Development and Comparison of Complementary Methods to Study Potential Skin and Inhalational Exposure to Pathogens During Personal Protective Equipment Doffing

Jennifer Therkorn,1 David Drewry,1 Jennifer Andonian,2 Lauren Benishek,3 Carrie Billman,7 Ellen R. Forsyth,1 Brian T. Garibaldi,2 Elaine Nowakowski,2 Kaitlin Rainwater-Lovett,1 Lauren Sauer,4,5 Maggie Schiffhauer,2 and Lisa L. Maragakis2,3,4; for the CDC Prevention Epicenters Program

1Applied Biological Sciences, Johns Hopkins Applied Physics Laboratory, Laurel, and 2Department of Hospital Epidemiology and Infection Control, Johns Hopkins Health System, 3Johns Hopkins University School of Medicine, and 4Johns Hopkins Bloomberg School of Public Health, Baltimore, Maryland

Background. Fluorescent tracers are often used with ultraviolet lights to visibly identify healthcare worker self-contamination after doffing of personal protective equipment (PPE). This method has drawbacks, as it cannot detect pathogen-sized contaminants nor airborne contamination in subjects’ breathing zones.

Methods. A contamination detection/quantification method was developed using 2-µm polystyrene latex spheres (PSLs) to investigate skin contamination (via swabbing) and potential inhalational exposure (via breathing zone air sampler). Porcine skin coupons were used to estimate the PSL swabbing recovery efficiency and limit of detection (LOD). A pilot study with 5 participants compared skin contamination levels detected via the PSL vs fluorescent tracer methods, while the air sampler quantified potential inhalational exposure to PSLs during doffing.

Results. Average PSL skin swab recovery efficiency was 40% ± 29% (LOD = 1 PSL/4 cm² of skin). In the pilot study, all subjects had PSL and fluorescent tracer skin contamination. Two subjects had simultaneously located contamination of both types on a wrist and hand. However, for all other subjects, the PSL method enabled detection of skin contamination that was not detectable by the fluorescent tracer method. Hands/wrists were more commonly contaminated than areas of the head/face (57% vs 23% of swabs with PSL detection, respectively). One subject had PSLs detected by the breathing zone air sampler.

Conclusions. This study provides a well-characterized method that can be used to quantitate levels of skin and inhalational contact with simulant pathogen particles. The PSL method serves as a complement to the fluorescent tracer method to study PPE doffing self-contamination.

Keywords. personal protective equipment doffing; exposure assessment; doffing self-contamination; inhalational exposure; methods development.

In September 2015, 2 healthcare workers at Texas Health Presbyterian Hospital became infected with Ebola virus disease (EVD) after providing care for a patient infected with Ebola virus [1, 2]. This case raised issues regarding whether the appropriate protections were provided to staff members, including training for safe and effective use of personal protective equipment (PPE), particularly the PPE doffing process. Proper PPE doffing is vital to reduce the risk for self-contamination and is an area of focus to ensure healthcare worker safety [3–7].

Given the risk of self-contamination during PPE doffing procedures, it is important to assess the likelihood of contamination at each point of the process. Furthermore, enhanced PPE ensembles for care of patients with high-consequence pathogens have multiple layers, adding to the complexity of doffing practices and increasing the risk of self-contamination [2, 6, 8–10]. Commercially available fluorescent products (liquid, lotion, or powder) used as tracers in combination with ultraviolet (UV) lights are frequently used as a study method to detect PPE doffing self-contamination [3, 6, 11, 12]. This approach provides feedback to healthcare workers as they can visualize where they have contaminated themselves [3]. However, this detection method has limitations. It has a relatively low sensitivity of detection, as eyesight alone cannot identify pathogen-sized fluorescent particles that may constitute contamination. Second, the commonly used fluorescent powders that dissolve into a liquid suspension may not fully or accurately represent the full spectrum of exposure risks.

Specifically, recent research has shown that there is the potential...
for reaerosolization of infectious particles from PPE during doffing [13], and this type of potential inhalational exposure has not been addressed by the fluorescent tracer method.

This study presents the development of a more sensitive method to investigate self-contamination using fluorescent, bacteria-sized (2 µm) polystyrene latex spheres (PSLs); additionally, for the first time, potential inhalational exposure for healthcare personnel during PPE doffing is investigated with the use of an air sampler to quantify PSLs in the doffer’s breathing zone. This approach is then directly compared to the traditional fluorescent tracer method in a pilot study to investigate self-contamination levels using the Centers for Disease Control and Prevention (CDC) PPE doffing guidelines for providers caring for patients with EVD or other high-consequence pathogens [14].

METHODS
This study encompasses 2 main components: (1) PSL contamination and detection methods development; and (2) a PPE doffing pilot study to evaluate the PSL method vs the fluorescent tracer method (Table 1). PSLs are a nonhazardous, inert substance. More background information on the approach and rationale for selection of the PSLs can be found in Drewry et al [13].

