

Diagnostics

Critical components of an effective FMD Response are appropriate sample collection and diagnostic testing, which are needed to detect and confirm the presence of FMD for critical reporting and surge capacity preparedness. These tools are necessary for investigations, surveillance, and tracebacks during an outbreak to determine the extent of disease spread, also will continue to be used after the outbreak to determine when transmission has stopped. Additionally, diagnostic surveillance serves as the basis for declaring stop movement at the beginning of an outbreak and disease freedom from FMD at the end.

Goals

During a suspected or actual FMD outbreak, the key goals of response are the following:

- ◆ Meet the surge requirements for diagnostic testing at specific intervals.
- ◆ Report all diagnostic test results to appropriate personnel and information management systems as soon as possible or in order of investigation priority level, after completion.

Foreign Animal Disease Investigation Policy and Procedures

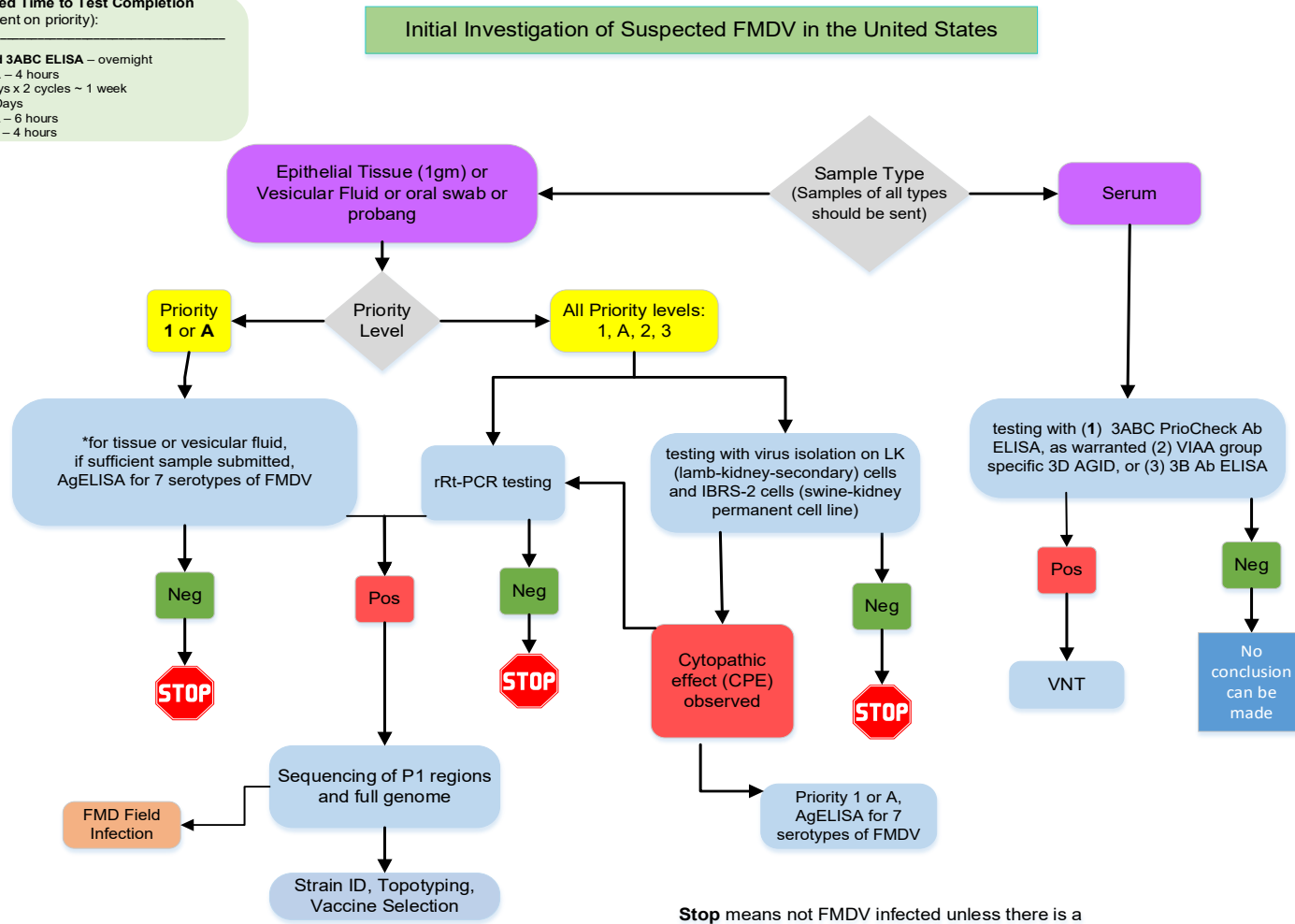
For information regarding FMD (or other foreign animal diseases) investigation policy and procedures see [Veterinary Services \(VS\) Guidance Document 12001.4](#) and the [Foreign Animal Disease Investigation Manual \(FAD PReP Manual 4-0\)](#).

