Research Paper

The inactivation of *Ascaris suum* eggs by short exposure to high temperatures

D. Naidoo, C. C. Appleton, C. E. Archer and G. L. Foutch

ABSTRACT

*Ascaris* sp. is the most prominent and resilient helminth of human health importance found in faecal sludge, making *Ascaris* sp. an ideal index organism for inactivation testing. Heat treatment destroys helminths, allowing for safe handling and possible reuse of sludge. Technology development focuses on rapid heating to minimize equipment size and cost. This study evaluates *Ascaris suum* eggs' viability with short heating time. *Ascaris* eggs were placed in a water bath at temperatures from 60 to 80 °C for various exposure times (5 seconds to 4 minutes) and were immediately processed and analysed via light microscopy. For all samples within these temperature and time ranges, less than 10% viable eggs were recovered. For 70, 75 and 80 °C, complete inactivation was observed for exposure time of 5 seconds and above.

**Key words** | *Ascaris suum*, heat treatment, inactivation, temperature, viability

INTRODUCTION

Approximately 2.3 billion people globally, and 695 million of the sub-Saharan Africa population, do not have access to improved sanitation (JMP 2017; Roche et al. 2017), with an estimated 1 billion performing open defecation (Tiwari et al. 2017). Approximately 2 billion people globally are infected with soil-transmitted helminths (Yadav & Mahato 2017). The inability of countries to provide water, sanitation and hygiene (WASH) results in 1.3 million deaths annually due to diarrhoeal diseases. Diarrhoeal disease is the cause of one in eight child mortality cases in children under five years of age worldwide (Kotloff 2017).

Diarrhoea manifests generally as a symptom of bacterial and viral infections but may also be a symptom of infection by parasitic worms (helminths), with *Ascaris lumbricoides* of greatest concern (Brownell & Nelson 2006). *Ascaris* eggs are the most resilient of all organisms found in sludge, as they can withstand harsh environmental conditions, such as desiccation, and can survive in both aerobic and anaerobic environments for up to seven years in the soil (Pecson & Nelson 2005). *Ascaris* spp. are therefore deemed fit as index organisms for parasitic contamination, as well as overall pathogenic contamination and inactivation (Sidhu & Toze 2009). If a treatment process can inactivate *Ascaris* spp. eggs, then it is very likely all other pathogens will be destroyed as well (Maya et al. 2012). Ethical approval for the collection of stool samples from human hosts (to obtain the human roundworm, *Ascaris lumbricoides*) is difficult. *Ascaris suum*, a parasite of pigs, is therefore sometimes used as a surrogate for *A. lumbricoides*, as the two species are morphologically...
indistinguishable from one another in all stages (Daugschies et al. 2013). A. suum eggs were therefore used in this study.

Ventilated pit latrines (VIP) are regarded as improved sanitation by the South African government. Once full, the pits are sealed or emptied (Bhagwan et al. 2008). Proper disposal and treatment of human excreta is important in reducing transmission of infectious diseases. Land application of treated sludge is becoming increasingly common, thereby encouraging resource reuse and recovery, alleviating environmental contamination and reducing human health impacts (Fewtrell & Bartram 2001).

Increased temperatures have been found to be effective and cost efficient for inactivation of faecal-sludge pathogens (Podichetty et al. 2014; Belcher et al. 2015). This study is an extension of a short communication by Naidoo & Foutch (2017) and aimed to fill the knowledge gap with regards to the effects of short exposures of Ascaris eggs to heat. It was part of a larger project funded by the Bill and Melinda Gates Foundation (BMGF), which focused on development of a prototype viscous heater for sludge treatment by heat exposure. Thomas et al. (2015) focused on a range of temperatures at exposure times >10 minutes. Brannen et al. (1975) and Thomas et al. (2015) reported Ascaris inactivation at 64 and 70 °C, respectively, for 1 minute exposure. No studies focus on inactivation of Ascaris eggs for exposure times less than a minute (Thomas et al. 2015). Test parameters for this study ranged from 60 to 80 °C, at exposure times of 5 seconds to 4 minutes, simulating the effects of the viscous heater during treatment. The isolated effects of heat on Ascaris eggs in water was investigated, rather than sludge, which might serve as a matrix to protect eggs against heat exposure.

