Detection of Foodborne Bacterial Pathogens from Individual Filth Flies

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Abstract

There is unanimous consensus that insects are important vectors of foodborne pathogens. However, linking insects as vectors of the pathogen causing a particular foodborne illness outbreak has been challenging. This is because insects are not being aseptically collected as part of an environmental sampling program during foodborne outbreak investigations and because there is not a standardized method to detect foodborne bacteria from individual insects. To take a step towards solving this problem, we adapted a protocol from a commercially available PCR-based system that detects foodborne pathogens from food and environmental samples, to detect foodborne pathogens from individual flies. Using this standardized protocol, we surveyed 100 wild-caught flies for the presence of Cronobacter spp., Salmonella enterica, and Listeria monocytogenes and demonstrated that it was possible to detect and further isolate these pathogens from the body surface and the alimentary canal of a single fly. Twenty-two percent of the alimentary canals and 8% of the body surfaces from collected wild flies were positive for at least one of the three foodborne pathogens. The prevalence of Cronobacter spp. on either body part of the flies was statistically higher (19%) than the prevalence of S. enterica (7%) and L. monocytogenes (4%). No false positives were observed when detecting S. enterica and L. monocytogenes using this PCR-based system because pure bacterial cultures were obtained from all PCR-positive results. However, pure Cronobacter colonies were not obtained from about 50% of PCR-positive samples, suggesting that the PCR-based detection system for this pathogen cross-reacts with other Enterobacteriaceae present among the highly complex microbiota carried by wild flies. The standardized protocol presented here will allow laboratories to detect bacterial foodborne pathogens from aseptically collected insects, thereby giving public health officials another line of evidence to find out how the food was contaminated when performing foodborne outbreak investigations.

Introduction

Insects play an important role in the transmission of food-related diseases because they can spread pathogens to food or food contact surfaces and utensils1. Among insects, flies, cockroaches, and ants exhibit behaviors that favor the spread of foodborne pathogens. These behaviors include an association with decaying matter, refuse and feces, endophily (entering buildings), and synanthropy (cohabiting with humans)2. Foodborne pathogens such as Salmonella spp., Listeria monocytogenes, Campylobacter spp., Escherichia coli O157:H7, and members of the genus Cronobacter (formerly Enterobacter sakazakii) have been reported to be transmitted by insects3-5. Synanthropic filth flies mechanically spread foodborne bacteria by transferring pathogens from their contaminated body surfaces. However, the presence of foodborne pathogens in the alimentary canal of flies can be up to three times greater than that observed on their body surfaces (body, head, legs, and wings)6. Foodborne pathogens can also remain in the fly’s alimentary canal for a greater length of time than on the body surface6,7 and in some instances, they are able to multiply, colonizing the fly’s digestive tract8,9. This increases the vector potential of flies because they can further spread foodborne pathogens through defecation and regurgitation10,11.

Nowadays, there are improved surveillance systems that are able to detect foodborne illness outbreaks more rapidly. While performing foodborne outbreak investigations, public health officials look for the food that may be the source(s) or vehicle(s) of infection. Investigators may also perform an environmental assessment of the facility (or facilities) involved to find out how the food was contaminated and may collect samples as part of the investigation12. Despite the vast amount of scientific literature concerning insects as carriers of foodborne pathogens, linking insects as vectors of the pathogen causing a particular foodborne illness outbreak has been challenging. This is mainly because insects are not being aseptically collected as part of environmental sampling programs during foodborne outbreak investigations. To include insects, particularly those that exhibit behaviors that favor the spread of foodborne pathogens, as part of an environmental sampling procedure, a standardized, rapid, sensitive and reliable protocol to detect foodborne pathogens from a single insect needs to be in place.

Traditional plating techniques for the detection of foodborne pathogens from insects are laborious and depend upon the competitive growth of the target bacteria in different culture media to overcome the rapid growth of the innate commensal microbiota of the insect. Most of the studies that have associated insects with bacterial pathogens have increased the sensitivity of the method by pooling together several insects rather than identifying the presence of pathogens on a per individual basis. Thus, those studies did not differentiate the body part of the insect where the pathogens were found13-18. The ability to identify whether foodborne pathogens are located on the body surface or in the alimentary...
canal of an individual insect is important as this may have epidemiological implications and may lead to different mitigation strategies. As mechanical vectors, flies that land on food for a short time may only transfer low levels of bacteria from their body surface, whereas those flies that regurgitate and defecate on the food increase the probability of transferring pathogens at potentially higher levels of infection. Consequently, it is important to estimate the prevalence of a foodborne pathogen per an individual insect and to differentiate the body part of that insect where the bacterial pathogen is located.