Testing for suspect FMD virus in the United States is done at the National Veterinary Services Laboratories—Foreign Animal Disease Diagnostic Laboratory (NVSL FADDL). Preliminary testing may be ongoing at a National Animal Health Laboratory Network (NAHLN) lab, but confirmation is only done by NVSL FADDL. If FMD virus (FMDV) is detected, then sequencing will be completed to reveal the strain, topotype, and to conduct possible vaccine matching.

Diagnostic Flowchart for Initial Investigation of FMD

Estimated Time to Test Completion
 (dependent on priority):

- VIAA and 3ABC ELISA – overnight
- 3B ELISA – 4 hours
- VI – 3 days x 2 cycles ~ 1 week
- VNT - 3 Days
- AgELISA – 6 hours
- rRT-PCR – 4 hours



Stop means not FMDV infected unless there is a circumstantial reason to request additional samples and conduct additional diagnostic testing.

Abbreviations	rRT-PCR: real-time reverse transcriptase polymerase chain reaction
Ag: antigen	VI: virus isolation
AGID: agar-gel immunodiffusion	VIAA: virus infection association antigen
ELISA: enzyme-linked immunosorbant assay	VNT: virus neutralization test

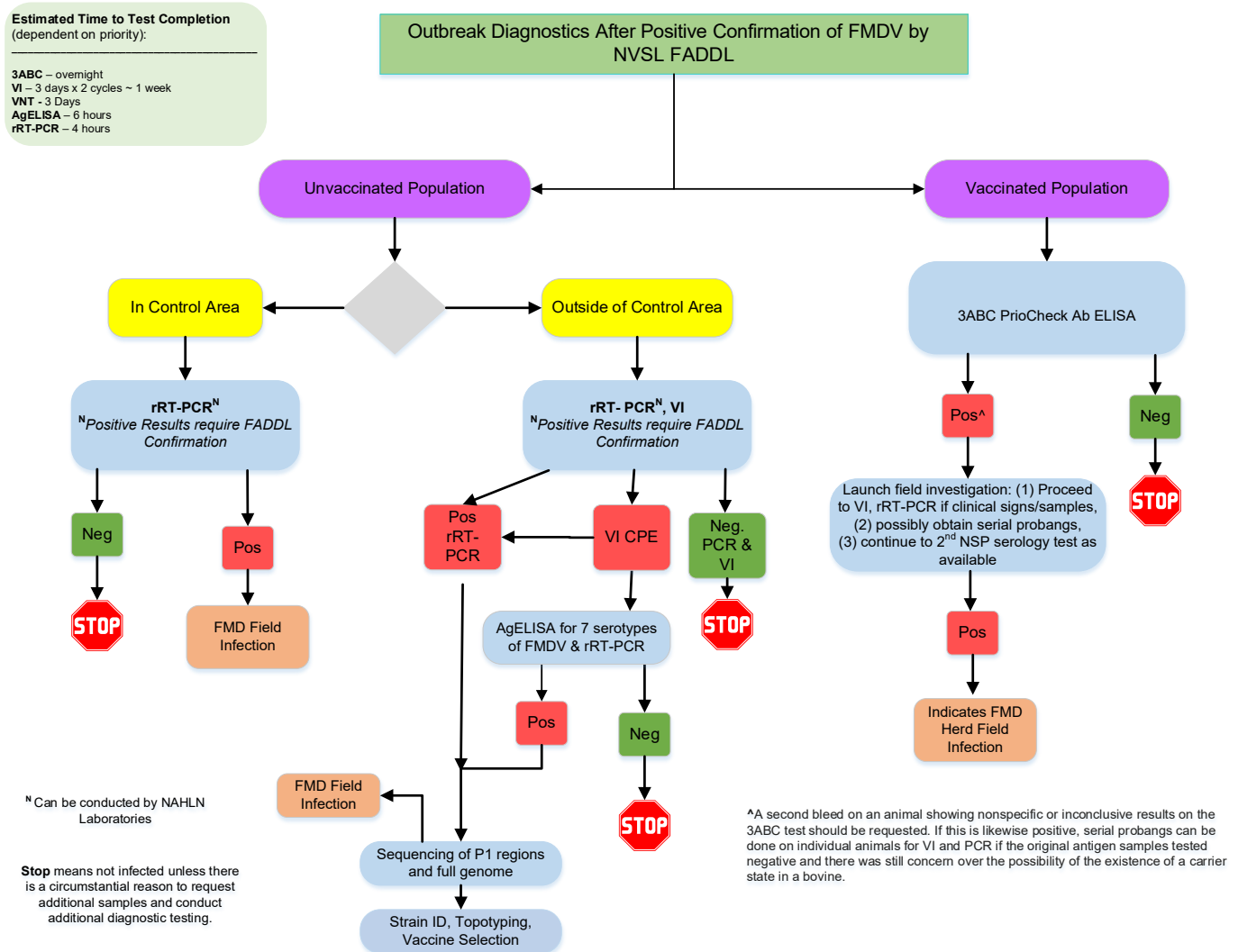
USDA APHIS Veterinary Services • National Preparedness and Incident Coordination (NPIC)

