Removal of *Cryptosporidium* and polystyrene microspheres from swimming pool water with sand, cartridge, and precoat filters

James E. Amburgey, Kimberly J. Walsh, Roy R. Fielding and Michael J. Arrowood

**ABSTRACT**

*Cryptosporidium* has caused the majority of waterborne disease outbreaks in treated recreational water venues in the USA for many years running. This research project evaluated some common US swimming pool filters for removing *Cryptosporidium* oocysts, 5-µm diameter polystyrene microspheres, and 1-µm diameter polystyrene microspheres. A 946 L hot tub with interchangeable sand, cartridge, and precoat filters was used at room temperature for this research. Simulated pool water for each experiment was created from Charlotte, NC (USA) tap water supplemented with alkalinity, hardness, chlorine, and a mixture of artificial sweat and urine. Precoat (i.e., diatomaceous earth and perlite) filters demonstrated pathogen removal efficiencies of 2.3 to 4.4 log (or 99.4–99.996%). However, sand and cartridge filters had average *Cryptosporidium* removals of 0.19 log (36%) or less. The combined low filter removal efficiencies of sand and cartridge filters along with the chlorine-resistant properties of *Cryptosporidium* oocysts could indicate a regulatory gap warranting further attention and having significant implications on the protection of public health in recreational water facilities. The 5-µm microspheres were a good surrogate for *Cryptosporidium* oocysts in this study and hold promise for use in future research projects, field trials, and/or product testing on swimming pool filters.

**Key words** | *Cryptosporidium*, diatomaceous earth, granular media filtration, precoat filtration, recreational water, swimming pools

**INTRODUCTION**

*Cryptosporidium* is a human protozoan parasite transmitted by the fecal-oral route while enclosed in a protective outer shell called an oocyst. The oocyst makes it highly-resistant to chlorine disinfection with Ct values on the order of 15,300 mg/L min (Shields *et al.* 2008b). *Cryptosporidium* caused more than 80% of reported waterborne disease outbreaks in treated recreational water venues (e.g., swimming pools) in the USA from 2005 to 2006 (Yoder *et al.* 2008). Cryptosporidiosis lasts an average of 12 days (with rare instances lasting as long as 4 weeks) in immunocompetent individuals with symptoms that can include: watery diarrhea, nausea, vomiting, fever, and abdominal cramping (Daniel 1996; Hoxie *et al.* 1997; Yoder & Beach 2010). Surveillance of cryptosporidiosis in the USA indicates that the reported incidence of infection has increased dramatically since 2004 (Yoder & Beach 2010). Both the number of reported cases and the number of individual outbreaks have shown overall upward trends since 2004 (Yoder *et al.* 2010). While it is difficult and expensive to assess the prevalence of protozoan parasites in public pools during normal non-outbreak conditions, a study of 160 filter backwash water samples from Atlanta, GA, USA showed that 13 (8.1%) were positive for the presence of *Giardia* or *Cryptosporidium* or both although the viability of the
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Accidental fecal release scenario

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Swimming pools are generally complex systems where a body of water experiences heterogeneous mixing and variable inputs of multiple contaminants. The filtration and disinfection of the water in a pool typically occurs as the water is recirculated through a treatment area at regular intervals that could be as high as 4–8 h on average (via a system of interconnected inlets and outlets spaced around the pool by the designer). A target pH (e.g., 7.2 to 7.6) and free chlorine level (e.g., 2 to 4 mg/L) are typically maintained in pool water (often by an automatic controller) to achieve some level of residual disinfection between treatment cycles. Unfortunately, the residual disinfectant is quite slow at inactivating chlorine-resistant pathogens like Cryptosporidium. Automatic control systems for filters exist, but most swimming pool filters are still operated manually. The overall quality and safety of the pool water depends on many variables (e.g., bather inputs, mixing dynamics, length of treatment cycle, disinfection process type(s), filtration effectiveness, as well as the training/performance of the pool operator), but the current study focused solely on the effectiveness of the filtration process for removal of particles of two specific sizes.