Even though the use of culture-independent methods to detect foodborne pathogens are increasingly being implemented, they have not been commercially used to detect foodborne pathogens from a single insect. Currently, there are validated molecular protocols that are commercially available for the rapid detection of foodborne pathogens from foods that are being used by industry and regulatory agencies. These methods include DNA-based systems for the detection of pathogens in a variety of food samples. Although molecular protocols are faster than traditional plating methods, enrichment of the sample is still required to obtain the sensitivity level of $10^5$ colony forming units (CFU) of the bacterial pathogen needed in polymerase chain reaction (PCR)-based methods. Additionally, isolation of pure bacterial colonies from PCR-positive samples is needed to confirm the pathogen using appropriate methods.

The aim of this protocol is to standardize a commercially available PCR-based system used to detect pathogens from food and environmental samples for the detection of foodborne bacteria from the body surface and the alimentary canal of a single fly and to further isolate those pathogens from the samples. The sensitivity of the protocol described here was first calibrated with lab-reared adult house flies (Musca domestica) that were experimentally fed with serial dilutions of each bacterial pathogen. The standardized protocol was subsequently used to survey 100 wild-caught flies for the presence of foodborne pathogens from their body surfaces and/or alimentary canals. This standardized protocol will allow public health laboratories to detect health threats posed by insects, allowing for the possibility of collecting them as part of the environmental sampling program when performing foodborne outbreak investigations.

### Protocol

1. **Collection of flies**

   1. Collect individual flies using sterile entomological sweep nets. Put the nets in a cooler and transfer them to the lab.

2. **Dissection of Flies**

   1. Immobilize aseptically collected flies by placing them at -20 °C for 5 - 7 min.

   2. Using sterile forceps place one fly in a sterile 2 ml tube containing 1 ml of pre-warmed (37 °C) buffered peptone water (BPW). Mix the tube gently by inversion for 2 min. It is essential that the whole body of the fly be in contact with the media so that the microflora present on the body surface (S) of the fly will be transferred to the BPW (BPW-S). Label the tube with a number and body part of the fly (i.e., 1S).

   **NOTE:** Please see Table of Specific Reagents/Equipment for a detailed description of materials and reagents mentioned in this protocol.

   3. Using sterile forceps remove the fly from the BPW-S media and transfer it to an empty and clean 2 ml tube to surface disinfect the fly. Incubate the tube containing the BPW-S media at 37 °C while performing the disinfection and dissection protocol.

      1. Surface-disinfect the fly by immersing it in 1 ml of 70% ethanol for 1 min, followed by a rinse with sterile distilled water before immersing it in 1 ml of freshly prepared 0.05% (v/v) bleach solution. Rinse 3 times with sterile distilled water. Transfer water from the last rinse to an autoclaved 2 ml tube.

      **NOTE:** Discard the liquid each time by using a 1,000 µl micropipette or by inverting the tube, making sure the fly remains inside the tube. Mix gently by inversion at each step of the surface-disinfection process.

      2. To evaluate the effectiveness of the disinfection process, transfer 100 µl of the water from the last rinse to a trypticase soy agar (TSA) plate and spread using a sterile L-shaped disposable spreader. Incubate the plate at 37 °C for 24 hr. After incubation, register the presence of any bacterial colonies.

      **NOTE:** The presence of bacterial colonies on the TSA plates indicates an inefficient surface-disinfection process. If this occurs, the presence of foodborne pathogens should only be reported on the body surface of the fly because cross-contamination between the body surface and the alimentary canal cannot be ruled out.

   4. After surface-disinfecting the fly, transfer it to a piece of autoclaved paper towel to remove excess water and then to a sterile 60 mm disposable Petri dish.

   5. Place the Petri dish under a dissecting scope and identify the fly to the species level using dichotomous keys for dipteran families. For Salmonella, use 1 ml of pre-warmed (42 °C) BPW. Incubate in a recirculating water bath at 42.5 °C for 22 - 24 hr. For secondary enrichment, transfer 100 µl of enriched BPW to 400 µl of pre-warmed (37 °C) brain heart infusion (BHI) broth previously placed in sterile cluster tubes. Incubate at 37 °C for 3 hr.

