Megasonic sonication for cost-effective and automatable elution of Cryptosporidium from filters and membranes

Abdelfateh Kerrouche¹,²,*, Marc P.Y. Desmulliez¹ and Helen Bridle²

¹) Heriot-Watt University, MicroSystems Engineering Centre (MISEC), Riccarton, Edinburgh, United Kingdom, EH14 4AS
²) Heriot-Watt University, Institute of Biological Chemistry, Biophysics and Bioengineering (IB3), Riccarton, Edinburgh, United Kingdom, EH14 4AS

* Corresponding author: Email addresses: a.kerrouche@hw.ac.uk / abdf.kerrouche@hotmail.co.uk

Tel.: +44 (0) 131 650 5814; fax: +44 (0) 131 650 6554

Abstract

Sample processing is a highly challenging stage in the monitoring of waterborne pathogens. This step is time-consuming, requires highly trained technicians and often results in low recovery rates of pathogens. In the UK but also in other parts of the world, Cryptosporidium is the only pathogen directly tested for in routine operational monitoring. The traditional sampling process involves the filtration of 1000L of water, semi-automated elution of the filters and membranes with recovery rates of about 30-40% typically. This paper explores the use of megasonic sonication in an attempt to increase recovery rates and reduce both the time required for processing and the number of labour-intensive steps. Results demonstrate that megasonic energy assisted elution is equally effective as the traditional manual process in terms of recovery rates. Major advantages are however offered in terms of reduction of the elution volume enabling the current centrifugation stage to be avoided. This saves time, equipment and staff costs and critically removes the step in the process that would be most
challenging to automate, paving the way thereby for highly effective automated solutions to pathogens monitoring.

Keywords: Cryptosporidium; elution; megasonic agitation; sonication; filtration; waterborne pathogens; monitoring.

1. Introduction

The presence of pathogens in drinking water is a major cause of disease outbreaks and endemic levels of illness, impacting upon productivity as well as quality of living (World Health Organisation, 2011; Hrudey et al., 2003). Water quality compromised by microbial contamination is also a concern for food producers and several disease outbreaks have been linked to the water utilised in food production (Söderström et al., 2008; Brugha et al., 1999). Although the labour-intensive monitoring of the water supply for the presence of pathogens can be expensive, such measures allow the reduction of the costs associated with disease outbreaks.

Cryptosporidium is a particularly problematic pathogen in this regard. This protozoan has a low infectious dose, a longevity of months in the water environment and a high resistance to disinfection by chlorination. Despite the removal of the regulatory requirement to directly test for the presence of Cryptosporidium in water, UK water utilities continue to perform regular, even daily, checks at many sites. Because of their low infectious dose, sample preparation is required to concentrate waterborne pathogens from a large volume of water, of the order of thousands of litres, to a small sample such as a few μLs to be used by detection devices (Bridle, 2013). Detection protocols such as the U.S. Environmental Protection Agency (EPA) method 1623.1 (Method 1623.1, 2012) or the UK Environment Agency Blue Book 2
publications (UK Environment Agency, 2010) stipulate a procedure for Cryptosporidium detection. This method consists of several steps involving filtration (1000L/24hrs), elution stage 1 (remove oocysts from filter into 1200mL), elution stage 2 (concentrate the eluate using a membrane to 50mL), centrifugation (centrifugation to 5mL), enrichment (immuno-magnetic separation IMS to separate oocysts from other particulate matter to 50μL) and detection (staining with fluorescent dyes followed by microscopic examination for identification). Most of these stages require a long time, large and/or specialised equipment or highly qualified staff.

Elution steps are critical in ensuring a high recovery rate of pathogens (Francy et al., 2013). Manufacturers of commercially available filters report rates in excess of 70%. However, personal communications with water utilities suggest that recovery rates do not often reach these levels. This is further confirmed by results of a variety of literature studies in which recovery rates on the order of 30% to 40% were repeatedly measured across a range of different water types (Polaczyk et al., 2008; Smith and Hill, 2009; Leskinen et al., 2010; Mull and Hill, 2009) or across a range of filters using lake water samples (Francy et al., 2013).

