Technical Data Sheet

APC-Cy™7 Mouse Anti-Human CD69

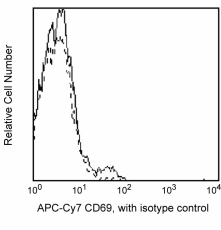
Product Information

| Material Number: |
|------------------|
| Alternate Name: |
| Size: |
| Vol. per Test: |
| Clone: |
| Isotype: |
| Reactivity: |
| Workshop: |
| Storage Buffer: |

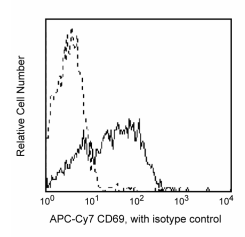
560912 Very Early Activation Antigen 25 tests 5 μl FN50 Mouse IgG1, κ QC Testing: Human IV A091 Aqueous buffered solution containing BSA and ≤0.09% sodium azide.

Description

Reacts with a 28/34 kDa dimeric glycoprotein expressed early during activation of lymphocytes and monocytes. FN50 monoclonal antibody labels NK cells and most lymphocytes of the follicular mantle and perifollicular/interfollicular zone as well as intragerminal center T cells of lymph nodes and tonsils.



Profile of resting peripheral blood lymphocytes analyzed by flow cytometry



Profile of PHA-stimulated (24 hour) peripheral blood lymphocytes analyzed by flow cytometry

Preparation and Storage

The monoclonal antibody was purified from tissue culture supernatant or ascites by affinity chromatography. The antibody was conjugated with APC-Cy7 under optimum conditions, and unconjugated antibody and free APC-Cy7 were removed. Store undiluted at 4°C and protected from prolonged exposure to light. Do not freeze.

Application Notes

| Application | | | |
|------------------|--|------------------|---------|
| Flow cytometry | | Routinely Tested | |
| Suggested Compar | ion Products | | |
| Catalog Number | Name | Size | Clone |
| 557873 | APC-Cy TM 7 Mouse IgG1, κ Isotype Control | 0.1 mg | MOPC-21 |

Product Notices

- 1. This reagent has been pre-diluted for use at the recommended Volume per Test. We typically use 1×10^{6} cells in a 100-µl experimental sample (a test).
- 2. Since applications vary, each investigator should titrate the reagent to obtain optimal results.
- 3. Please refer to www.bdbiosciences.com/pharmingen/protocols for technical protocols.
- 4. For fluorochrome spectra and suitable instrument settings, please refer to our Fluorochrome Web Page at www.bdbiosciences.com/colors.

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| of any patents. BL use of our produc product or as a co written authoriza For Research Use (| D Biosciences will ne ts. Purchase does n mponent of anoth- tion of Becton Dick Only. Not for use in | ot be held responsi ot include or carry er product. Any us inson and Compan diagnostic or thera | ble for patent infrin any right to resell or e of this product oth y is strictly prohibite apeutic procedures. | gement or other vio r transfer this produc per than the permitte d. | e the above product in violation lations that may occur with the ct either as a stand-alone ed use without the express 11 BD |

- 5. Please observe the following precautions: Absorption of visible light can significantly alter the energy transfer occurring in any tandem fluorochrome conjugate; therefore, we recommend that special precautions be taken (such as wrapping vials, tubes, or racks in aluminum foil) to prevent exposure of conjugated reagents, including cells stained with those reagents, to room illumination.
- 6. APC-Cy7 tandem fluorochrome emission is collected in a detector for fluorescence wavelengths of 750 nm and higher.
- 7. APC-Cy7 is a tandem fluorochrome composed of Allophycocyanin (APC), which is excited by laser lines between 595 and 647 nm and serves as an energy donor, coupled to the cyanine dye Cy7TM, which acts as an energy acceptor and fluoresces at 780 nm. BD Biosciences Pharmingen has maximized the fluorochrome energy transfer in APC-Cy7, thus maximizing its fluorescence emission intensity, minimizing residual emission from APC, and minimizing required electronic compensation in multilaser-laser flow cytometry systems. Note: Although every effort is made to minimize the lot-to-lot variation in residual emission from APC, it is strongly recommended that every lot be tested for differences in the amount of compensation required and that individual compensation controls are run for each APC-Cy7 conjugate.
- 8. Warning: Some APC-Cy7 conjugates show changes in their emission spectrum with prolonged exposure to formaldehyde. We recommend that you analyze fixed samples within four hours.
- 9. Cy is a trademark of Amersham Biosciences Limited. This conjugated product is sold under license to the following patents: US Patent Nos. 5,486,616; 5,569,587; 5,569,766; 5,627,027.
- 10. This product is subject to proprietary rights of Amersham Biosciences Corp. and Carnegie Mellon University and made and sold under license from Amersham Biosciences Corp. This product is licensed for sale only for research. It is not licensed for any other use. If you require a commercial license to use this product and do not have one return this material, unopened to BD Biosciences, 10975 Torreyana Rd, San Diego, CA 92121 and any money paid for the material will be refunded.
- 11. Caution: Sodium azide yields highly toxic hydrazoic acid under acidic conditions. Dilute azide compounds in running water before discarding to avoid accumulation of potentially explosive deposits in plumbing.
- 12. Source of all serum proteins is from USDA inspected abattoirs located in the United States.

References

Beavis AJ, Pennline KJ. Allo-7: a new fluorescent tandem dye for use in flow cytometry. Cytometry. 1996; 24(4):390-395. (Biology)

Knapp W, Dorken B, Rieber EP, et al, ed. *Leucocyte Typing IV*. New York: Oxford University Press; 1989:1-1208. (Clone-specific) Roederer M, Kantor AB, Parks DR, Herzenberg LA. Cy7PE and Cy7APC: bright new probes for immunofluorescence. *Cytometry*. 1996; 24(3):191-197. (Biology) Schlossman S, Boumell L, et al, ed. *Leucocyte Typing V*. New York: Oxford University Press; 1995. (Biology)