Vector/Pathogen/Host Interaction, Transmission

External Surface Disinfection of the Lesser Mealworm (Coleoptera: Tenebrionidae)

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J. Med. Entomol. 43(5): 916–923 (2006)

ABSTRACT

Understanding the dynamics of movement of bacteria within the environment and between species is crucial to unraveling the epidemiology of bacterial diseases and to developing biosecurity measures to prevent dissemination. Many arthropods, some beneficial and some detrimental, inhabit poultry houses. The lesser mealworm, Alphitobius diaperinus (Panzer) (Coleoptera: Tenebrionidae), is a pest commonly found in poultry litter that can harbor pathogens involved in both human and animal health issues. Current farm management practices perpetuate persistent infestations contributing to the dispersal of beetles and pathogens. To study the dissemination of bacteria by this beetle, we require the ability to differentiate internal from external sources of bacteria carried by the beetle. In this study, we tested previously described methods to externally disinfect beetles and found disinfectant efficacies between 40 and 98%. The irregular surface of the insect posed a challenge to cleansing procedures because the surface offered many recesses able to sequester bacteria. Complete bacterial disinfection was achieved with a serial treatment of ethanol and hydrogen peroxide or hydrogen peroxide/peracetic acid.

KEY WORDS
lesser mealworm, external disinfection, Alphitobius diaperinus, bacteria, poultry litter beetle

Lesser mealworms, Alphitobius diaperinus (Panzer) (Coleoptera: Tenebrionidae), at all life stages, inhabit manure and feed in commercial poultry operations and are one of the most abundant insect species recovered from broiler chicken and turkey litter samples (Pfeiffer and Axtell 1980, Stafford et al. 1988, Axtell and Arends 1990, Rueda and Axtell 1997). Lesser mealworms are omnivorous scavengers that feed on manure, spilled chicken feed, cracked eggs, chicken carcasses, house fly maggots, and detritus (Pfeiffer and Axtell 1980, Axtell and Arends 1990, Rueda and Axtell 1997). In turn, these beetles are often fodder for chickens, wild birds, and opportunistic rodents. In addition, the beetles are inadvertently dispersed to neighboring residences by the spreading of beetle-containing manure on nearby fields (Armitage 1986). They are a prime candidate for participating in the transmission of bacteria among fauna inhabiting the poultry house environment and have previously been implicated in the transmission of several disease agents, including bacterial pathogens (De las Casas et al. 1968, 1973, 1976; Despins et al. 1994; McAllister et al. 1994, 1995, 1996; Hald et al. 1998; Gray et al. 1999).

Beetle morphology can reduce the effectiveness of surface sterilization by preventing adequate access to bacteria. The insect integument and presumably the fecal material adhering to the exoskeleton provide refuge to bacterial organisms. Flexible joints, wings, and spiracles occurring between hard plates offer anchorage to microbes, whereas structures such as the elytra and the cuticle serve as protective covers (Chapman 1982a). In addition, ectodermal invaginations, such as sutures or sulci, may act to shield bacterial organisms from access and displacement during disinfectant procedures (Chapman 1982b). This study assesses the efficacy of previously described and newly developed methods to disinfect the external surface of beetles. We have devised a specific method resulting in complete surface bacterial disinfection for use in future studies to explore the environmental transfer of bacteria by this beetle.

Studies exploring the transfer of bacteria by litter beetles have been reported, but there is limited assessment of how a transferred bacterium is harbored by a beetle. This task requires the ability to differentiate bacteria carried externally or internally. Although several previous studies described protocols to disinfect the surface of beetles, no data were presented validating the efficacy of the techniques (De las Casas et al. 1968, 1972; Harein and De las Casas 1968; Harein et al. 1970, 1972; McAllister et al. 1994, 1995, 1996; Hald et al. 1998; Gray et al. 1999).

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Materials and Methods

Beetles. The Southern Plains Agricultural Research Center (SPARC) starter colony of *A. diaperinus* was a generous gift from a colony originally isolated from a poultry farm located in Wake County, North Carolina, and maintained by Dr. D. W. Watson (North Carolina State University, Raleigh, NC). The SPARC colony was initiated and has remained in production since 2004. Beetles were reared in 1,000-ml wheat bran (Morrison Milling Co., Denton, TX) in plastic containers (15 by 15 by 30 cm) with screen tops and held at 30°C under a photoperiod of 8:16 (L:D) h. Additional within each cage, a 6-cm² sponge moistened with 30% (Morrison Milling Co., Denton, TX) in plastic containers (15 by 15 by 30 cm) with screen tops and held at 30°C under a photoperiod of 8:16 (L:D) h. Additional within each cage, a 6-cm² sponge moistened with deionized water (dH₂O) and a 0.5-cm-thick slice of a medium-sized apple were replenished twice per week, and 30 ml of fishmeal (Omega Protein, Inc., Hammond, LA) was added to the wheat bran once per week.