   6. Using autoclaved fine tip forceps gently pull the anus and the whole alimentary canal (A) out of the fly and aseptically transfer it to another disposable Petri dish.

   **NOTE:** Please see Table of Specific Reagents/Equipment for a detailed description of materials and reagents mentioned in this protocol.

3. **Primary and Secondary Enrichment**

   1. Label all primary and secondary enrichment tubes containing media according to the sample number and the body part of the fly.

   2. Under a sterile hood, transfer 300 µl of BPW-S (surface) to sterile 2 ml tubes containing the following media:

      1. **For Salmonella**, use 1 ml of pre-warmed (42 °C) BPW. Incubate in a recirculating water bath at 42.5 °C for 22 - 24 hr. For secondary enrichment, transfer 100 µl of enriched BPW to 400 µl of pre-warmed (37 °C) brain heart infusion (BHI) broth previously placed in sterile cluster tubes. Incubate at 37 °C for 3 hr.
2. **For Cronobacter**, use 1 ml of pre-warmed (37 °C) BPW with novobiocin (10 mg/L; Wallace, M., personal communication). Alternatively, use 1 ml of the R & F Enterobacter sakazakii enrichment broth with supplement (vancomycin and ceftazidim) as primary enrichment. Incubate at 37 °C for 22 - 26 hr. For secondary enrichment, transfer 100 µl of enriched BPW with novobiocin to 400 µl of pre-warmed (37 °C) BHI broth previously placed in sterile cluster tubes. Incubate at 37 °C for 3 hr.

3. **For L. monocytogenes**, use 1 ml of freshly prepared RT 24 Listeria enrichment broth (24 LEB) with selective supplement. Incubate at 37 °C for 44 ± 5 hr. No secondary enrichment is required for the detection of *L. monocytogenes*.

4. Repeat steps 3.2.1 - 3.2.3 above using the tube labeled as BPW-A.

4. **Preparation of the PCR-Based System for Amplification and Detection of the Target Foodborne Pathogen**

Steps 4-8 use a commercial PCR cycler/detector system, a computer workstation, and ready-to-use kits to screen for *Salmonella* (*Salmonella* 2 standard assay kit), *Cronobacter* species (*E. sakazakii* standard assay kit), and *Listeria monocytogenes* (*L. monocytogenes* 24E assay kit). Standard assays use PCR end-point detection. Each kit contains PCR-ready tablets with an intercalating dye that emits a fluorescence signal when binding to double-stranded DNA. The signal is captured during the detection phase of the PCR system program, generating a melting curve that is interpreted by the software as positive or negative.

1. Prepare reagents and equipment as specified by the manufacturer’s protocol per each target foodborne pathogen. **NOTE**: The protocols for detecting *Salmonella* and *Cronobacter* require a one-step lysis procedure whereas the protocol for detecting *L. monocytogenes* requires a two-step lysis procedure (see sections 5 and 6, respectively).

2. Turn on the automated heating block selecting the specific program for the target pathogen. Alternatively, if the heating blocks are manual, set temperatures to 37 °C (for *Salmonella*, *Cronobacter* spp., and *L. monocytogenes*) or to 55 ± 2 °C (for *L. monocytogenes* part 2 of lysis – see step 6.2) and 95 ± 3 °C.

3. Make sure that the cooling blocks have been refrigerated O/N, otherwise chill them at 2 - 8 °C for at least 2 hr.

4. Using the computer software of the PCR-based detection system, create a rack file following manufacturer’s instructions.

5. Label and arrange cluster tubes containing the lysis reagent in the rack, according to the rack file.

6. Initialize the PCR-based detection system instrument.

5. **Perform Lysis for the Detection of Salmonella and Cronobacter**

1. Prepare the lysis reagent by adding 150 µl of protease to one 12 ml bottle of lysis buffer.

2. Transfer 200 µl of lysate from each of the previously labeled cluster tubes. **NOTE**: Cluster tubes containing the lysis reagent can be stored at 2 - 8 °C for up to 2 weeks.

3. Using long pipette tips, transfer 20 µl of secondary enriched samples (see steps 3.2.1 and 3.2.2) to corresponding cluster tubes containing 200 µl of lysis reagent. Use new pipette tips for each sample. **NOTE**: Keep tubes from primary and secondary enrichment in the refrigerator (at RT, <4 °C) for further confirmation analysis of PCR-positive/ negative samples.

