

Recombinant SARS-CoV S1 Subunit protein

Catalog Number: ATGP4012

PRODUCT INFORMATION

Expression system

HEK293

Domain

14-667aa

UniProt No.

P59594

NCBI Accession No.

NP_828851.1

Alternative Names

E2 glycoprotein precursor, Spike glycoprotein, S glycoprotein, E2, Peplomer protein, Severe acute respiratory Syndrome-related Coronavirus, SARS, SRAS-CoV, SARS-CoV1, spike protein S1

PRODUCT SPECIFICATION

Molecular Weight

73.7kDa(660aa)

Concentration

0.25mg/ml (determined by Absorbance at 280nm)

Formulation

Liquid. In Phosphate-Buffered Saline (pH 7.4) containing 10% glycerol

Purity

> 90% by SDS - PAGE

Endotoxin level

< 1 EU per 1ug of protein (determined by LAL method)

Biological Activity

Measured by its binding ability in a functional ELISA with Human ACE-2 (CAT# ATGP3963)

Tag

His-Tag

Application

SDS-PAGE, Bioactivity

Storage Condition

Can be stored at +2C to +8C for 1 week. For long term storage, aliquot and store at -20C to -80C. Avoid repeated freezing and thawing cycles.

BACKGROUND

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Description

Severe Acute Respiratory Syndrome Coronavirus (SARS-CoV), Middle Eastern Respiratory Syndrome Coronavirus (MERS-CoV), and the recently identified novel Coronavirus (SARS-CoV-2) belong to the Coronaviridae family, genus Betacoronavirus, that has been related to important epidemiological outbreaks. SARS-CoV emerged in 2003 as a significant threat to human health. SARS-CoV has four structural proteins, known as the S (spike), E (envelope), M (membrane), and N (nucleocapsid) proteins. The spike protein, responsible for allowing the virus to attach to and fuse with the membrane of a host cell and is a large type I transmembrane protein containing two subunits, S1 and S2. S1 mainly contains a receptor binding domain (RBD), which is responsible for recognizing the cell surface receptor. S2 contains basic elements needed for the membrane fusion. The S protein plays key parts in the induction of neutralizing-antibody and T-cell responses, as well as protective immunity. It attaches the virion to the cell membrane by interacting with host receptor, initiating the infection. A metalloproteinase, angiotensin-converting enzyme 2 (ACE-2), has been identified as a functional receptor for SARS-CoV through interaction with a receptor binding domain (RBD) located at the C-terminus of S1 subunit. Recombinant SARS-CoV spike S1 subunit fused to His-tag at C-terminus, was expressed in HEK293 cell and purified by using conventional chromatography techniques.

Amino acid Sequence

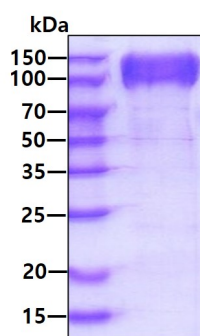
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KSNVVRGWVF GSTMNNKSQS VIIIINNSTNV VIRACNFELC DNPFFAVSKP MGTQTHMIF DNAFNCTFEY ISDAFSLDVS  
EKSGNFKHLR EFVFKNKDGF LYVYKGYQPI DVVRDLPSGF NTLKPIFKLP LGINITNFRA ILTAFSPAQD IWGTSAAAYF  
VGYLKPTTFM LKYDENGIT DAVDCSQNPL AELKCSVKSF EIDKGIYQTS NFRVVPDGDV VRFNPITNLC PFGEVFNATK  
FPSVYAWERK KISNCVADYS VLYNSTFFST FKCYGVSATK LNDLCFSNVY ADSFVVKGDD VRQIAPGQTG VIADYNYKLP  
DDFMGCVLAW NTRNIDATST GNYNKYRYL RHGKLRPFER DISNVPFSPD GKPCTPPALN CYWPLNDYGF YTTTGIGYQP  
YRVVLSFEL LNAPATVCGP KLSTDLIKNQ CVNFNFNGLT GTGVLTPSSK RFQPFQFGR DVSDFTDSVR DPKTSEILDI  
SPCAFGGVSV ITPGTNASSE VAVLYQDVNC TDVSTAIHAD QLTPAWRIYS TGNNVFQTQA GCLIGAEHVD TSYECDPIG  
AGICASYHTV SLLR<HHHHHH>
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General References

Kukla M., et al, (2020) J Clin Med. 9:1420.
Ayoub A., et al, (2020) J Clin Virol. 129:104521.
Tortorici, M.A. and D. Veessler (2019). Adv. Virus Res. 105:93-116.
Li F, et al, (2005) Science. 309:1864-1868.
Struck AW, et al, (2012) Antiviral Res. 94:288-296.

DATA

SDS-PAGE



3ug by SDS-PAGE under reducing condition and visualized by coomassie blue stain

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Biological Activity

SARS-CoV S1 Subunit is coated at 5 ug/ml (100 ul/well) can bind Human ACE-2 (CAT# ATGP3963) in a Functional ELISA assay.