PSL Protocols: Skin Swabbing and PSL Quantification by Microscopy
To establish the limit of detection for the PSL method, known quantities of PSLs were pipetted onto and then sampled from porcine skin coupons using swabs, and the full procedural recovery efficiency was estimated (ie, from skin swabbing to elution of PSLs from the swab tip). Given the potential for procedural loss of PSLs from skin swabbing, this experiment with skin coupons allowed for determination of the correction factor to use in the pilot study doffing experiments to scale up the observed numbers of PSLs recovered from skin swabs to estimate the actual total number of PSLs present on skin.

The process used 4500 g of porcine skin, a human skin substitute [17–19], with a skin thickness of about 1.5 cm. The skin was cut into rectangular-shaped coupons (50 cm²) using a scalpel and cleaned with 70% ethanol followed by a disinfecting wipe (Oxivir wipes, Diversey, Charlotte, North Carolina). The disinfectants were allowed to dry on the skin coupons, and then they were put in plastic zip bags and frozen (−20°C) until use. The coupons were removed from the freezer and thawed at 4°C for 24 hours prior to use.

The PSL solution was serially diluted to achieve concentrations of 10²–10⁶ PSL/mL. Using the methods of Therkorn et al [20], the 5 PSL concentrations were spiked (100 µL) onto the skin coupons, with each dilution spiked on 5 different skin samples (25 total coupon samples). For each of the spiked PSL concentrations, there were 5 control standards prepared where 100 µL was pipetted directly into 1 mL of filtered, deionized (DI) water.

After allowing the spiked droplets to dry for 5 hours in a biosafety cabinet, each skin coupon was swabbed to collect PSLs. Each coupon was swabbed with 1 sterile foam tipped swab

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**Table 1. Comparison of Polystyrene Latex Spheres and Fluorescent Tracer Methods for Studying Self-contamination During Personal Protective Equipment Doffing**

| Method description | PSL Method | Fluorescent Tracer Method |
|--------------------|------------|---------------------------|
| Method description | 2-µm PSLs (G0200, Thermo Fisher Scientific, Waltham, Massachusetts) diluted in water | Fluorescent powder slurry (Glitter Bug, Brevis Corporation, Salt Lake City, Utah) mixed in a viscous suspension of water and oil |
| Suspension composition | PSLs diluted 1:10 in filtered, deionized water (10⁷ PSL/mL) | Glitter Bug powder (75 mg/mL) mixed in grapeseed oil and water (1:6 ratio oil to water) |
| Application of contaminant to healthcare worker study subjects | 25 mL of PSL suspension in 3-jet Collison nebulizer (Mesa Laboratories, Butler, New Jersey); 4 min of continuous aerosol generation while healthcare worker turned 90° every 60 sec | 1000 mL of mixture in a pesticide hand sprayer (RL Flo-Master 2000 mL capacity, Lowell, Michigan); 5 sweeping passes of sprayer from head to feet on front and back of healthcare worker |
| Simulated contamination type | Representative of pathogen dispersal via respiratory secretions, such as coughing and sneezing [13, 15] | Representative of a wet patient and gross liquid contamination [8]; the grapeseed oil was included to represent a viscous bodily fluid |
| Sampling method | From skin: sterile foam-tipped swabs (13-cm handle, scored with thumb stop, Puritan Medical Products, Guilford, Maine) premoistened in filtered, deionized water with 0.05% Triton-X 100 | Not applicable |
| Detection method | PSL counting via epifluorescent microscopy | Visual inspection with a ultraviolet lamp |
| Quantification method | Skin: number of PSLs per cm² of skin on each swabbed body part | Number and relative size of contamination spots detected on skin and scrubs |

Abbreviations: PPE, personal protective equipment; PSL, polystyrene latex sphere.
premoistened in an elution suspension of filtered deionized water with 0.05% Triton-X 100 as previously recommended in similar research [21–23]. Each swab was assigned a 15-mL conical tube containing 5 mL of the elution suspension. Swabs were dipped into the elution suspension, and excess fluid was expressed from the swab by pressing against the inside of the tube. The moistened swab was then swept across the entire surface of the skin coupon 4 times in the following directions: horizontal, vertical, diagonal from top right to bottom left, and opposite diagonal direction [24]. The swab was turned each time the sweeping direction changed to expose all surfaces of the swab. Then, the swabs were placed back into the elution fluid in the conical tube and stored at 4°C until processing. As recommended, only 1 person conducted all of the swabbing to minimize sampling variability [25–27]. To ensure there was no contamination introduced onto the coupons in any of the procedures, 3 clean skin coupons were swabbed to serve as blanks.

