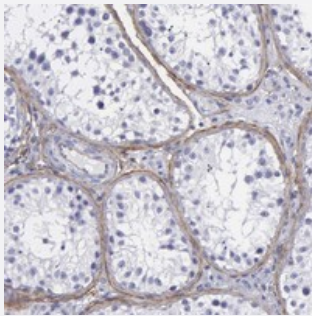


FBLN5 polyclonal antibody

Catalog # PAB20009 Size 100 uL

Applications



Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections)

Immunohistochemical staining of human testis with FBLN5 polyclonal antibody (Cat # PAB20009) shows distinct positivity in capsular cells.

Specification

Product Description	Rabbit polyclonal antibody raised against recombinant FBLN5.
Immunogen	Recombinant protein corresponding to amino acids of human FBLN5.
Sequence	QAQCTNGFDLDRQSGQCLDIDECRTIPEACRGDMMCVNQNGGYLCIPRTNPVYRGPYSNPYSTPYSGPYAAAPPLSAPNYPTISRPLICRFGYQMDESNCVDVDECATDSHQCNPQTQICINTEGGYTCSCTDGYWLLEGQCLDIDEC
Host	Rabbit
Reactivity	Human
Form	Liquid
Purification	Antigen affinity purification
Isotype	IgG
Recommend Usage	Immunohistochemistry (1:50-1:200) Western Blot (1:250-1:500) The optimal working dilution should be determined by the end user.
Storage Buffer	In PBS, pH 7.2 (40% glycerol, 0.02% sodium azide)

Storage Instruction

Store at 4°C. For long term storage store at -20°C.
Aliquot to avoid repeated freezing and thawing.

Note

This product contains sodium azide: a POISONOUS AND HAZARDOUS SUBSTANCE which should be handled by trained staff only.

Applications

- Western Blot
- Immunohistochemistry (Formalin/PFA-fixed paraffin-embedded sections)

Immunohistochemical staining of human testis with FBLN5 polyclonal antibody (Cat # PAB20009) shows distinct positivity in capsular cells.

Gene Info — FBLN5

Entrez GeneID[10516](#)**Protein Accession#**[Q9UBX5](#)**Gene Name**

FBLN5

Gene Alias

ARMD3, DANCE, EVEC, FIBL-5, FLJ90059, UP50

Gene Description

fibulin 5

Omim ID[123700](#) [219100](#) [604580](#) [608895](#)**Gene Ontology**[Hyperlink](#)**Gene Summary**

The protein encoded by this gene is a secreted, extracellular matrix protein containing an Arg-Gly-Asp (RGD) motif and calcium-binding EGF-like domains. It promotes adhesion of endothelial cells through interaction of integrins and the RGD motif. It is prominently expressed in developing arteries but less so in adult vessels. However, its expression is reinduced in balloon-injured vessels and atherosclerotic lesions, notably in intimal vascular smooth muscle cells and endothelial cells. Therefore, the protein encoded by this gene may play a role in vascular development and remodeling. Defects in this gene are a cause of autosomal dominant cutis laxa, autosomal recessive cutis laxa type I (CL type I), and age-related macular degeneration type 3 (ARMD3). [provided by RefSeq]

Other Designations

developmental arteries and neural crest epidermal growth factor-like|urine p50 protein

Disease

- [Aortic Aneurysm](#)
- [Atherosclerosis](#)
- [Calcinosis](#)
- [Coronary Artery Disease](#)
- [Diabetes Mellitus](#)
- [Disease Progression](#)
- [Genetic Predisposition to Disease](#)
- [Macular Degeneration](#)
- [Tobacco Use Disorder](#)
- [Vision](#)