

# Datasheet for ABIN7519711

## **BMPR2 Protein (Fc Tag)**



### Overview

Quantity:	10 μg
Target:	BMPR2
Origin:	Human
Source:	HEK-293 Cells
Protein Type:	Recombinant
Biological Activity:	Active
Purification tag / Conjugate:	This BMPR2 protein is labelled with Fc Tag.

#### Product Details

Product Details	
Purpose:	Active Recombinant Human BMPR-2 Protein
Sequence:	SQNQERLCAF KDPYQQDLGI GESRISHENG TILCSKGSTC YGLWEKSKGD INLVKQGCWS HIGDPQECHY EECVVTTTPP SIQNGTYRFC CCSTDLCNVN FTENFPPPDT TPLSPPHSFN RDETI
Specificity:	Ser27-Ile151
Purity:	> 95 % by SDS-PAGE.
Sterility:	0.22 µm filtered
Endotoxin Level:	< 1 EU/µg of the protein by LAL method.
Biological Activity Comment:	Measured by its binding ability in a functional ELISA. Immobilized Human BMP2 at 1 μg/mL (100 μL/well) can bind Human BMPR2 with a linear range of 19.5 ng/ml-1.46 μg/ml.

## **Target Details**

Target:	BMPR2
Alternative Name:	BMPR-2 (BMPR2 Products)
Background:	Description: The bone morphogenetic protein type II receptor (BMPR-II, or BMPR2), a receptor
	for the transforming growth factor (TGF)-beta/bone morphogenetic protein (BMP) superfamily
	Reduced expression or function of BMPR2 signaling leads to exaggerated TGF-beta signaling
	and altered cellular responses to TGF-beta. In endothelial cells, BMPR2 mutation increases the
	susceptibility of cells to apoptosis. BMPR2 transduces BMP signals by forming heteromeric
	complexes with and phosphorylating BMP type I receptors. The intracellular domain of BMPR2
	is both necessary and sufficient for receptor complex interaction. It had been identified that
	BMPR2 plays a key role in cell growth. Its mutations lead to hereditary pulmonary hypertension
	and knockout of Bmpr-II results in early embryonic lethality. The C-terminal tail of BMPR2
	provides binding sites for a number of regulatory proteins that may initiate Smad-independent
	signalling. BMPR2 mutations were predicted to alter the BMP and TGF-b1/SMAD signalling
	pathways, resulting in proliferation rather than apoptosis of vascular cells, and greatly increase
	the risk of developing severe pulmonary arterial hypertension. BMPR2 gene result in familial
	Primary pulmonary hypertension (PPH) transmitted as an autosomal dominant trait, albeit with
	low penetrance. Heterozygous germline mutations of BMPR2 gene have been identified in
	patients with familial and sporadic PPH, indicating that BMPR2 may contribute to the
	maintenance of normal pulmonary vascular structure and function. Tctex-1, a light chain of th
	motor complex dynein, interacts with the cytoplasmic domain of BMPR2 and demonstrate tha
	Tctex-1 is phosphorylated by BMPR-II, a function disrupted by PPH disease causing mutations
	within exon 12. BMPR2 and Tctex-1 co-localize to endothelium and smooth muscle within the
	media of pulmonary arterioles, key sites of vascular remodelling in PPH.
	Name: BMPR2, BMPR-II, BMPR3, BMR2, BRK-3, POVD1, PPH1, T-ALK, bone morphogenetic
	protein receptor type-2,BMPR-II,BMPR3,BMR2,BRK-3,POVD1,PPH1,T-ALK
Gene ID:	659
JniProt:	Q13873-1
Pathways:	Growth Factor Binding
Application Details	

Restrictions: For Research Use only

## Handling

Format:	Lyophilized
Reconstitution:	Centrifuge the vial before opening. Reconstitute to a concentration of 0.1-0.5 mg/mL in sterile
	distilled water. Avoid votex or vigorously pipetting the protein. For long term storage, it is
	recommended to add a carrier protein or stablizer (e.g. 0.1 % BSA, 5 % HSA, 10 % FBS or 5 %
	Trehalose), and aliquot the reconstituted protein solution to minimize free-thaw cycles.
Buffer:	Lyophilized from a 0.22 µm filtered solution of PBS, pH 7.4.
Storage:	-20 °C,-80 °C
Storage Comment:	Store the lyophilized protein at -20°C to -80°C for long term. After reconstitution, the protein
	solution is stable at -20°C for 3 months, at 2-8°C for up to 1 week.