Microbial quality and suitability of roof-harvested rainwater in rural villages for crop irrigation and domestic use

Mosimanegape Jongman and Lise Korsten

ABSTRACT

The study aimed at assessing the microbiological quality and suitability of roof-harvested rainwater (RHRW) for crop irrigation and domestic use. In total, 80 rainwater tanks (246 samples) across three rural villages (Ga-Molepane, Jericho and Luthngele) were visited. Culture-based techniques were used to isolate bacterial microbes and identities were confirmed using matrix-assisted laser desorption/ionization time of flight (MALDI-TOF-MS). Uncultured fungal populations were also identified using pyrosequencing. Salmonella spp. (3%), Listeria monocytogenes (22%), total coliforms (57.7%), Escherichia coli (30.5%), Enterococcus spp. (48.8%), Pseudomonas spp. (21.5%) were detected in RHRW samples after rainfall. Fungal sequences belonging to species known to cause fever, coughing and shortness of breath in humans (Cryptococcus spp.) were identified. This study indicates that RHRW quality can be affected by external factors such as faecal material and debris on rooftops. The use of untreated RHRW could pose a potential health risk if used for irrigation of crops or domestic use, especially in the case of a relative high population of immunocompromised individuals. This study does not dispute the fact that RHRW is an alternative irrigation water source but it recommends treatment before use for domestic purposes or for watering crops.

Key words | human pathogens, MALDI-TOF-MS, microbial quality, pyrosequencing, roof-harvested rainwater

INTRODUCTION

Approximately 11% of the world's population is without access to quality clean water. Despite all the development and infrastructure, an estimated 1,800 million people will still lack access to safe water by 2025 (UN 2012). Due to current water shortages, the use of potable municipal drinking water for irrigation in South Africa is not encouraged (Gemmell & Schmidt 2012). As a result, microbial quality of alternative water sources has received increased attention not only in South Africa but worldwide. Harvested rainwater is a realistic alternative source that can be used for irrigation and drinking purposes. Roof-harvested rainwater (RHRW) has been used for potable and non-potable purposes in many countries (Ahmed et al. 2014). Despite public health perception, rainwater poses a low risk to human health (Ahmed et al. 2014; Dobrowsky et al. 2014; Chidamba & Korsten 2015a). Various microorganisms, including human pathogens, could be present in dust, faeces of birds, insects and sometimes small mammals present on rooftops (Ahmed et al. 2010). During rainfall, faeces and other debris harbouring pathogens could be washed into rainwater harvesting tanks thus degrading its quality (Pachepsky et al. 2011). Pathogens have been reported in RHRW (Lye 2002; Lee et al. 2012; Ahmed et al. 2014) including bacteria, viruses and parasites, many of which could lead to waterborne illnesses (Hageskal et al. 2009).
The most frequently studied group of microorganisms with respect to microbial quality of water is bacteria. The presence of potentially pathogenic microorganisms, such as *Listeria monocytogenes*, *Campylobacter* spp., *Salmonella* spp., *Escherichia coli* (E. coli), *Cryptosporidium* spp. and *Aeromonas* spp. in RHRW has been reported (Ahmed et al. 2012; De Kwaadsteniet et al. 2013). Fungi are usually ignored with respect to microbial quality of water, mainly because many disease outbreaks related to use of contaminated water are caused by pathogenic bacteria (Hageskal et al. 2009). However, many fungal species have been detected in water including *Fusarium oxysporum* Schltdl., *Fusarium moniliforme* Sheld., *Aspergillus flavus* Link, *Penicillium chrysogenum* Thom., *Penicillium coryophilum* Dierckx, *Acremonium strictum* Gams, *Phialophora bubakii* Schol. and *Phoma exigua* Sacc. (Monchy et al. 2011); some of these are pathogenic to humans or are known plant pathogens. Hageskal et al. (2009) stated that *Aspergillus fumigatus* Fresen is one of the most important fungal pathogens detected in water that could cause infections in immunocompromised individuals. The presence of pathogenic fungi in water has been considered as a chronic and possibly underestimated problem (Hageskal et al. 2009). Plant pathogens play an important role in infecting irrigated crops during the seedling stage (Bell et al. 2006), growing conditions or post-harvest (Prusky 1996). No study has reported on the link between RHRW and crop disease or decay potential due to the presence of plant pathogens.