4. Prepare negative controls by adding 20 µl of sterile BHI media to cluster tubes containing 200 µl of lysis reagent.

5. Prepare positive controls by adding 20 µl of O/N bacterial cultures (grown in BHI) of any known *Salmonella* or *Cronobacter* strain to cluster tubes containing 200 µl of lysis reagent.

6. Cap cluster tubes and secure tightly using the capping tool.

7. Place the rack of cluster tubes in the automated heating block after selecting the specific program for the target pathogen. Alternatively, incubate cluster tubes at 37 ± 2 °C for 20 min, followed by incubation at 95 ± 3 °C for 10 min. Finally, transfer the cluster tubes to cooling blocks (2 - 8 °C) for 5 min. **NOTE**: Cluster tubes containing the lysate can be stored at -20 °C for up to 2 weeks.

6. **Perform Lysis for the Detection of L. monocytogenes**

1. Prepare 1.8 ml of sterile deionized water to the bottle of fully thawed lysing agent 1. **NOTE**: Store lysing agent 1 at 2 - 8 °C until ready to use. After opening and diluting, store at RT (20 - 30 °C) for up to 6 months.

2. Combine lysing agents 1 and 2 in a 4:1 ratio (40 µl of diluted lysing agent 1 and 10 µl of lysing agent 2 per each sample). Transfer 50 µl of the combined lysing agents to cluster tubes. Use the mixture within 4 hr.

3. Add 500 µl of primary enriched sample (see step 3.2.3) to the cluster tube containing the 50 µl of the combined lysing agents.

4. Prepare a negative control by adding 500 µl of sterile 24 LEB to 50 µl of the combined lysing agents.

5. Prepare a positive control by adding 500 µl of O/N *L. monocytogenes* culture grown in 24 LEB to 50 µl of the combined lysing agents.

6. Cap the cluster tubes, mix gently and place in heating block at 37 ± 1 °C for 30 min. **NOTE**: Keep tubes from primary enrichment in the refrigerator for further confirmation analysis of PCR-positive/negative samples.

2. Perform part 2 of lysis as follows:

   1. Prepare the lysis reagent as instructed in steps 5.1 and 5.2.

   2. Using long pipette tips transfer 20 µl of part one lysate to cluster tubes containing 200 µl of lysis reagent. Use new pipette tips for each sample.

   3. Cap cluster tubes and secure tightly using the capping tool.

   4. Incubate cluster tubes at 37 °C for 44 ± 5 hr. No secondary enrichment is required for the detection of *L. monocytogenes*.

   5. Cap the cluster tubes, mix gently and place in heating block at 37 ± 1 °C for 30 min. **NOTE**: Keep tubes from primary enrichment in the refrigerator for further confirmation analysis of PCR-positive/negative samples.

   6. Cap the cluster tubes, mix gently and place in heating block at 37 ± 1 °C for 30 min. Finally, transfer the cluster tubes to cooling blocks (2 - 8 °C) for 5 min.
7. Hydrate PCR-Ready Tablets

1. Select a chilled (4 °C) PCR cooling block and place a PCR tube rack over the insert.
2. Place corresponding PCR tubes containing the PCR-ready tablets (included with each kit) for the target foodborne pathogen in the holder, according to the rack file.
3. Using the decapping tool, carefully remove the caps from PCR-tubes. Discard the caps and verify that each tube contains a tablet.
4. Transfer 50 µl (for *Salmonella* and *Cronobacter*) or 30 µl (for *L. monocytogenes*) of lysate to specific PCR tubes. Use new optical caps and secure tightly onto the PCR tubes using the capping tool.

NOTE: After adding the lysate to PCR-ready tablets, samples must remain chilled at 2 - 8 °C until loaded into the PCR-based detection system. The PCR tubes can be centrifuged at 2,500 x g for a few seconds to assure that the full volume is in the bottom of the tube.
5. Load the PCR tubes into the PCR cycler/detection system instrument by opening the instrument drawer.
6. Place the rack of PCR tubes into the wells in the drawer and check that the tubes are seated correctly.
7. Close the drawer and initiate the program as described by the manufacturer’s protocol.

NOTE: The PCR-based instrument has preset cycling parameters for each foodborne pathogen.
8. Verify that the PCR cycling status bar displays a blue bar indicating that the amplification portion of the program is running.