Disinfecting Agents. Hydrogen peroxide (H₂O₂) (Sigma-Aldrich, St. Louis, MO), 95% ethanol (EtOH) (EMD Chemicals, Gibbstown, NJ), Tween 80 (Amresco, Solon, OH), and sodium hypochlorite (NaOCl) (Sigma-Aldrich) were diluted to working concentrations with sterile dH₂O. The 7.35% H₂O₂/0.23% peracetic acid was a commercially available formulation, SporGon (Decon Labs, Inc., Bryn Mawr, PA).

These agents were used individually or in combination to formulate and evaluate 11 disinfection protocols designed for comparison to previously described techniques or to test new combinations for increased efficacy. The protocols were grouped into EtOH (three), NaOCl (four), and H₂O₂ (four)-based protocols as described below. All protocols were evaluated on individual beetles held and disinfected in 1.5-ml tubes covered with a sterile barrier film (Parafilm; Sigma-Aldrich).

For each protocol, the tube in which the beetle was immersed was covered by the barrier film, inverted three times, and sonicated for 2 min at 40 kHz (model 8851-34, Cole Parmer, Vernon Hills, IL) before the next step. The beetle was transferred into a new sterile tube for each successive immersion or rinse. Evaporation was performed by transferring the beetle into a sterile tube, which was placed into a sterile, biosafety hood to allow the EtOH to evaporate. Sonication was used to assist in dislodging of the bacteria and to ensure uniformity of the agitation procedure for comparison of protocols.

EtOH-Based Protocols. (A) Each beetle was immersed in 95% EtOH and then rinsed in sterile dH₂O for 30 s followed by sonication for 2 min; the rinse was repeated. (B) Each beetle was immersed in 70% EtOH and the EtOH evaporated for 5 min. (C) Each beetle was immersed in 95% EtOH, and the EtOH evaporated for 5 min.

NaOCl-Based Protocols. (D) Each beetle was immersed in 2% NaOCl/10% Tween 80 and then rinsed in sterile dH₂O for 30 s followed by sonication for 2 min; the rinse was repeated. (E) Each beetle was immersed in 2% NaOCl/Tween 80, allowed to soak an additional 8 min, and then immersed in 70% EtOH. The beetle was then rinsed in sterile dH₂O for 30 s followed by sonication for 2 min; the rinse was repeated twice. (F) Each beetle was immersed in 2% NaOCl/Tween 80, allowed to soak an additional 8 min, and then immersed in 95% EtOH and the EtOH evaporated for 5 min. (G) Each beetle was immersed in 95% EtOH, followed by immersion in 2% NaOCl/Tween 80, and allowed to soak an additional 8 min. The beetle was then immersed in 70% EtOH and the EtOH evaporated for 5 min.

H₂O₂-Based Protocols. (H) Each beetle was immersed in 20% H₂O₂, followed by immersion in 5% EtOH. (I) Each beetle was immersed in 20% H₂O₂, followed by immersion in 95% EtOH. (J) Each beetle was immersed in 95% EtOH and the EtOH evaporated for 5 min, followed by immersion in 20% H₂O₂. (K) Each beetle was immersed in 95% EtOH and the EtOH evaporated for 5 min, followed by immersion in 7.35% H₂O₂/0.23% peracetic acid.