Filtration practices

Relatively little is known about the capabilities of common US swimming pool filters to remove waterborne pathogens. Research has shown that ‘high-rate’ (i.e., 25–49 m/h filter loading rate) swimming pool sand filters can only parasites was not reported (Shields et al. 2008a). In a study of 803 Oklahoma children, 58% of adolescents (ages 14 to 21) were seropositive for C. parvum, which indicates prior infection by the pathogen (Ford 1999). The true burden of cryptosporidiosis is not known with certainty, but recent estimates have ranged from 300,000 to 748,000 cases annually in the USA (Yoder & Beach 2007; Beach 2011). Multiple sources have indicated that weaker subpopulations (e.g., very young children, elderly people, pregnant women, and the immunocompromised) could die from cryptosporidiosis (Daniel 1996; Hoxie et al. 1997; Ford 1999). A quantitative risk assessment model of Cryptosporidium in swimming pools recently confirmed there is a ‘significant public health risk’ (Pintar et al. 2010). As in the drinking water industry, the burden for safety often falls primarily on physical removal (i.e., filtration).

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Pool water treatment overview

Swimming pools are generally complex systems where a body of water experiences heterogeneous mixing and variable inputs of multiple contaminants. The filtration and disinfection of the water in a pool typically occurs as the water is recirculated through a treatment area at regular intervals that could be as high as 4–8 h on average (via a system of interconnected inlets and outlets spaced around the pool by the designer). A target pH (e.g., 7.2 to 7.6) and free chlorine level (e.g., 2 to 4 mg/L) are typically maintained in pool water (often by an automatic controller) to achieve some level of residual disinfection between treatment cycles. Unfortunately, the residual disinfectant is quite slow at inactivating chlorine-resistant pathogens like Cryptosporidium. Automatic control systems for filters exist, but most swimming pool filters are still operated manually. The overall quality and safety of the pool water depends on many variables (e.g., bather inputs, mixing dynamics, length of treatment cycle, disinfection process type(s), filtration effectiveness, as well as the training/performance of the pool operator), but the current study focused solely on the effectiveness of the filtration process for removal of particles of two specific sizes.

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consistently deliver 0.11 to 0.28 log (22 to 48%) removal of Cryptosporidium oocysts and/or a microsphere surrogate without coagulation (Amburgey et al. 2007, 2008, 2009a, b; Croll et al. 2007). These levels of removal appear inadequate to prevent outbreaks of cryptosporidiosis, which is supported by surveillance data on outbreaks investigated each year in the US. However, drinking water treatment research has shown that granular media filters can remove in excess of 6 log (99.9999%) of Cryptosporidium oocysts (States et al. 2002). Although sand filters are similar for swimming pool and drinking water treatment facilities, there are two important differences between drinking water treatment and pool water treatment practices. First, drinking water treatment facilities perform chemical coagulation prior to filtration, which greatly enhances the ability of the filters to remove the pathogens by reducing the natural electrostatic repulsion that exists between negatively charged pathogens and negatively charged filter media in water (Hendricks 2006). Second, the filter loading rates (flow per unit of surface area of filter) are typically at least 4 to 5 times lower in US drinking water filters than in pools (e.g., 5 to 10 m/h for drinking water versus 37 to 49 m/h in pools).

Diatomaceous earth (DE) is also used in drinking water treatment. However, the two important differences between sand filters used for drinking water and pool water treatment (stated previously) do not apply to DE filtration. The filter loading rates used for DE filters in both industries are similar, and neither industry necessarily practices coagulation as pretreatment for DE filters (Logsdon, 2008; NSPF, 2009). Unlike sand filters, DE filters rely dominantly on the size exclusion principle to prevent pathogens from passing through the tiny pores in the DE media. This is why sand filter media depths are a minimum of 0.25 m (and commonly called ‘depth filters’) while DE media depths are only 1.6–3.2 mm because they are ‘surface filters’. Drinking water research has shown Cryptosporidium removal in excess of 6 log for DE filters (Ongerth & Hutton 1997; Ongerth & Hutton 2001). Nevertheless, two important differences still exist between the DE filtration practices used in the drinking water and pool water industries. The grades of DE used in drinking water treatment are of a finer grade, which is typically measured as ‘permeability’. Typical permeability values of the DE media used for swimming pools is 4–5 μm², but the permeability of the DE media used in drinking water treatment is typically less than 1 μm² (Hendricks 2006). The finer-grained precoat media has proportionally smaller pores between the grains and removes smaller particles more effectively. Secondly, the amount of DE media loaded into each filter per unit surface area of the media support material is typically two-fold greater (at 1 kg/m²) in the drinking water industry (Logsdon 2006). More media per unit surface area creates a thicker layer of DE ‘cake’ that particles must pass through to avoid removal.