Reports on the role of irrigation water in the contamination of produce, subsequently leading to foodborne illness outbreaks are increasing (Allende & Monaghan 2015). The transfer of any pathogen in water used for irrigation onto crops could be risky if produce is consumed fresh. Studies have linked the presence of pathogens on fresh produce to contaminated water (Pachepsky et al. 2011; Du Plessis et al. 2013). The presence and subsequent transfer of fungal pathogens from irrigation water onto fresh produce has not been fully investigated.

Traditional culturing assays underestimate microbial diversity due to methodological constraints and limitations including the ones used for fungal populations. Previous studies on fungal analysis of water used only culturing methods. Disadvantages of culturing fungi include (1) inability to grow non-sporulating fungi and (2) diverse nutrient requirements (Hageskal et al. 2009). As a result, reliance on molecular methods has received global recognition. The 454 pyrosequencing platform is a recent type of next generation sequencing technology. This technique is a good diagnostic tool for detection and identification of microbial pathogens. Despite being an expensive technique, the 454 pyrosequencing technique is rapid and flexible with 99% accuracy (Liu et al. 2015). Therefore, this study aims to assess the microbiological quality of RHRW for irrigation and domestic purposes using culturable and non-culturable approaches for bacteria and fungi, respectively. In addition, the presence of potential human pathogens (bacterial and fungal) was also assessed as part of the microbial population.

**MATERIALS AND METHODS**

**Study sites and sample collection**

Sampling was conducted at three rural villages in South Africa: Ga-Molepane (Limpopo Province), Jericho (North West Province) and Luthengele (Eastern Cape Province). All villages experience serious water shortages, hence the communities rely on RHRW. Ga-Molepane and Luthengele villages have no boreholes and households depend entirely on RHRW during the wet season supplemented with stream water during dryer months. Domesticated animals use the stream as a water source. Jericho village has boreholes which often are not adequate to supply the community with enough water for domestic use. At Luthengele village, leafy green vegetables (mainly cabbage) are grown but RHRW is not used for watering. In total, 246 RHRW samples were collected from water tanks during the 2013–2014 rainy seasons (September to May). The strategy was to sample before and after the onset of the main rainy season. Randomly selected households (80) were used for this study. Each site was visited twice in 2013 and repeated in 2014 following the same strategy, before and after onset of the rainy season and sampling from the same households. Sanitary factors, i.e., overhanging trees, animal faecal material and other debris were identified on the rooftops. Samples were collected from the outlet taps located close to the base of the water tanks, in sterilized 2
litre (L) bottle containers. Prior to collecting water samples, the tap was sterilized with 70% ethanol, and water was allowed to run for 30 to 60 s to flush out stagnant water from the tap (Ahmed et al. 2010). After collection, the samples were stored on ice in cooler boxes during transportation. All samples were brought back to Pretoria University Plant Pathology Laboratories, and processed within 24 hr.

**Enumeration and isolation of bacteria**

Microbes in water were concentrated by filtering 750 mL of water through 0.45-μm pore-sized (47 mm diameter) nitrocellulose membranes (Sartorius Stedim Biotech, Gottingen, Germany) by vacuum filtration (Shi et al. 2015), and membranes aseptically transferred into 9 mL of sterile tryptic soy broth (TSB) (Merck, Johannesburg, South Africa). The tubes were vortexed vigorously for 2 to 5 minutes (min) to detach the bacteria from the membranes. After serial dilutions, 100 microlitre (μL) aliquots were plated out in triplicates onto Reasoner’s 2A agar (R2A) (Merck) for heterotrophic plate counts. Membrane filters in 9 mL TSB were incubated at 37 °C for 24 hr with agitation (150 rpm). Afterwards, 0.1 mL of the pre-enriched broth was inoculated into 10 mL Rappaport-Vasidallis and Listeria enrichment broth for selective enrichment of *Salmonella* (Gomba et al. 2016) and *L. monocytogenes*, respectively. Rappaport-Vasidallis was incubated at 42 °C for 24 hr while *Listeria* enrichment broth was incubated at 37 °C for 18-24 hr. A loopful of each selective enrichment culture with growth was streaked onto xylose lysine deoxycholate agar (for *Salmonella* spp.) and Oxford-Listeria selective agar (for *L. monocytogenes*) plates.

