KEY PRINCIPLES OF BIOSECURITY

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ABSTRACT

The goal of our research program is to put science into biosecurity protocols. We realize that bigger issues such as siting, pig proximity, and aerosol transmission offer risks that we cannot control in many cases. Thus, our research has focused on the details within production units that we can control - specifically the role of people as mechanical vectors in transmitting porcine pathogens. These details are important because we most likely track pathogens among groups of pigs before we observe the clinical signs of an outbreak.

INTRODUCTION

Biosecurity and sanitation practices are implemented on many pork production units to prevent the introduction of pathogens to the herd or to groups of pigs within the herd. These protocols must take into consideration a multitude of risks for pathogen introduction. Many decisions regarding biosecurity protocols on pork production units are currently based on producer and veterinary experience and opinion, not on scientific research. Not knowing the extent to which biosecurity measures need to be implemented to prevent the transmission of porcine pathogens is an important problem, because, until that information is known, pork producers will run one of two risks:

- Expenditure of time and money on unnecessary biosecurity measures, or
- Insufficient biosecurity measures that place the pig population at risk for economically devastating disease outbreaks.

The argument often presented is that all biosecurity measures, even if not effective, are important because implementation of biosecurity protocols sensitizes personnel to biosecurity issues. The biosecurity mind-set of the personnel is thought to enable workers to pay more attention to details that, if performed sloppily, might place the herd at risk of infection. I wholeheartedly agree that we must encourage our colleagues to pay attention to these details in their work. As most biosecurity procedures have not been validated, we must do the best we can with the information that we have to date. However, I feel that a dangerous premise is set when we recommend procedures that have been scientifically shown to be ineffective, just to give the 'perception' that we are doing everything possible to prevent breaches in biosecurity. Encouraging people to perform biosecurity tasks that are known to be worthless damages our credibility. One would not ask personnel to vaccinate a herd for pseudorabies using a modified live vaccine that had been mixed and then stored for 2 weeks at 90°F just to give the perception that by vaccinating the pigs, they were doing everything possible to
prevent an outbreak of pseudorabies. Eventually, employees and clients will recognize the hoax and your future recommendations will not be heard.

**BOOT BATHS**

Farm personnel use boot baths with the goal of preventing mechanical transmission of pathogens among groups of pigs. However, in the authors' experience, boot bath maintenance on most facilities is poor, and frequently boot baths are grossly contaminated with organic matter. People commonly avoid stepping into boot baths or simply step through the bath without stopping to clean their boots.

Literature on boot bath use is scarce and usually limited to the authors’ opinions on proper procedure. Phenolic detergents have been recommended for use in boot baths (Quinn, 1991). Effective utilization of boot baths consisted of cleaning boots in a preliminary bath filled with dilute detergent, followed by immersion of clean boots to a depth of 15 cm, for at least 1 minute, in a second bath filled with detergent. The author advocated that large units prepare new boot baths daily or when visibly contaminated and small units prepare new boot baths every 3 days (Quinn, 1991).

We recently evaluated Cidex Formula 7*, Nolvasan®, Chlorox®, Betadine Solution, 1Stroke Environ®, Rocal-D Plus, and Virkon®S utilizing various boot bath protocols (Amass et al., 2000; Amass et al., 2001). Basic principles of proper boot bath use learned from these experiments include:

- Scrubbing visible manure from boots enhances removal of significant numbers of bacteria. Simply walking through a boot bath will not reduce bacterial counts. Standing in a boot bath for up to 2 minutes without scrubbing off the manure did not significantly reduce bacterial counts except when a cost-prohibitive disinfectant (Cidex formula 7*) was used.
- Scrubbing visible manure off in a water bath is as efficacious as scrubbing manure off in a bath of the disinfectants tested as far as reducing bacterial counts. Although not tested, detergents may make manure removal easier.
- Scrubbing off manure in a clean disinfectant boot bath (1Stroke Environ®) reduces the bacterial count more than scrubbing boots in a contaminated boot bath.
- Boots that have been scrubbed free of manure and then soaked in Rocal-D Plus for 5 or more minutes meet the standard for disinfection.
- Boots that have been scrubbed free of manure and then dipped in Virkon®S meet the standard for disinfection the majority of the time.
- Boots that have been scrubbed in Virkon®S for 30 seconds meet the standard for disinfection; however, a clean tub of Virkon®S must be used each time.