Pool filters are commonly designed to keep swimming pools looking clear and pleasing to the eye (i.e., turbidity removal), which is not the same as for effective pathogen removal. The US swimming pool industry has traditionally relied on disinfectants, such as free chlorine, to control the spread of waterborne diseases. However, chlorine is inefficient at inactivating Cryptosporidium, so the burden of microbial safety falls on filtration. The drinking water industry also relied heavily on chlorine until chlorine-resistant pathogens forced changes in the 1980s (for Giardia) and the 1990s and beyond (for Cryptosporidium). These chlorine-resistant pathogens forced the drinking water industry to put considerable regulatory emphasis on filtration optimization to achieve physical removal of these pathogens. US drinking water regulations have become increasingly stringent on pathogen removal in recent years in order to safeguard public health. The swimming pool industry could be forced to take the same approach. However, the average level of training and human resources devoted to swimming pool operations tends to be considerably less than for drinking water operations, which necessitates simple yet effective filtration options. Constantly optimizing coagulation and closely monitoring filtered water turbidity might not be the most practical approach for the majority of US swimming pools.

**Research objectives**

The primary objective of this research was to evaluate some common swimming pool filtration technologies for removing Cryptosporidium oocysts, 5-μm polystyrene microspheres, and 1-μm polystyrene microspheres from a simulated pool water. The filter types chosen were sand, cartridge, and precoat. Standard commercially available filters and media were used in these experiments. Knowing the key differences between drinking water and pool water.
practices for both sand and precoat filters, a limited number of additional experiments were conducted in an effort to achieve better pathogen removal performance with the sand and precoat filters.

**METHODS**

**Experimental setup (model pool)**

A 946 L commercial hot tub was used at room temperature (20 °C) without aeration to serve as the pool (or water tank) for this research. The original pump and filter were removed and replaced by a commercially available (to the pool industry) 2.2 kW centrifugal pump and a set of three interchangeable filters (i.e., sand, cartridge, and precoat) connected via a 51 mm diameter PVC pipe loop measuring approximately 6.5 m in length. A schematic of the experimental setup is shown in Figure 1, and Table 1 contains detailed information about the materials used in this research. The flow was measured with a digital paddle-wheel flow meter and controlled with a 51 mm diameter PVC ball valve.

### Filter descriptions

The sand filter contained approximately 35 cm of sand (with an effective size of 0.49 mm and uniformity...
coefficient of 1.5), but only about 25 cm of the sand was necessarily used for filtration as the remainder was below the surface of the laterals. A cross-sectional drawing of the 0.18 m² sand filter appears in Figure 2. The surface areas of the precoat and cartridge filters were 2.2 and 22.3 m², respectively. The three types of precoat media used in this study were standard pool grade DE with a permeability of 4.4 μm², fine DE with a permeability of 1.2 μm², and fine perlite with a permeability of 1.5 μm². The manufacturer-supplied filter cartridges were used in the cartridge filter, but no pore size rating was given in the product literature. All filters were from the same manufacturer and similar in size, shape, and construction materials despite the very different filtration technologies and surface areas being utilized.