In this paper we explore the use of a novel physical approach to filter and membrane elution, namely the use of megasonic sonication as a replacement to manual processes of filter elution. In the last few years, megasonic wave assisted cleaning systems have been widely used to clean various types of objects possessing complex surface geometries such as electronic devices, semiconductor wafers or component parts (Kaufmann et al., 2008; Busnaina et al., 1995; Helbig et al., 2008). In megasonic assisted agitation, an piezoelectric transducer, placed inside a tank, produces high frequency sound waves, typically over 1 MHz, that propagate through the liquid. Each point along the sound wave oscillates between a maximum and a minimum pressure. When the minimum pressure is below the vapour
pressure of the liquid, bubbles are formed. As the pressure increases to the maximum
pressure, the bubbles implode decreasing local turbulence at the implosion sites (Chitra et al.,
2004). Megasonic waves propagate at a higher frequency than ultrasonic waves. Smaller
bubbles with less resulting cavitation energy are created, resulting in a gentler elution and
potentially avoiding destruction of the pathogens (Al-Sabi et al., 2011).

Studies on the effect of the sonication of filters using ultrasound were performed to elute
bacteria from filters for safe drinking water (Mendez et al., 2004) or from food samples
(Ruban et al., 2011).

The effects of ultrasound with different sonication power and time durations on waterborne
protozoa *Cryptosporidium* and *Giardia* were studied. The results showed that changes in
parasite characteristics became visible (the shells were broken) when sonication time was
extended (Al-Sabi et al., 2011). A study investigated the effect of underwater ultrasound on
the viability of *Cryptosporidium* oocysts and demonstrated that more than 90% of the
dispersed *Cryptosporidium* oocysts could be deactivated in few minutes of continuous
sonication (Ashokkumar et al., 2003). However, the deactivation of oocysts by this method is
undesirable if one wishes to preserve the viability of the pathogens for further determination
of their infectivity. Additionally, DNA degradation could be incompatible with the molecular
tools currently under development (Bridle et al., 2014). In contrast, through the minimisation
of the time required for bubble growth, megasonic sonication offers a way to elute
undamaged and potentially viable oocysts from filters and membranes. This paper presents,
for the first time, the use of megasonic sonication for pathogen elution and evaluates its
qualities in terms of recovery rates, pathogen viability, processing, time required and potential
for automation.
2. Materials and methods

2.1. Standard elution protocol

The standard elution procedure as recommended in the U.S. Environmental Protection Agency (EPA) method 1623.1 (Method 1623.1, 2012) or the UK Environment Agency Blue Book publications (UK Environment Agency, 2010) is used by the water utility company, Scottish Water, which assisted in the microscopic evaluation of oocysts following the different elution protocols. The Filta-Max sponge filter from the IDEXX company, is first removed from the filter housing and placed into a washing station which encompasses a concentrator unit. In this washing station the filter is rinsed twice with 600mL of Phosphate-Buffered Saline with Tween® 20 (PBS T) for about 20 minutes although the duration of the rinsing time depends on the water sample. The wash solution is then passed through a membrane placed at the bottom of the concentrator placed on a magnetic stirrer attached to a hand pump to generate a vortex in the suspension within the concentrator. This magnetic stirring maximises the amount of particulates held in suspension throughout the filtration process, and should prevent oocysts from strongly attaching themselves to the membrane. After the liquid has reached a stable rotational velocity, the sample is drained away through the membrane using a vacuum below 40KPa. The membrane is then removed and placed inside a polythene bag containing 5 to 10mL of PBST. Once the bag is sealed, the surface of the membrane is rubbed between thumb and forefinger for 70 ± 10 seconds until the membrane appears to be clean. Finally, the eluent liquid is removed using a plastic Pasteur pipette and added to a 50mL centrifuge tube with the concentrate fraction obtained from the rinsed stirrer bar. The addition of 5-10mL of PBST and rubbing is repeated a second time and the volume in the centrifuge tube made up to 50mL. The 50mL was then passed onto centrifugation, immunomagnetic separation and microscopy for detection and
enumeration of oocysts. Two elution stages can be distinguished from the above procedure: one from the sponge filter where 1.2L of PBST is used for further sample concentration, the other from the membrane whereby 50mL of PBST is employed.

Both stages were studied in this article. In the case of the sponge filters, 1000 litres of uncontaminated water were spiked with 100 oocysts and filtered through the sponge filter over 24 hours. Recovery rates were then measured by carrying out the rest of the traditional process. In the case of the membranes, 100 oocysts in 1mL of water were passed directly through the membrane and recovery rates were determined by undertaking the rest of the standard procedure.

