Evaluation of Hand Sanitizer for On-Site Disinfection of Mouthguards

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Submission: February 21, 2019; Published: March 28, 2019

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Abstract

The oral cavity may be an important and overlooked reservoir of systemic infection, and athletic appliances may constitute a potentially important and under-recognized risk factor for infection. Oral appliances, e.g. toothbrushes, mouthguards and dentures, can transmit organisms responsible for both local and systemic disease [1]. Home-based sanitation procedures that are effective for cleaning toothbrushes, e.g. rinsing with soap and water, mouth rinses, and microwave irradiation, are not convenient for sanitizing mouthguards at the athletic site.

Oral appliances become colonized by microorganisms after a certain time in the mouth [2]. Gram-positive cocci, gram-positive bacilli, gram-negative cocci, gram-negative bacilli, yeasts, and molds were isolated from 81 athletic mouthguards submitted for microbial analysis [3]. The two most frequently gram-positive cocci isolated from athletic mouthguards were Staphylococcus aureus and Staphylococcus epidermidis, prominent skin pathogens that are also associated with endocarditis, pericarditis, pneumonia, osteomyelitis, food intoxication, and athletic equipment contamination [4].

The oral cavity may be an important and overlooked reservoir of systemic infection, and athletic appliances may constitute a potentially important and under-recognized risk factor for infection [5]. The possible disease transmission by contaminated athletic mouthguards, one to a leg abscess and another to lung infection, has been documented in two young football players [6]. The oral cavity is commonly colonized with S. aureus which may be associated with a wide range of serious infections, especially methicillin-resistant S. aureus (MRSA) [7]. Athletes have been the subject of an increasing number of reported MRSA cases [8].

Methicillin-resistant S. aureus is an important nosocomial pathogen, which is often transmitted from the colonized site to the site of subsequent infection and is responsible for considerable morbidity and mortality. Two types of MRSA are identified in the literature: hospital-associated MRSA and community-associated MRSA. An increase in community associated MRSA (aka close-quarter MRSA) outbreaks has been noted in basketball, football, rugby, volleyball, and wrestling athletes. and some strains of S. aureus are the most common cause of cultured skin abscesses in athletes [9]. Among athletes, MRSA may be spread from skin to skin contact with open abrasions and contaminated persons, from surface to skin contact with contaminated sports equipment (e.g. mouthguards), or by sharing personal items [9].

Good personal hygiene is the key to prevention and control of community associated MRSA outbreaks. Such practices among athletes include the following: frequent hand washing, covering abrasions or seeping wounds, disallowing athletes with open wounds in whirlpools or saunas, discouraging shared personal items, requiring showers after all practices and games, wearing sandals in showers, isolating athletes who have infections, and washing protective gear after each use [10]. Recommended infection control measures include regular and thorough cleaning of equipment, however when antimicrobial treatment is recommended, specific products are not identified.

Athletic mouthguards are difficult to clean and provide retentive sites for microorganisms. Although adequate brushing or
scrubbing with a dentifrice is an effective means of controlling the biofilm, inappropriate home-care quality and frequency are factors that compromise the efficacy of the mechanical control of biofilm on mouthguards [11]. In addition, many athletic mouthguards are often left in the locker room, and while in use at the athletic site, are often transferred from the mouth to hands, and temporarily stored or carried at distant sites (e.g. helmets, shorts, sleeves, socks etc.). Alternative means of disinfecting mouthguards at the athletic site may include the use of hand sanitizer. Therefore, the purpose of this study was to assess this alternative method of disinfecting samples of acrylic from surface inoculates of specific bacteria.

