

 Telephone:
 + 1 (509) 334-5815

 Fax:
 + 1 (509) 332-5356

 E-mail:
 vmrd@vmrd.com

 Web site:
 http://www.vmrd.com

Certificate of Analysis

BOVINE RESPIRATORY SYNCYTIAL VIRUS (BRSV)

FA Substrate Slide

CATALOG NO.: SLD-IFA-BRSV SIZE: 12 well LOT: P080808-001 EXPIRATION: 04 April 2031

AGENT: Bovine Respiratory Syncytial Virus (BRSV) Cell Culture Substrate: MDBK cells Virus Strain: TN-511



QUALITY CONTROL METHOD: Indirect FA using VMRD BRSV Positive Control (catalog no.PC-IFA-BRSV), BRSV Negative Control (catalog no. NC-IFA-BRSV), and Anti-Bovine IgG_{1,2} FITCPolyclonal Conjugate (catalog no. CJ-F-BOVG-AP-1ML or CJ-F-BOVG-AP-10ML).Specific Reaction:2-4+ fluorescence with the positive control and negative with the negative control. No background.Other Reactions or Comments:0-25 infected cells per high power field.

PATTERN OF FLUORESCENCE: Granular cytoplasmic fluorescence with some small syncytia.

INTENDED USE: Useful in detecting viral antibody by IFA. May be used to differentiate antibody class (IgG or IgM) with suitable quality fluoresceinated second antibody conjugate.

STABILITY: Foil-pouch sealed slides are stable when stored at -20°C. Avoid self-defrosting freezers.

DESCRIPTION: Slides are virus-infected cell cultures grown on the surface of Teflon-masked slides. They are supplied fixed and unstained in moisture-free foil pouches. All wells contain 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.

FOR IN VITRO LABORATORY USE ONLY.

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RECOMMENDED STAINING PROCEDURE FOR INDIRECT FA:

- 1. Warm slide to room temperature before removing from foil pouch.
- Place 50 μl diluted serum on the designated wells. Dilute serum in serum diluting buffer, pH 7.2 (catalog no. 210-93-SB).
- 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. 210-90-RB) 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 50 µl 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 <u>back and edges</u> 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. 210-92-MF) 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 50 µl of 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. 210-90-RB) and then soak for 10 minutes in FA rinse buffer, pH 9.0.
- 5. Drain slide and dry <u>back and edges</u> 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. 210-92-MF) 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	
- DI/dH ₂ O	Q.S. to 1 liter

* This recipe makes 1 liter. If you need less, adjust recipe accordingly. Store at 4°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
- NaHCO ₃	
- NaCl	8.5 gm
- DI/dH ₂ O	Q.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.