MATERIALS AND METHODS

A. suum eggs were purchased from Excelsior Sentinel Inc. (USA) and stored at 4 °C until needed. Egg stock solutions were prepared from the initial egg samples with approximately 550 eggs per 1 mL of solution. A 34-litre heating water bath controlled experimental exposure temperatures. Eggs were exposed to heat treatments in 15 mL polypropylene test tubes preheated in the water bath by adding 13 mL of boiling water per tube and immersing these tubes into the water via a modified polystyrene rack. A thermocouple and data logger monitored temperature over time. Eggs were analysed via light microscopy, before and after incubation at 25 °C for 28 days in a cooling incubator.

Experimental temperatures were 60, 65, 70, 75 and 80 °C, and exposure times 5, 10, 15, 30 and 45 seconds, and 1, 2, 3 and 4 minutes. For 80 °C, shorter exposure times of 1–10 seconds were also tested. The water bath was preheated prior to each experiment. The thermocouple was placed into the tube prior to exposure to ensure steady-state temperature (Figure 1). For each treatment, a 1 mL aliquot of egg stock solution was placed into a 15 mL test tube and used to assess developmental states of the eggs before exposure to heat. This allowed counting of the total number of eggs per 1 mL of solution. One mL aliquots of egg stock solutions (approximately 550 eggs per 1 mL) were pipetted into the test tubes using a 1 mL Pasteur pipette and time recorded as T0. Eggs were exposed for the respective times, after which the tube was immediately removed from the water bath. Samples were returned to room temperature to avoid prolonged heat exposure, by emptying the contents of the test tube (i) into beakers containing 40–50 mL iced water at approximately 9 °C or (ii) onto a 20 μm sieve within a bowl containing tap water at room temperature (Figure 1(d)). Control samples were not heated but were cooled and processed similarly to treated samples.

Eggs were processed following Moodley et al. (2008) and Pebsworth et al. (2012), modified by Naidoo et al. (2016). Ammonium bicarbonate (made up with deionized water to form 99% ammonium bicarbonate, NH₄HCO₃ by Sigma) was used to rinse the test tubes and beaters that previously contained treated samples. The tube or beaker contents were rinsed thoroughly over a 20 μm sieve. The deposit was transferred into a plastic 15 mL test tube using a Pasteur pipette and centrifuged at 1,310 × g (3,000 rpm) for 5 minutes using a bench-top centrifuge (with a swing-out rotor). The supernatant was discarded, and the egg pellet was analysed under a light microscope at 100× and 400× magnification.

Egg viability was assessed by pipetting two drops of egg suspension onto a plain glass microscope slide and covering with a 40 × 22 mm coverslip. The eggs were categorized as either potentially viable or non-viable based on morphology.
Potentially viable (PV) eggs included those that were undeveloped (at different cell stages), embryonated with a visibly motile larva, and embryonated containing an immobile larva (Naidoo et al. 2016). Non-viable (NV) eggs included those that were dead (globules inside the egg shell), embryonated with a necrotic larva, and mechanically broken (Naidoo et al. 2016). Samples were analysed before and after incubation such that treated samples were immediately analysed, washed back into the test tube after counting, incubated for 28 days at 25 °C and then re-analysed after incubation. This process was repeated for all temperatures and exposure times and for controls. Eggs were further analysed via oil immersion microscopy (1,000×, 1,500× and 2,000×) in order to evaluate heat damage in greater detail (three-gear focusing using a magnification changer allowed for magnifications greater than 1,000×).

Normality of data was tested using a one-sample Kolmogorov–Smirnov test using IBM SPSS Statistics (version 24, IBM Corp., Armonk, NY, USA) and transformed accordingly using arcsine transformation, then analysed statistically. A nested analysis of variance (ANOVA) was run on the data using R (version 3.0.2, R Core Team 2016). The Shapiro–Wilk test was used to test for normality and a Levene’s test was used to test for homogeneity of
variance of the studentized residuals from the ANOVA. Tukey’s multiple comparisons post-hoc test was used to compare means of egg viability across and between different treatments. Percentage viability was calculated as follows:

\[
\text{Percentage viable eggs recovered} = \frac{\text{Total viable eggs recovered}}{\text{Total viable eggs inoculated}} \times 100(1)
\]

An independent samples t-test was run on percentage viable eggs recovered, comparing processing with iced or tap water. A second independent samples t-test was run on percentage of viable eggs recovered before and after incubation for each replicate of each treatment (IBM SPSS v.24). The criterion set for this study for successful inactivation was ≤ < 10% viable eggs recovered after treatment. This is similar to a study by Ayçiçek et al. (2001) and Naidoo et al. (2016), which considered 90% mortality of eggs as significant.