Pathogen and microsphere seeding

Inline feed of the oocyst/microsphere suspensions was made possible by a digital peristaltic pump feeding directly into the PVC pipe just upstream of the centrifugal pump. The oocyst/microsphere suspensions were made in a 1-L glass Erlenmeyer flask of simulated pool water and stirred continuously with a magnetic stirrer and Teflon®-coated stir bar prior to and during the experiments. The inline feed system allowed for feeding the oocysts/microspheres into the system without stopping and restarting the precoat filters during each experiment, which had been observed to hinder the filter performance in pre-trial runs.

Filter operation

The sand and cartridge filters were operated differently because the stop/start process was not expected to alter their performance once the flow stabilized. For the sand and cartridge filter experiments, the flow of water was stopped just prior to beginning the oocyst/microsphere seeding, and valves were repositioned to redirect the filtered water to the drain piping thereby preventing recirculation. Preliminary experiments (data not shown) revealed that the low removals associated with the sand and cartridge filters allowed rapid accumulation of particles in the spa water with recirculation, which caused the influent number of oocysts/microspheres to increase rapidly and potentially alter the accuracy of calculations as well as the filter performance itself. Sand filters were backwashed with simulated pool water prior to experiments to prevent the pore water from changing chemically during the experiments while the other two types of filters were simply drained completely and filled with simulated pool water at the beginning of the experiment.

All filters were pressure filters (as opposed to gravity or vacuum filters). The filter loading rates were chosen at the maximum rated value for each filter in order to assess the worst-case scenarios for each filter in terms of Cryptosporidium removal. The spa system was capable of pumping water at up to 227 L/min. The precoat and cartridge filters were operated at 227 L/min for filter loading rates of 6.1 and 0.61 m/h, respectively. Water was pumped through the sand filter at 144 L/min for a filter loading rate of 49 m/h. All filters were designed and approved by the National Sanitation Foundation (Ann Arbor, MI, USA) to operate at the prescribed flow rates according to their product labels. The DE or perlite media was added to the precoat filter (in slurry form) in the amount of 0.49 kg/m² of filter surface area. However, one exception was that the amount of fine DE was increased to 1.5 kg/m² in the second experiment only.

Coagulant application

The coagulant, when used, was poured directly into the spa following an initial dilution in 10 L of simulated pool water and allowed to recirculate and mix for a period of time prior to seeding oocysts and/or microspheres. The minimum recommended dosage on the bottle (1.6 mg/L as product) was used without any optimization or zeta potential measurements. This product was chosen because of its history and prevalence in the US swimming pool market. Currently, coagulant use is not required with any type of filter. Use is
at the discretion of the pool operator, which often means it is
not used at all by a given facility. When a coagulant is used, it
is often in response to a real or perceived water quality pro-
blem. Coagulant usage guidelines are not well-established,
and the directions on the bottle tend to be vague about the
manner of application and frequency of use. While the ingre-
dients and formulation information are proprietary, the
primary active ingredient is a cationic polymer.

Simulated pool water

Simulated pool water was created for each experiment from
946 L of Charlotte, NC (USA) tap water supplemented with
sodium bicarbonate to an alkalinity of 150 mg/L as CaCO₃,
with calcium chloride to a hardness of 250 mg/L as CaCO₃,
with sodium hypochlorite to a free chlorine concentration of
2 mg/L, with hydrochloric acid to a pH of 7.5, and with a
mixture of artificial sweat and urine described in Table 2
to a final total organic carbon (TOC) concentration of
approximately 20 mg/L as C. This TOC concentration was
chosen to match the campus pool at UNC Charlotte,
which is near the upper end of the 3.0 to 23.6 mg/L average
TOC concentration range observed in a survey of 23 indoor
pools in the Southeastern USA (Kanan 2010).

