## Brilliant Violet 421™ anti-human CD200R

Catalog # / 2246570 / 100 tests

Size: 2246565 / 25 tests

Clone: OX-108

Isotype: Mouse IgG1, ĸ

Human CD200R full length fusion Immunogen:

protein

Reactivity: Human, Non-human primate

Preparation: The antibody was purified by affinity

> chromatography and conjugated with Brilliant Violet 421™ under optimal conditions. The solution is free of unconjugated Brilliant Violet 421™

and unconjugated antibody.

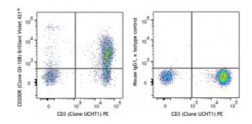
Phosphate-buffered solution, pH 7.2, Formulation:

containing 0.09% sodium azide and

BSA (origin USA).

Workshop Number: **HCDM** listed

**Concentration:** Lot-specific



Human perpipheral blood lymphocytes were stained with CD3 (clone UCHT1) PE and CD200R (clone OX-108) Brilliant Violet 421™ (left) or Mouse IgG1, κ Brilliant Violet 421TM isotype

control (right).

## **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 μl per million cells in 100 μl staining volume or 5 µl per 100 µl of whole blood.

Brilliant Violet 421™ excites at 405 nm and emits at 421 nm. The standard bandpass filter 450/50 nm is recommended for detection. Brilliant Violet 421<sup>™</sup> is a trademark of Sirigen Group Ltd.

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**Application** 

Notes:

Additional reported applications (for the relevant formats) include: The 5E8/CXCR2 antibody is useful for immunofluorescent staining and flow

cytometric analysis of CXCR2 expression.

**Application** References:

1. Wright GJ, et al. 2003. J. Immunol. 171:3034.

**Description:** CD200R, also known as OX2R, is a membrane glycoprotein with up to 70% of

its weight derived from N-linked glycosylation. CD200R is expressed primarily by monocytes and neutrophils, but also by other leukocytes including T cells and mast cells. The interaction between CD200 and CD200R may contribute to pathways that suppress and limit macrophage

induced inflammatory damage in tissues.

Antigen References:

Wright GJ, et al. 2003. J. Immunol. 171:3034.
Gorczynski R, et al. 2004. J. Immunol. 172:7744.