Time constraints make proper use of boot baths within production units difficult. However, spending time and money to implement boot bath procedures on a farm without using them correctly is a waste of resources. Although going through the motions of stepping in a boot bath has benefits of increasing employee awareness of biosecurity and maintaining a clean
workplace, this insufficient biosecurity measure as tested in this study may place the pigs at risk for infection because contaminated boots are being used by personnel.

In conclusion, boot stations with hoses and brushes will facilitate manure removal. Disinfectants should be selected based on efficacy, cost, ease of use, and environmental friendliness. Manure should be removed from boots before placing them in a boot bath or else a new clean boot bath needs to be prepared each time boots are cleaned. The intention of this research was not to have everyone stop cleaning boots, but instead, to encourage the use of effective footwear cleaning protocols.

PEOPLE

People-flow into and within production units comprises a large component of biosecurity; however little research is available to support common policies regarding people movement. Sellers et al. (1970), sampled people who had contacted animals infected with FMDV. More FMDV was isolated from the nose than the mouth of these people. Virus was isolated from the nose of one person at 28 hours, but was not isolated after 48 hours. Nose blowing or washing was not effective in eliminating the virus, and cloth or industrial masks were not effective in preventing inhalation of the virus. Transfer of the virus between people was documented after persons in contact with infected animals spoke to unexposed colleagues in a box for 4 minutes. One year later, Sellers et al. (1971) reported that FMDV could be transferred by human beings, from infected pigs, to susceptible cattle. Results from Seller's work appear to be the origin for the "48 hour rule" used by many producers even though different viruses and bacteria may be harbored for longer or shorter periods by humans. Wentworth et al. (1997) recorded transmission of SIV to human caretakers. In this study, pig-to-human transmission occurred despite the use of Animal Biosafety Level 3 containment practice (coveralls, boots, goggles, gloves, hairnets, and dust masks).

In contrast, Goodwin (1985) reported that the culture of breath and hair samples from a person exposed to pigs experimentally infected with M. hyopneumoniae did not result in reisolation of M. hyopneumoniae. Additionally, we could not detect pig-to-human transmission of S. suis using throat swab samples collected from farm personnel who were working in close daily contact with infected pigs (Amass et al., 1998).

Our investigations (Amass et al., 2000) of people as mechanical vectors for PRRSV were less conclusive. Although people did not transmit PRRSV from pigs with acute PRRS to uninfected pigs under the conditions of our study, there was some evidence that people could be contaminated with PRRS viral RNA after contact with infected pigs. PRRS viral RNA was detected in saliva and fingernail rinse samples of 2 of 10 people immediately after exposure to PRRSV-inoculated pigs, on a third person (fingernail rinse) at 5 hours, and a fourth person (nasal swab) at 48 hours after exposure to infected pigs. Further studies should address these findings using virus isolation instead of nRT-PCR to determine if the PRRSV RNA found on people is infectious.
Our studies of people as mechanical vectors of TGEV demonstrated that people could act as mechanical vectors and spread TGEV to healthy pigs; however, handwashing and changing outerwear after exposure to infected pigs was sufficient to prevent transmission (Alvarez et al., 2001).

Thus, it would appear that the risk of transmitting diseases back-and-forth between human beings and swine varies with the pathogen. Quantification of the risk of transmission of common porcine pathogens and individual strains of these pathogens on an individual basis is essential.

CONCLUSIONS

Further research is needed to validate biosecurity methods used in pork production. Once effective biosecurity procedures are defined, producers and veterinarians can develop protocols for production units commensurate with the greatest risks for that farm, keeping in mind that removal of visible manure is central to all biosecurity efforts whether the contaminated surface is a boot, clothing, truck or skin.

LITERATURE CITED


