

INTENDED USE

CD59-FITC is a monoclonal antibody (MAb) labelled with fluorescein isothiocyanate (FITC) designed for flow cytometry use as a direct immunofluorescence reagent in the identification and enumeration of CD59 antigen-expressing cells.

SUMMARY AND EXPLANATION

Flow Cytometry (FC) is a powerful tool in analytical and quantitative characterization of cells which provides rapid and multiparametric analysis of heterogeneous cell populations on a cell-by-cell basis. Flow cytometry is performed on cell suspension after incubating it with fluorescent-labelled antibodies directed against specific cellular proteins. Positive cells relative fluorescence intensity indicates the amount of antibody bonded to specific cell sites providing information about antigen expression ⁽¹⁾.

Membrane-associated C-regulatory proteins are the major protection against Complement attack. CD59 (Protectin), a 19 kDa GPI-linked membrane glycoprotein is one of these regulatory proteins binding to the complement components C8 and C9 preventing the destruction of autologous cells. CD59 has a broadly distributed expression on all lymphocytes, monocytes, granulocytes, erythrocytes and in a variety of non-haematological cells. CD59 modulates T cell adhesion and plays a role in T cell activation ^(2, 3).

CD59 deficiency was already detected in clinical inflammatory disorders. Cells from patients with the haematological disorder paroxysmal nocturnal haemoglobinuria (PNH) are defective for GPI anchor synthesis and hence lack surface expression of CD59 and are highly susceptible to the lytic effects of Complement attack. For this reason CD59 together with CD55, another GPI-anchored protein, have been systematically used in diagnosis of PNH ^(4, 5).

PROCEDURE PRINCIPLES

Flow cytometry is an innovative technology that can evaluate simultaneously different characteristics for a single cell. Flow cytometers use hydrodynamic focusing to individually present cells to one or more laser beams. As cells are intercepted by light a set detectors recover signals of two different kinds: those generated by dispersed light (FSC/SSC), which mainly reflect cell size and internal complexity, and those related to fluorochromes light emission when cells are labelled. Recovered signals are then amplified by a series of linear and logarithmic amplifiers and converted in electrical signals large enough to be plotted graphically.

Fluorochrome-labelled monoclonal antibodies bind specific antigens, therefore cell populations carrying this antigen will be detected when the reagent is added to a sample and passed through a flow cytometer.

Detection of targeted populations can be hindered by erythrocytes presence. This problem can be avoided by their removal using a red blood cell lysing solution previous to sample acquisition. Quicklysis™ (CYT-QL-1) erythrocyte lysing solution without fixatives is recommended since it requires no washing steps. Quicklysis™ use minimises sample handling and avoids centrifuge processed which are usually associated to cell losses ^(6, 7).

CD59 positive cells are expressed as a percentage of different leucocytes subsets or erythrocytes present in a sample. Since every flow cytometer has different operating characteristics each laboratory must determine its optimal operating procedure.

REAGENT COMPOSITION

The purified monoclonal CD59 Antibody conjugated with fluorescein isothiocyanate (FITC) is supplied in phosphate-buffered saline (PBS) containing 0.1% sodium azide.

Clone: MEM-43

Isotype: IgG2a.

Amount per 1 mL vial: 200 tests (5 µL MAb per determination).

Reagent is considered non-sterile.

STORAGE CONDITIONS

The reagent is stable when stored at 2-8 °C until expiration date shown on label. The reagent should not be frozen or exposed to direct light during storage or during cell incubation. Reagent vial should be kept dry and once open stored in vertical position to avoid any possible spillage.

WARNINGS AND RECOMMENDATIONS

1. For research use only.
2. This product is supplied ready to use, any modification by dilution or addition of other compounds should be validated by the user.
3. The reagent is stable until its expiry date when properly stored. Do not use it after expiration date shown on label. If product is stored in conditions different from those recommended, such conditions must be validated by the user.
4. Alteration in reagent appearance, such as precipitation or discoloration indicates instability or deterioration. In such cases, the reagent should not be used.
5. It contains 0,1% sodium azide (CAS-Nr. 26628-22-8) as a preservative but care should anyway be taken to avoid microbial contamination of reagent and reduce incorrect results that can probably arise from that contamination.
 - Sodium azide (NaN₃) is harmful if swallowed (R22). If swallowed, seek medical advice immediately and show this container or label (S46).
 - Wear suitable protecting clothing (S36).
 - Contact with acids liberates very toxic gas (R32).
 - On disposal, flush with large amounts of water to prevent azide build-up in metal plumbing since explosive conditions may develop.
6. All patient specimens and materials are considered biohazards and should be handled as if capable of transmitting infection ⁽⁸⁾. Disposal should be made according to the established legal precautions. It is also recommendable the use of appropriate protective gloves and clothing when handling this product. Product use should be made by personnel qualified to perform the described procedures. Avoid sample contact with skin or mucous membranes. Wash immediately with abundant water if skin contact has occurred.