The PSLs were eluted from the swabs using a 2-minute vortex followed by a 10-minute ultrasonication (M1800, Branson Ultrasonics, Danbury, Connecticut) [20, 28]. PSL detection and counting were conducted using a modified version of the Collection of Airborne Microorganisms on Nucleopore filters, Estimation and Analysis method [20, 29] via epifluorescence microscopy (Nikon Eclipse Ni-E, Nikon Instruments, Melville, New York). A 1-mL volume of elution suspension containing PSLs was filtered through a black polycarbonate filter (GTMP, 0.2 μm, 25 mm, Merck Millipore, Cork, Ireland). The filter was allowed to dry and then fixed onto a microscope slide. At least 40 random field views were counted and averaged, and this averaged number was multiplied by the total number of field views to estimate the number of PSLs across the entire filter [29]. To determine the PSL recovery efficiency and limit of detection, the results were then calculated as the recovered number of PSLs per swab relative to the actual spiked number of PSLs as determined from the prepared standards.

**Air Sampling for PSLs in the Subjects’ Breathing Zone**

An additional sampling method was employed to assess the possibility of exposure to aerosolized PSLs during the donning process. The Button Sampler, a commonly used personal breathing zone sampler in occupational and environmental health studies [30–33], was used to measure the concentration of airborne PSLs in the subjects’ breathing zone during PPE donning. This sampler closely follows the convention for sampling inhalable airborne particles [34]. The Button Sampler was clipped onto the shirt collar of the scrubs underneath the PPE, and the air pump of the sampler was attached to the powered air purifying respirator (PAPR) belt. Care was taken to ensure that the tubing from the Button Sampler to its pump did not interfere with the PPE placement and did not change the wearer experience. The sampler pump was turned on for collection prior to the subject entering the room for contamination. The sampler on/off button was easy to push through the PPE so as not to disturb the ensemble. The pump ran continuously throughout contamination and donning to measure potential inhalational exposures during the process. It was turned off by the trained observer after the final step in the donning process. The filter inside the Button Sampler where the PSLs were captured was recovered with clean forceps and stored in a 50-mL conical tube for each subject. The PSLs were eluted from the Button Sampler filters and enumerated using the same methods as described above for the swabs.

**Pilot Test**

A pilot test was conducted to compare the results of the PSL method to the fluorescent tracer method. The CDC’s PPE donning and doffing guidelines for care of patients with EVD were used for this pilot test [14]. Table 2 describes the elements of the PPE ensemble. Five subjects were recruited to participate who had no prior experience with the PPE and doffing guidelines. The study protocols were approved by the Johns Hopkins School of Medicine Institutional Review Board (approved 26 January 2018, IRB00084233). All participants provided written informed consent as required by the IRB protocol.

All subjects were provided paper scrubs and hospital-approved footwear and asked to remove any accessories. Each subject wiped his or her face with a skin cleansing wipe to remove makeup and excess skin oil (Gentle Skin Cleansing Cloths, Cetaphil, Galderma, Fort Worth, Texas). The study then proceeded as follows (detailed description below): (1) study subjects were checked by investigators for background fluorescence contamination on skin and scrubs in a dark room under UV light; (2) the Button Sampler was attached to the shirt collar of scrubs; (3) PPE was donned by subjects; (4) subjects were contaminated with both the fluorescent tracer and the PSLs; (5) PPE was doffed by subjects; (6) subjects were checked for fluorescent tracer contamination by UV light; and (7) subjects were swabbed for PSLs and Button Sampler filters were retrieved. To minimize any potential for unintended cross-contamination of subjects during the experiment, separate rooms were used for PPE donning, subject contamination, PPE doffing, and skin swabbing, with a designated walking path between these rooms (Figure 1).

After subjects donned their PPE according to the CDC guidelines for the care of patients with EVD, they were then contaminated with the 2 different materials: fluorescent tracer slurry followed by nebulized PSLs. Table 1 provides details on these 2 methods, including suspension composition and equipment used for spraying the contamination. Participants were instructed to stand with palms facing forward on a marked spot on the floor in the contamination room 0.5 m away from where the fluorescent tracer slurry was sprayed. A research technician ensured that the slurry was well mixed and then sprayed the subject with a pesticide hand sprayer by using a left to right...
sweeping motion (5 total passes) moving from the subject's head to feet. The subject was then instructed to turn 180°, and this contamination process was repeated on the subject's back. The fluorescent tracer slurry was developed after trying different combinations of commercially available oils, fluorescent products, and sprayers; the goal was the selection of a slurry that sprayed consistently, adhered to the PPE surface without runoff, and could be visualized under UV light.

