abcam

Product datasheet

APC Anti-CD105 antibody [SN6], prediluted ab155367

1 References 1 Image

Overview

Product name APC Anti-CD105 antibody [SN6], prediluted

Description APC Mouse monoclonal [SN6] to CD105, prediluted

Host species Mouse

Conjugation APC. Ex: 645nm, Em: 660nm

Tested applications
Suitable for: Flow Cyt
Species reactivity
Reacts with: Human

Immunogen Full length native protein (purified) corresponding to CD105.

Positive control U937 cells.

General notes

The Life Science industry has been in the grips of a reproducibility crisis for a number of years.

Abcam is leading the way in addressing this with our range of recombinant monoclonal antibodies and knockout edited cell lines for gold-standard validation. Please check that this product meets

your needs before purchasing.

If you have any questions, special requirements or concerns, please send us an inquiry and/or contact our Support team ahead of purchase. Recommended alternatives for this product can be

found below, along with publications, customer reviews and Q&As

Properties

Form Liquid

Storage instructions Shipped at 4°C. Store at +4°C.

Storage buffer pH: 7.20

Preservative: 0.09% Sodium azide Constituents: 0.2% BSA, 99% PBS

Purity Protein G purified

Clonality Monoclonal

Clone numberSN6IsotypelgG1Light chain typekappa

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Applications

The Abpromise guarantee

Our Abpromise quarantee covers the use of ab155367 in the following tested applications.

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

Application	Abreviews	Notes
Flow Cyt		Use 5µl for 10 ⁶ cells. ab37391 - Mouse monoclonal lgG1, is suitable for use as an isotype control with this antibody.

Target

Function Major glycoprotein of vascular endothelium. May play a critical role in the binding of endothelial

cells to integrins and/or other RGD receptors.

Tissue specificity Endoglin is restricted to endothelial cells in all tissues except bone marrow.

Involvement in diseaseDefects in ENG are the cause of hereditary hemorrhagic telangiectasia type 1 (HHT1)

[MIM:187300, 108010]; also known as Osler-Rendu-Weber syndrome 1 (ORW1). HHT1 is an autosomal dominant multisystemic vascular dysplasia, characterized by recurrent epistaxis, muco-cutaneous telangiectases, gastro-intestinal hemorrhage, and pulmonary (PAVM), cerebral (CAVM) and hepatic arteriovenous malformations; all secondary manifestations of the underlying vascular dysplasia. Although the first symptom of HHT1 in children is generally nose bleed, there

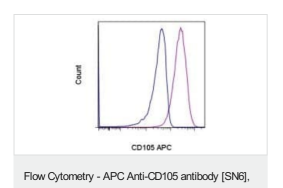
is an important clinical heterogeneity.

Cellular localization

prediluted (ab155367)

Membrane.

Images



Flow cytometric analysis of U937 cells with mouse IgG1 kappa Isotype Control APC (blue histogram) or ab155367 (purple histogram). Total viable cells were used for analysis.

Please note: All products are "FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES"

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