

Product code	65-165
Size	100 µg
Storage	-20°C
Concentration	1.0 mg/ml
Buffer	PBS- with 50% glycerol
Purity	Purified IgG fraction with protein A from hybridoma cell culture medium
Immunogen	Human Influenza B Virus strain Nagasaki/1/87, one of the strains of B/Victoria group.
Isotype	mouse IgG2ак
Reactivity	According to Ref.1 during epidemic in Osaka 1996-97, 10B8 antibody reacted with HA protein
	of all Influenza B virus isolates belonging to Victoria group tested (73 strains) and none of
	clinical 27 isolates belonging to Yamagata group as examined by PAP staining. It also reacts
	with Victoria group vaccine strains; Shangdong/7/1997, Malasia/2506/2004. However, note
	that HA changes during passages and may change reactivity to this antibody. By western
	blotting, reactivity with B/Malasia/2506/2004 and B/Massachusetts/2/2012 was tested
	positive. No cross reactivity with any strains of influenza A virus.
Special notes	N/A
Application	1. Western blotting (1/500~1/1,000 dilution)
	2. Immunofluorescent and Immunocytochemical staining (1/100~1/200 dilution)
	3. Immunoprecipitation (1/200 dilution)
	4. Neutralization of infectivity (NT) (assay dependent)
	5. Hemagglutination Inhibition (HI) (assay dependent)
	6. ELISA (assay dependent)
Background	Hemagglutinin (HA) binds to sialic acid-containing receptors on the cell surface, bringing
	about the attachment of the virus particle to the cell. Plays a major role in the determination
	of host range restriction and virulence. Class I viral fusion protein. Responsible for
	penetration of the virus into the cell cytoplasm by mediating the fusion of the membrane of
	the endocytosed virus particle with the endosomal membrane. Low pH in endosomes induce
	an irreversible conformational change in HA2, releasing the fusion hydrophobic peptide.
	Several trimers are required to form a competent fusion pore.
	Post-translational modification <sup>i</sup>
	HAo is consists of 584 amino acids with molecular mass of 63,275. In natural infection,
	inactive HA is matured into HA1 and HA2. By the sequence similarity it is indicated to be
	palmitoylated.
Data Link	UniProtKB <u>P03460</u> Influenza B/Lee/1940 HA protein.
Please note: All products are FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC	
PROCEDURES. NOT FOR MILITARY USE.	

## Anti-Influenza A Virus Nucleoprotein antibody, mouse monoclonal (10B8)



Data Images: 65-165 Anti-Influenza B Virus HA antibody, mouse monoclonal (10B8)



Fig.1 Immunofluorescence assay of MDCK (canine kidney ) cells infected with Influenza B virus, using anti-Influenza B virus HA antibody (clone 10B8).

Samples were taken at 24 hours post-infection. Anti-Influenza B Virus HA antibody (clone 10B8) efficiently detected HA in the B/Malasia/2506/2004 virus (Victorial group) but not in B/Florida/4/2006 virus (Yamagata group) infected MDCK cells. The cells were fixed with 4% paraformaldehyde in PBS- and permeabilized with 0.1% Triton X-100 in PBS. The bound antibody was visualized by a further reaction with an Alexa Fluor 488-conjugated secondary antibody.

Images on the left are mock-infected MDCK cells as negative control.





- 1. B/Mie/1/1993
- 2. B/Johannes Burg/5/1999
- 3. B/Florida/4/2006
- 4. B/Lee/1940
- 5. B/Florida/4/2006



- 6. Shandong/7/97
- 7. B/Malasia/2506/2004
- 8. B/Massachusetts/2/2012

First antibody was used at 1/500 dilution and as 2<sup>nd</sup> antibody, HRP-conjugated goat anti-mouse IgG antibody was used at 1/10,000 dilution. Positions of marker proteins are indicated in kDa on the left. **Clone 10B8 recognizes an epitope on HA2 region**.

 $\ensuremath{\textbf{References}}$  : This product has been used in the following publication

- 1.Nakagawa N. et al. Rapid detection and identification of two lineages of influenza B strains with monoclonal antibodies. J Virol Methods. 1999;79:113-2 ICC, IP
- Nakagawa N. et al. Heterogeneity of Influenza B Virus Strains in One Epidemic Season Differentiated by Monoclonal Antibodies and Nucleotide Sequences. <u>J Clin Microbiol.</u> 2000;38:3467-9. HI, NT
- 3. Nakagawa N. et al. Variation of the Conserved Neutralizing Epitope in Influenza B Virus Victoria Group Isolates in Japan. <u>J Clin Microbiol.</u> 2005 ;43:4212-4. **HI**
- 4. Nakagawa N. et al. Discovery of the neutralizing epitope common to influenza B virus victoria group isolates in Japan. J Clin Microbiol. 2006;44:1564-6. **HI**