

## Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD32a-High Expression)

Catalog Number: CK-003

Components	Catalog No.	Size
Human 4-1BB (Luc) HEK293 Reporter Cell	CHEK-ATF073	$2 \times (1 \text{ vial contains} \sim 5 \times 10^{6} \text{ cells})$
CHO/Human CD32a Stable Cell Line (High Expression)	SCCHO-ATP061H	$2 \times (1 \text{ vial contains} \sim 5 \times 10^{6} \text{ cells})$

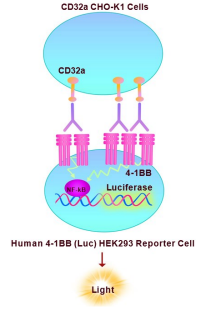
#### • Description

The Cell-based Kit consists of two engineered cell lines, Human 4-1BB (Luc) HEK293 Reporter Cell (Cat.No.CHEK-ATF073) and CHO/Human CD32a Stable Cell Line (High Expression) (Cat.No.SCCHO-ATP061H).

The Human 4-1BB (Luc) HEK293 Reporter Cell was engineered to not only express NF-κB signaling response element, but also express the receptor full length human 4-1BB (Gene ID: 3604), which can drive luciferase expressing systems by 4-1BB ligand/ agonist antibody stimulation. The CHO/Human CD32a Stable Cell Line was engineered to express full length human CD32a receptor (Gene ID: 2212), with different levels of CD32a expression (High, Medium, Low), which can be used to test agonist antibody whether in a CD32a-dependent manner to strengthen the agonistic activity. When co-cultured with Human 4-1BB (Luc) HEK293 Reporter Cell and anti-4-1BB agonist antibody, the anti-4-1BB antibody can be crosslinked, thereby strengthening 4-1BB pathway-activated luminescence.

### • Application

• Screen for Anti-human 4-1BB antibodies whether in a CD32a-dependent manner to strengthen the agonistic activity





# • Cell Line Profile of Human 4-1BB (Luc) HEK293 Reporter Cell

Cell line	Human 4-1BB (Luc) HEK293 Reporter Cell
Host Cell	HEK293
Property	Adherent
Complete Growth Medium	DMEM + 10% FBS
Selection Marker	Puromycin (2 μg/mL) + Hygromycin (20 μg/mL)
Incubation	37°C with 5% CO <sub>2</sub>
Doubling Time	22-24 hours
Transduction Technique	Lentivirus

# • Cell Line Profile of CHO/Human CD32a Stable Cell Line (High Expression)

Cell line	CHO/Human CD32a Stable Cell Line (High Expression)	
Host Cell	СНО	
Property	Adherent	
Complete Growth Medium	F-12K + 10% FBS	
Selection Marker	Hygromycin (20 μg/mL)	
Incubation	37°C with 5% CO <sub>2</sub>	
Doubling Time	22-24 hours	
Transduction Technique	Lentivirus	



## • Cell Culture of Human 4-1BB (Luc) HEK293 Reporter Cell

### Materials Required

- DMEM medium (Gibco, Cat.No.11965-092)
- Fetal bovine serum (CellMax, Cat.No.SA211.02)
- Puromycin (InvivoGen, Cat.No.ant-pr-5b)
- Hygromycin B (Invitrogen, Cat.No.10687010)
- Complete Growth Medium: DMEM + 10% FBS
- Culture Medium: DMEM + 10% FBS, Hygromycin (20 μg/mL), Puromycin (2 μg/mL)
- Freeze Medium: 90% FBS, 10% (V/V) DMSO
- T-75 Culture flask (Corning, 430641)
- Cryogenic storage vials (SARSTEDT, 72.379.007)
- Thermostat water bath
- Centrifuge
- Luna cell counter (Logos Biosystems, LUNA-II)
- CO<sub>2</sub> Incubator (Thermo, 3111)
- Biological Safety Cabinet (Thermo, 1389)

#### Recovery

- 1. Thaw the vial by gentle agitation in a 37°C water bath. To reduce the possibility of contamination, keep the cap out of the water. Thawing should be rapid (approximately 2 minutes).
- 2. Remove the vial from the water bath as soon as the contents are thawed, and decontaminate by spraying with 70% ethanol. All the operations from this point on should be carried out under strict aseptic conditions.
- 3. Transfer the vial contents to a centrifuge tube containing 4.0 mL complete growth medium and spin at approximately 1000 rpm for 5 minutes.
- 4. Resuspend cell pellet with 5 mL complete growth medium and transfer the cell suspension into T-75 flask containing 10-15 mL of pre-warmed complete growth medium.
- 5. Incubate at 37°C with 5% CO<sub>2</sub> incubator until the cells are ready to be split.



#### Subculture

- 1. Remove and discard culture medium.
- 2. Wash the cells once with sterile PBS.
- 3. Add 3 mL of 0.25% trypsin to cell culture flask. Place the flask at 37°C for 5-7 minutes, until 90% of the cells have detached.
- 4. Add 6.0 to 8.0 mL of culture medium and aspirate cells by gently pipetting.
- 5. Add appropriate aliquots of the cell suspension to new culture vessel.
- 6. Incubate at 37 ℃ with 5% CO<sub>2</sub> incubator.

**Subcultivation Ratio:** A subcultivation ratio of 1:6 to 1:10 is recommended.

Medium Renewal: Every 2 to 3 days.

