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## 14-520ACL: TIM-3 Stable Cell Line

**Application:** Functional Assay

# **Description**

TIM-3 Stable Cell Line is a stably transfected CHO-K1 cell line which expresses human T-cell immunoglobulin and mucindomain containing-3 (TIM-3, also known as HAVCR2 and CD366).

Sequence data: hTIM-3 (accession number NP 116171)

MFSHLPFDCVLLLLLLLTRSSEVEYRAEVGQNAYLPCFYTPAA
PGNLVPVCWGKGACPVFECGNVVLRTDERDVNYWTSRYWLNGDFRKGDVSLTIENVTL
ADSGIYCCRIQIPGIMNDEKFNLKLVIKPAKVTPAPTRQRDFTAAFPRMLTTRGHGPA
ETQTLGSLPDINLTQISTLANELRDSRLANDLRDSGATIRIGIYIGAGICAGLALALI
FGALIFKWYSHSKEKIQNLSLISLANLPPSGLANAVAEGIRSEENIYTIEENVYEVEE
PNEYYCYVSSRQQPSQPLGCRFAMP

### **Product Info**

Amount: 1 Vial

**Content:** Each vial contains 2 ~ 3 x 10^6 cells in 1 ml of 90% FBS + 10% DMSO

**Storage condition :** Immediately upon receipt, store in liquid nitrogen.

## **Application Note**

### Application:.

Screen for antibodies of human TIM-3 through Flow Cytometry.

#### **Culture conditions:**

Cells should be grown at  $37^{\circ}$ C with 5% CO<sub>2</sub> using DMEM medium (w/ L-Glutamine, 4.5g/L Glucose and Sodium Pyruvate) supplemented with 10% heat-inactivated FBS and 1% Pen/Strep, plus 10  $\mu$ g/ml of Blasticidin.

It is recommended to quickly thaw the frozen cells upon receipt or from liquid nitrogen in a  $37^{\circ}$ C water-bath, transfer to a tube containing 10 ml of growth medium without Blasticidin, spin down cells, resuspend cells in pre-warmed growth medium without Blasticidin, transfer resuspended cells to T25 flask and culture in  $37^{\circ}$ C-CO<sub>2</sub> incubator.

Leave the T25 flask in the incubator for  $1\sim2$  days without disturbing or changing the medium until cells completely recover viability and become adherent. Once cells are over 90% adherent, remove growth medium and passage the cells through trypsinization and centrifugation. At first passage, switch to growth medium containing Blasticidin. Cells should be split before they reach complete confluence.

To passage the cells, detach cells from culture vessel with Trypsin/EDTA, add complete growth medium and transfer



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to a tube, spin down cells, resuspend cells and seed appropriate aliquots of cells suspension into new culture vessels. Subcultivation ration = 1:10 to 1:20 weekly. To achieve satisfactory results, cells should not be passaged over 16 times.

### **LIMITED USE RESTRICTIONS:**

THIS PRODUCT IS SOLELY FOR IN VITRO RESEARCH USE ONLY. NOT FOR DIAGNOSTIC OR THERAPEUTIC USE.

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This product is <u>solely for Internal Research Purposes</u> and <u>not for Commercial Purposes</u>. Commercial Purposes include, but are not limited to (1) use of the cell line in manufacturing; (2) use of the cell line to provide a service, information or data; (3) use of the cell line for therapeutic, diagnostic or prophylactic purposes; or (4) resale of the cell line whether or not such cell lines are resold for use in research. <u>The buyer cannot sell, give or otherwise transfer this product to a third party.</u>

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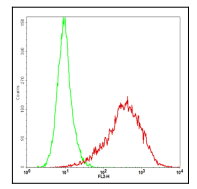


Fig-1: Detection of human TIM-3 in the CHO-K1/TIM-3 stable cell line by Flow Cytometry using monoclonal anti-human TIM-3 antibody (ABEOMICS, Cat. No. 10-4076). A goat anti-mouse PE-conjugated secondary antibody was used. Parental CHO-K1 cells (Green); CHO-K1/TIM-3 stable cell line (Red).