### 2.2. Elution with megasonic sonication

A transducer from the Company Sonosys with a frequency of 2MHz and an output power of 1200 Watts was employed to investigate the elution with megasonic energy assisted agitation (Sonosys. 2015). The encapsulated transducer made of stainless steel was positioned at the bottom side of an existing tank as shown in Figure 1. The sponge filters were added to a large plastic bag with up to 1.2L of PBST whereas the membranes were added to the bag utilised in the traditional approach with up to 50mL volumes of PBST.

### 2.3. Assessment of oocysts viability

An excystation assay was performed accordingly to protocol. Briefly a sample of 1 million oocysts in 40µL of Hanks Buffered Salt Solution (HBSS) were added to 50µL of trypsin at pH=3 and incubated in a water bath for 60mins at 37°C followed by re-suspension in 90µL HBSS using 10µL sodium bicarbonate and 10µL sodium deoxycholate at ~pH=8 for 40mins at the same temperature. An aliquot of the excysted solution was placed on a microscope
slide and counted under differential interference contrast microscopy for a minimum of 250
counts per sample (Blewett 1989a and Blewett 1989b). Three replicates of both the control and
the solution treated with megasonic energy were counted. The latter solution was exposed to
megasonic agitation for 120 minutes a week before the excystation assay took place. All
samples were stored in the fridge during that time.

2.4. Reagents and equipment

Spiked samples of Cryptosporidium parvum oocysts counted on the flow cytometer (BD
Influx™ cell sorter) were generously provided by Scottish Water. The oocystswere purchased
from the company Creative Science, spin out company from the Moredun Institute, which
produced and isolated these oocysts. Oocysts used for the experiments were prepared about
two months before tests took place and were stored in the fridge. The filters utilised are Filta-
Max Filter Modules from IDEXX (Idexx. 2015) and all other reagents were from Cellabs Pty
Ltd.

Figure 1: Experimental set-up for the elution using megasonic sonication. The sponge filters,
seen at the top of the figure have a doughnut shape when fully expanded and are enclosed in
a plastic bag. The membrane is seen in a smaller bag on the bottom left of the figure. The
megasonic transducer, seen as a black square, is placed at the bottom of the bath filled with
water.
3. Results

3.1. Influence of sonication time during elution on oocysts recovery rate

3.1.1. Sonication of membranes

One of the challenges with optimising elution protocols and maximising recovery rates is the large number of operational sampling parameters that can impact upon the results. These include water type, choice of elution solution, volume and flow rate of elution, spiking volumes/quantities and differences in other sample processing steps and detection methods.

The first parameter investigated in this study was the duration of the sonication and its impact on the recovery rate achieved. As detailed in the “Materials and method” section, this experiment utilised membranes and recovery rates were determined using centrifugation, immunomagnetic separation (IMS) and microscopy. Figure 2 shows the recovery rate of the membranes eluted with megasonic energy as a function of the duration of the elution. The graph clearly demonstrates a time-dependence within the first twenty minutes of elution below which, increasing elution time enhances the recovery rate. After this duration, the recovery rate reaches a plateau at around 45%, which is a rate similar to the control membranes eluted via the traditional method. This result indicates that 20 mins is sufficient to maximise recovery rates.

Figure 2: Recovery rates using elution with megasonic energy assisted agitation.

50mL elution volumes, 100 oocysts spiked into 1mL were passed through the membrane using the traditional set-up. One experiment was carried out at 2, 4, 6, 8, 10 and 120 minutes and two experiments were carried out at 20, 30, 40, 50 and 60 minutes.
3.1.2. Sonication of the IDEXX filters and membranes

Figure 3 shows comparison between controlled tests carried out without megasonic agitation as in the normal procedure and tests with megasonic agitation. In the case of the filters, the control samples underwent traditional process using 1200mL of PBST in the first stage and 50 mL in the second stage followed by centrifugation, IMS and microscope detection of the stained oocysts. For the megasonic samples, the sponge filters were eluted inside the megasonic bath using 1200mL of PBST for 20 minutes and then traditional membrane elution was used for the 2nd stage. The control sample for the membranes underwent traditional process and involving manual rubbing of the membrane. The megasonic sample was eluted into 50mL of PBST for 20 minutes.

