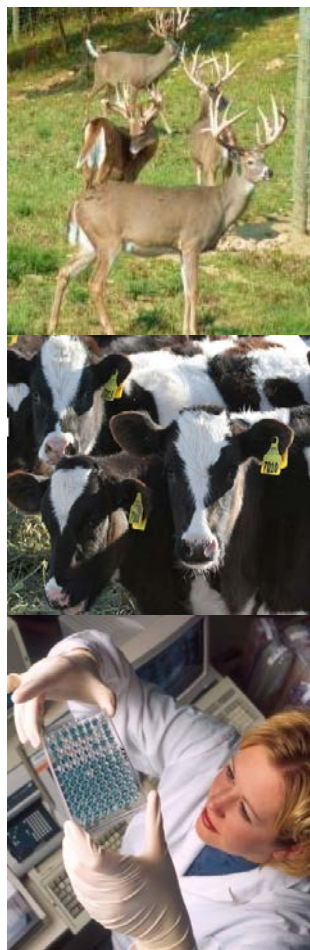


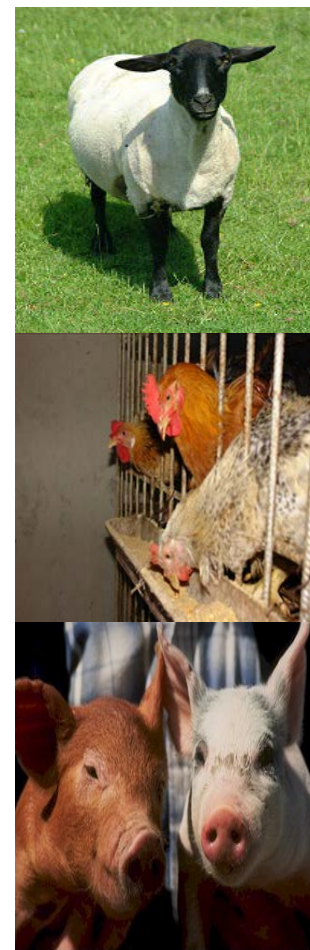
Veterinary Services



Update on *Theileria equi* genotyping at NVSL

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EP genotyping at NVSL

- NVSL's approach
- Progress
- Next steps
- Potential pitfalls

Whole genome sequencing

- NVSL has developed expertise in next-gen sequencing and bioinformatics
 - TB and *Brucella* programs
 - Organism typing moving towards WGS
- Can we successfully sequence the larger genome of *T. equi* using same approach?

Progress

- Proof of concept
 - *In vitro*-derived *T. equi*
 - Ct value 29.1
 - Captured 97% of the *T. equi* genome at a coverage of 200x
 - Sequencing results were 30% *T. equi* and 70% equine
- Sample inventory
 - Approx. 600 whole blood samples archived

Next steps

- Build database of sequences
 - Compare sequence data to known epi info
 - Compare WGS to other published genotyping studies
- Evaluate methods of concentrating *T. equi* DNA
 - Minimize horse DNA
 - Decreased cost
 - Increased sensitivity

Next steps (cont.)

- Archived vs. fresh samples
 - Archived stored in EDTA, significant lysis of cells
 - Two different approaches for DNA concentration
 - Affinity purification for archived samples
 - WBC depletion for fresh samples
 - Enrich by *in vitro* culture?

Potential pitfalls

- Sensitivity
 - Can't be cultured/isolated like bacteria
 - Low parasitemia in chronic carriers
 - Extraction of sufficient amount of *T. equi* for sequencing
- Exchange of genetic information

Questions?