### Cryopreservation

- 1. Remove and discard spent medium.
- 2. Detach cells from the cell culture flasks with 0.25% trypsin.
- 3. Centrifuge at 1000 rpm for 5 min at RT to pellet cells.
- 4. Resuspend the cell pellets with culture medium and count viable cells.
- Centrifuge at 1000 rpm for 5 min at RT and resuspend cells in freezing medium to a concentration of 5×10<sup>6</sup> to 1×10<sup>7</sup> cells/mL.
- 6. Aliquot into cryogenic storage vials. Place vials in a programmable cooler or an insulated box placed in a -80°C freezer overnight, then transferring to liquid nitrogen storage.

#### Storage

- **Product format:** Frozen
- Storage conditions: Liquid nitrogen immediately upon receipt



# • Cell Culture of CHO/Human CD32a Stable Cell Line (High Expression)

### Materials Required

- F-12K Nutrient Mixture (Gibco, Cat.No.21127-022)
- Fetal bovine serum (CellMax, Cat.No.SA211.02)
- Hygromycin B (Invitrogen, Cat.No.10687010)
- Complete Growth Medium: F-12K + 10% FBS
- Culture Medium: F-12K + 10% FBS, Hygromycin (20 µg/mL)
- Freeze Medium: 90% FBS, 10% (V/V) DMSO
- T-75 Culture flask (Corning, 430641)
- Cryogenic storage vials (SARSTEDT, 72.379.007)
- Thermostat water bath
- Centrifuge
- Luna cell counter (Logos Biosystems, LUNA-II)
- CO<sub>2</sub> Incubator (Thermo, 3111)
- Biological Safety Cabinet (Thermo, 1389)

#### Recovery

- 1. Thaw the vial by gentle agitation in a 37°C water bath. To reduce the possibility of contamination, keep the cap out of the water. Thawing should be rapid (approximately 2 minutes).
- 2. Remove the vial from the water bath as soon as the contents are thawed, and decontaminate by spraying with 70% ethanol. All the operations from this point on should be carried out under strict aseptic conditions.
- 3. Transfer the vial contents to a centrifuge tube containing 4.0 mL complete growth medium and spin at approximately 1000 rpm for 5 minutes.
- 4. Resuspend cell pellet with 5 mL complete growth medium and transfer the cell suspension into T-75 flask containing 10-15 mL of pre-warmed complete growth medium.
- 5. Incubate at 37°C with 5% CO<sub>2</sub> incubator until the cells are ready to be split.



#### Subculture

- 1. Remove and discard culture medium.
- 2. Wash the cells once with sterile PBS.
- 3. Add 3 mL of 0.25% trypsin to cell culture flask. Place the flask at 37°C for 5-7 minutes, until 90% of the cells have detached.
- 4. Add 6.0 to 8.0 mL of culture medium and aspirate cells by gently pipetting.
- 5. Add appropriate aliquots of the cell suspension to new culture vessel.
- 6. Incubate at 37 ℃ with 5% CO<sub>2</sub> incubator.

**Subcultivation Ratio:** A subcultivation ratio of 1:6 to 1:10 is recommended.

**Medium Renewal:** Every 2 to 3 days.

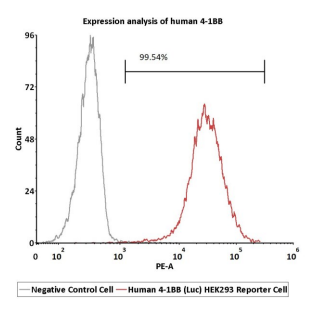
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#### Storage

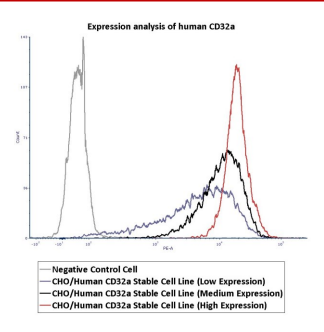
- Product format: Frozen
- Storage conditions: Liquid nitrogen immediately upon receipt

# • Receptor Assay



**Fig1.** Expression analysis of human 4-1BB on Human 4-1BB (Luc) HEK293 Reporter Cell by FACS. Cell surface staining was performed on Human 4-1BB (Luc) HEK293 Reporter Cell or negative control cell using PElabeled anti-human 4-1BB antibody.





Catalog No.	Stable Cell Line	MFI for CD32a (PE)
SCCHO-ATP061L	CHO/Human CD32a Stable Cell Line (Low Expression)	5196.88
SCCHO-ATP061M	CHO/Human CD32a Stable Cell Line (Medium Expression)	11336.11
SCCHO-ATP061H	CHO/Human CD32a Stable Cell Line (High Expression)	18309.90

**Fig2. Expression analysis of human CD32a on CHO/Human CD32a Stable Cell Line by FACS.** Cell surface staining using PE-labeled anti-human CD32a antibody was performed on CHO/Human CD32a Stable Cell Line with different expression levels: CHO/Human CD32a Stable Cell Line (Low Expression); CHO/Human CD32a Stable Cell Line (Medium Expression); CHO/Human CD32a Stable Cell Line (High Expression).

### • License Disclosure

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# • Related Products

<u>Products</u>	Cat.No.
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD32a-Low Expression)	CK-001
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD32a- Medium Expression)	CK-002
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD16a (158V)-Low Expression)	CK-004
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD16a (158V)-Medium Expression)	CK-005
Cell-based Screening Kit for Anti-human 4-1BB Antibody (HumanCD16a (158V)-High Expression)	CK-006
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD32b-Low Expression)	CK-007
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD32b-Medium Expression)	CK-008
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD32b-High Expression)	CK-009
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD64-Low Expression)	CK-010
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD64-Medium Expression)	CK-011
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human CD64-High Expression)	CK-012
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human PD-L1-Low Expression)	CK-013
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human PD-L1-Medium Expression)	CK-014
Cell-based Screening Kit for Anti-human 4-1BB Antibody (Human PD-L1-High Expression)	CK-015