The data in Figure 3 were analysed statistically using a one-way Analysis of Variance (ANOVA) test to examine whether there was a significant effect of the use of megasonic energy in the recovery rate of the oocysts during the elution of filters and membranes. The analysis was carried out using Microsoft Excel program for Windows 8 package. The F-ratios was F=0.606 for the filters and F=0.01 for the membranes, both ratios being less than the critical F-ratio, $F_{\text{crit}} (0.05,1,4) = 7.709$, indicating thereby that the analysis fails to reject the null hypothesis of major difference in the achieved recovery rates between the control and the elution carried out using megasonic assisted agitation. There is therefore no significant difference in terms of enhanced recovery rate.

Figure 3: Comparison between controlled tests without megasonic agitation as in the normal procedure and tests with megasonic agitation for both filters and membranes.

Results obtained for an average of 3 replicates, spiked with 100 oocysts.
3.2. Performance of megasonic elution at different volumes

Filters were placed inside a plastic bag with different volumes of PBST to study the performance of megasonic elution. Table 1 shows that the recovery rate increases with the volume of PBST at the 1st stage of the elution.

Table 1: Recovery rate of filter sonicated for 40 minutes using different volumes of PBST. Filters were spiked with 100 oocysts.

Membranes were also placed inside a plastic bag with different volumes of PBST to study the performance of megasonic elution at different volumes. Table 2 shows that the recovery rate falls slightly, from 66% to 53%, when the volume of PBST in the 2nd stage elution is decreased from 50mL to 15mL.

Table 2: Recovery rate of membrane sonicated for 20 minutes using different volumes of PBST. Membranes were spiked with 100 oocysts.

4. A full procedure for megasonic elution

The previous results investigated the impact of megasonic elution for each of the different filtration stages, demonstrating that 20 minutes of megasonic elution is sufficient to match recovery rates achieved by the existing protocol. Thus the main advantage of utilising megasonic elution is in replacing the existing elution method with an easy to use, automatable approach. Additionally, the use of megasonic elution reduces operator variability and should increase the reproducibility of the results in terms of recovery rates of the pathogens. A key finding of this work is that the volume of elution solution can be
reduced such that megasonic elution would allow the centrifugation stage of the traditional process to be skipped as 10mL would be sufficient to achieve the same recovery rate as in the traditional method for the membrane alone as described in Figure 4. This is confirmed by a single factor Analysis of Variance which shows no significant difference between the recovery rate of oocysts with megasonic energy for both stages and the control tests as the F-ratio is F=2.41 which is less than F_{crit}(0.05,1.4)= 7.709.

Figure 5 presents a timeline of the existing and proposed elution methods using megasonic agitation. In addition to removing a stage that is challenging to automate (centrifugation), the sonicated elution for one sample saves approximately 15 minutes and about 600mL of PBST. Although centrifugation could simultaneously process 20 samples, time saving scales with the number of samples as the membrane rubbing cannot be scaled up without increasing the number of operators. However, all membranes could be processed in one step for the megasonic elution approach. Thus the time savings for 20 samples would become 1 hour 50 minutes.

5. Impact of megasonic sonication on oocyst viability

Destruction of oocysts during the elution procedure is obviously undesirable; additionally inactivation of oocysts would prevent any subsequent determination of infectivity. Therefore
experiments were carried out to check the impact of megasonic energy on oocysts. However, an excystation assay is considered a more reliable means of assessing oocyst viability and this test was subsequently performed. The results are shown in Table 3, clearly showing no difference in excystation rate (percentage excystation) or in the sporozoite/shell ratio between the control sample and the sample exposed to elution by megasonic sonication. The excystation assay was performed a week after the megasonic exposure to confirm that the megasonic agitation did not have influence oocyst viability via a slow acting mechanism, which might not have been observed had the excystation assay been undertaken immediately after megasonic exposure.

Table 3: Results of the excystation assay

6. Conclusions

This article reports the first investigation of a novel physical approach for the elution of filters and membranes used in waterborne pathogen monitoring. It is demonstrated that the sonication of filters and membranes using a megasonic transducer preserves the viability of oocysts and achieves recovery rates similar to the established sampling procedure. The key advantages of this novel method lie in (1) the decrease of the volume of the reagents required, (2) the reduction of the manual intervention needed, (3) the reduction of time and resources and (4) the potential for automation. In order to fully document the economic impact of this new elution method, a robust cost of ownership (COO) assessment should be undertaken. Whereas the performance of this method has been demonstrated with Cryptosporidium, an extension of the use of sonication for elution to other pathogens is obviously desirable. The next steps are to work towards an automated filtration/elution
system based on megasonic sonication assisted elution. As centrifugation would no longer be
required, this set-up could then easily be integrated with automated IMS and detection
protocols for a fully automated solution to waterborne pathogen monitoring.

