abcam

Product datasheet

Human BMI1 knockout HEK-293T cell line ab266514

2 Images

Overview

Product name	Human BMI1 knockout HEK-293T cell line		
Parental Cell Line	HEK293T		
Organism	Human		
Mutation description	Knockout achieved by using CRISPR/Cas9, Homozygous: 543 bp deletion in exon 2		
Passage number	<20		
Knockout validation	Sanger Sequencing, Western Blot (WB)		
Tested applications	Suitable for: WB		
Biosafety level	2		
General notes	Recommended control: Human wild-type HEK293T cell line (<u>ab255449</u>). Please note a wild- type cell line is not automatically included with a knockout cell line order, if required please add recommended wild-type cell line at no additional cost using the code WILDTYPE-TMTK1.		
	Cryopreservation cell medium: Cell Freezing Medium-DMSO Serum free media, contains 8.7% DMSO in MEM supplemented with methyl cellulose.		
	Culture medium: DMEM (High Glucose) + 10% FBS		
	Initial handling guidelines: Upon arrival, the vial should be stored in liquid nitrogen vapor phase and not at -80°C. Storage at -80°C may result in loss of viability.		
	 Thaw the vial in 37°C water bath for approximately 1-2 minutes. Transfer the cell suspension (0.8 mL) to a 15 mL/50 mL conical sterile polypropylene centrifuge tube containing 8.4 mL pre-warmed culture medium, wash vial with an additional 0.8 mL culture medium (total volume 10 mL) to collect remaining cells, and centrifuge at 201 x g (rcf) for 5 minutes at room temperature. 10 mL represents minimum recommended dilution. 20 mL represents maximum recommended dilution. 		
	 3. Resuspend the cell pellet in 5 mL pre-warmed culture medium and count using a haemocytometer or alternative cell counting method. Based on cell count, seed cells in an appropriate cell culture flask at a density of 2x10⁴ cells/cm². Seeding density is given as a guide only and should be scaled to align with individual lab schedules. 4. Incubate the culture at 37°C incubator with 5% CO₂. Cultures should be monitored daily. 		
	Subculture guidelines:		
	All seeding densities should be based on cell counts gained by established methods. A guide seeding density of 2x10 ⁴ cells/cm ² is recommended.		
	A partial media change 24 hours prior to subculture may be helpful to encourage growth, if required.		

Cells should be passaged when they have achieved 80-90% confluence. This product is subject to limited use licenses from The Broad Institute and ERS Genomics Limited, and is developed with patented technology. For full details of the limited use licenses and relevant patents please refer to our **limited use license** and **patent pages**.

We will provide viable cells that proliferate on revival.

Properties

Number of cells	1 x 10 ⁶ cells/vial, 1 mL	
Adherent /Suspension	Adherent	
Tissue	Kidney	
Cell type	epithelial	
STR Analysis	Amelogenin X D5S818: 8, 9 D13S317: 12, 14 D7S820: 11 D16S539: 9, 13 vWA: 16, 19 TH01: 7, 9.3 TPOX: 11 CSF1PO: 11, 12	
Antibiotic resistance	Puromycin 1.00µg/ml	
Mycoplasma free	Yes	
Storage instructions	Shipped on Dry Ice. Store in liquid nitrogen.	
Storage buffer	Constituents: 8.7% Dimethylsulfoxide, 2% Cellulose, methyl ether	

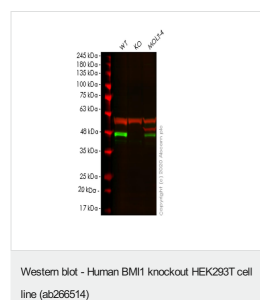
Target Function Component of the Polycomb group (PcG) multiprotein PRC1 complex, a complex required to maintain the transcriptionally repressive state of many genes, including Hox genes, throughout development. PcG PRC1 complex acts via chromatin remodeling and modification of histones; it mediates monoubiquitination of histone H2A 'Lys-119', rendering chromatin heritably changed in its expressibility. In the PRC1 complex, it is required to stimulate the E3 ubiquitin-protein ligase activity of RNF2/RING2. **Sequence similarities** Contains 1 RING-type zinc finger. **Post-translational** Monoubiquitinated (By similarity). May be polyubiquitinated; which does not lead to proteasomal modifications degradation. **Cellular localization** Nucleus. Cytoplasm.

Applications

The Abpromise guarantee Our <u>Abpromise guarantee</u> covers the use of ab266514 in the following tested applications.

The application notes include recommended starting dilutions; optimal dilutions/concentrations should be determined by the end user.

Application	Abreviews	Notes
WB		Use at an assay dependent concentration. Predicted molecular weight: 36 kDa.



All lanes : Anti-Bmi1 antibody [EPR3745(2)] (<u>ab126783</u>) at 1/1000 dilution

Lane 1 : Wild-type HEK293T cell lysate Lane 2 : BMI1 knockout HEK293T cell lysate Lane 3 : MOLT-4 cell lysate

Lysates/proteins at 20 µg per lane.

Secondary

All lanes : Goat anti-Rabbit lgG H&L (IRDye® 800CW) preadsorbed (<u>ab216773</u>) at 1/10000 dilution

Predicted band size: 36 kDa Observed band size: 37 kDa

Lanes 1-3: Merged signal (red and green). Green - <u>ab126783</u> observed at 37 kDa. Red - loading control <u>ab7291</u> observed at 50 kDa.

<u>ab126783</u> Anti-Bmi1 antibody [EPR3745(2)] was shown to specifically react with Bmi1 in wild-type HEK293T cells. Loss of signal was observed when knockout cell line ab266514 (knockout cell lysate <u>ab256850</u>) was used. Wild-type and Bmi1 knockout samples were subjected to SDS-PAGE. <u>ab126783</u> and Anti-alpha Tubulin antibody [DM1A] - Loading Control (<u>ab7291</u>) were incubated at room temperature for 2.5 hours at 1 in 1000 dilution and 1 in 20000 dilution respectively. Blots were developed with Goat anti-Rabbit IgG H&L (IRDye[®] 800CW) preadsorbed (<u>ab216773</u>) and Goat anti-Mouse IgG H&L (IRDye[®] 680RD) preadsorbed (<u>ab216776</u>) secondary antibodies at 1 in 20000 dilution for 1 hour at room temperature before imaging.

Homozygous: 543 bp deletion in exon2

Sanger Sequencing - Human BMI1 knockout HEK293T cell line (ab266514)

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