NOTE: For standard PCR assays, the processing time of the full program (amplification and detection) takes approximately 3 - 3.5 hr to complete.

8. Review Results

1. After processing is complete, follow the screen prompts from the PCR-based system instrument to remove samples and review results.
2. If the target foodborne pathogen is present in the sample (either the surface or the alimentary canal of the fly) the well is red with a ‘plus’ sign (positive). If the pathogen is absent, the well is green with a ‘minus’ sign (negative).
3. If the well is yellow with a red bar across the center, it indicates a signal error.

9. Isolation of Bacterial Pathogens from PCR-Positive Results

1. Select tubes from the primary (for *L. monocytogenes*) or secondary (for *Salmonella* and *Cronobacter*) enrichment of those samples that were PCR-positive. Also, randomly select 3 - 5% of samples that were PCR-negative and proceed as follows:

2. For *Salmonella*:
   1. Add 100 µl of the secondary enrichment media to 10 ml of Rappaport-Vassiliadis (RV) medium and to 1 ml of tetrahionate (TT) broth. Incubate tubes at 42.5 °C in a recirculating water bath for 22 - 24 hr.
   2. After incubation, streak a 3 mm loopful (10 µl) of each, RV and TT media on bismuth sulfite (BS) agar, xylose lysine desoxycholate (XLD) agar, and Hektoen enteric (HE) agar. Incubate plates at 35 ± 1 °C for 22 - 24 hr.
   3. After incubation, examine plates for the presence of typical *Salmonella* colonies on each media. If no isolated colonies can be obtained after several sub-culturing steps, consider the sample as negative and report as a false positive for the PCR-based system.

NOTE: For typical *Salmonella* colonies on specific media see 22. Select five presumptive typical *Salmonella* colonies and subculture them on BS, XLD, or HE until pure cultures of isolated/single colonies are obtained.
4. Select one pure colony and identify presumptive *Salmonella* by using biochemical commercial tests such as the VITEK 2 identification card or API biochemical identification system, following the manufacturer’s instructions.

3. For *Cronobacter*:
   1. Streak a 3 mm loopful (10 µl) of the secondary enrichment media on two plates of chromogenic culture media such as R&F *Enterobacter sakazakii* (Cronobacter) chromogenic plating medium, and/or ChromID Sakazakii Agar. Incubate plates at 35 °C for 22 - 24 hr.
   2. After incubation, examine plates for the presence of typical *Cronobacter* colonies (blue-black to blue-gray). Select 5 presumptive *Cronobacter* colonies and subculture them onto R&F *Enterobacter sakazakii* (Cronobacter) chromogenic plating medium, ChromID Sakazakii Agar, or TSA until pure cultures of isolated/single colonies are obtained.

NOTE: If no isolated colonies can be obtained after several sub-culturing steps, consider the sample as negative and report as a false positive for the PCR-based detection system.
3. Select one pure colony and identify presumptive *Cronobacter* by using biochemical commercial tests such as the VITEK 2 identification card or API 20E biochemical identification system, following the manufacturer’s instructions.

4. For *L. monocytogenes*:
   1. Streak a 3 mm loopful (10 µl) of the primary enrichment media on two plates of Brilliance Listeria agar (BLA). Incubate plates at 36 ± 1 °C for 22 - 26 hr.
   2. After incubation, examine plates for the presence of presumptive *L. monocytogenes* (blue-green) colonies. Select 5 presumptive *L. monocytogenes* colonies and subculture them on BLA until obtaining pure cultures of isolated/single colonies. Re-incubate negative plates at 36 ± 1 °C for an additional 22 - 26 hr.
   3. Select one pure colony and identify presumptive *L. monocytogenes* by using commercial biochemical tests such as the VITEK 2 identification card or API Listeria biochemical identification system, following the manufacturer’s instructions.

5. Presumptive foodborne pathogens isolated from insects should be further confirmed and serotyped in a reference laboratory.

