

Product datasheet

info@arigobio.com

ARG70022 anti-Factor IX antibody

Package: 250 μg Store at: -20°C

Summary

Product Description Goat Polyclonal antibody recognizes Factor IX

Tested Reactivity Hu

Tested Application IHC-P, WB

Host Goat

Clonality Polyclonal

Isotype IgG

Target Name Factor IX

Species Human

Immunogen Human Factor IX purified from plasma.

Conjugation Un-conjugated

Alternate Names Coagulation factor IX; HEMB; FIX; PTC; Plasma thromboplastin component; F9 p22; THPH8; EC

3.4.21.22; P19; Christmas factor

Application Instructions

Application table	Application	Dilution
	IHC-P	Assay-dependent
	WB	Assay-dependent
Application Note	* The dilutions indicate recommended starting dilutions and the optimal dilutions or concentrations should be determined by the scientist.	

Properties

Form Liquid

Purification Affinity purification with immunogen.

Buffer 10 mM HEPES (pH 7.4), 150 mM NaCl and 50% (v/v) Glycerol.

Stabilizer 50% (v/v) Glycerol

Storage instruction For continuous use, store undiluted antibody at 2-8°C for up to a week. For long-term storage, aliquot

and store at -20°C. Storage in frost free freezers is not recommended. Avoid repeated freeze/thaw cycles. Suggest spin the vial prior to opening. The antibody solution should be gently mixed before use.

Note For laboratory research only, not for drug, diagnostic or other use.

Bioinformation

Gene Symbol

www.arigobio.com arigo.nuts about antibodies 1/2

Gene Full Name coagulation factor IX

Background This gene encodes vitamin K-dependent coagulation factor IX that circulates in the blood as an inactive

zymogen. This factor is converted to an active form by factor XIa, which excises the activation peptide and thus generates a heavy chain and a light chain held together by one or more disulfide bonds. The role of this activated factor IX in the blood coagulation cascade is to activate factor X to its active form through interactions with Ca+2 ions, membrane phospholipids, and factor VIII. Alterations of this gene, including point mutations, insertions and deletions, cause factor IX deficiency, which is a recessive X-linked disorder, also called hemophilia B or Christmas disease. Alternative splicing results in multiple transcript variants encoding different isoforms that may undergo similar proteolytic processing.

[provided by RefSeq, Sep 2015]

Function Factor IX is a vitamin K-dependent plasma protein that participates in the intrinsic pathway of blood

coagulation by converting factor X to its active form in the presence of Ca(2+) ions, phospholipids, and

factor VIIIa. [UniProt]

Calculated Mw 52 kDa

PTM Activated by factor XIa, which excises the activation peptide (PubMed:9169594, PubMed:1730085). The

propeptide can also be removed by snake venom protease (PubMed:20004170, PubMed:20080729).

The iron and 2-oxoglutarate dependent 3-hydroxylation of aspartate and asparagine is (R)

stereospecific within EGF domains.

Predominantly O-glucosylated at Ser-99 by POGLUT1 in vitro. Xylosylation at this site is minor. [UniProt]

Cellular Localization Secreted [UniProt]