4700 River Road Unit 41 • Riverdale, MD 20737

Diagnostics During a FMD Outbreak

After NVSL FADDL confirmation of FMD on a premises (index case), subsequent [swab samples](#) for rRT-PCR may be sent to NAHLN laboratories. Incident Command will provide specific instructions regarding the direction and collection of samples, which is likely to change as the outbreak evolves. In all cases 1) NVSL FADDL will confirm the index case, 2) presumptive positive samples (on a rRT-PCR) from outside an established Control Area will be tested and confirmed by NVSL FADDL, and 3) NVSL FADDL will receive samples routinely from inside the Control Area to monitoring for the disease status in susceptible species.

Diagnostic Flowchart after Positive Confirmation of FMD in the United States



^N Can be conducted by NAHLN Laboratories

Stop means not infected unless there is a circumstantial reason to request additional samples and conduct additional diagnostic testing.

^AA second bleed on an animal showing nonspecific or inconclusive results on the 3ABC test should be requested. If this is likewise positive, serial probangs can be done on individual animals for VI and PCR if the original antigen samples tested negative and there was still concern over the possibility of the existence of a carrier state in a bovine.

Diagnostic Surge Capacity & Reporting Notification

In an FMD outbreak, additional resources—such as personnel and materials—may be needed for sample collection. Additional capacity may also be required for laboratory sample testing. NAHLN labs have the capability to conduct rRT-PCR tests. Ideally, NAHLN laboratories will also have the capability to conduct 3ABC ELISA serology testing and to differentiate between infected and vaccinated animals (DIVA testing) on a herd basis if required in the outbreak. Surge capacity can help facilitate a rapid response as well as continuity of business for non-infected premises. Should NVSL FADDL or the State NAHLN laboratory become overwhelmed by diagnostic testing requirements, NAHLN labs from across the country may provide surge capacity. Individual laboratories have

independent protocols on how to manage personnel if a surge is required. See the NAHLN [website](#) for list of approved NAHLN labs.

- ◆ Cases of clinical illness that are found to be presumptive positive by NVSL-FADDL, based on the current case definition, will be reported to the affected States, other States, Tribal Nations, industry, other Federal agencies, trading partners, and the World Organization for Animal Health (OIE).
- ◆ Appropriate Federal-State-Tribal-industry response and containment measures will be initiated during FMD investigations.
- ◆ For more information reporting and notification, see APHIS VS Guidance Document 12001 and the *FAD Investigation Manual (FAD PReP Manual 4-0)* which can be found at <http://www.aphis.usda.gov/fadprep>