing medium (10% DMSO in FBS) when cells reach 80% confluency. Cells have been demonstrated to be stable for at least 15 passages; BPS Bioscience recommends preparing frozen stocks so cells are not used beyond passage 20.</u>

### Mycoplasma Testing

This cell line has been screened using the MycoAlert<sup>™</sup> Mycoplasma Detection Kit (Cat. #LT07-118) to confirm the absence of Mycoplasma contamination. MycoAlert Assay Control Set (Cat. #LT07-518) was used as a positive control.

# Application

The Foxp3 promoter Jurkat reporter cell line is suitable for monitoring the transcription activity of Foxp3 in response to stimulant, and establishing cell-based screens for inhibitors that target specific Foxp3-stimulating molecules. This reporter cell line has been tested and validated using phorbol 12-myristate 13-acetate (PMA) with ionomycin (**Figure 2**). BPS Bioscience does not recommend starving the cells overnight in serum-free medium prior to stimulation.

# Materials Required but Not Supplied for Cell Culture

- Thaw Medium 2 (BPS Bioscience #60184)
- Growth Medium 2B (BPS Bioscience #79530)



# Materials Required but Not Supplied for Cellular Assay

- Assay Medium 2B (BPS Bioscience #79619)
- PMA (Fisher, Cat # BP685-1)
- Ionomycin (Fisher, Cat # BP25271)
- ONE-Step<sup>™</sup> Luciferase Assay System (BPS Bioscience #60690)
- Luminometer
- 96-well tissue culture-treated white clear-bottom assay plate

# **Application References**

1. Mantel, P.Y., *et al.* (2006) Molecular Mechanisms Underlying FOXP3 induction in Human T cells. *J. Immunology* **176(6)**:3593-602.

# Assay Protocol

 In a white opaque 96- well plate, seed cells at ~1 x10<sup>4</sup> cells/well (100 μl per well) in Assay Medium 2B (BPS #79619)

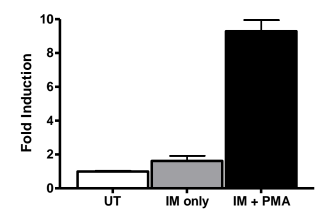
**Assay Medium 2B (BPS #79619):** RPMI 1640 medium (Thermo Fisher, Cat. #A1049101) supplemented with 1% Penicillin/Streptomycin (Hyclone #SV30010.01).

Cells should be growing at log phase at time of seeding. Using cells from over-confluent culture (ie. stationary growth) will significantly reduce signal output.

- Prepare fresh working solution of PMA (Fisher, Cat # BP685-1) at 100 ng/ml (from 100 μg/ml stock in DMSO) and ionomycin (Fisher, Cat # BP25271) at 1.67 μg/ml (from 1 mg/ml stock in DMSO) in PBS.
- Immediately treat cells with 10 μl of working solution of PMA (ie. 10 ng/ml, final concentration) and ionomycin (167 ng/ml, final concentration) for 24 hours at 37°C with 5% CO<sub>2</sub> (Figure 2).For maximum signal, the final concentration should be <u>167 ng/ml</u> ionomycin and <u>0.15-30 ng/ml</u> of PMA in 100 μL of medium (Figure 3).
- 4. Add ONE-Step<sup>™</sup> Luciferase Assay System (BPS Bioscience, Cat. #60690) to each well, according to recommended protocol.
- 5. Read luminescence using a luminometer. Normalize luminescence to wells that contain only medium to obtain the Relative Luminescence Units (RLUs).

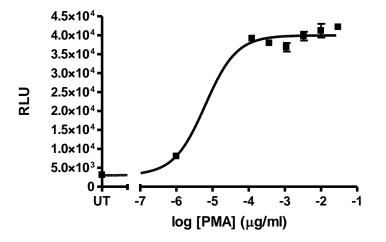


### **Quality Assurance and Functional Analysis**

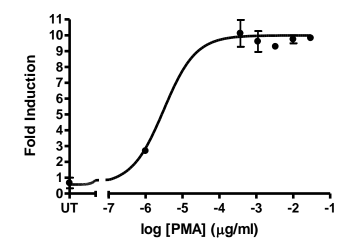


### Figure 2. Foxp3 Jurkat reporter responses to a combination of PMA and ionomycin.

Foxp3 Jurkat reporter cells were seeded on a white opaque 96- well plate at approximately 1 x104 cells/well (100  $\mu$ l per well) in serum free RPMI medium. Cells were treated with 167 ng/ml ionomycin (IM only), with 167 ng/ml IM and 10 ng/ml PMA (IM+PMA) or untreated (UT) at 37°C with 5% CO2. Fold induction = relative luminescence normalized to untreated cells; n=4.

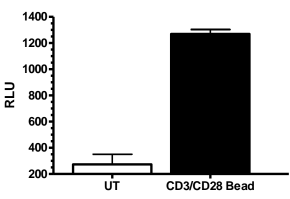


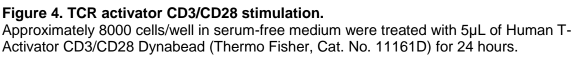




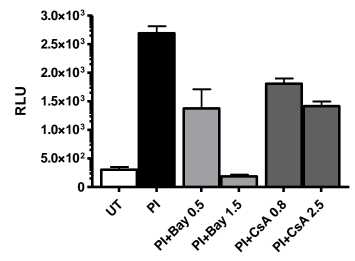
### Figure 3. Foxp3 Jurkat reporter activity in response to PMA and ionomycin.

Approximately 1 x10<sup>4</sup> cells/well (100  $\mu$ l per well) in serum free RPMI medium were untreated (UT) or treated with 167ng/ml of ionomycin with 0- 30 ng/ml of PMA for 24h. (Top) Clone 1D8; RLU = relative luminescence normalized to RPMI. (Bottom) Clone 1C9; Fold induction = RLU of treated cells with respect to untreated cells; n=3. Error bar = standard deviation (SD).









# Figure 5. Inhibition of Foxp3 reporter by BAY11-7082 and Cyclosporin A.

Approximately 8000 cells/ well in SF- medium were left untreated (UT), treated with 30ng/ml of PMA and 167ng/ml Ionomycin (PI), PI with 0.5 or 1.5  $\mu$ M Bay 11-7082 (Bay; Sigma Cat. No. B5556) (PI + Bay), or PI with 0.8 or 2.5  $\mu$ M Cyclosporin A (CsA; Sigma Cat. No. 30024) for 24 hours. n =3. RLU = Relative Luminescence Unit.

# **References:**

- 1. Tone Y et.al. (2008) Nat Immunology. 9: 194-202
- 2. Liu R et.al. (2015) Cancer Res. 75: 1703-1713
- 3. Soligo M. et al. (2011) Eur J. Immunology. 41: 503-513



### Vector

Human Foxp3 promoter-Luciferase was cloned into the MCS of pCDNA3.1<sup>™</sup> (+) vector (Invitrogen, Cat. #V79020).

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### **Related Products**

Product	<u>Cat. #</u>	<u>Size</u>
ONE-Step <sup>™</sup> Luciferase Assay System	60690-1	10 ml
ONE-Step <sup>™</sup> Luciferase Assay System	60690-2	100 ml
ONE-Step <sup>™</sup> Luciferase Assay System	60690-3	1 L
Thaw Medium 2	60184	100 ml
Growth Medium 2B	79530	500 ml
Assay Medium 2B	79619	100 ml