7. Reagent use with incubation times or temperatures different from those recommended may cause erroneous results. Any changes in procedure must be validated by the user.

PROCEDURE

Material included

CD59-FITC can be used for 200 determinations (5 µL MAb per determination).

Material required but not included

- 488 nm ion argon laser-equipped flow cytometer and appropriate computer hardware and software.
- Test tubes suitable for the used flow cytometer. Usually 6 mL tubes (12x 75 mm) with a rounded bottom are used.
- Automatic pipette (100µL) and tips.
- Micropipette and tips.
- Chronometer.
- Vortex Mixer.
- Isotypic control reagent.
- Quicklysis™ lysing solution.
- Wash buffer (phosphate buffered saline (PBS) containing 0.1% sodium azide).
- Pasteur pipette or vacuum system.

Preparation

Whole blood sample must be taken aseptically by venipuncture^(9, 10) in a sterilized tube containing an appropriate anticoagulant (use of EDTA is recommended). Flow cytometry analysis require one hundred (100) µL whole blood sample per tube, assuming a normal range of approximately 4 to 10 x 10³ leucocytes per µL. Samples with high white blood cell count should be diluted with PBS to obtain an approximate cell concentration of 1 x 10⁴ cells/µL. Store blood samples at 18-22°C until they are to be tested. It is advisable to test blood samples within 24 hours after their extraction. Samples with suspended cell aggregates or haemolysed should be rejected.

For PNH studies, CD55 and CD59 expression on leucocytes should be performed following a lyse-wash-and then stain direct immunofluorescence technique⁽¹¹⁾. For other studies a stain and the lyse method is recommended.

1. Mix 100µL of peripheral blood with 2 mL of Quicklysis™ erythrocyte lysing solution and incubate sample in the dark at room temperature for 10 minutes.
2. In order to wash out leftover lysing solution, centrifuge for 5 min at 540g, discard supernatant and resuspend cell pellet in 2mL of PBS for a second centrifugation step (5 min at 540g). Discard supernatant and resuspend cell pellet.
3. Add 5µL of CD59-FITC.
To evaluate non-specific binding of the antibody, an appropriated isotype control tube can be prepared.
3. Incubate in the dark at room temperature for 15 minutes.
4. Acquisition in the flow cytometer should be performed within the first four hours after sample preparation. If samples are not acquired immediately after preparation, they should be stored in the dark at 2-8°C. Instrument calibration must be done according to manufacturers' advice. Before acquiring samples, adjust threshold or discriminator to minimise debris and ensure that populations of interest are included. Also, previous to acquisition samples should be mixed on a vortex (low speed) to reduce cell aggregation.

*Note: The use of other lysing solutions may require elimination of lysed red blood cells. Follow lysing solution manufacturer recommended protocol.

Flow cytometry analysis

Confirm that the cytometer is correctly aligned and standardised for light dispersion and fluorescent intensity. Compensation should be set following cytometer manufacturer instructions.

Verify CD59 positive cells comparing the parameter CD59 with the relevant parameters (e.g. CD14 for monocytes, CD16 for neutrophils, etc.).

Results are commonly reported as a percentage of CD59 positive cells to different leucocytes subsets count or erythrocyte count present.