All media were incubated at 37 °C for 24-48 hr except R2A plates (25 °C for 24-72 hr) and *Salmonella* enrichment broth. All media used were obtained from Merck. After incubation, colonies recovered using R2A were counted, recorded and expressed as colony-forming units per mL.

**Indicator system**

Densities of total coliforms, *E. coli*, *Pseudomonas* and *Enterococci* were determined with Colilert-18, Pseudolert-18 and Enterolert-18 chromogenic substrate tests kits and Quantitray 2000 trays (Idexx, Westbrook, Maine, USA), as per the manufacturer’s instructions. For highly contaminated samples, 1 mL was homogenized in 99 mL sterile water to satisfy the 100 mL requirement for quantification using Quantitray 2000 trays. The manufacturer’s most probable number (MPN) data table was used to generate microbial density estimates per 100 mL.

After incubation (37 °C for 18-24 hr), the backing material of each Quantitray was disinfected with 70% ethanol and a sterile razor blade used to pierce the backing material of three filters 750 mL of water to satisfy the 100 mL requirement for quantification. An aliquot of 1 ml was homogenized in 99 mL sterile water per the manufacturer’s instructions. For highly contaminated samples, 1 mL was homogenized in 99 mL sterile water to satisfy the 100 mL requirement for quantification using Quantitray 2000 trays. The manufacturer’s most probable number (MPN) data table was used to generate microbial density estimates per 100 mL.

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**Matrix-assisted laser desorption ionization identification**

Presumptive isolates were purified on NA prior to MALDI-TOF-MS analysis. The automated process from MALDI-TOF-MS measurement to species identification occurs without user intervention (Pinto et al. 2011; Chidamba & Korsten 2015b). Using default settings, the manufacturer’s MALDI Biotyper 5.0 software (Bruker Daltonics, Germany) analyzed spectra of the isolates and compared it with reference spectra in the database. A logarithmic score is generated corresponding to similarity of spectral patterns and interpreted as per manufacturer’s instructions. Results are conveyed as log (score) values ranging from 0 to three levels. Scores designate high confidence level species identification (≥2,300), species identification (≥2,000), genus identification (1,700-1,999) and no identification (<1,700).

**DNA extraction**

Genomic DNA (gDNA) was isolated directly from filter paper, with a ZM fungal/bacterial DNA miniprep™ kit (Zymo Research Corporation, Inqaba Biotech catalogue # D6005) as per the manufacturer’s specifications. The DNA concentration was determined with the Qubit 2.0...
Fluorometer (Lifescience Technology, Johannesburg, South Africa). Isolated gDNA was stored at −20°C and kept on ice during further experimental procedures.

**Pyrosequencing analysis**

DNA samples were sent to Inqaba Labs (Pretoria) for tag encoded pyrosequencing of a portion of the fungal internal transcribed spacer (ITS) region using the Roche GS FLX + 454 pyrosequencer (GATC Biotech, Konstanz, Germany). The fungal ITS region was amplified using primer set ITS1F (CTTGGTCATTTAGAGGAAGTAA) and ITS4R (TCCTCCGCTTATGGATATGC) (McHugh et al. 2014). Bacterial pyrosequencing is not presented in this paper but has been presented in another paper (Chidamba & Korsten 2015b). The matrix-assisted laser desorption/ionization time of flight (MALDI-TOF-MS) database restricted fungi identification, hence we relied on 454 fungal pyrosequencing to identify isolates.