Experimental procedure and sampling

Two experiments were performed for each set of conditions.
For each experiment, duplicate samples were collected from
the filter influent and effluent pipes. Sampling began approxi-
mately 1 min after the start of an experiment. The hydraulic
detention time of the pipe loop from the point of microsphere
injection through the pump and sand filter to filter effluent
sample point was calculated to be 35 s at a flow rate of
144 L/min. So, roughly two hydraulic detention times
passed prior to the collection of the first sample and four
detention times between replicate sets of samples. A delay
of one hydraulic detention time was used between collection
of the influent and effluent paired samples in an attempt to
sample the same water volume before and after
filtration. The hydraulic detention time of the system including the
hot tub was only 7.2 min, so the duration of the microsphere
seeding experiments was limited to less than 6 min each for
the sand filter configuration. The cartridge and precoat
filter setups used a higher flow rate and proportionally
shorter hydraulic retention times (25 s) with entire exper-
iments lasting no more than 4 min each. So, it was not
possible to look at performance changes or headloss develop-
ment rates versus time of filtration as part of this study.

Microspheres and Cryptosporidium

Fluorescent-green carboxylate-modified polystyrene micro-
spheres (4.869 μm, std. dev. 0.246 μm) were fed simultane-
ously with Cryptosporidium oocysts, but the smaller
diameter microspheres (0.951 μm, std. dev. 0.010 μm)
were used in separate experiments. A 1 L microsphere sus-
pension was fed inline during the experiment to achieve
the target filter influent concentration of approximately
15–150 oocysts/microspheres per mL of water. Influent
samples of 50 mL were collected in sterile 50 mL conical-
bottomed plastic centrifuge tubes, and the volume of the
effluent samples varied from 50 mL to 1 L with the larger
samples collected in glass media bottles. A maximum of
approximately 10⁸ heat-inactivated (55 °C for 30 min) Cryp-
tosporidium parvum oocysts (Iowa isolate; purified
according to Arrowood & Donaldson (1996)) and/or YG flu-
orescent carboxylate-modified polystyrene microspheres
were used in each experiment. The microspheres also
served as an internal control for oocyst staining procedure
since the microspheres do not have to be stained to be
seen under an epifluorescence microscope. Further, micro-
spheres are not susceptible to excystation (breaking open)

Table 2: Artificial sweat and urine formula

<table>
<thead>
<tr>
<th>Component</th>
<th>Concentration (mg/L)</th>
<th>Concentration (mM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetic acid</td>
<td>12.6</td>
<td>0.21</td>
</tr>
<tr>
<td>Ammonium chloride</td>
<td>48.4</td>
<td>0.90</td>
</tr>
<tr>
<td>Creatinine</td>
<td>1.8</td>
<td>0.016</td>
</tr>
<tr>
<td>Lactic acid</td>
<td>32.8</td>
<td>0.36</td>
</tr>
<tr>
<td>Potassium chloride</td>
<td>4.0</td>
<td>0.053</td>
</tr>
<tr>
<td>Sodium chloride</td>
<td>59.6</td>
<td>1.02</td>
</tr>
<tr>
<td>Sodium phosphate (monobasic)</td>
<td>4.2</td>
<td>0.030</td>
</tr>
<tr>
<td>Sodium sulfite</td>
<td>2.0</td>
<td>0.016</td>
</tr>
<tr>
<td>Urea</td>
<td>18.7</td>
<td>0.31</td>
</tr>
</tbody>
</table>
were stored at 4°C with antibiotics prior to use and obtained from Dr Michael Arrowood’s laboratory at the Centers for Disease Control and Prevention in Atlanta, GA, USA.

Sample analysis

Samples were filtered through 3-μm polycarbonate track-etched (PCTE) filters in 25-mm glass microanalysis filter funnels by a regulated 3-place vacuum manifold. Sample volumes analyzed were adjusted to obtain between 10 and 150 oocysts and/or microspheres per sample. The staining method used for Cryptosporidium involved placing 600 μL of Cryptosporidium-specific monoclonal antibody (obtained from Dr. Arrowood’s lab) on top of the filter for 30 min at room temperature, rinsing twice with 750 μL of 1× PBS (pH 7.4), applying 500 μL of a 1:150 dilution of a fluorescein isothiocyanate (FITC)-labeled goat anti-mouse antibody in DABCO solution (Freer 1984) and a 25-mm square glass cover slip for enumeration under epifluorescence microscope at 100× (for microspheres) or 250× (for oocysts) total magnification. Separate counts were performed for microspheres and oocysts whenever they were used together. The raw counts per mL from the filter influent and effluent sample pairs were used to calculate the percent and log removals. The fluorescent filter set used in the microscope had a 450–490-nm excitation wavelength range, a 510-nm dichroic filter, and a 520-nm emission filter. The PCTE filters with 3-μm pores were used for the 5-μm microspheres and oocytes, which were assayed simultaneously. A 0.65-μm pore size PCTE filter was used for the experiments using only the 1-μm microspheres. Samples were stored at 4°C prior to analysis.