Experimental Design. Three replications of each experimental wash protocol were conducted using 30 beetles per protocol per replication. To collect a sample of the resident bacteria present on the outer surface of the beetles before external disinfection (PRE-wash), each beetle was immersed in 1 ml of tryptic soy broth (TSB; Difco, Sparks, MD) at room temperature. The beetle was removed and immediately subjected to one of the experimental disinfection protocols (see above). An aliquot of 0.1 ml of PRE-wash TSB was serially diluted, spread on blood agar plates (Becton Dickinson, Sparks, MD), and incubated overnight at 37°C. Bacterial load, expressed as colony-forming units (CFU), was determined after an 18–24-h incubation. After each disinfection protocol was completed, surviving residual bacteria were sampled (POST-wash) by again immersing the beetle in 1 ml of TSB at room temperature. The beetle was removed, and an aliquot of 0.1 ml of POST-wash TSB was serially diluted, spread on blood agar plates, and incubated overnight at 37°C. CFU were determined after an 18–24-h incubation. To ensure detection of bacterial contamination below the plating threshold of 10 CFU, the remaining 0.9 ml of PRE-wash and POST-wash TSB was enriched by incubation overnight at 37°C (ENRICH). After incubation, an aliquot of 0.1 ml of TSB was spread on a blood agar plate and incubated overnight at 37°C. The presence or absence of bacteria was recorded after 18–24-h incubation at 37°C. CFU data were analyzed by logistic regression and enrichment data were analyzed by exact logistic regression in PROC LOGISTIC (SAS Institute, Cary, NC; Agresti 2002). Representative colonies of bacteria present were collected from PRE- and POST-wash blood plates for identification by growth on selective media or by ribotyping.

For PRE-TSB control, 60 beetles were taken from the same cage and split into two groups of 30 beetles. From one group, beetles were placed into individual tubes containing TSB. These served as PRE-wash samples for this control study to ensure that the tested beetles were contaminated with bacteria. Beetles
from the second group, without prior immersion in TSB, were subjected to protocol K as described above.

**Bacterial Isolation and Preliminary Identification**

**Culture Methods.** Individual bacteria from mixed cultures were grown in TSB and then subcultured onto blood agar plates for selection of isolated colonies. Individual isolates were identified by growing them on a bank of selective and differential media and then comparing the results from the unknown isolate to data for known species (Atlas 1997). To identify aerobic bacteria, 10-µl aliquots were streaked onto Brilliant Green agar (BGA; Becton Dickinson), MacConkey (Becton Dickinson), m Enterococcus (ME; Becton Dickinson), Rogosa (Becton Dickinson), CHROMagar E. coli, and CHROMagar Orientation (CHROMagar, Paris, France) plates, and then incubated at 37°C for 24 h. Each initial colony selection was streaked onto fresh media to ensure cultural purity. A bacterial lawn of each pure isolate culture was pre-streaked onto fresh media to ensure cultural purity. A pattern be an 85% or greater match to an existing database (containing patterns in the bacterial database provided by Qualicon, Inc.).

**Ribotype Characterization.** Bacterial isolates from blood agar plates were incubated at 37°C for 24 h to allow a bacterial lawn to form. Samples were collected while in log phase growth. Bacteria were suspended in a neutral pH buffer (Qualicon, Inc., Wilmington, DE), heated at 90°C for 10 min, combined with two additional lytic enzymes (Qualicon, Inc.), and analyzed according to manufacturer’s instructions using the restriction enzymes EcoRI or PvuII. The RiboPrinter microbial characterization system (Qualicon, Inc.) characterizes the 5, 16, and 23S rRNA and flanking regions of a bacterial sample by using specified restriction enzymes. The resulting rRNA pattern (RiboPrint pattern) was automatically characterized and identified by comparing the pattern to reference patterns in the bacterial database provided by Qualicon, Inc. (containing >6000 isolates) and a custom, food animal-specific database (containing >400 isolates) developed by C.S. Identification requires that the ribopattern be an 85% or greater match to an existing static ribopattern. When ribotype identification match was <85%, isolate identification was confirmed using API Staph, API 20E, API 20NE, API 20 Strep, ID 32 STAPH (Analytical Profile Index, bioMerieux, Inc., Durham, NC) manual identification test strips.

