# abcam

## Product datasheet

# Recombinant Human Von Willebrand Factor protein (Tagged) ab152801

1 References 1 图像

描述

产品名称 重组人Von Willebrand Factor蛋白(Tagged)

纯度 >= 80 % Purified via GST Tag.

Glutathione Sepharose

表达系统 Wheat germ Accession P04275

**蛋白长度** Protein fragment

无动物成分 No

性质 Recombinant

种属 Human

序列 MGAQDEEEGIQDLDGLLVFDKIVEVTLLNLPWYNEETEGQRG

**EMTAPKSP** 

 ${\tt RAKIRGTLCAEGTRGRSSTARCSLFGSDFVNTFDGSMYSFAG}$ 

**YCSYLLAG** 

GCQKRSFSIIGDFQNGKRVSLSVYLGEFFDIHLFVNGTVTQG

**DQRVSMPY** 

ASKGLYLETEAGYYKLSGEAYGFVARIDGSGNFQVLLSDRYF

**NKTCGLCG** 

NFNIFAEDDFMTQEGTLTSDPYDFANSWALSSGEQWCERASP

PSSSCNIS SGEMQKVGVDWPGCTWMVCDFWI

预测分子量 56 kDa including tags

**氨基酸** 1 to 273

标签 GST tag N-Terminus

技术指标

Our Abpromise guarantee covers the use of ab152801 in the following tested applications.

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

应**用** SDS-PAGE

Western blot

ELISA

1

形式 Liquid

**补充说明** This recombinant protein is a short type (isoform) of VWF.

#### 制备和贮存

稳**定性和存储** Shipped on dry ice. Upon delivery aliquot and store at -80℃. Avoid freeze / thaw cycles.

00.8:Ha

Constituents: 0.31% Glutathione, 0.79% Tris HCI

#### 常规信息

功能 Important in the maintenance of hemostasis, it promotes adhesion of platelets to the sites of

vascular injury by forming a molecular bridge between sub-endothelial collagen matrix and platelet-surface receptor complex GPlb-IX-V. Also acts as a chaperone for coagulation factor VIII, delivering it to the site of injury, stabilizing its heterodimeric structure and protecting it from

premature clearance from plasma.

组织特异性 Plasma.

疾病相关 Defects in VWF are the cause of von Willebrand disease (VWD) [MIM:277480]. VWD defines a

group of hemorrhagic disorders in which the von Willebrand factor is either quantitatively or qualitatively abnormal resulting in altered platelet function. Symptoms vary depending on severity and disease type but may include prolonged bleeding time, deficiency of factor VIII and impaired platelet adhesion. Type I von Willebrand disease is the most common form and is characterized by partial quantitative plasmatic deficiency of an otherwise structurally and functionally normal Willebrand factor; type II is associated with a qualitative deficiency and functional anomalies of the

Willebrand factor; type Ill is the most severe form and is characterized by total or near-total absence of Willebrand factor in the plasma and cellular compartments, also leading to a profound

deficiency of plasmatic factor VIII.

序列相似性 Contains 1 CTCK (C-terminal cystine knot-like) domain.

Contains 4 TIL (trypsin inhibitory-like) domains.

Contains 3 VWFA domains. Contains 3 VWFC domains. Contains 4 VWFD domains.

结**构域** The von Willebrand antigen 2 is required for multimerization of vWF and for its targeting to

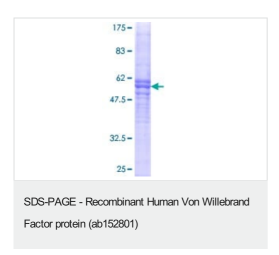
storage granules.

翻译后修饰 All cysteine residues are involved in intrachain or interchain disulfide bonds.

N- and O-glycosylated.

细胞定位 Secreted. Secreted > extracellular space > extracellular matrix. Localized to storage granules.

图片



12.5% SDS-PAGE analysis of ab152801 stained with Coomassie Blue.

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