## RESULTS

Eggs that appeared undamaged and remained at the one-celled stage after treatment were scored as undeveloped pre-incubation, as embryo death could only be confirmed post-incubation. The percentage of viable eggs recovered pre-incubation is therefore much greater than post-incubation, especially for treatment combinations that did not yield visible egg damage. The p-values are not only indicators of significant differences between percentage of viable eggs recovered, but also the level of damage and the general morphological states between exposure times. Samples exposed for 5 seconds at 70, 75 and 80 °C resulted in the most significant shift in egg state, post-incubation (from undeveloped to developed) and were used as the baseline for comparison of successive exposure times (the least damage could be seen after 5 seconds). The general ANOVA (Table 1) indicated that temperature, independently, and in conjunction with secondary variables such as exposure time, processing with iced or tap water and point of analysis, had a significant effect on percentage of viable eggs recovered (p < 0.001).

The first level of nestedness compares each temperature (primary variable) against other temperatures and two controls for the percentage of viable eggs recovered after treatment. No significant difference was observed between control 1 (iced water) and control 2 (tap water) (p = 0.760), indicating that the processing method had no effect on percentage viability (note: controls were not heat treated but were processed by the same method as treated samples). There was, however, a significant difference in viable eggs recovered between each test temperature combination from 60 °C to 80 °C (p < 0.001), as well as between controls and heat-treated samples (p < 0.001).

The second nested level compares the effect of each test temperature (primary variable) on the percentage of viable eggs recovered after treatment across various exposure times (secondary variable). Samples were compared across temperatures at each exposure time. When comparing the controls with 60 °C, no significant difference in development was seen up to 15 seconds (p > 0.05), however exposure times of 50 seconds and above showed a significant decline in development (Figure 2(a)). There was a significant difference in development up to 15 seconds (p < 0.001) between 60 and 65 °C (comparing Figure 2(a) and 2(b)). From 50 seconds onwards, there were no significant differences between each exposure time (p > 0.05), and a similar trend was seen for 65 °C. Viable egg recovery

| Primary variable         | F(DF) | p-Value | Variable combinations   | p-Value |
|--------------------------|-------|---------|-------------------------|---------|
| Temperature              | F(6, 552)=1705.8 | <0.001 | Temperature/Exposure time | <0.001 |
|                          |       |         | Temperature/Exposure time/Processing | <0.001 |
|                          |       |         | Temperature/Exposure time/Processing/Analysis | <0.001 |

*Variable combinations represent the nestedness of the statistical design. Processing refers to the method used (iced or tap water) and analysis refers to whether eggs were examined before or after incubation.
had dropped to 0% after 4 minutes’ exposure, although at 60 °C, minimal development was still evident. At 60 °C, percentages of viable eggs recovered post-incubation were mostly <10% per treatment, and almost 0% at 3 and 4 minutes (Figure 2(a)). Little visible egg damage was evident pre-incubation, thus they were initially classified as viable (undeveloped eggs). When considered together with the low viability percentages shown in Figure 2(a), successful inactivation at 60 °C can be attributed to exposures of ≥45 seconds, which met the criterion for this study.

Significant egg viability differences were observed between 5 seconds and every time interval from 10 seconds to 4 minutes (5:10 seconds, \( p = 0.002 \); 5:15 seconds up to 4 minutes, \( p < 0.001 \)), indicating an increase in visible damage after 5 seconds when exposed to 65 °C (Figure 2(b)). Between exposures of 5 and 10 seconds, the percentage of viable eggs dropped to approximately 10% (post-incubation), and almost zero after 15 seconds onwards. There was no significant change in the state of eggs after treatment from 10 seconds onwards and between each time interval from 10 seconds onwards and between each time interval.
(p > 0.05), confirming that successful inactivation (≤10% viable eggs recovered) occurred after exposures of 10 seconds and longer, meeting the criterion of this study. Complete die-off was, however, not achieved fully for either 60 or 65 °C as isolated cases of egg development were evident even after 4 minutes of exposure.