Materials and Methods

Bacterial cultivation and preparation of cells. *Staphylococcus epidermidis* (aka *S. aureus*), *Lactobacillus casei*, and *Escherichia coli* were purchased from the American Type Culture Collection, Manassas, Va. *S. epidermidis* ATCC 12228 was grown on/in brain heart infusion agar and broth. *L. casei* (aka *L. rhamnosus*) ATCC 7469 was grown on/in Lactobacillus agar and broth. *E. coli* ATCC 10798 was grown on/in trypticase soy agar/broth. The purchased freeze dried cells were reconstituted in their respective broths and then streaked on agar plates. Colonies were examined for purity and single colonies were transferred to broth and grown to log phase (4hrs) at which time they were used experimentally.

Experimental set-up

All experiments were conducted in a laminar control hood (Environmental Air Control, Inc., Hagerstown, Md). The surface of the hood was disinfected with Clorox wipes prior to conducting the experiments. All tubes, broths, agars and pipet tips were sterilized by autoclaving at 121 C for 20 min.

Materials to be tested

Purell™ and Clorox were purchased from local pharmacies. Clorox was diluted to 2%.

Preparation of acrylic specimens

A standard hole puncher was used to make circular discs from a hard-soft acrylic mouth guard (Figure 1). Disc sizes were polished so that they were 6mm wide in diameter, 2.5mm thick and weighed 1g. The cut discs were sterilized by autodaving at 121 C for 20 min.

Experimental Procedure for Evaluating the Effect of Hand Sanitizer on Mono-species Biofilms of *S. epidermidis*, *E. coli*, and *L. casei*.

The effect of a 20 second exposure to Purell™ on single species biofilm vitality was measured like the method of Welch et al. [12] as follows:

**Formation of biofilms:** Mono-species biofilms could form on the surface of the discs by incubating them in cells growing through log phase for 4hrs at 37 C with agitation. A disc immersed in sterile broth (no cells) was used as negative control. (For quality control, biofilms attached to 10 separate discs were stained with crystal violet [a dye that is specifically taken up by gram positive bacteria]. Distaining with 30% acetic acid confirmed that all discs retained similar numbers of bacteria).

**Rinsing:** After the experimental discs were removed using sterile procedures, planktonic cells were removed by rinsing with 5ml sterile saline, vortexing for 10sec in 1ml sterile saline, and rinsing again with 5ml sterile saline.

**Treatment:** Discs with adherent cells were then dipped into sterile Eppendorf tubes containing 1ml of the following: sterile saline (positive control), Purell™, or 2% sodium hypocholorite (Clorox) for 20 seconds. All discs were then rinsed once with 1ml sterile saline.

**Allowing biofilm cells to recover:** Discs were placed into 10ml replenished media. All four tubes of broth containing the discs (designated as working solutions) were then incubated at 37 C for 24 hours. Growth curves were generated for all four specimens, i.e. negative control, positive control, disc dipped into Purell™, and disc dipped into Chlorox. Distaining with 30% acetic acid confirmed that all discs retained similar numbers of bacteria.

**Assessing biofilm cell viability:** In addition to growth curves, after 4hrs incubation, samples from each working solution were diluted in ten-fold steps for the purpose of estimating bacterial cell numbers. From the dilutions, 0.1ml was spread evenly across the surface of their respective agars. All agar plates were incubated for 48hrs. at 37 C, after which all visible colonies were counted. The number of colonies forming units (CFU) were determined by counting segments of the plates like the procedure described Loeshe et al. [13]. All experiments were performed in triplicate.

**Statistical analysis:** The Student t-test was used to compare means of colony firming units between positive control samples and experimental Purell™ samples.

Results

Figure 2 shows the growth curves for *S. aureus* generated by cells attached to acrylic discs inserted into replenished media. Similar curves were found for *L. casei* and *E. coli* (data not shown).
Discs treated with Purell™ showed diminished growth patterns, with significantly less surviving colony forming units, as seen in Figures 3-5. The results of the surface application experiments for all three organisms are summarized in Table 1. As compared to positive control, surface application of hand sanitizer (Purell™) for 20 sec significantly inhibited growth of all three bacteria (p < 0.05). Three trials also indicated that mixing of 2% sodium hypochlorite (Clorox) with the bacterial inoculum on the surface of discs for 20 seconds completely inhibited growth of all three organisms.