Following contamination with the fluorescent tracer, subjects were then contaminated with the PSLs using a Collison nebulizer. Subjects were instructed to stand on another marked spot on the floor in the contamination room located 0.5 m from the outlet of the nebulizer. The nebulizer was on a stand placed at 1.5 m height. Subjects were instructed to face toward the nebulizer with palms forward, and then PSL nebulization began by supplying air to the aerosol generator (6 L/minute, 20 lbs per square inch gauge). The subjects were then asked to turn 90° every 60 seconds to ensure all surfaces of their external PPE were exposed to the nebulized PSLs. Following the 4-minute nebulization time, 2 high-efficiency particulate air filtration units (dry filter unit-1000, Lockheed Martin Integrated Systems, Gaithersburg, Maryland) were turned on operating at a combined 5000 L/minute and allowed to run for 5 minutes to fully purge the air of the contamination room prior to the subject exiting; the contamination room was approximately 25 m³ volume. An aerodynamic particle sizer (TSI, Shoreview, Minnesota) was included in the contamination room to monitor the particle number size distribution of the nebulized PSLs and the background levels of air contamination in between subjects. The average peak PSL concentration during contamination was

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**Table 2. Elements of the Personal Protective Equipment Ensemble Tested in the Pilot Study**

| PPE Item        | Product Name                                      | Manufacturer                                      |
|-----------------|---------------------------------------------------|--------------------------------------------------|
| Gown            | SmartGown impervious surgical gown                | Cardinal Health (Dublin, Ohio)                   |
| Gloves (outer)  | Biogel Skinsense (synthetic [nonlatex] polychloroprene surgical glove) | Mölnlycke (Gothenburg, Sweden)                   |
| Gloves (inner)  | Synthetic (nonlatex) polychloroprene surgical glove (synthetic [nonlatex] polychloroprene surgical underglove) | Mölnlycke                                          |
| Isolation gown  | MediChoice overhead poly-coated gown              | Owens and Minor (Richmond, Virginia)             |
| Boot covers     | Hi Guard regular full coverage boot, universal size | Kimberly Clark (Irvine, Texas)                   |
| Belt-mounted high efficiency PAPR | Air-Mate Assembly 231-01-30 | 3M (Maplewood, Minnesota) |
| Tape            | Duct tape                                          | 3M                                               |
| PAPR hood       | White Respirator Hood BE-10–3 (regular, Tychem double shroud) | 3M                                               |

Abbreviation: PAPR, powered air purifying respirator; PPE, personal protective equipment.

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**Figure 1.** Map of the study area indicating subject walking path and separate rooms for each activity: (1) personal protective equipment (PPE) donning; (2) contamination; (3) PPE donning and visible (Glitter Bug) contamination check; and (4) skin swabbing for polystyrene latex spheres. Abbreviations: PPE, personal protective equipment; PSLs, polystyrene latex spheres.
216 PSL/L air ± 488 PSL/L air. Past studies have shown that coughing can produce up to about $10^5$ particles/L air with a mean size by particle number of 1–2 μm [35]. Therefore, the PSL contamination in the present study is not outside of a realistic range for respiratory particulate that could be generated by a patient. For an illustration of the nebulized PSL particle number size distribution during subject contamination, see Supplementary Figure 1.

The subjects then doffed the PPE according to the CDC guidelines for care of patients with EVD including the use of a trained observer and doffing assistant [14]. The same individuals served in these roles for each participant, and the doffing assistant changed PPE between each participant to minimize the risk for subject cross-contamination. After doffing, the room was fully darkened, and a UV light was used to locate the areas for fluorescent tracer contamination. These were recorded on a standard sheet for each participant indicating locations and relative sizes of contaminated skin and scrub areas (Supplementary Figure 2). Finally, the subjects proceeded to the final station for PSL skin swabbing. The locations for skin swabbing (Table 3) were informed by subject matter expertise in healthcare, with past research indicating the most likely area for doffing self-contamination to be the hands, and the most common activity to result in self-contamination to be during removal of the PAPR hood [5, 7, 9, 11, 36, 37]. For more details on the derivation of skin surface area per swab location, see Supplementary Table 1.

RESULTS

PSL Protocols: Skin Swabbing and PSL Quantification by Microscopy

The overall average recovery efficiency for swabbing the PSLs from the porcine skin coupons was 40% ± 29% (Table 4). The precision, as indicated by the coefficient of variation (CV), ranged from 21% to 91% (Table 4). The CV tended to decrease as the spiked PSL number increased.

The theoretical limit of detection (LOD) for the PSL swabbing can be described as follows [24]:

$$\text{LOD} = \frac{1 \text{ PSL} \times \left(\frac{\text{total eluted volume}}{\text{filtered volume}}\right)}{\text{Swab recovery efficiency (\%)} \times \text{Swabbed skin area (cm}^2\text{)}}$$

It is assumed that at least 1 PSL is required for detection as indicated in the equation above. For the present experiments, 1 mL of the 5-mL eluted swab suspension was filtered through the black polycarbonate filter. Assuming an average swab recovery efficiency of 40% and a swabbed surface area of 50 cm² per porcine coupon, this results in a theoretical LOD of 0.25 PSL/cm², or about 13 PSL per skin coupon. As illustrated in Figure 2, the number of PSLs recovered was found to be linear in relation to the number of PSLs applied to the skin coupon, with variability in swabbing recovery generally decreasing as the number of spiked PSLs increased. The high variability for the lowest number of spiked PSLs (~10) and the inability to detect PSLs on all 5 spiked coupons for this number support the validity of the theoretical LOD calculated above.