Quality control measures

The spa system was thoroughly cleaned between experiments with a minimum of three drain-and-fill rinses with recirculation at 227 L/min, and samples were collected prior to seeding in each experiment to measure any potential carryover between experiments. A control experiment was used to determine whether or not the oocysts and microsphere were destroyed by the pump and/or lost due to surface attachment within the system. All statistical calculations were performed on the percent removal data (not the log-manipulated values) with commercially available software (Microsoft Office Excel 2007; Microsoft, Redmond, WA, USA).

RESULTS AND DISCUSSION

Particle challenge tests were performed for three categories of filters (i.e., sand, cartridge, and precoat). Subcategories included sand filtration with and without coagulant as well as precoat filtration with three individual filter media products. For each of the preceding filtration scenarios, particle removal was determined separately for Cryptosporidium oocysts, 5-μm microspheres, and 1-μm microspheres. Data from individual experiments are summarized in Table 3.

Sand and cartridge filter removals of Cryptosporidium

Percent Cryptosporidium removals and log Cryptosporidium removals under varying conditions are shown in Figures 3 and 4, respectively. Cryptosporidium removals averaged 0.16 log (31%) for the sand filter operated at 49 m/h and 0.19 log (36%) for the cartridge filter operated at 0.61 m/h. Sand filter removals nearly doubled to a mean of 0.41 log (61%) with the addition of a coagulant to the system. The coagulant dose was not optimized, which might explain the relatively low removals in comparison with drinking water research results or the precoat media used in this study. Charge neutralization is a process that requires the number of positive charges (from the coagulant) to balance the negative surface charges on the particles. The improvement with coagulant addition indicates promise, but further trials with alternate coagulant dosages were beyond the scope of this project. Furthermore, the coagulant was added and recirculated for a minimum of 4 h prior to adding the oocysts. This practice seemed consistent with what might actually occur before a fecal accident in a swimming pool intermittently employing a coagulant, but this method of coagulation is
significantly different than that practiced in drinking water production. In drinking water treatment, adding a continuous feed of coagulant immediately following oocysts/microspheres (just prior to filtration) is expected, but alternate coagulant feeding scenarios were not explored in the present study. These two preceding coagulant addition scenarios could be termed ‘passive’ and ‘active’ treatment, respectively.

**Precoat filter removals of Cryptosporidium**

Precoat filter removals of *Cryptosporidium* oocysts were significantly higher than for the other two types of filters as shown in Figure 4. The standard grade of DE used in swimming pools removed 2.3 log (99.6%) of oocysts. Two finer-grained (lower permeability) types of precoat media each