**Table 1:** Effect of Hand Sanitizer on Biofilm Vitality

| Organism   | Treatment    | CFU at 10⁴ Dilution*   |
|------------|--------------|------------------------|
| **S. aureus** | None (+ control) | 105.1 ± 9.3            |
|            | Purell™      | 9.3 ± 1.2              |
| **L. casei** | None (+ control) | 45.4 ± 3.6             |
|            | Purell™      | 4.8 ± 0.8              |
| **E. coli**  | None (+ control) | 127.5 ± 6.7            |
|            | Purell™      | 28.4 ± 3.9             |

*Data is presented as median ± standard deviation of 3 trials; - control (disc with no cells) and treatment with 2% NaOCl = 0 CFU

**Discussion**

Athletic mouthguards become colonized by microorganisms after a certain period in the mouth. Previous studies have reported that these appliances become contaminated by several oral microorganism that may be involved in oral and systemic diseases [14]. Although dentists usually recommend at-home disinfection of dentures by immersion in antimicrobial agents for a certain length of time [15] data is lacking concerning the effectiveness of antimicrobials on bacteria attached to these appliances. In addition, in contrast to dentures, there is no emphasis on disinfection of mouthguards either at home or at the arena.

Bacteria with pathogenic potential, *L. casei*, *S. aureus*, and *E. coli*, were selected for this study. *L. casei* is characteristically known to lead to the progression of caries lesions, and bactereemia, meningitis, and endocarditis have been reported, particularly in immunocompromised patients [16]. *S. aureus* causes several diseases, such as pneumonia, sepsis, abscesses, infective endocarditis, and osteomyelitis and MRSA [7]. *E. coli* is an important cause of diarrhea, urinary tract infections, and septicemia [17].

Immersion of discs into sodium hypochlorite 2% (Clorox, bleach, NaOCl) for 20 sec was considered the gold standard for disinfection from bacteria. NaOCl is widely used as the main root canal irritant because of its broad antimicrobial activity in endodontics. It is also a low-cost method to disinfect an appliance [4]. Da Silva et al. [18] investigated the effectiveness of different solutions for disinfecting acrylic resin specimens contaminated with *Streptococcus mutans*, *S. aureus*, and *E. coli*. They found 2% NaOCl to be...
the best antimicrobial agent against the tested microorganisms. This result is supported by Salvia et al. [19], however, although the use of NaOCl, is an effective and low cost methods of sterilization, it is not applicable to use in the locker room or on the playing field.

In this study, we evaluated the efficacy of an alternative protocol-application of Purell™ - for disinfection of mouthguard specimens compared with the gold standards of chlorox application. Disinfection means to eliminate most harmful microorganisms from surfaces or objects. Purell™ reduced the levels of all bacteria but did not eliminate the microorganisms completely. The efficacy of Purell in this instance, i.e. it disinfects but does not sterilize, may be explained by the fact that the active ingredient, alcohol, is dispersed in a viscous carrier glycerol, which may not allow contact with all the organisms on the specimen. Also, the concentration of ethanol, at 29%, is low. Ethanol destroys the membranes of most bacteria on contact. In the same way, the discs were only immersed for 20 seconds, equivalent to the time used for hand sanitation. Longer immersions may eliminate the bacteria altogether. Purell™ is used almost universally in public facilities such as schools, clinics, and hospitals. It evaporates rapidly, leaves no after taste, and is not toxic to soft tissues [20]. In conclusion, this study supports the use of hand sanitizer for the disinfection of athletic mouthguards that is convenient to use at the athletic site. This in vitro study demonstrates the susceptibility of S. aureus, E. coli, and L. casei to hand sanitizer (Purell™) and demonstrates that it disinfects the surfaces of acrylic specimens of mouthguards.

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