**Results**

A comparison of external disinfection protocols demonstrated large variation in treatment efficacy (Table 1). External disinfection using 95% EtOH followed by a dH2O rinse (protocol A), disinfected less than one-half of the beetles. However, allowing external surface evaporation after treatment with EtOH instead of immediately performing a dH2O rinse, significantly improved disinfection by 61% (protocol C versus A; *P* < 0.0001). Using 2% NaOCl/10% Tween 80 followed by a dH2O rinse (protocol D) disinfected slightly more than one-half of the beetles. Adding a 70% EtOH treatment to the protocol significantly improved disinfection by 36% (protocol E versus D; *P* < 0.0003). Removing the dH2O rinse completely and adding a higher strength 95% EtOH treatment, which was subsequently allowed to evaporate, also significantly improved disinfection by 40% (protocol F versus D; *P* < 0.002). Prior treatment of 95% EtOH with NaOCl/Tween 80 improved disinfection by 19% (protocol G versus F; *P* < 0.03). However, the addition of a second 95% EtOH treatment before the NaOCl/Tween 80 and 95% EtOH treatments did not significantly increase efficacy (protocol G versus F). A dH2O rinse to wet the beetle followed by 20% H2O2 (protocol H) disinfected almost all of the beetles. The substitution of a 95% EtOH, which was subsequently allowed to evaporate, before treatment with 20% H2O2 or 7.35% H2O2/0.23% peracetic acid, disinfected 100% of the beetles (protocols J and K). However statistically nonsignificant the increase between protocols H versus J or K (2%), it gave us the outcome we desired, which was consistently reproducible complete disinfection of culturable, aerobic bacteria.

The measuring of resident bacteria before executing the disinfection protocol required a pretreatment of beetles by immersion in TSB. As this would not be a step normally included in the external disinfection protocol, we demonstrated that this initial immersion in TSB did not affect the efficacy of the disinfection.

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**Table 1. Comparison of the efficacy of lesser mealworm surface disinfection protocols**

| Disinfection Protocol | CFU | Enrichment |
|-----------------------|-----|------------|
|                       | PRE-Wash Mean ± SE | POST-wash Mean ± SE | % Clean beetles ± SE |
| EtOH-based protocols  |     |            |                           |
| A 95% EtOH-H2O        | 10.99 ± 0.00 | 9.14 ± 0.07 | 48.52 ± 0.07 |
| B 70% EtOH-Evaporation | 3.99 ± 0.25 | 14.23 ± 0.50 | 60.19 ± 0.07 |
| C 95% EtOH-Evaporation | 1.27 ± 0.01 | 1.27 ± 0.01 | 78.26 ± 0.01 |
| NaOCl-based protocols |     |            |                           |
| D NaOCl-H2O           | 13.22 ± 0.00 | 222.16 ± 2.63 | 66.72 ± 0.00 |
| E NaOCl-70% EtOH-H2O  | 2.98 ± 0.10 | 0.00 ± 0.02 | 90.87 ± 0.04 |
| F NaOCl-95% EtOH-Evaporation | 1.68 ± 0.01 | 0.00 ± 0.01 | 90.32 ± 0.01 |
| G 95% EtOH-NaOCl-95% EtOH | 2.69 ± 0.56 | 0.00 ± 0.00 | 95.15 ± 0.00 |
| H2O2-based protocols  |     |            |                           |
| H H2O2-H2O2          | 1.69 ± 0.56 | 0.00 ± 0.01 | 97.78 ± 0.00 |
| I H2O2-95% EtOH-Evaporation | 1.18 ± 0.56 | 16.36 ± 0.46 | 91.43 ± 0.03 |
| J 95% EtOH-Evaporation-H2O2 | 1.19 ± 0.56 | 0.00 ± 0.00 | 100.00 ± 0.00 |
| K 95% EtOH-Evaporation-H2O2/peracetic | 1.52 ± 0.56 | 0.00 ± 0.00 | 100.00 ± 0.00 |

*Average of three replicates, 30 beetles per replicate.*

*POST-wash is significantly different from PRE-wash (*P* < 0.001).
protocol by testing protocol K (7.35% H$_2$O$_2$/0.23% peracetic acid) without PRE-wash immersion in TSB. A comparison between beetles exposed to prior TSB immersion and those disinfected without prior immersion showed no significant difference ($P < 1.0$) in wash efficacy. This demonstrated that the PRE-wash TSB immersion had no affect on the outcome of the disinfectant protocol.

The resident bacterial load of each beetle was measured individually before exposure to a disinfection protocol (Fig. 1). A frequency distribution of the log$_{10}$ CFU of the results indicated that bacterial load was variable on individual beetles. Bacterial load ranged from 0 CFU on 20 beetles to 2.5 $\times$ 10$^6$ CFU on one beetle, averaging 7.6 $\times$ 10$^3$– 9.0 $\times$ 10$^4$ CFU, with a median of 1.2 $\times$ 10$^3$ CFU.