Pilot Test: Skin Swabbing and Button Sampler Results

All 5 of the subjects completed the full study protocol with an average doffing time of 8 minutes ± 2 minutes. Figure 3 illustrates the locations and relative sizes and numbers of contamination spots for each subject for the 2 types of contaminations tested in the pilot study. There are 3 apparent trends demonstrated by Figure 3: (1) All subjects were contaminated. The locations that were contaminated across every subject included the hands, face/head, and legs. (2) Only 2 of 5 subjects had overlapping detection of both types of contamination on a wrist (subject 1) and back/front of hands (subject 5). For all other subjects, the PSL method enabled a more sensitive means of skin contamination detection that was not detectable by the fluorescent tracer method alone. (3) The PSL method resulted in the identification of more skin contamination than the fluorescent method, while the fluorescent tracer method enabled detection of contamination on scrubs and shoes in addition to skin.

Figure 4 illustrates the number of PSLs recovered from all of the skin swabbing locations for each subject; results are divided into 2 graphs (head/face vs hands/wrists) to better illustrate trends in contamination. Across all subjects, the hands/wrists were more commonly contaminated than areas of the head/face with 57% (17/30 swabs) vs 23% (7/30 swabs) of swabs resulting in PSL detection, respectively. For those swabs where PSLs were detected, the numbers ranged from 1 PSL/cm² to 1390 PSL/cm² on head/face, and 1 PSL/cm² to 192 PSL/cm² on hands/wrist. This highlights that the study subjects varied greatly in individual tendencies for self-contamination.

While all subjects had varying levels of PSL skin contamination, only subject 4 had PSLs detected in the breathing zone as indicated by the Button Sampler; given that subject 4 had relatively low levels of skin PSL contamination, there does not appear to be a clear relationship between skin contamination and breathing zone presence of PSLs (Figure 5).

DISCUSSION

The results obtained in the present study for porcine skin swabbing recovery efficiencies of PSLs (40% ± 29%) are similar to the results reported for previous studies investigating the recovery efficiency of Bacillus anthracis spores from stainless steel coupons using premoistened macro-foam swabs. Spiking spore numbers of $10^8$ to $10^7$ and eluting swabs with a 2-minute vortex, average recovery efficiencies have been reported ranging from 16% to 64% with overall averages typically in the range of 25%–45% [22, 24, 38]. The precision for the PSL recovery efficiency (CV = 21%–91%) was also similar to previous studies investigating the same range of spiked B. anthracis spore numbers from steel coupons.
### Table 3. Locations for Skin Swabbing, Estimated Total Skin Surface Areas, and Respective Limits of Detection for Polystyrene Latex Spheres

| Swab Location\(^a\) | Description | Approximate Skin Surface Area\(^b\), cm\(^2\) | Theoretical Limit of Detection for Swabbed Area\(^c\), No. of PSLs |
|----------------------|-------------|----------------------------------|--------------------------------------------------|
| Forehead             |             | 51.4                             | 13                                               |
| Right and left cheek |             | 31.9                             | 8                                                |
| Chin                 |             | 13.7                             | 3                                                |
| Right and left outer rim of ear | | 6.1                             | 2                                                |
| Right and left inner wrist | | 40.6                             | 10                                               |
| Right and left back of hand (fingers not included) | | 74.1                             | 19                                               |
| Index to thumb       |             | 18.2                             | 5                                                |

Abbreviation: PSL, polystyrene latex sphere.

\(^a\)Where right and left samples are indicated, these samples were taken with a separate swab for right and left.

\(^b\)For more details on the derivation of skin surface area per swab location, see Supplementary Table 1.

\(^c\)Theoretical limit of detection (LOD) calculated using equation 1 and results presented in Table 3 indicating the overall recovery efficiency of PSL skin swabbing to be 40%, resulting in an LOD of 0.25 PSL/cm\(^2\).

