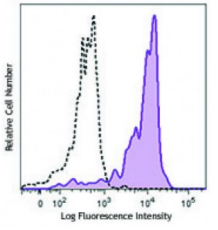
Product Data Sheet

PE anti-human CD191 (CCR1)

| Catalog # / Size: | 2414520 / 100 tests 2414515 / 25 tests |
|-----------------------|--|
| Clone: | 5F10B29 |
| Isotype: | Mouse IgG1, κ |
| Immunogen: | Human CCR1 transfected cells |
| Reactivity: | Human |
| Preparation: | The antibody was purified by affinity chromatography and conjugated with PE under optimal conditions. The solution is free of unconjugated PE and unconjugated antibody. |
| Formulation: | Phosphate-buffered solution, pH 7.2, containing 0.09% sodium azide and 0.2% (w/v) BSA (origin USA). |
| Concentration: | Lot-specific |



Human peripheral blood monocytes were stained with CD191 (CCR1) PE (clone 5F10B29, filled histogram) or mouse IgG1, κ PE isotype control (open histogram).

Applications:

| Applications: | Flow Cytometry |
|------------------------|--|
| Recommended Usage: | Each lot of this antibody is quality control tested by immunofluorescent staining with flow cytometric analysis. For flow cytometric staining, the suggested use of this reagent is 5 microL per million cells or 5 microL per 100 microL of whole blood. It is recommended that the reagent be titrated for optimal performance for each application. |
| Application Notes: | This clone does not cross-react with human CCR4, CCR5, CCR6, CCR7, or CCR8. |
| Description: | CD191, also known as CCR1, is a 41 kD, G-protein coupled receptor expressed predominantly by monocytes. CCR1 is also expressed by a subset of T cells and eosinophils. CCR1 positive cells can migrate in response to a CCL3 and CCL5 gradient. CCR1 knock-out studies suggest that this molecule plays an important role in inflammation and susceptibility to viruses and parasites. |
| Antigen References: | 1. Su SB, <i>et al.</i> 1996. <i>J. Leuko. Biol.</i> 60:658. 2. Su S, <i>et al.</i> 1997. <i>Blood.</i> 90:605. 3. Ayehunie S, <i>et al.</i> 1997. <i>Blood.</i> 90:1379. |

4. Gerard C, et al. 1997. J. Clin.

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