LIMITATIONS

- Blood samples should be stored at 18-22°C and tested within 24 hours after they were obtained.
- It is advisable to acquire stained samples as soon as possible to optimise results. Non-viable cells may show unspecific staining. Prolonged exposure of whole blood samples to lytic reagents may cause white cell destruction and targeted population cell loss.
- When using whole blood procedures some red blood cells may not lyse, for instance if there are nucleated red blood cells or if abnormal protein concentration and haemoglobinopathies are observed. This may cause misleading results since unlysed red blood cells are counted as leucocytes.
- Results obtained by flow cytometry may be erroneous if cytometer laser is misaligned or if gates are incorrectly set.
- Each laboratory should establish a normal range for CD59+ cells using its own test conditions.
- Certain patients may present special problems due to altered or very low number of a certain cellular population.
- Cells separated from whole blood by means of density gradients may not have the same relative concentration as in whole blood. This may be relatively insignificant in individuals with normal white blood cell counts. In leucopenic patients, the selective loss of specific subsets may affect determination accuracy.
- Knowledge of antigen normal expression pattern and its relation to other relevant antigens is paramount to carry out an adequate analysis^(2, 11).
- Abnormal states of health are not always represented by abnormal percentages of certain leucocyte populations. An individual in an abnormal state of health may show the same leucocyte percentage as a healthy person.

QUALITY CONTROL

- Pipettes precision and cytometer calibration should be verified to obtain optimal results.
- In multicolour panels, fluorochromes emit in wavelengths that can show certain spectral overlap which must be corrected by electronic compensation. Optimal compensation levels can be established by analysing cells from healthy individuals stained with mutually exclusive monoclonal antibodies conjugated with appropriate fluorochromes.
- Non-specific binding of the antibody can be evaluated using an appropriated isotype control tube.







REFERENCES

1. Orfao A, González de Buitrago JM. La citometría de flujo en el laboratorio clínico. Sociedad española de bioquímica clínica y patología Molecular 1995.
2. Zhang H, Yu J, Chen S, Morgan BP, Abagyan R, Tomlinson S. Identification of the Individual Residues that Determine Human CD59 Species Selective Activity. J. of Biolog. Chemistry 274: 10969–10974 (1999)
3. Zalman LS, Brothers MS, Muller-Eberhard HJ. Isolation of homologous restriction factor from human urine. Immunochemical properties and biologic activity. J Immunol 143:1943–1947 (1989)
4. Gorter A, Meri S. Immune evasion of tumor cells using membrane-bound complement regulatory proteins. Immunol Today 20: 576-582(1999)
5. Lida Y, Takeda J, Miyata T, Inoue N, Nishimura J, Kitani T, Maeda K, and Kinoshita T. Characterization of genomic PIG-A gene: a gene for glycosylphosphatidylinositol-anchor biosynthesis and paroxysmal nocturnal hemoglobinuria. Blood 83: 3126–3131 (1994)
6. Menéndez P, et al. Comparison between a lyse-and-then-wash method and a lyse-non-wash technique for the enumeration of CD34+ hematopoietic progenitor cells. Cytometry (Comm. Clin. Cytometry) 34: 264-271 (1998)
7. Gratama JW, Menéndez P, Kraan J, Orfao A. Loss of CD34+ hematopoietic progenitor cells due to washing can be reduced by the use of fixative-free erythrocyte lysing reagents. J Immunol. Methods 239: 13-23 (2000)
8. Protection of Laboratory Workers from occupationally acquired infections. Second edition; approved guideline (2001). Villanova PA: National Committee for Clinical Laboratory Standards; Document M29-A2.
9. Procedures for the collection of diagnostic blood specimens by venipuncture- approved standard; Fifth edition (2003). Wayne PA: National Committee for Clinical Laboratory Standards; Document H3-A5.
10. Clinical applications of flow cytometry: Quality assurance and immunophenotyping of lymphocytes; approved guideline (1998). Wayne PA: National Committee for Clinical Laboratory Standards; Document H42-A.
11. Richards SJ, Barnett D. The role of flow cytometry in the diagnosis of paroxysmal nocturnal hemoglobinuria in the clinical laboratory. Clin Lab Med 27: 577-90 (2007).

WARRANTY

This product is warranted only to conform to quantity and label specifications. There are no warranties that extend beyond the description on product label. Cytognos' sole liability is limited to either product replacement or refund of the purchase price.

EXPLANATION OF SYMBOLS

	Use by (use by YYYY-MM)
	Storage temperature limitation
	Consult instruction for use
RUO	For research use only
	Batch code
	Code number
	Manufacturer

PRODUCED BY CYTOGNOS SL

Polígono La Serna, Nave 9
37900 Santa Marta de Tormes
Salamanca (Spain)
Phone: + 34-923-125067
Fax: + 34-923-125128

Ordering information: admin@cytognos.com
Technical information: support@cytognos.com

www.cytognos.com