

Product datasheet

Recombinant Human Ephrin B1 protein (denatured) ab111630

[1 Image](#)

Description

Product name	Recombinant Human Ephrin B1 protein (denatured)
Purity	> 90 % SDS-PAGE.
Expression system	Escherichia coli
Accession	<u>P98172</u>
Protein length	Protein fragment
Animal free	No
Nature	Recombinant
Species	Human
Sequence	MGSSHHHHHHSSGLVPRGSHMLAKNLEPVSWSSLNPKF LSGKGLVIYPKI GDKLDIICPRAEAGRPYEEYKLYLVRPEQAAACSTVLDPNV LVTCNRPEQ EIRFTIKFQEFSPNYMGLEFKKHHDYYTSTSNGSLEGLNR EGGVCRT TMKIIMKVGQDPNAVTPPEQLTTSRPSKEADNTVKMATQAP GSRGSLGDSD GKHETVNQEEKSGPGASGGSSGDPDGGFFNSK
Predicted molecular weight	25 kDa including tags
Amino acids	28 to 237
Tags	His tag N-Terminus

Specifications

Our **Abpromise guarantee** covers the use of **ab111630** in the following tested applications.

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

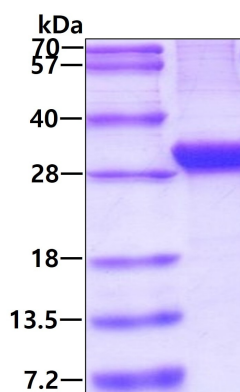
Applications SDS-PAGE

Form Liquid

Preparation and Storage

Stability and Storage	<p>Shipped at 4°C. Upon delivery aliquot and store at -20°C or -80°C. Avoid repeated freeze / thaw cycles.</p> <p>pH: 8.00</p> <p>Constituents: 2.4% Urea, 0.32% Tris HCl, 5% Glycerol (glycerin, glycerine)</p>
General Info	
Function	Binds to the receptor tyrosine kinases EPHB1 and EPHA1. Binds to, and induce the collapse of, commissural axons/growth cones in vitro. May play a role in constraining the orientation of longitudinally projecting axons.
Tissue specificity	Heart, placenta, lung, liver, skeletal muscle, kidney, pancreas.
Involvement in disease	Defects in EFNB1 are a cause of craniofrontonasal syndrome (CFNS) [MIM:304110]; also known as craniofrontonasal dysplasia (CFND). CFNS is an X-linked inherited syndrome characterized by hypertelorism, coronal synostosis with brachycephaly, downslanting palpebral fissures, clefting of the nasal tip, joint anomalies, longitudinally grooved fingernails and other digital anomalies.
Sequence similarities	Belongs to the ephrin family.
Post-translational modifications	Inducible phosphorylation of tyrosine residues in the cytoplasmic domain.
Cellular localization	Membrane.

Images



15% SDS-PAGE showing ab111630 at approximately 25.3kDa (3µg).

SDS-PAGE - Recombinant Human Ephrin B1
protein (denatured) (ab111630)

Please note: All products are "FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES"

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