The species of bacteria carried by the beetles was relatively consistent. Although beetles were reared in separate cages, they used feed and bran from identical sources and were placed into a common cage as adults before use in experiments; thus, similar contamination profiles were found. The beetles were contaminated with 12 bacterial species, representing nine genera: Acinetobacter spp. (Brisou and Prévot; 97% API match), Aerococcus spp. (Williams; 100% API match), Bacillus spp. (Meyer and Gottheil; 97% ribotype match), Bacillus fusiformis (Cohn; 91% ribotype match), Enterobacter cloacae (Jordan; 94% ribotype match), Klebsiella oxytoca (Flügge; 94% ribotype match), Micrococcus lutetus (Schloëf; 91% ribotype match), Pseudomonas aeruginosa (Schroeter; 85% ribotype match), Staphylococcus gallinarum (Devrieze; 94% ribotype match), Streptococcus kloosii (Schleifer; 92% ribotype match), Streptococcus xylosis (Schleifer and Kloos; 91% ribotype match), and Streptococcus spp. (Rosenbach; 92% API match). The bacteria showed varying sensitivities to the disinfection protocols (Table 2). Acinetobacter spp. and S. gallinarium seemed particularly resistant to EtOH- and NaOCl-based washes. M. luteus was also resistant to 70% EtOH and NaOCl washes, but it did not survive a 95% EtOH wash. Bacillus spp. and Enterobacter cloacae resisted sequential washing in dH$_2$O and H$_2$O$_2$, but they were removed by sequential washing in 95% EtOH and H$_2$O$_2$. Although we did not identify the taxa, we noted that sequential washing in 95% EtOH and H$_2$O$_2$ also disinfected fungi present on the PRE-wash plates (data not shown).

**Discussion**

Litter beetles, especially the lesser mealworm, have become serious pests within the poultry brooder and laying industry. Because of their mobility, feeding habits, and prey potential, these beetles are implicated as mechanical vectors for diseases (e.g., Mareks disease, avian influenza, bacterial diseases, fowl pox, coccidiosis, and New Castle disease). The insect offers surfaces to support bacterial, fungal, and viral organisms (De las Casas et al. 1968, 1972, 1973; Harien et al. 1970; McAllister et al. 1994, 1995). These beetles have high reproductive rates and are difficult to control. Ultimately, they are portrayed as a reservoir source contributing to the persistence and transmission of pathogens among individual birds within a poultry facility (Harien et al. 1970, 1972; Brown et al. 1992; Hald et al. 1998).

Previous studies have attempted to surface sterilize beetles to establish the carriage of pathogens internally (De las Casas et al. 1968, 1972; Harien and De las Casas 1968; Harien et al. 1970, 1972; McAllister et al.
Table 2. Comparison of surface disinfection protocols by removal of specific bacterial species

| Bacterial spp. | A 95% EtOH | B 70% EtOH | C 95% EtOH | D NaOCl | E NaOCl | F NaOCl | G 95% EtOH | H H2O2 | I 95% EtOH | J 95% EtOH | K 95% EtOH |
|----------------|----------|----------|----------|--------|--------|--------|----------|--------|----------|----------|----------|
| Acinetobacter spp. | + | + | + | + | + | + | + | + | + | + | + | + |
| Bacillus spp. | + | - | + | + | + | - | + | + | - | - | + | - |
| Bacillus subtilis | + | + | + | + | + | + | + | + | + | + | + | + |
| Enterococcus faecalis | + | + | + | + | + | + | + | + | + | + | + | + |
| Klebsiella oxytoca | + | + | + | + | + | + | + | + | + | + | + | + |
| Micrococcus luteus | + | + | + | + | + | + | + | + | + | + | + | + |
| Pseudomonas aeruginosa | + | + | + | + | + | + | + | + | + | + | + | + |
| Staphylococcus gallinarum | + | + | + | + | + | + | + | + | + | + | + | + |
| Staphylococcus aureus | + | + | + | + | + | + | + | + | + | + | + | + |
| Staphylococcus xylosus | + | + | + | + | + | + | + | + | + | + | + | + |
| Streptococcus spp. | + | + | + | + | + | + | + | + | + | + | + | + |

*Representative bacteria, selected from PRE- and POST-wash blood agar plates, identified using selective culture methods and ribotype characterization.

+ bacteria was present; blank space, bacteria was not present; —, bacteria removed by corresponding disinfection protocol.
E. coli for competence of lesser mealworms for disinfected by this method (protocol B). The reservoir of disinfection techniques was determined by subsequent individual exposure of each beetle to nutrient broth and removal of contaminated specimens from the study. We determined that washing in 2% NaOCl/Tween 80 followed by dH$_2$O was only 66.72% effective (protocol D). Similarly, De las Casas et al. (1968) determined that ~50% of the pupae subjected to these disinfection procedures were still contaminated. In our study, the simple addition of an EtOH rinse step after disinfection by NaOCl/Tween 80 increased effectiveness to an average of 92.33% (protocol E and F).