When comparing across temperatures between 60 and 70 °C, 75 and 80 °C, significant differences in egg viability were noted at all test exposure times, indicating a decline in egg development (thus increased effectiveness of treatment of the eggs) for each successive exposure time (Figure 2(a)-2(e)). The same can be said for comparisons between 65 and 75 °C and between 65 and 80 °C exposures (p < 0.001). As seen in Figure 2(c), visible damage occurred only at 2 minutes’ exposure to 70 °C, however significant die-off occurred from 5 seconds’ exposure onwards. A similar level of damage to the eggs occurred after treatment at 70 °C across all exposure times (p > 0.05 between each time interval). At 75 °C, a significant difference in egg viability was observed between 5 seconds and all subsequent exposure times from 10 seconds to 4 minutes (p < 0.001), indicating an increase in visible damage from ≥10 seconds’ exposure (Figure 2(d)).

No significant differences in egg viability were observed between 5 seconds and all other exposure time from 10 seconds to 4 minutes (p > 0.05), indicating that visible damage occurred from the first exposure time point (5 seconds) when exposed to 80 °C (Figure 2(e)). Further analysis of exposure to 80 °C for shorter periods (1–10 seconds) showed that visible damage to the eggs was only apparent after exposures of 4–5 seconds (Figure 2(f)). Although eggs appeared healthy and undamaged after exposure to 80 °C for 1–10 seconds, most failed to develop further after incubation. Some development was evident for eggs exposed for up to 2 seconds. Even so, development was arrested at stages of cleavage between two cells and the gastrula.

Results from the independent samples t-test between the two processing methods (cooling by iced and tap water) indicated no significant difference in the percentage of viable eggs recovered after treatment (p > 0.05). The third level of comparison of the nested ANOVA (temperature/exposure time/processing method) was therefore not outlined in further detail.

**DISCUSSION**

Table 2 summarizes the results from the inactivation trials and indicates which exposure time was most effective in inactivating eggs for each test temperature. It highlights the exposure time at which damage occurred but was not visible (where further development did not occur post-incubation), visible damage occurred (globular eggs seen as indicated by Naidoo et al. (2016) to be explored in a second article) and complete die-off was achieved (where no viable eggs were recovered after treatment). For each column, a decreasing trend can be observed, indicating that with increasing temperature, the time required for damage (both non-visible and visible) and egg inactivation to occur, decreases. For 60 and 65 °C, die-off did not reach a 0% level, with few eggs able to develop, although viability percentages after incubation were negligible.

The current study indicated that heat treatment is a successful inactivation method when applied over a range of temperatures. This study also contributes towards filling the literature gap that exists for short temperature exposures and the resulting effect on the viability of Ascaris eggs. Exposure to each of the test temperatures was successful on its own, and eventually reached either negligible viability percentages (60 and 65 °C) or complete die-off (0% viability at 70, 75 and 80 °C), after exposure for up to 4 minutes.

Table 2 | Summary of the Ascaris egg viability results for each test temperature of the current study

| Temperature (°C) | Damage not visible | Visible damage | Complete die-off |
|------------------|--------------------|----------------|------------------|
| 60 °C            | 30 seconds         | 3 minutes      | –                |
| 65 °C            | 15 seconds         | 5 minutes      | –                |
| 70 °C            | 5 seconds          | 2 minutes      | 15 seconds       |
| 75 °C            | 5 seconds          | 1 minute       | 10 seconds       |
| 80 °C            | 5 seconds          | 5 seconds      | 5 seconds        |
| 80 °C – Shorta   | 1 second           | 4 seconds      | 4 seconds        |

*aShort refers to the repeated 80 °C experiment with exposure times between 1 and 10 seconds.

*bComplete die-off was confirmed by microscopic analysis post-incubation.
As already mentioned, exposures at 60 and 65°C resulted in almost 100% inactivation, except for a few eggs that developed after incubation. Popat et al. (2010) investigated inactivation of *A. suum* by anaerobic digestion at thermophilic temperatures (thermophilic refers to the ability of an organism, in this case bacteria, to survive temperatures ranging from 46 to 108°C). It was reported that there was a reduction in egg viability after 2 hours of heat treatment at 51.1–55°C and that the composition of the sludge might play a role in the effectiveness of heating.