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Table 1: Recovery rate of filter sonicated for 40 minutes using different volumes of PBST. Filters were spiked with 100 oocysts. (n = 3 trials)

<table>
<thead>
<tr>
<th>Volume of PBST (mL)</th>
<th>Mean recovery rate (%)</th>
<th>Standard deviation (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>50</td>
<td>4.66</td>
<td>2.05</td>
</tr>
<tr>
<td>100</td>
<td>16.66</td>
<td>2.86</td>
</tr>
<tr>
<td>500</td>
<td>25</td>
<td>1.63</td>
</tr>
<tr>
<td>600</td>
<td>31.33</td>
<td>1.69</td>
</tr>
<tr>
<td>1200</td>
<td>38.62</td>
<td>2.62</td>
</tr>
</tbody>
</table>
Table 2: Recovery rate of membrane sonicated for 20 minutes using different volumes of PBST. Membranes were spiked with 100 oocysts. *(n =3 trials)*

<table>
<thead>
<tr>
<th>Volume of PBST (mL)</th>
<th>Mean recovery rate (%)</th>
<th>Standard Deviation (SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>50.33</td>
<td>2.05</td>
</tr>
<tr>
<td>20</td>
<td>61</td>
<td>1.63</td>
</tr>
<tr>
<td>50</td>
<td>65.66</td>
<td>1.24</td>
</tr>
</tbody>
</table>
Table 3: Results of the excystation assay

<table>
<thead>
<tr>
<th>Sample</th>
<th>Excystation percentage</th>
<th>Sporozoite/shell ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control ((n =1 \text{ trial}))</td>
<td>97</td>
<td>2.4</td>
</tr>
<tr>
<td>Megasonic ((n =3 \text{ trials}))</td>
<td>Mean value = 96</td>
<td>Mean value = 2.26</td>
</tr>
<tr>
<td></td>
<td>SD = 1.11</td>
<td>SD = 0.36</td>
</tr>
</tbody>
</table>
Figure 1: Experimental set-up for the elution using megasonic sonication. The sponge filters, seen at the top of the figure have a doughnut shape when fully expanded and are enclosed in a plastic bag. The membrane is seen in a smaller bag on the bottom left of the figure. The megasonic transducer, seen as a black square, is placed at the bottom of the bath filled with water.
Figure 2: Recovery rates using elution with megasonic energy assisted agitation. 50mL elution volumes, 100 oocysts spiked into 1mL were passed through the membrane using the traditional set-up. One experiment (n=1) was carried out at 2, 4, 6, 8, 10 and 120 minutes and two experiments (n=2) were carried out at 20, 30, 40, 50 and 60 minutes.
Figure 3: Comparison between controlled tests without megasonic agitation as in the normal procedure and tests with megasonic agitation for both filters and membranes. Results obtained for an average of 3 replicates, spiked with 100 oocysts.
Figure 4: Recovery rates of controlled tests without megasonic and full megasonic elution using 600ml in the first stage and only 10ml in the 2\textsuperscript{nd} stage elution and detection process without centrifugation.
<table>
<thead>
<tr>
<th>Time</th>
<th>Existing method</th>
<th>Megasonic elution</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Place the filter module into the filter housing</td>
<td>Unscrew the filter, place the sponges inside the plastic bag and put the bag into the megasonic bath</td>
</tr>
<tr>
<td></td>
<td>Wash the sample with PBST 1200 ml 20-30 minutes</td>
<td>600 ml PBST</td>
</tr>
<tr>
<td></td>
<td>Concentrate the sample through a membrane</td>
<td>Conserve the sample through a membrane</td>
</tr>
<tr>
<td></td>
<td>Rub the membrane and put the sample from the bag to a 50 ml centrifuge tube. Add 5-10 ml PBST 2 minutes</td>
<td>Place the membrane inside the plastic bag and put the bag inside the megasonic bath 5-10 ml PBST 20 minutes</td>
</tr>
<tr>
<td>2 minutes</td>
<td>Centrifugation 50 ml Tube 30 minutes</td>
<td>IMS separation Max: 10 ml volume</td>
</tr>
<tr>
<td></td>
<td>IMS separation Max: 10 ml volume</td>
<td>To detection</td>
</tr>
</tbody>
</table>

*Figure 5: Schematic timeline describing the savings in terms of process time and volume of PBST of the megasonic elution compared to the traditional elution method.*