NOTE: Cluster tubes containing the lysate can be stored at -20 °C for up to 2 weeks.
Representative Results

This protocol was first calibrated on a set of lab-reared house flies that were experimentally fed for 24 hr with liquid fly food (2% powder milk) containing serial dilutions (10^2 – 10^8 CFU/ml) of *C. sakazakii*, *S. enterica*, *L. monocytogenes*, or *C. jejuni* (*n* = 21 for each bacterial pathogen). Enrichment media as well as incubation times and temperatures were adjusted for each foodborne pathogen until the PCR-based system was able to detect the lowest levels of bacteria (10^2 CFU/ml) from the body surface and the alimentary canal of a single experimentally fed fly. Using the enrichment media and conditions described in the protocol section, the PCR-based system detected *C. sakazakii*, *S. enterica*, and *L. monocytogenes* from the body surface of 100% of flies fed with bacterial inocula >10^6 CFU/ml (Figure 1A). When flies were fed with 10^2 CFU/ml, the percentage of detection of *C. sakazakii*, *S. enterica*, and *L. monocytogenes* from their body surface was 100%, 66%, and 33%, respectively (Figure 1A). The PCR-based system also detected these three foodborne pathogens from the alimentary canal of flies fed with all bacterial concentrations at percentages ≥33% (Figure 1B). However, the detection of *C. jejuni* was only achieved when lab-reared flies were experimentally fed with liquid food containing the highest bacterial inoculum (10^6 CFU/ml). Hence, *C. jejuni* was excluded from the group of foodborne pathogens that could be tested from individual synanthropic filth flies using this PCR-based detection system.

With this standardized protocol, we were able to determine the prevalence of *Cronobacter* spp., *S. enterica*, and *L. monocytogenes* from the body surface and/or the alimentary canal of 100 wild flies that were individually and aseptically caught from the dumpster area of ten urban restaurants located in the metropolitan area of Washington, D.C. Collected filth flies were representative of at least six species including *M. domestica* (47%), *Lucilia cuprina* (33%), *L. sericata* (14%), *Cochliomyia macellaria* (2%), *Sarcophaga haemorrhoidalis* (2%), and *Ophyra leucostoma* (1%). One fly was identified only to family level (Anthomyiidae; 1%). The surface-disinfection protocol was effective at avoiding cross-contamination between the body parts of the fly because no bacterial growth was observed on TSA plates for water from the last disinfection rinse of each individual fly. Thus, a distinction could be made between foodborne bacteria present on the body parts of each fly.

No false positives were detected from samples of the body surface and the alimentary canal of individual flies when using this commercial PCR-based system for the detection of *S. enterica* and *L. monocytogenes*, and the confirmation of viable pathogens on agar plates was in agreement with PCR-positive results. However, it was not possible to isolate pure cultures of *Cronobacter* spp. from all PCR-positive samples. Hence, the detection of this pathogen by the PCR-based system showed false positives from the body surface (50%; 9/18) and the alimentary canal (48%; 16/33) of single wild-caught flies. Randomly selected PCR-negative samples that were plated on specific media, confirmed the absence of the foodborne pathogens. Therefore, no false negatives were detected from any of the samples when using this commercial PCR-based system to detect *Cronobacter* spp., *S. enterica*, or *L. monocytogenes*.

Only those PCR-positive samples where the pathogen was isolated and confirmed were considered positive and included for statistical analysis. The overall prevalence of foodborne pathogens in the alimentary canal of wild-caught filth flies was significantly higher than on the body surface (χ^2 = 6.8772, df = 1, *p* = 0.0087). 22% of the alimentary canals and 6% of the body surfaces of collected wild flies were positive for at least one of the three foodborne pathogens (Figure 2). Overall, the prevalence of *Cronobacter* spp. on either the body surfaces or alimentary canals of collected flies was statistically higher (19%; Fisher’s exact test *p* = 0.0165) than the prevalence of *S. enterica* (7%) and *L. monocytogenes* (4%). However, no statistical differences were observed when performing pairwise comparisons between the body parts of the flies for each bacterial pathogen (Figure 3; Fisher’s exact test *p* = 0.1464, *p* = 0.1184, and *p* = 0.6212 for *Cronobacter* spp., *S. enterica*, and *L. monocytogenes*, respectively). None of the flies were positive for all three pathogens evaluated. However, three of the flies (two *L. cuprina* and one *L. sericata*) carried *Salmonella* spp. and *L. monocytogenes* on the surface or in the alimentary canal.
Figure 1. Detection levels of *Cronobacter sakazakii*, *Salmonella enterica*, *Listeria monocytogenes*, and *Campylobacter jejuni* from (A) the body surface and (B) the alimentary canal of individual lab-reared house flies fed with liquid food containing different bacterial inocula (n = 21 for each bacterial pathogen, n = 3 per each bacterial concentration).
Figure 2. Percentage of body surfaces and alimentary canals of individual flies found positive for any of the target foodborne pathogens.

