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LARGE Protein (AA 1-756) (rho-1D4 tag)





Go to Product page

Overview

Quantity:	1 mg
Target:	LARGE
Protein Characteristics:	AA 1-756
Origin:	Human
Source:	Insect Cells
Protein Type:	Recombinant
Purification tag / Conjugate:	This LARGE protein is labelled with rho-1D4 tag.
Application:	Western Blotting (WB), SDS-PAGE (SDS), ELISA, Crystallization (Crys)

Product Details

Sequence:

MLGICRGRRK FLAASLSLLC IPAITWIYLF SGSFEDGKPV SLSPLESQAH SPRYTASSQR
ERESLEVRMR EVEEENRALR RQLSLAQGRA PSHRRGNHSK TYSMEEGTGD SENLRAGIVA
GNSSECGQQP VVEKCETIHV AIVCAGYNAS RDVVTLVKSV LFHRRNPLHF HLIADSIAEQ
ILATLFQTWM VPAVRVDFYN ADELKSEVSW IPNKHYSGIY GLMKLVLTKT LPANLERVIV
LDTDITFATD IAELWAVFHK FKGQQVLGLV ENQSDWYLGN LWKNHRPWPA LGRGYNTGVI
LLLLDKLRKM KWEQMWRLTA ERELMGMLST SLADQDIFNA VIKQNPFLVY QLPCFWNVQL
SDHTRSEQCY RDVSDLKVIH WNSPKKLRVK NKHVEFFRNL YLTFLEYDGN LLRRELFGCP
SEADVNSENL QKQLSELDED DLCYEFRRER FTVHRTHLYF LHYEYEPAAD STDVTLVAQL
SMDRLQMLEA ICKHWEGPIS LALYLSDAEA QQFLRYAQGS EVLMSRHNVG YHIVYKEGQF
YPVNLLRNVA MKHISTPYMF LSDIDFLPMY GLYEYLRKSV IQLDLANTKK AMIVPAFETL
RYRLSFPKSK AELLSMLDMG TLFTFRYHVW TKGHAPTNFA KWRTATTPYR VEWEADFEPY
VVVRRDCPEY DRRFVGFGWN KVAHIMELDV QEYEFIVLPN AYMIHMPHAP SFDITKFRSN

KQYRICLKTL KEEFQQDMSR RYGFAALKYL TAENNS

Sequence without tag. Tag location is at the discretion of the manufacturer. If you have a special request, please contact us.

Characteristics:

- Made in Germany from design to production by highly experienced protein experts.
- Human LARGE Protein (raised in Insect Cells) purified by multi-step, protein-specific process to ensure crystallization grade.
- State-of-the-art algorithm used for plasmid design (Gene synthesis).

This protein is a made to order protein and will be made for the first time for your order. Our experts in the lab will ensure that you receive a correctly folded protein.

The big advantage of ordering our made-to-order proteins in comparison to ordering custom made proteins from other companies is that there is no financial obligation in case the protein cannot be expressed or purified.

In the unlikely event that the protein cannot be expressed or purified we do not charge anything (other companies might charge you for any performed steps in the expression process for custom-made proteins, e.g. fees might apply for the expression plasmid, the first expression experiments or purification optimization).

When you order this made-to-order protein you will only pay upon receival of the correctly folded protein. With no financial risk on your end you can rest assured that our experienced protein experts will do everything to make sure that you receive the protein you ordered.

The concentration of our recombinant proteins is measured using the absorbance at 280nm.

The protein's absorbance will be measured in several dilutions and is measured against its specific reference buffer.

The concentration of the protein is calculated using its specific absorption coefficient. We use the Expasy's protparam tool to determine the absorption coefficient of each protein.

Purification:

Three step purification of membrane proteins expressed in baculovirus infected SF9 insect cells:

- 1. Membrane proteins are fractioned by ultracentrifugation and subsequently solubilized with different detergents (detergent screen). Samples are analyzed by Western blot.
- 2. The best performing detergent is used for solubilization and the proteins are purified via their rho1D4 tag via two rho1D4 antibody columns: one DTT resistant, the other one not. Eluate fractions are analyzed by Western blot.
- 3. Protein containing fractions of the best purification are subjected to second purification step through size exclusion chromatograph. Eluate fractions are analyzed by SDS-PAGE and Western blot.

Purity:

>95 % as determined by SDS PAGE, Size Exclusion Chromatography and Western Blot.

Sterility:

0.22 µm filtered

Product Details Protein is endotoxin-free. Endotoxin Level: Grade: Crystallography grade Target Details LARGE Target: Alternative Name: LARGE (LARGE Products) Background: Bifunctional glycosyltransferase with both xylosyltransferase and beta-1,3glucuronyltransferase activities involved in the biosynthesis of the phosphorylated O-mannosyl trisaccharide (N-acetylgalactosamine-beta-3-N-acetylglucosamine-beta-4-(phosphate-6-)mannose), a carbohydrate structure present in alpha-dystroglycan (DAG1) (PubMed:22223806). Phosphorylated O-mannosyl trisaccharid is required for binding laminin Glike domain-containing extracellular proteins with high affinity and plays a key role in skeletal muscle function and regeneration. LARGE elongates the glucuronyl-beta-1,4-xylose-beta disaccharide primer structure initiated by B3GNT1/B4GAT1 by adding repeating units [-3-Xylose-alpha-1,3-GlcA-beta-1-] to produce a heteropolysaccharide (PubMed:25279699). {ECO:0000269|PubMed:15661757, ECO:0000269|PubMed:15752776, ECO:0000269|PubMed:21987822, ECO:0000269|PubMed:22223806, ECO:0000269|PubMed:25138275, ECO:0000269|PubMed:25279697, ECO:0000269|PubMed:25279699}. Molecular Weight: 89.2 kDa Including tag. UniProt: 095461 **Application Details Application Notes:** In addition to the applications listed above we expect the protein to work for functional studies as well. As the protein has not been tested for functional studies yet we cannot offer a gurantee though. Comment: In cases in which it is highly likely that the recombinant protein with the default tag will be

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receive your protein of interest.

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Restrictions:

insoluble our protein lab may suggest a higher molecular weight tag (e.g. GST-tag) instead to

increase solubility. We will discuss all possible options with you in detail to assure that you

Handling

Format:	Liquid
Buffer:	100 mM NaCL, 20 mM Hepes, 10% glycerol. pH value is at the discretion of the manufacturer.
Handling Advice:	Avoid repeated freeze-thaw cycles.
Storage:	-80 °C
Storage Comment:	Store at -80°C.
Expiry Date:	Unlimited (if stored properly)

Images

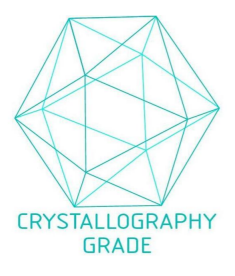


Image 1. "Crystallography Grade" protein due to multi-step, protein-specific purification process