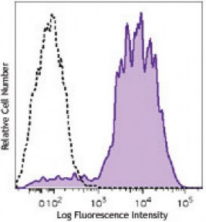
Product Data Sheet

Brilliant Violet 711[™] anti-mouse/human CD11b

Catalog # / Size:	1106205 / 125 μl 1106210 / 50 μg	
Clone:	M1/70	we Cell Number
Isotype:	Rat IgG2b, κ	
Immunogen:	C57BL/10 splenocytes	
Reactivity:	Human	
Preparation:	The antibody was purified by affinity chromatography and conjugated with Brilliant Violet 711 [™] under optimal conditions. The solution is free of unconjugated Brilliant Violet 711 [™] and unconjugated antibody.	Relat
Formulation:	Phosphate-buffered solution, pH 7.2, containing 0.09% sodium azide and BSA (origin USA).	C! w M
Concentration:	microg sizes: 0.2 mg/ml microL sizes: lot-specific	sh Ce



C57BL/6 mouse bone marrow cells were stained with CD11b (clone M1/70) Brilliant Violet 711[™]. Data shown was gated on the myeloid cell population.

Applications:

- Applications: Flow Cytometry
- **Recommended** Usage: Each lot of this antibody is quality control tested by immunofluorescent staining with flow cytometric analysis. For immunofluorescent staining using the microg size, the suggested use of this reagent is ≤ 0.4 microg per million cells in 100 microL volume. For immunofluorescent staining using the microL size, 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.

Brilliant Violet 711[™] excites at 405 nm and emits at 711 nm. The bandpass filter 710/50 nm is recommended for detection, although filter optimization may be required depending on other fluorophores used. **Be sure to verify that your cytometer configuration and software setup are appropriate for detecting this channel.** Refer to your instrument manual or manufacturer for support. Brilliant Violet 711[™] is a trademark of Sirigen Group Ltd.

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Application Notes:
 Additional reported applications (for relevant formats of this clone) include: immunoprecipitation^{1,4}, *in vitro* blocking^{3,9,12}, depletion^{2,8}, immunofluorescence microscopy^{6,7,10}, and immunohistochemistry of acetone-fixed frozen sections^{5,11-13} and paraffin sections²⁸. The LEAF™ purified antibody (Endotoxin <0.1 EU/µg, Azide-Free, 0.2 µm filtered) is recommended for functional assays (Cat. No. 101231). For *in vivo* studies or highly sensitive assays, we recommend Ultra-LEAF™ purified antibody (Cat. No. 101248) with a lower endotoxin limit than standard LEAF™ purified antibodies (Endotoxin <0.01 EU/microg).

Application References:	 Springer T, <i>et al.</i> 1978. <i>Eur. J. Immunol.</i> 8:539. (IP) Ault K and Springer T. 1981. <i>J. Immunol.</i> 126:359. (Deplete) Springer TA, <i>et al.</i> 1982. <i>Immunol. Rev.</i> 68:171. (Block) Ho MK and Springer TA. 1983. <i>J. Biol. Chem.</i> 258:2766. (IP) Flotte TJ, <i>et al.</i> 1983. <i>Am. J. Pathol.</i> 111:112. (IHC) Noel GJ, <i>et al.</i> 1990. J. Clin. Invest. 85:208. (IF) Allen LA and Aderem A. 1996. <i>J. Exp. Med.</i> 184:627 (IF) D'Amico A and Wu L. 2003. <i>J. Exp. Med.</i> 198:293. (Deplete) Brickson SJ, <i>et al.</i> 2003. <i>Appl Physiol.</i> 95:969. (Block) Clatworthy MR and Smith KG. 2004. <i>J. Exp. Med.</i> 199:717. (IF) Hata H, <i>et al.</i> 2002. <i>J. Immunol.</i> 168:3088. (IHC) Zhang Y, <i>et al.</i> 2003. <i>J. Exp. Med.</i> 197:121. (Block, FC) Iwasaki A and Kelsall BL. 2001. <i>J. Immunol.</i> 166:4884 (IHC, FC) Tailleux L. 2003. <i>J. Exp. Med.</i> 197:121. (Block, FC) Olver S, <i>et al.</i> 2006. <i>J. Immunol.</i> 176:2872. (FC) PubMed Ponomarev ED, <i>et al.</i> 2006. <i>J. Immunol.</i> 176:1402. (FC) Dzhagalov I, <i>et al.</i> 2007. <i>Nature Immunol.</i> 8:753. Rasmussen JW, <i>et al.</i> 2008. <i>J. Immunol.</i> 180:609. PubMed Napimoga MH, <i>et al.</i> 2008. <i>J. Immunol.</i> 180:609. PubMed Kim DD, <i>et al.</i> 2008. <i>Blood</i> 112:1109. PubMed Kim DD, <i>et al.</i> 2008. <i>Blood</i> 112:480. PubMed Korian LA, <i>et al.</i> 2009. <i>Cancer Res.</i> 69:3086. (FC) PubMed Norian LA, <i>et al.</i> 2010. <i>J. Immunol.</i> 184:573. PubMed Norian LA, <i>et al.</i> 2010. <i>J. Immunol.</i> 184:573. PubMed Norian LA, <i>et al.</i> 2010. <i>J. Immunol.</i> 184:573. PubMed Korian LA, <i>et al.</i> 2010. <i>J. Immunol.</i> 184:573. PubMed Korian LA, <i>et al.</i> 2010. <i>J. Immunol.</i> 184:573. PubMed Korian LA, <i>et al.</i> 2010. <i>J. Immunol.</i> 184:573. PubMed
Description:	CD11b is a 170 kD glycoprotein also known as α M integrin, Mac-1 α subunit, CR3, and Ly-40. CD11b is a member of the integrin family, primarily expressed granulocytes, monocytes/macrophages, dendritic cells, NK cells, and subsets and B cells. CD11b non-covalently associates with CD18 (β 2 integrin) to f Mac-1. Mac-1 plays an important role in cell-cell interaction by binding its ligation.

ed on s of T form Mac-1. Mac-1 plays an important role in cell-cell interaction by binding its ligands ICAM-1 (CD54), ICAM-2 (CD102), ICAM-4 (CD242), iC3b, and fibrinogen.

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1. Barclay A, et al. 1997. The Leukocyte Antigen FactsBook Academic Press. Antigen

- 2. Springer TA. 1994. Cell 76:301.
- References:
- 3. Coxon A, et al. 1996. Immunity 5:653.