Brannen et al. (1975) and Brandon (1978) both investigated the effects of heat exposure on the embryonation of *Ascaris lumbricoides* eggs, in a controlled-temperature water bath, similar to that used in the current study. Brannen et al. (1975) reported a near complete inhibition of egg development when exposed to 55°C for 4 minutes and <1% embryonation of eggs when exposed to 64°C for 1 minute. Brandon (1978) reported that 1 hour of exposure to 55°C was sufficient for reducing the number of viable *Ascaris* eggs to near negligible amounts. Both studies support the findings of the current study, but only at the lower test temperatures, showing a similar relationship between exposure to heat and egg inactivation (Figure 2).

Steer & Windt (1978) investigated the effects of composting on *Ascaris* eggs. They concluded that temperatures of 65°C should be maintained for 70 days within the composting system for successful inactivation of eggs. The results of the current study challenge the recommendations of Steer & Windt (1978), as exposure to 65°C for 30 seconds was sufficient for negligible inactivation percentages (Table 2 and Figure 2(b)). It can be argued that the different methods of heat exposure in the two studies resulted in the discrepancy between the reported inactivation rates.

Thomas et al. (2015) investigated the relationship between *A. suum* egg inactivation and the effects of heat and shear stress/force independently and combined. The methods for testing heat treatment were similar to the current study. Human-faecal simulants were spiked with *A. suum* eggs and smeared along the inside of 7 mL polyethylene vials (Thomas et al. 2015). The vials were sealed and wrapped inParafilm to prevent water entering, and were fully submerged in a water bath for 60 seconds at 40°C. Treatments increased in 10°C intervals up to 100°C. Cooling to room temperature was done by immersing the heated vials in a 27°C water bath for 60 seconds. At 47, 51 and 55°C, the embryonation rates of eggs were 94.7, 91.1 and 89.7%, respectively. This aligns with the current study, which found that eggs were able to continue development at temperatures up to 65°C when exposed to heat for up to 2 minutes (viability is visible in Figure 2(b), even if negligible). Low temperature treatment therefore requires longer exposure times.

At 70°C, 5 seconds’ exposure was sufficient to halt further embryo development, with 100% inactivation occurring from 15 seconds onwards. Thomas et al. (2015) stated that for temperatures of 70°C and above, almost no viable eggs were recovered after treatment. This supports the data presented in the current study, where egg development, if any, at temperatures from 70°C onwards, approached zero viable egg recovery (Figure 2(c)). Belcher et al. (2015) published the first study on *Ascaris* eggs based on the viscous heater, that serves as a preamble to the current study which provides important foundation work on the effects of temperature, independently, on *Ascaris* eggs. A baseline for egg inactivation was determined by feeding seeded sludge into the heater. Representative samples were taken for each inlet pump and rotating core speed combination, at increasing and plateau temperatures, in order to determine egg viability. The operational speeds also determined residence time for specific generated temperatures. The findings of Belcher et al. (2015) were in line with Thomas et al. (2015), by recording 90% egg inactivation at temperatures above 70°C, further supporting the findings of the current study. At 75 and 80°C and ≤10 seconds of heat exposure, 100% egg inactivation was achieved (Figure 2(d)–2(f)). In most cases, all structural and morphological integrity of both the egg and the contained embryo were lost. Belcher et al. (2015) also reported 100% egg inactivation when exposed to 85°C with residence times under 1 minute, which was in line with the present study’s findings of 100% inactivation at 80°C for 4 seconds.

Pecekon & Nelson (2003) investigated the effects of temperature, exposure time, pH and ammonia on the inactivation of *Ascaris* eggs. Eggs were stored at different pH levels, ammonia concentrations and temperatures for 24 hours. Between a threshold temperature of 44 to 48°C, eggs went from low to complete inactivation. In the absence of ammonia, 99% inactivation was reported at 48°C and pH 7, 9 and
When the ammonia concentration was held constant at 1,000 ppm with pH 7 at 48 °C, pH 9 at 44 °C and pH 11 at 44 °C, 99% inactivation was achieved. At lower pH values, the presence of ammonia had a negligible effect on inactivation rates, as a temperature of 48 °C was still required for 99% inactivation. At higher pH values, the presence of ammonia reduced the temperature required for 99% inactivation, indicating that the addition of ammonia is a valuable secondary treatment option in combination with temperature. This is because the alteration of eggshell permeability by heat may allow for easier penetration of the ammonia (or possibly other chemical treatments) (Barrett 1976).

