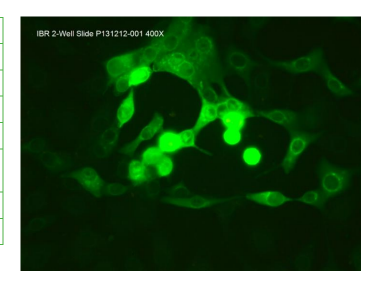


CERTIFICATE OF ANALYSIS

Bovine Herpesvirus Type 1 (BHV-1/IBR)

FA Control Slide

Catalog No.:	SLD-FAC-IBR
Size:	2 Well
Well Capacity:	50 μΙ
Lot:	P131212-001
Expiration:	06 January 2018
Agent:	Bovine Herpesvirus Type 1 (BHV-1/IBR)
Strain:	Kentucky
Cell Culture Substrate:	MDBK cells



Description:

Wells contain virus-infected cell cultures grown on the surface of Teflon-masked slide. They are supplied fixed and unstained in moisture-free foil pouches. Each slide contains one positive and one negative cell culture well. The positive well contains both positive and negative cells. The positive cells usually total no more than 30% of the cells in the well so that good contrast may be seen.

Quality Control Method:

Direct FA using BHV-1/IBR FA conjugate (catalog no. CJ-F-IBR-1ML or 10ML).

Specific Reaction: 3-4+ fluorescence on the positive well and negative on the negative well

with no background. There are 0 to 20 positive cells and/or plagues per

high-power field.

Other Comments: NA

Pattern Of Fluorescence:

Primarily undifferentiated cytoplasmic with some nuclear fluorescence, especially in rounded cells and degenerating cells in plaques with acellular centers.

Intended Use:

For positive and negative control of direct or indirect FA tests for BHV-1/IBR.

Storage:

Store sealed in foil pouch at -20°C. Avoid self-defrosting freezers.

References: NA

Recommended Staining Procedure for Indirect FA:

- 1. Warm slide to room temperature before removing from foil pouch.
- 2. Place diluted serum on the designated wells. Dilute serum in serum diluting buffer, pH 7.2 (catalog no. FASDB-100ML) however if high background due to anti-bovine IgG activity is present it may be advisable to use SSDB-100ML.
- 3. Incubate slide in humid chamber at 37°C for 30 minutes.
- 4. Using a wash bottle, gently rinse slide briefly in FA rinse buffer, pH 9.0 (catalog no. FARB-4X) and then soak for 10 minutes in FA rinse buffer, pH 9.0.
- 5. Drain slide and dry around wells by pressing blotter (included in pouch) to front surface. Place labeled anti-IgG or IgM on the wells.
- 6. Incubate as in step 3.
- 7. Rinse as in step 4.
- 8. Drain slide and dry back and edges with a paper towel. Do not allow stained surface to dry. Do not rinse with water.
- 9. Mount with mounting fluid [glycerol/FA rinse buffer, pH 9.0, (50/50)] (catalog no. FAMF-10ML) and view with good quality fluorescence microscope at 100X-250X. Confirmation may be made at 400X.

Recommended Staining Procedure for Direct FA:

- 1. Warm slide to room temperature before removing from foil pouch.
- 2. Place direct FA conjugate on the designated wells.
- 3. Incubate slide in humid chamber at 37°C for 30 minutes.
- 4. Using a wash bottle, gently rinse slide briefly in FA rinse buffer, pH 9.0 (catalog no. FARB-4X) and then soak for 10 minutes in FA rinse buffer, pH 9.0.
- 5. Drain slide and dry back and edges with a paper towel. Do not allow stained surface to dry. Do not rinse with water.
- 6. Mount with mounting fluid [glycerol/FA rinse buffer, pH 9.0, (50/50)] (catalog no. FAMF-10ML) and view with good quality fluorescence microscope at 100X-250X. Confirmation may be at 400X.

Serum Diluting Buffer (pH 7.2):*

-	Na ₂ HPO ₄	1.19 gm
-	NaH ₂ PO ₄	0.22 gm
-	NaCl	8.55 gm
-	BSA	10.0 gm
	DI/dH ₂ O	

^{*}This recipe makes 1 liter. If you need less, adjust recipe accordingly. Store at 2-7°C. Add 0.09% NaN₃ if diluted serum is not going to be used within one week.

4X FA Rinse Buffer (pH 9.0):

-	Na ₂ CO ₃ 11.4 gm
-	NaHCO333.6 gm
-	NaCl8.5 gm
-	DI/dH ₂ OQ.S. to 1 liter

Final pH should be 9.0-9.5. This is a 4X concentrate and should be diluted 1/4 with DI/distilled water for use as a working buffer. Keep in a tightly stoppered container at room temperature. MOUNTING FLUID is made by mixing glycerol and FA rinse buffer, pH 9.0, in equal proportions.