Lesser mealworms have been surface sterilized against infectious bursal disease virus with 10% H$_2$O$_2$ followed by rinsing in dH$_2$O (McAllister et al. 1995). In our study, disinfection with 10% H$_2$O$_2$ was <100% effective against bacteria (data not shown). An increase to 20% H$_2$O$_2$ (protocol H) was found to be 97.8% effective against bacteria. The addition of a 95% EtOH treatment before a 20% H$_2$O$_2$ wash resulted in consistent, complete surface bacterial disinfection.

H$_2$O$_2$ is often used as an antimicrobial or bleaching agent. However, it is an unstable compound that quickly neutralizes itself by reverting to oxygen and water. Stability depends upon many factors, but solutions of H$_2$O$_2$ which are kept in dark, inert containers that are completely free of contamination are relatively stable. Commercial solutions, however, usually contain minute amounts of impurities, which can cause decomposition; therefore, stabilizers are sometimes added (Goor 1989, Hess 1995, CHEMINFO 2005). Concentrated H$_2$O$_2$ solutions can also react exothermically with solvents. Although H$_2$O$_2$ proved to be the best cleansing agent, its instability in pure form and exothermic reactivity at high concentrations made it problematic to use; therefore, we investigated a commercially available substitute. The substitute consisted of a less concentrated formulation of 7.35% H$_2$O$_2$ with the addition of 0.23% peracetic acid, a stabilizer, a surfactant and a corrosion inhibitor. Our results demonstrate that this formulation was as effective as 20% H$_2$O$_2$ treatment.

The thin epicuticle secreted on the outside of the cuticle serves to minimize loss of insect body water, contributing to their success in a terrestrial environment (Bursell and Clements 1967). The epicuticle consists of several layers, including a superficial wax or lipid layer of long-chain hydrocarbons and esters of fatty acids and alcohols (Chapman 1982a, Lockey 1988). This deterrent to desiccation also presumably diminishes the effectiveness of surface disinfection by H$_2$O$_2$ or NaOCl treatment due to its hydrophobic nature. We found that prior treatment with EtOH enhanced the efficacy H$_2$O$_2$ and NaOCl treatments. Because lipids are soluble in organic solvents, it is likely that EtOH treatment either diminished the waxy layer or wetted the exterior sufficiently to allow a subsequent agent improved access to the beetle.
surface and invaginations, and the microorganisms residing there.

Surface disinfections that included EtOH or NaOCl were detrimental to the beetles and often resulted in their death. H$_2$O$_2$ treatment seemed less deleterious because the beetles usually survived the treatment. For our purposes and for many of the studies discussed, survival of the specimens was not required. However, for future studies where survival is desired, a modification in the protocols to a protocol less adverse to the survival of the beetles while still maintaining efficacy as a disinfectant will be necessary.

In summary, this study assessed the efficacy of previously described and newly developed methods to disinfect the external surface of A. diaperinus. Although many of the bacterial disinfecting procedures examined cleaned >90% of the beetles, we endeavored to find a protocol that disinfected 100% of the insects. When investigating bacterial transfer, complete exoskeleton disinfection is important to discern conclusively an internal source of contamination. Several previously reported studies used protocols that purportedly disinfected the surface of beetles; however, we could find no data validating the efficacy of the described techniques. Although it is possible that the insects harbor bacterial organisms internally, insufficiently sterilized exoskeletons can contaminate subsequent homogenized specimens. Therefore, it would be unclear whether the bacteria being measured were actually being harbored internally. We found that a combination of treatment with 95% EtOH, allowed to evaporate, followed by a 20% H$_2$O$_2$ or 7.35% H$_2$O$_2$/0.23% peracetic acid wash removed all culturable, aerobic bacteria from the external surface of the adult lesser mealworms.

Acknowledgments

We thank Jesus Esquivel and Sharon Mowery for assistance in establishing the beetle colony; Gretchen Jones, Kate Andrews. Jason Leger, and Melanie Sandera for technical assistance; and Sara Duke for assistance with statistical analysis.

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Received 13 December 2005; accepted 27 February 2006.