CONCLUSIONS AND RECOMMENDATIONS

- All test temperatures (between 5 seconds and 4 minutes) met the criterion for this study, i.e., inactivation was considered successful if the recovery of viable eggs after treatment and incubation was ≤10% (Figure 2(a)–2(f)).
- Complete die-off within the tested exposure time range was noted for 70 °C, 75 °C and 80 °C, however treatment at 60 °C and 65 °C allowed for development of a few eggs after incubation.
- Incubation of heat-treated samples is required when egg damage is not visible, in order to confirm die-off.
- The results of the current study therefore show that residence times of as low as 4 seconds at 80 °C may be recommended for successful inactivation when using the viscous heater.

Field application of treatment processes, as well as treatment in the indigenous medium and not a saline suspension may result in differing inactivation levels. As explained by Jebri et al. (2015) and Mun et al. (2009), the suspension medium of the eggs determines the extent of heat treatment. Buttar et al. (2015) reported that sludge or any similar faecal simulant may act as insulation for eggs exposed to heat, suggesting that higher temperatures and prolonged exposure times might be needed for testing eggs in vitro. The role of the suspension medium during heating needs to be investigated. Further work is also required to test the actual effects of heat treatment of the eggs using the viscous heater.

ACKNOWLEDGEMENTS

The authors of this article would like to thank the Bill & Melinda Gates Foundation (OPP1141554) for funding throughout this project.