**Table 3 | Removal data for individual experiments for each type of filter and particle**

<table>
<thead>
<tr>
<th></th>
<th>Cryptosporidium % removal</th>
<th>Cryptosporidium Log removal</th>
<th>5-μm Microspheres % removal</th>
<th>5-μm Microspheres Log removal</th>
<th>1-μm Microspheres % removal</th>
<th>1-μm Microspheres Log removal</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sand</strong></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Exp. 1-1</td>
<td>31</td>
<td>0.16</td>
<td>36</td>
<td>0.17</td>
<td>39</td>
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*Indicates precoat media was loaded at 1.5 Kg/m².*
produced higher removals of oocysts than the standard DE media. Perlite and a finer grade of DE removed oocysts with an average removal efficiency of 3.2 and 4.3 log, respectively. The filtration rate remained constant at 6.1 m/h throughout all of the precoat filtration experiments, but the amount of precoat media increased in one experiment. The mass of standard DE and perlite was identical in all experiments (i.e., 1.1 kg or 0.5 kg/m²), but the lower density of perlite produced a precoat layer of approximately double the thickness of the DE precoat layer. Half of the data points with fine grade of DE were produced at 0.5 kg/m², but the other half were produced with 1.5 kg/m² of DE. The thickness of the precoat layer did appear to make some difference in the Cryptosporidium removal as shown in Figure 5, but the number of replicate samples (n = 2) was insufficient for statistical analysis. The dominant factor impacting Cryptosporidium removal was likely the permeabilities of the precoat media, which were 4.4, 1.5, and 1.2 μm² for the standard DE, perlite, and fine DE media, respectively. A series of heteroscedastic, 2-tailed Student’s t-tests showed that the differences in oocyst removals between each of the media/filter types were statically significant (α = 0.05) with the exception of the sand filter versus the cartridge filter. For the statistical analyses, data from all four of the fine DE experiments were grouped together.

**Five-micron microsphere (Cryptosporidium surrogate) removals**

The microspheres were used as a potential non-infectious surrogate for Cryptosporidium oocysts as they have nearly identical size, shape, and density. If the microspheres behave like Cryptosporidium oocysts in pools, then they could be used to safely evaluate any full-scale pool filtration system under normal operating conditions without posing any risk to bathers. The percent removals and log removals of the 5-μm microspheres are also shown in Figures 3 and 4, respectively. The Cryptosporidium removal results in Figures 3 and 4 are very similar to the microsphere removals. A series of heteroscedastic, 2-tailed Student’s t-tests did not indicate any statistically significant differences (α = 0.05) between the oocyst and 5-μm microsphere removals for any of the experiments with identical treatment. So, the microspheres appeared to be a good surrogate for Cryptosporidium oocysts under all test conditions in these filters. Polystyrene microspheres have also been found to be a good surrogate for oocysts in multiple drinking water research studies (Dai & Hozalski 2003; Emelko et al. 2003; Amburgey et al. 2005; Brown & Emelko 2009).

**One-micron microsphere removals**

The percent removals and log removals of the 1-μm microspheres are shown in Figures 3 and 4, respectively. The 1-μm microspheres were used in this study to evaluate how each type of filtration performed at removing particles even smaller than Cryptosporidium oocysts. These experiments sought to test the perceived limits of each filtration scenario. Mean sand and cartridge filter removals dropped
below 0.12 log (25%) and approached the control (no filter) removal rate of 0.05 log (10%).

Mean removals tended to be lower across each filtration condition with the exception of the sand filter-coagulant combination where the mean removal remained almost constant. The sand filter with coagulant actually showed slightly higher average removals than the standard DE or perlite for the 1-μm microspheres. There was one unplanned difference in the experimental procedure between the oocyst/5-μm microsphere experiments and the 1-μm microsphere experiments using the sand filter with coagulant that could have contributed to relative increase in removal efficiency between sand filter with coagulant versus the other filtration scenarios for the 1-μm removal experiments. The coagulant was added and recirculated for approximately 30 min prior to the 1-μm removal experiments instead of at least 4 h as was done in the oocyst/5-μm microsphere experiments. The shorter interval between coagulant addition and the beginning of the seeding experiment could have increased the removal efficiency of the coagulant/sand filter system, which could indicate that this was a time-dependent treatment process for this system. It could be important that the ratio of surface area of the sand grains to the volume of water in this research setup was much larger than in a typical swimming pool, which could have exerted an increased coagulant demand on the system due to the natural negative surface charge of sand grains in water. The preceding design issue, if significant, calls the real-world applicability of the coagulation experiments into question, but not the nature of the coagulation results, which was a general improvement over no coagulant. However, the impact of this system’s surface area ratios would only be expected to influence the coagulation trials since none of the other trials were adding chemicals that would potentially alter the charge of any surface (i.e., the other experiments were purely physical removal without a chemical component).