**Pyrosequencing data processing and analysis**

Pyrosequencing data analysis for selected samples (Tanks 1–5) was performed using QIIME as established in previous studies (McHugh et al. 2014), with modifications. Selection of samples for analysis was based on population diversity. Sequences with less than 200 bp and quality score less than 25 were removed from the pyrosequencing-derived data sets using the UCLUST pyrosequencing pipeline initial processing (Cleary et al. 2011). The web-server SnoWMAN 1.15 (http://snowman.genome.tugraz.at) (Stocker et al. 2011) was used with the following settings for the taxonomic-based analysis: analysis type: UCLUST pipeline; chimera filtering reference database: Chimerarefdbs_09_Feb_2012_aligned; Mothur alignment reference database: fungalITS_24-Sep-2013-current_prokMSA_aligned; Mothur chimera checking reference database: chimerarefdbs_39-Feb-2012_16S_aligned and taxonomic reference database: greengenes_24-Mar-2010; rarefaction method: RDP; taxonomy: RDP; confidence threshold: 80%; include taxa covering more than 1%.

Operational taxonomic units for rarefaction analysis were clustered at 97% (species level), 95% (genus level) and 80% (phylum level) similarity cut-offs (Yang et al. 2014). The calculation of ACE (abundance-based coverage estimator), Evenness, Shannon, Chao1 and Species Richness diversity indices and the generation of principal coordinate analysis plots were generated using the open source software package QIIME (http://qiime.sourceforge.net), designed for combining heterogeneous experimental data sets, analysis of high throughput community sequencing data and obtaining new insights about various microbial communities (Caporaso et al. 2010).

**Statistical analysis**

Statistical analyses were conducted with SAS statistical software (version 9.3). Analysis of variance tests were performed and the least significance difference test used to derive statistical differences ($P > 0.05$) of microbial loads among all sampling areas.

**RESULTS**

**Microbial quality of RHRW**

Out of 246 samples, average total coliforms (57.7%), *E. coli* (50.5%), *Enterococcus* spp. (48.8%), *Pseudomonas* spp. (21.5%) and total aerobic bacteria (100%) were detected after rainfall. Prevalence of total coliforms, *Enterococcus* and *Pseudomonas* from all sites was not significantly different before and after rainfall. Before the rainy season, total coliforms were more prevalent in water from Ga-Molepane (43%), followed by Luthengele (35%) and Jericho (31%). Total coliforms were more prevalent in water samples from Ga-Molepane (73.7%) while the least prevalence was detected in samples from Luthengele after rainfall (Table 1). For season 1, concentrations of *E. coli* isolated from Ga-Molepane were significantly different from *E. coli* isolated from Jericho, but not significantly different from Luthengele *E. coli* isolates. Higher levels of *E. coli* were detected in water samples from Ga-Molepane (31.6%), followed by Luthengele (31.1%) and Jericho (29.2%) after rainfall. At least 45 and 51% of water samples from Luthengele represented *Enterococcus* spp. before and after rainfall, respectively, while 21.3% accounted for *Pseudomonas* spp. in RHRW from Jericho after rainfall. In general, water samples from Ga-Molepane had the highest concentrations...
of faecal coliforms, E. coli, Enterococcus spp. and Pseudomonas spp. (Table 1). Total aerobic bacteria ranged from a geometric mean of 5.26 (Ga-Molepane) to 5.91 (Jericho) log CFU/mL (Table 1). Means for aerobic plate counts from Ga-Molepane, Jericho and Luthengele were significantly different from one another, with samples from Jericho recording the highest and samples from Ga-Molepane recording the lowest.

Salmonella spp. and L. monocytogenes were present in RHRW from Ga-Molepane, Jericho and Luthengele villages. In total, 2 and 29 (before rainfall), 5 and 43 (after rainfall) RHRW samples tested positive for Salmonella spp. and L. monocytogenes, respectively (Table 2). Generally, the prevalence of L. monocytogenes and Salmonella spp. increased after rainfall at all sampling sites (Table 2). The prevalence of L. monocytogenes was 18.2 (before rainfall) and 31.1% (after rainfall) in RHRW samples from Jericho. Salmonella spp. was not detected before rainfall in water from Ga-Molepane and Jericho but was present after rainfall. The number of Salmonella positive samples increased (from 0 (before rainfall) to 2 (after rainfall)) in RHRW samples from Ga-Molepane.