Figure 3. Prevalence of *Cronobacter* spp., *Salmonella enterica*, and *Listeria monocytogenes* from the body surface and the alimentary canal of synanthropic wild-caught flies. The *p* values reported are from pairwise comparisons between the body surface and the alimentary canal for each bacterial pathogen (Fisher’s exact test, *p* value < 0.05 indicates statistical significance). Copyright © American Society for Microbiology, Journal of Applied and Environmental Microbiology 78 (22):7891-902, 2012. doi: 10.1128/AEM.02195-12.

**Discussion**

Previous studies that have detected foodborne pathogens from wild insects have used a great variety of protocols that might not include the necessary information to accurately assess the food-related risk of the presence of a single fly in foods or food-related environments. Here, we demonstrated that using this standardized protocol, it is possible to detect and isolate *Cronobacter* spp., *S. enterica*, and *L. monocytogenes* from the body surface and the alimentary canal of single flies caught in the wild. Because insects may carry low numbers of the target foodborne pathogen and high numbers of other indigenous microbiota, this protocol requires primary (and sometimes secondary) enrichment of the samples in specific culture media to increase the sensitivity of detection of the target foodborne pathogen. Results from the PCR-based detection system were obtained within approximately 30 hr (for the detection of *Cronobacter* spp. and *S. enterica*) and 48 hr (for...
the detection of *L. monocytogenes*) after initially processing the samples. Thus, this protocol is reliable as well as rapid and sensitive enough to screen a single fly for the presence of foodborne pathogens.

Confirmation of PCR-positive results and isolation of viable bacteria is part of the standard operating procedure of many laboratories. In addition, for epidemiology purposes, pure bacterial cultures from PCR-positive samples are required to further confirm and serotype the foodborne pathogen using biochemical, immunological, or genetic methods. Although no false positives were observed when detecting *S. enterica* and *L. monocytogenes* from the body parts of single wild-caught flies, using this protocol, we found up to a 50% rate of false positives for *Cronobacter* spp. This suggests that the PCR-based detection system for the genus *Cronobacter* may cross-react with other *Enterobacteriaceae* present among the highly complex microbiota carried by flies. Thus, isolation and purification of pure colonies of the genus *Cronobacter* from PCR-positive samples require more selective plating than the other pathogens evaluated.

This protocol has primarily been standardized to screen individual wild-caught flies for the presence of *Cronobacter* spp., *S. enterica*, and *L. monocytogenes* using a commercial PCR-based detection system. However, this protocol was also easily adapted to screen body parts of single flies for the presence of other foodborne pathogens such as enterohemorrhagic *E. coli* O157:H7 (using either the *E. coli* O157:H7 MP standard assay kit or the *E. coli* O157:H7 real-time assay kit) and the shiga-toxigenic *E. coli* (STEC) group (using the real-time STEC suite), obtaining sensitivities >80% (unpublished data). Also, this protocol can potentially be adapted to detect foodborne pathogens from other insects that are known vectors of diseases (cockroaches and ants), but more research in this area is needed.

Foodborne illness outbreak investigations are very dynamic and comprise a multi-step process that may vary according to the specific situation and the local environment being investigated12,27. These investigations are important because they provide immediate public health protection by preventing future illnesses. Additionally, these investigations can elucidate new mechanisms by which foodborne microorganisms are spread, and raise important questions that lead to new areas for research26. Investigative techniques as well as standardized, rapid, and sensitive protocols are necessary for detecting foodborne pathogens from individual insects. This standardized protocol opens the opportunity to aseptically collect insects like flies, which can vector the foodborne bacterial pathogen, as part of an environmental sampling program. The epidemiological information that can be gained from this would be of use in constructing an accurate picture of the mechanisms of transmission of foodborne pathogens by insects (i.e., length of exposure time: a fly by landing versus flies landing, defecating, and regurgitating).

Finally, even though the commercial PCR-based detection system described here is practical to use and simplifies PCR amplification and visualization of a genus-level amplicon, it is by no means the only appropriate system. The lysate from enriched samples could alternatively be used to screen for the presence of foodborne pathogens by using publically available species-specific primer pairs. However, detection sensitivity should be demonstrated prior to their use.

Disclosures

The use of specified instrumentation is not an endorsement by the U.S. Food and Drug Administration. The authors certify that there is no conflict of interest with any financial organization regarding the material discussed in this article.

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