As shown in Figure 5, the precoated filter removals were considerably lower at 0.37 to 1.3 log for the 1-μm microspheres than for the larger 5-μm microspheres and oocysts (2.2 to 4.8 log). Coagulant use may hold promise for increasing the filtration removals of all three types of filters though demonstrated herein only for the sand filter. Hendricks (2006) stated that coagulant use improved the removal of smaller particles in DE filters, but there was a corresponding increase in the rate of headloss accumulation. The removal of the 1-μm microspheres was highest for the fine DE media (with a mean of 86% or 0.85 log for two loading levels used for the fine DE) as shown in Figure 3 and 4. Only with the finest grade of DE at triple the standard loading rate, were any of the filters tested capable of removing greater than 1 log (90%) of the 1-μm microspheres. Removals of 1-μm microspheres were 0.61–0.67 log (75–79%) and 1.3–1.4 log (95–96%) for the standard and tripled media loading rates for fine DE, respectively.

CONCLUSIONS

The results show that swimming pool sand filters currently thought to be used by the majority of the pools in the USA
(as well as cartridge filters) have limited effectiveness for pathogen removal with overall removals averaging less than 0.19 log (36%) for Cryptosporidium oocysts and both sizes of microspheres under the conditions studied. Based on this finding, it seems logical that the pool industry would want to support new research to identify techniques for improving sand filter performance to better safeguard the health of bathers. Coagulation prior to sand filtration is one technique that warrants further study since it is required prior to sand filtration in drinking water treatment operations throughout the world as well as in the majority of European public swimming pool facilities. While there are no obvious barriers to coagulation prior to sand filtration, there is a significant lack of comprehensive data regarding the current filter design and operating practices in the USA. The size and depth of filter media, filter loading rates, and backwashing practices are known to have significant impacts on filter performance, but accurate values for these parameters are not currently available to regulators, researchers, pool designers, or even filter manufacturers. A survey of US aquatic facilities could establish a baseline of current practices and help identify (through future research) the best strategies for improving filter performance and protecting public health.

Diatomaceous earth and perlite produced higher levels of pathogen removal with mean removals ranging from 2.3 log (99.4%) to 4.4 log (99.996%) with the higher pathogen removals obtained in experiments with finer grades of DE and perlite. The performance of the finer grades of precoat media was further enhanced by increasing the thickness of the media layer in the filter. The 1-μm microsphere removals by the precoat filters were considerably lower than for the 5-μm microsphere removals with removals averaging 0.37 to 0.85 log. The impact of stopping and restarting a precoat filter without changing the media and pathogen distribution within the filter media is a topic that appears to warrant further investigation. The effects of integrity and cleanliness of precoat media support materials over time on pathogen removal is still unknown. The use of precoat media on top of the sand in a sand filter might be a way to quickly improve the performance of existing sand filters, but this approach would require long-term full-scale studies at multiple locations to gain widespread acceptance.

The 5-μm microspheres appeared to be an adequate surrogate for Cryptosporidium in terms of filter removals for all three types of filters and under all tested conditions in simulated swimming pool water. This finding could be important for future research studies, new product testing protocols, as well as for full-scale performance evaluations of swimming pool filters. While the risk factors associated with Cryptosporidium outbreaks extend beyond swimming pool filtration removal efficiency, there does appear to be significant room for improvement in this area.

Based on the low average removal efficiency ≤0.19 log (36%) average removal efficiency of sand and cartridge filters for Cryptosporidium, the prevalence of these filters in existing recreational water facilities, and the highly chlorine-resistant properties of Cryptosporidium oocysts; further safeguarding public health in recreational water facilities in the USA could be quite challenging. The epidemiological data (both endemic and outbreak) suggest that this problem is not going away on its own. The leaders in the pool industry should take an active role in fully assessing this problem, identifying potential solutions, and making the critical decisions required to resolve this problem.

ACKNOWLEDGEMENTS

EP Minerals generously provided funding for this research as well as the precoat filter media. Pentair Water sized and donated the centrifugal pump, three pool filters, and the PVC valves used to direct water into and out of the filters.

DISCLAIMER

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

REFERENCES