Matrix-assisted laser desorption time of flight

Bacterial isolates from all sampling sites, as identified by MALDI-TOF-MS, were tabulated (Table 3). Of the total, opportunistic pathogens (Staphylococcus spp. (Niazi et al. 2010) (13%), Enterococci spp. (Shioya et al. 2011) (7%) and Pseudomonas spp. (de Bentzmann & Plésiat 2011) (3%)) and known potential human pathogens (L. monocytogenes (6%) and Salmonella spp. (4%)) were identified.

Of the total bacteria isolated from Ga-Molepane, E. coli prevalence was 15%, the highest compared to other sites. Various species belonging to Enterococci were identified. These included E. faecalis (only Ga-Molapane village) and E. faecium (all sites) (Table 3). Non-human pathogenic environmental bacterial isolates, among them, Acinetobacter spp. and Anaerococcus spp., were also detected (results not shown).

Fungal pyrosequencing

At phylum level classification, fungal sequences were classified into five phyla (Figure 1). These include Ascomycota,
Basidiomycota, Chytridiomycota, Glomeromycota and Zygomycota. Ascomycota dominated the data set (32.1%), while 45.7% of the sequences were not classified to any known fungal phyla. Of the identified sequences 17, 4.8 and 0.09% were classified to Basidiomycota, Chytridiomycota and Glomeromycota, respectively (Figure 1). There were notable differences in fungal diversity in the selected RHRW samples. Fungal sequences (95%) from Tank 1 were not identified while all sequences in Tank 4 were identified. Ascomycota was identified in four out of the five tanks (Tanks 2, 3, 4 and 5). The highest concentration of Chytridiomycota (70%) was identified in Tank 2. Only two fungal phyla (Basidiomycota and Zygomycota) were identified in Tank 1 RHRW (Figure 2). Tanks 2 and 3 had more diverse fungal populations with five identified sequences represented in both RHRW tanks.

Classification at species level revealed a diverse fungal population. Sordariomycetes species and Davidiella tassiana were more predominant in Tank 3 water samples (29.3% and 18.1%, respectively) than other water samples (Figure 2). Sequences of species in Tank 3 (13.9%) remained unclassified. Other species also identified included Cryptococcus spp., Dothideomycetes spp., Tremellales spp. and Knufia perforans Sterflinger.

**DISCUSSION**

In this study, microbial quality of RHRW was tested for the presence of bacterial indicators (total coliforms, *E. coli* and *Enterococci*), opportunistic (*Pseudomonas* spp.) and potential pathogenic microorganisms (*L. monocytogenes* and *Salmonella* spp.), and fungal organisms. To assess the suitability of RHRW for irrigation and domestic use conventional culture-based methods were used. There were significant variations of *E. coli* and aerobic plate

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**Table 2** | Prevalence of human pathogenic bacteria in RHRW

<table>
<thead>
<tr>
<th>Site</th>
<th>Salmonella Before rain</th>
<th>L. monocytogenes Before rain</th>
<th>Salmonella After rain</th>
<th>L. monocytogenes After rain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ga-Molepane</td>
<td>0 (0)</td>
<td>2 (10.5)</td>
<td>2 (10.5)</td>
<td>4 (21.1)</td>
</tr>
<tr>
<td>Jericho</td>
<td>0 (0)</td>
<td>8 (18.2)</td>
<td>1 (2.2)</td>
<td>14 (31.1)</td>
</tr>
<tr>
<td>Luthengele</td>
<td>2 (3.4)</td>
<td>19 (32.2)</td>
<td>2 (3.4)</td>
<td>25 (41.6)</td>
</tr>
</tbody>
</table>

**Table 3** | Frequency of selected significant dominant bacterial isolates from RHRW as identified by matrix-assisted laser desorption time of flight mass spectrometry