### Table 4. Results Summary From Porcine Skin Swabbing to Recover and Detect Polystyrene Latex Spheres

| Spiked No. of PSLs | Recovery, %, Average ± 1 SD | Coefficient of Variation, % | No. (%) of Skin Coupons With PSLs Detected |
|--------------------|-----------------------------|-----------------------------|-------------------------------------------|
| \(9 \times 10^5\)  | 60 ± 55                     | 91                          | 3/5 (60)                                   |
| \(4 \times 10^7\)  | 24 ± 18                     | 72                          | 5/5 (100)                                  |
| \(5 \times 10^3\)  | 49 ± 10                     | 21                          | 5/5 (100)                                  |
| \(4 \times 10^4\)  | 33 ± 17                     | 51                          | 5/5 (100)                                  |
| \(4 \times 10^5\)  | 32 ± 12                     | 36                          | 5/5 (100)                                  |

Abbreviations: PSL, polystyrene latex sphere; SD, standard deviation.
(21%–91%) [38]. However, it should be noted that, as opposed to steel coupons, the porcine skin coupons used in the present study were inherently flawed with minor imperfections, such as skin wrinkles and divots [39], likely adding to the sampling variability. Regardless, the use of skin coupons is a better representation of actual skin swabbing performed in the pilot test.

As the PSLs simulate a microorganism-sized particle, the PSL method described herein allows for quantification of the actual total number of potential pathogens that could be expected to be present on skin after doffing [13]. The PSL contamination method simulates a realistic level of contamination that a healthcare worker might experience from a coughing patient with a respiratory pathogen [35]. This has implications not only for evaluating contamination from doffing practice, but for assessing possible areas of high risk for leading to infection in a real patient care environment. Recent studies have also highlighted the importance of including a contamination technique that allows for quantitative measures of doffing self-contamination, with a means to relate this information to real-world potential for infection.
These studies have incorporated nonpathogenic viruses (bacteriophages MS2 and Φ6) into study protocols as part of the fluorescent tracer mix to compare virus contamination results to the fluorescent tracer [5, 37, 40]. However, this type of protocol incorporating nonpathogenic viruses may be logistically impractical in all clinical environments. While every subject was contaminated to some extent, only 2 of 5 subjects had overlapping detection of both types of viruses.

**Figure 4.** Skin swabbing results for polystyrene latex spheres (PSLs) detected across subjects and swab locations on the head/face vs hands/wrists after doffing of personal protective equipment. Results are presented as number of PSLs per cm² of swabbed skin, as estimated using anthropomorphic data (Supplementary Table 1). Abbreviation: PSLs, polystyrene latex spheres.
contamination (ie, PSL and fluorescent tracer) on a body part. The PSL method did enable a more sensitive means of skin contamination detection, but this new method cannot be used to universally scan scrubs for contamination; therefore, it serves as a complement to the fluorescent tracer method to study self-contamination during PPE doffing. The wide variability in the tendencies for the study subjects to contaminate themselves highlights the individualized aspect of this type of research. PPE doffing educational and training programs should therefore include individual feedback to healthcare workers to reduce overall levels of contamination and to identify and provide additional training for individuals who are at higher risk for self-contamination [9].

The finding that one of the study subjects had PSLs in their breathing zone during doffing warrants further investigation. The occurrence of PSL air contamination may be attributed to the doffing style used by this subject, and this will be explored further in a larger, future study. With an airborne PSL concentration of $9.7 \times 10^5$ PSL/L air, and assuming a mean moderate intensity, short-term inhalation rate of 27 L/minute for adults during PPE doffing. The wide variability in the tendencies for the study subjects to contaminate themselves highlights the individualized aspect of this type of research. PPE doffing educational and training programs should therefore include individual feedback to healthcare workers to reduce overall levels of contamination and to identify and provide additional training for individuals who are at higher risk for self-contamination [9].

The finding that one of the study subjects had PSLs in their breathing zone during doffing warrants further investigation. The occurrence of PSL air contamination may be attributed to the doffing style used by this subject, and this will be explored further in a larger, future study. With an airborne PSL concentration of $9.7 \times 10^5$ PSL/L air, and assuming a mean moderate intensity, short-term inhalation rate of 27 L/minute for adults [41] with an average doffing time of 8 minutes, this would result in exposure to >$2 \times 10^8$ PSLs. However, this is assuming that the PSL concentration was consistent throughout doffing. A limitation of the present study is that the use of only 1 Button Sampler throughout the contamination and doffing procedures to measure breathing zone PSLs results in an inability to distinguish when these processes the exposures may have occurred. Regardless, the potential for a high level of inhalational exposure during PPE doffing warrants further investigation. Future work will include a more thorough investigation of the potential for PSLs to be present in the doffer’s breathing zone by a series of filter samplers that turn on sequentially during PPE doffing to identify the doffing steps that introduce the most risk.

These samplers will also be placed in a grid around the doffer to characterize the PSL concentration throughout the space with fans to ensure a well-mixed study room.

CONCLUSIONS

Overall, this study provides a well-characterized method that can be used to quantitate levels of skin and inhalational contact with simulant pathogen particles. Cross-comparison of the results obtained from this new PSL method vs the traditional fluorescent tracer method suggests that future studies should use both contamination methods as complementary approaches to allow for more thorough, comprehensive, and objective contamination assessment. Further research will be required to better understand the presence of airborne particles in the breathing zone during PPE doffing, and how that may or may not represent true healthcare worker exposure.

