BioLegend®

PE anti-human CD45

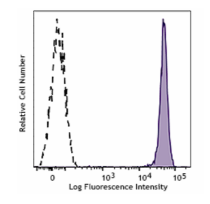
Analyte Specific Reagent. Analytical and performance characteristics are not established.

Catalog# / Size	982322 / 500 µL
Clone	HI30
Workshop	IV N816
Other Names	LCA, T200
Isotype	Mouse IgG1, κ
Description	CD45 is a 180-240 kD single chain type I membrane glycoprotein also known as leukocyte common antigen (LCA) and T200. It is a tyrosine phosphatase expressed on the plasma membrane of all hematopoietic cells, except erythrocytes and platelets. CD45 is a signaling molecule that regulates a variety of cellular processes including cell growth, differentiation, cell cycle, and oncogenic transformation. CD45 plays a critical role in T and B cell antigen receptor-mediated activation by dephosphorylating substrates including p56Lck, p59Fyn, and other Src family kinases. CD45 non-covalently associates with lymphocyte phosphatase-associated phosphoprotein (LPAP) on T and B lymphocytes. CD45 has been reported to bind galectin-1 and to be associated with several other cell surface antigens including CD1, CD2, CD3, and CD4.

Product Details

Reactivity	Human					
Formulation	Phosphate-buffered solution, pH7.2, 0.09% sodium azide, 0.2% (w/v) BSA (origin USA), an stabilizer.					
Preparation	The antibody was purified by affinity chromatography and conjugated with PE under optimal conditions.					
Concentration	10 μg/mL					
Storage & Handling	The antibody solution should be stored undiluted between 2°C and 8°C, and protected from prolonged exposure to light. Do not freeze .					
Application	Suggested for Flow Cytometry					
Disclaimer	WARNINGS AND PRECAUTIONS					
	 Use appropriate personal protective equipment and safety practices per universal precautions when working with this reagent. Refer to the reagent safety data sheet. This antibody contains sodium azide. Follow federal, state and local regulations to dispose of this reagent. Sodium azide build-up in metal wastepipes may lead to explosiv conditions; if disposing of reagent down wastepipes, flush with water after disposal. All specimens, samples and any material coming in contact with them should be considered potentially infectious and should be disposed of with proper precautions and in accordance with federal, state and local regulations. Do not use this reagent beyond the expiration date stated on the label. Do not use this reagent if it appears cloudy or if there is any change in the appearance of the reagent as these may be an indication of possible deterioration. Avoid prolonged exposure of the reagent or stained cells to light. 					

Product Data



Typical results from human peripheral blood lymphocytes stained either with HI30 PE used at 5 μ L/test (filled histogram) or with an isotype control (open histogram).

Symbols Glossary*

Symbol	Meaning	Symbol Title	Symbol No.	Symbol	Meaning	Symbol Title	Symbol No.
REF	Catalog number	Catalogue number	5.1.6	Í	Indicates the need for the user to consult the instructions for use.	Consult instructions for use	5.4.3
X	Indicates the temperature limits to which the medical device can be safely exposed.	Temperature limit	5.3.7	漱	Indicates a medical device that needs protection from light sources.	Keep away from sunlight	5.3.2
X	Indicates the upper limit of temperature to which the medical device can be safely exposed.	temperature	5.3.6	Σ	Indicates the date after which the medical device is not to be used.	Use-by date	5.1.4
•••	Indicates the medical device manufacturer.	Manufacturer	5.1.1	EC REP	Indicates the authorized representative in the European Community.	Authorized representative in the European Community	5.1.2
LOT	Indicates the manufacturer's batch code so that the batch or lot can be identified.	Batch code	5.1.5	IVD	Indicates a medical device that is intended to be used as an in vitro diagnostic medical device.	<i>In vitro</i> diagnostic medical device	5.5.1

* Symbol information is from EN ISO 15223-1:2016 Medical devices – Symbols to be used with medical device labels, labelling and information to be supplied – Part 1: General requirements

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