REFERENCES

Ayçiçek, H., Yarsan, E., Sarimehmetoğlu, H. O., Tanyüksel, M., Girginkardeşler, N. & Özyurt, M. 2001 Efficacy of some disinfectants on embryonated eggs of Toxocara canis. Turkish Journal of Medical Science 31, 35–39.
Barrett, J. 1976 Studies on the induction of permeability in Ascaris lumbricoides eggs. Parasitology 73 (1), 109–121.
Belcher, D., Foutch, G. L., Smay, J., Archer, C. & Buckley, C. A. 2015 Viscous heating effect on deactivation of helmint eggs in ventilated improved pit sludge. Water Science and Technology 72 (7), 1119–1126.
Bhagwan, J. N., Still, D., Buckley, C. & Foxon, K. 2008 When last did we look down the pits? WISA Paper. Water Institute of South Africa, Midrand, 11.
Brandon, J. R. 1978 Parasites in Soil/Sludge Systems. Department of Energy, Sandia Laboratories, available from the National Technical Information Service, Springfield, VA, USA.
Brennen, J. P., Garst, D. M. & Langley, S. 1975 Inactivation of Ascaris lumbricoides Eggs by Heat, Radiation, and Thermoradiation (No. SAND–75–0165). Sandia Labs, Albuquerque, NM and Livermore, CA, USA.
Brownell, S. A. & Nelson, K. L. 2006 Inactivation of single-celled Ascaris suum eggs by low-pressure UV radiation. Applied and Environmental Microbiology 72 (2), 2178–2184.
Buttar, B. S., Nelson, M. L., Busboom, J. R., Hancock, D. D., Walsh, D. B. & Jasmer, D. P. 2015 Effect of heat treatment on viability of Taenia hydatigena eggs. Experimental Parasitology 113, 421–426.
Daugschies, A., Bangoura, B. & Lendhner, M. 2013 Inactivation of exogenous endoparasite stages by chemical disinfectants: current state and perspectives. Parasitology Research 112, 917–952.
Fewtrell, L. & Bartram, J. 2001 Guidelines, Standards and Health: Assessment of Risk and Risk Management for Water-Related Infectious Disease. World Health Organization, Geneva, Switzerland.
Jebri, S., Hmaied, F., Jofre, J., Mendez, J., Barkallah, I. & Hamdi, M. 2015 Effect of gamma irradiation on bacteriophages used as viral indicators. Water Research 47 (11), 3673–3678.
Joint Monitoring Programme (JMP) 2017 Progress on Drinking Water, Sanitation and Hygiene – 2017 Update and SDG Baseline. Geneva, Switzerland.
Kotloff, K. L. 2017 The burden and etiology of diarrheal illness in developing countries. Pediatric Clinics 64 (4), 799–814.
Maya, C., Torner-Morales, F. J., Lucario, E. S., Hernandez, E. & Jiménez, B. 2012 Viability of six species of larval and non-larval
helminth eggs for different conditions of temperature, pH and dryness. Water Research 46, 4770–4782.
Moodley, P., Archer, C., Hawksworth, D. & Leibach, L. 2008 Standard Methods for the Recovery and Enumeration of Helminth ova in Wastewater, Sludge, Compost and Urine-Diversion Waste in South Africa: Report to the Water Research Commission. Water Research Commission, South Africa.
Mun, S., Cho, S. H., Kim, T. S., Oh, B. T. & Yoon, J. 2009 Inactivation of Ascaris eggs in soil by microwave treatment compared to UV and ozone treatment. Chemosphere 77 (2), 285–290.
Naidoo, D. & Foutch, G. L. 2017 The time-temperature relationship for the inactivation of Ascaris eggs. Journal of Water Sanitation and Hygiene for Development 8 (1), 123–126.
Naidoo, D., Archer, C., Louton, B. & Rodda, N. 2016 Testing household disinfectants for the inactivation of helminth eggs on surfaces and in spills during pit latrine emptying. Water SA 42 (4), 560–570. https://dx.doi.org/10.4314/wsa.v42i4.06.
Pebsworth, P. A., Archer, C. E., Appleton, C. C. & Huffman, M. A. 2012 Parasite transmission risk from geophagic and foraging behaviour in chacma baboons. American Journal of Primatology 74 (10), 940–947.
Pecson, B. M. & Nelson, K. L. 2003 The effects of exposure time, temperature, pH, and ammonia concentration on the inactivation rate of Ascaris eggs. Proceedings of the Water Environment Federation 2003 (10), 534–539.
Pecson, B. M. & Nelson, K. L. 2005 Inactivation of Ascaris suum eggs by ammonia. Environmental Science & Technology 39 (20), 7909–7914.
Podichetty, J. T., Islam, M. W., Van, D., Foutch, G. L. & Johannes, A. H. 2014 Viscous heating analysis of simulant faeces by computational fluid dynamics and experimentation. Journal of Water, Sanitation and Hygiene for Development 4 (1), 62–71.
Popat, S. C., Yates, M. V. & Deshusses, M. A. 2010 Kinetics of inactivation of indicator pathogens during thermophilic anaerobic digestion. Water Research 44 (20), 5963–5972.
Roche, R., Bain, R. & Cumming, O. 2017 A long way to go – estimates of combined water, sanitation and hygiene coverage for 25 sub-Saharan African countries. PloS One 12 (2), e0171783.
Sidhu, J. P. & Toze, S. G. 2009 Human pathogens and their indicators in biosolids: a literature review. Environment International 35 (1), 187–201.
Steer, A. G. & Windt, C. N. 1978 Composting and fate of Ascaris lumbricoides ova. Water SA 4 (3), 129–132.
Thomas, J. E., Podichetty, J. T., Shi, Y., Belcher, D., Dunlap, R., McNamara, K., Reichard, M. V., Smay, J., Johannes, A. J. & Foutch, G. L. 2015 Effect of temperature and shear stress on the viability of Ascaris suum. Journal of Water Sanitation and Hygiene for Development 5 (3), 402–411.
Tiwari, A., Russpatrick, S., Hoehne, A., Matimelo, S. M., Mazimba, S., Nkhata, L., Osbert, N., Winters, A., Winters, B. & Larsen, D. A. 2017 Assessing the impact of leveraging traditional leadership on access to sanitation in rural Zambia. The American Journal of Tropical Medicine and Hygiene 97 (5), 1355–1361.
Yadav, S. N. & Mahato, S. 2013 Study on intestinal helminth parasites in school children of Rangeli Municipality of Morang District in Eastern Nepal. American Journal of Health Research 5 (2), 50–53.