Supplementary Data

Supplementary materials are available at Clinical Infectious Diseases online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

Notes

Acknowledgments. The authors thank contributing members of the Johns Hopkins EpiCenter team for their assistance in this project, and all of our volunteer participants.

Disclaimer. The findings and conclusions in this report are those of the authors and do not necessarily represent the official position of the Centers for Disease Control and Prevention.

Financial support. This work was supported by funding from the Prevention Epicenters Program of the Centers for Disease Control and Prevention (cooperative agreement U54CK000447).

Supplement sponsorship. This article appears as part of the supplement “Personal Protective Equipment for Preventing Contact Transmission of Pathogens: Innovations from CDC’s Prevention Epicenters Program,” sponsored by the CDC’s Prevention Epicenters Program.

Potential conflicts of interest. D. D. and K. R. L. have a patent pending (“Method for Release and Tracking of Aerosol Particles Within a Hospital Environment”). All other authors report no potential conflicts of interest. All authors have submitted the ICMJE Form for Disclosure of Potential Conflicts of Interest. Conflicts that the editors consider relevant to the content of the manuscript have been disclosed.

References

1. Frieden TR, Damon I, Bell BP, Kenyon T, Nichol S. Ebola 2014—new challenges, new global response and responsibility. N Engl J Med 2014; 371:1177-80.

2. Courage KH. How did a Dallas nurse catch Ebola? Available at: https://www.scientificamerican.com/article/how-did-a-dallas-nurse-catch-ebola/. Accessed 1 February 2019.

3. Almrid H, Koganti S, Tomas ME, Cadnum JI, Jenison A, Donkeye CT. A pilot study to assess use of fluorescent lotion in patient care simulations to illustrate pathogen dissemination and train personnel in correct use of personal protective equipment. Antimicrob Resist Infect Control 2016; 5:40.

4. Beam EL, Gibbs SG, Boulter KC, Beckerette ME, Smith PW. A method for evaluating health care workers' personal protective equipment technique. Am J Infect Control 2011; 39:415-20.

5. Gasanova LM, Teal LJ, Sickbert-Bennett EE, et al; CDC Prevention Epicenters Program. Assessment of self-contamination during removal of personal protective equipment for Ebola patient care. Infect Control Hosp Epidemiol 2016; 37:1156-61.

6. Chughtai AA, Chen X, Macintyre CR. Risk of self-contamination during doffing of personal protective equipment. Am J Infect Control 2018; 46:1329-34.
7. Mumma JM, Durwo FT, Ferguson AN, et al; Centers for Disease Control and Prevention. Epicenters Program, Division of Healthcare Quality Promotion. Human factors risk analyses of a doffing protocol for Ebola-level personal protective equipment: mapping errors to contamination. Clin Infect Dis 2018; 66:950–8.
8. Zellmer C, Van Hooft S, Sadrar N. Variation in health care worker removal of personal protective equipment. Am J Infect Control 2015; 43:750–1.
9. Kang J, O’Donnell JM, Coliaianne B, Bircher N, Ren D, Smith KJ. Use of personal protective equipment among health care personnel: results of clinical observations and simulations. Am J Infect Control 2017; 45:17–23.
10. Mitchell R, Roth V, Gravel D, et al; Canadian Nosocomial Infection Surveillance Program. Are health care workers protected? An observational study of selection and removal of personal protective equipment in Canadian acute care hospitals. Am J Infect Control 2013; 41:240–4.
11. Tomas ME, Kundrapu S, Thota P, et al. Contamination of health care personnel during removal of personal protective equipment. JAMA Intern Med 2015; 175:1904–10.
12. Zamora JE, Murdoch J, Simchison B, Day AG. Contamination: a comparison of 2 personal protective systems. CMAJ 2006; 175:249–54.
13. Drewry D, Sauer L, Shaw-Saliba K, et al. A novel approach to identify potential provider and environmental contamination on a clinical biocoutainment unit using aerosolized pathogen simulants. Health Secur 2017; 16:83–91.
14. Centers for Disease Control and Prevention. Guidance on personal protective equipment (PPE) to be used by healthcare workers during management of patients with confirmed Ebola or persons under investigation (PUIs) for Ebola who are clinically unstable or have bleeding, vomiting, or diarrhea in U.S. hospitals, including procedures for donning and doffing PPE. Available at: https://www.cdc.gov/vhf/ebola/healthcare-us/ppguidance.html. Accessed 28 September 2018.
15. Therkorn J, Drewry D III, Pilholski T, et al. Impact of air-handling system ex-fault on dissemination pattern of simulant pathogen particles in a clinical biocoutainment unit. Indoor Air 2019; 29:143–55.
16. Aizenberg V, Grinshpun SA, Willeke K, Smith J, Baron PA. Performance characteristics of the button personal inhalable aerosol sampler. AIHAJ 2000; 61:398–404.
17. Summerfield A, Meurens F, Ricklin ME. The immunology of the porcine skin and its value as a model for human skin. Mol Immunol 2015; 64:16–21.
18. Ranamukhaarachchi SA, Lehnert S, Ricklin ME, et al. A micromechanical comparison of human and porcine skin before and after preservation by freezing for medical device development. Sci Rep 2016; 6:32074.
19. Jacobs U, Kaiser M, Toll R, et al. Porcine ear skin: an in vitro model for human skin. Skin Res Technol 2007; 13:19–24.
20. Therkorn J, Thomas N, Calderon L, Scheinbeim J, Mainelis G. Design and development of a passive bioaerosol sampler using polarized ferroelectric polymer film. J Aerosol Sci 2017; 105:128–44.
21. Selwyn S, Ellis H. Skin bacteria and skin disinfection reconsidered. Br Med J 1972; 1:136–40.
22. Rose L, Jensen B, Peterson A, Banerjee SN, Srdulino MJ. Swab materials and Bacillus anthracis spore recovery from nonporous surfaces. Emerg Infect Dis 2004; 10:1023–9.
23. McGinley KJ, Webster GE, Leyden JJ. Regional variations of cutaneous propionibacteria. Appl Environ Microbiol 1978; 35:62–6.
24. Hoolges LR, Rose LJ, O’Connell H, Arudondo MJ. National validation study of a swab protocol for the recovery of Bacillus anthracis spores from surfaces. J Microbiol Methods 2010; 81:141–6.
25. Michel CE, Roper KG, Divena MA, Lee HH, Taylor HR. Correlation of clinical trachoma and infection in Aboriginal communities. PLoS Negl Trop Dis 2011; 5:e986.
26. Simpkins C, Skays MV, Hero J-M. Swabber effect: Swabbing technique affects the detectability of Batrachochytrium dendrobatidis. Herpetol Rev 2014; 45:1–5.
27. Familiar López M, Rebollar EA, Harris RN, Vredenburg VT, Hero JM. Temporal variation of the skin bacterial community and Batrachochytrium dendrobatidis infection in the terrestrial cryptic frog Phylora leveridgei. Front Microbiol 2017; 8:2535.
28. Burton NC, Adhikari A, Grinshpun SA, Hornung R, Reponen T. The effect of filter material on bioaerosol collection of Bacillus subtilis spores used as a Bacillus anthracis simulant. J Environ Monit 2005; 7:475–80.
29. Palmgren U, Ström G, Blomquist G, Malmberg P. Collection of airborne microorganisms on Nuclepore filters, estimation and analysis—CAMNEA method. J Appl Bacteriol 1986; 61:401–6.
30. Adhikari A, Martuzevicius D, Reponen T, et al. Performance of the Button personal inhalable sampler for the measurement of outdoor aeroallergens. Atmos Environ 2003; 37:4723–33.
31. Sagona JA, Shalat XL, Wang Z, et al. Comparison of particulate matter exposure estimates in young children from personal sampling equipment and a robotic sampler. J Expo Sci Environ Epidemiol 2017; 27:299–305.
32. Lee T, Grinshpun SA, Martuzevicius D, et al. Relationship between indoor and outdoor bio-aerosols collected with a button inhalable aerosol sampler in urban homes. Indoor Air 2006; 16:37–47.
33. Bidinger E, Grinshpun SA, Willeke K, Hamed A, Tabakoff W, Airflow and particle velocities near a personal aerosol sampler with a curved, porous aerosol sampling surface AU—Aizenberg, Vitaly. Aerosol Sci Technol 1998; 28:247–58.
34. Aizenberg V, Reponen T, Grinshpun SA, Willeke K. Performance of Air-O-Cell, Burkard, and Button samplers for total enumeration of airborne spores. AIHAJ 2000; 61:855–64.
35. Yang S, Lee GW, Chen CM, Wu CC, Yu KP. The size and concentration of droplets generated by coughing in human subjects. J Aerosol Med 2007; 20:484–94.
36. Lai JY, Guo YP, Or PP, Li Y. Comparison of hand contamination rates and environmental contamination levels between two different glove removal methods and distances. Am J Infect Control 2011; 39:104–11.
37. Kwon JH, Burnham CD, Reske KA, et al. Assessment of healthcare worker protocol deviations and self-contamination during personal protective equipment donning and doffing. Infect Control Hosp Epidemiol 2017; 38:1077–83.
38. Hoolges LR, Rose LJ, Peterson A, Noble-Wang J, Arudondo MJ. Evaluation of a macrofoam swab protocol for the recovery of Bacillus anthracis spores from a steel surface. Appl Environ Microbiol 2006; 72:4429–30.
39. Genzer J, Groenewold J. Soft matter with hard skin: from skin wrinkles to templating and material characterization. Soft Matter 2006; 2:310–23.
40.相似的参考文献部分。