<table>
<thead>
<tr>
<th>Species name</th>
<th>Ga-Molepane</th>
<th>Jericho</th>
<th>Luthengele</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Aeromonas caviae</em></td>
<td>1</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td><em>Aeromonas jandaei</em></td>
<td>3</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td><em>Aeromonas veroni</em></td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td><em>Bacillus cereus</em></td>
<td>5</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td><em>Bacillus pumilus</em></td>
<td>2</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td><em>Enterobacter amnigenus</em></td>
<td>2</td>
<td>2</td>
<td>7</td>
</tr>
<tr>
<td><em>Enterobacter asburiae</em></td>
<td>5</td>
<td>7</td>
<td>1</td>
</tr>
<tr>
<td><em>Enterobacter cancerogenus</em></td>
<td>1</td>
<td>0</td>
<td>3</td>
</tr>
<tr>
<td><em>Enterococcus faecalis</em></td>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td><em>Enterococcus faecium</em></td>
<td>5</td>
<td>15</td>
<td>9</td>
</tr>
<tr>
<td><em>Enterococcus durans</em></td>
<td>9</td>
<td>7</td>
<td>12</td>
</tr>
<tr>
<td><em>Enterococcus mundii</em></td>
<td>6</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td><em>Escherichia vulneris</em></td>
<td>7</td>
<td>1</td>
<td>11</td>
</tr>
<tr>
<td><em>E. coli</em></td>
<td>15</td>
<td>11</td>
<td>9</td>
</tr>
<tr>
<td><em>Klebsiella pneumonia</em></td>
<td>4</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td><em>Listeria monocytogenes</em></td>
<td>9</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td><em>Morganellamorganii</em></td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td><em>Pantoa agglomerans</em></td>
<td>6</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><em>Pseudomonas otidis</em></td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td><em>Pseudomonas taetrolens</em></td>
<td>1</td>
<td>6</td>
<td>5</td>
</tr>
<tr>
<td><em>Salmonella</em> spp.</td>
<td>3</td>
<td>4</td>
<td>6</td>
</tr>
<tr>
<td><em>Staphylococcus auricularis</em></td>
<td>3</td>
<td>12</td>
<td>4</td>
</tr>
<tr>
<td><em>Staphylococcus capitis</em></td>
<td>1</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>

*Values are in percentages calculated from the total isolates identified.
counts between the three rural villages sampled. A previous study reported similar variations in the microbiological quality of RHRW (Evans et al. 2006a). The study further reported extensive disparities in the level of water pollution and bacterial composition between sites, and over time. Results of our study indicate the presence of potential human pathogenic bacteria in RHRW from all the sites tested.

Salmonella spp. (3%) and L. monocytogenes (22%) were detected in RHRW samples from all three sampling areas. Ahmed et al. (2015) detected Salmonella spp. but not L. monocytogenes in RHRW. A previous study indicated that drinking RHRW containing Salmonella spp. caused diarrhoea and abdominal pains (De Kwaadsteniet et al. 2013). Based on observations during this study, RHRW was being contaminated with rooftop faecal material and dust. The source of this contamination was not specifically studied in this paper but it has been observed and previously described in other studies (Evans et al. 2006b; Ahmed et al. 2012).

The rainfall patterns of Jericho and Ga-Molepane are similar and the rainfall events occur seasonally. Hence, E. coli concentration in RHRW is significantly higher immediately after rainfall events. Luthengele area, on the other hand, experiences rain events more frequently. As a result, there is little time for accumulation of faecal material and other debris on rooftops. This could partially explain E. coli concentrations not being significantly different before and after rain events. The low coliforms and E. coli counts detected in RHRW before the start of the rainy season may suggest that the rooftops are more consistently washed down by pouring rainwater and are therefore not consistently highly contaminated. However, the presence of Enterococci spp., L. monocytogenes and Salmonella spp. suggests otherwise. The mere presence of L. monocytogenes in water does not mean infection is possible upon consumption. However, the pathogen can proliferate in storage tanks leading to high quantities suitable for the onset of listeriosis (Morobe et al. 2009; Golberg 2015).

In a recent study, low counts of E. coli were observed but detection of faecal indicator bacteria (Enterococci spp.) suggested the water may be contaminated (Chidamba & Korsten 2015a). Although the concentration of E. coli most likely decreased with time, Enterococci and other pathogens persisted owing to their resilience (Ahmed et al. 2012). This study therefore finds RHRW a potential hazard if used for domestic purposes or to irrigate crops due to the presence of well-known human pathogens. However, it is worth noting that using polluted irrigation water does not always result in contaminated crops.

Basic systems of roof-water harvesting are used in the sampling sites visited. The system is deprived of first flush diverters and filtration apparatus (Chidamba & Korsten 2015a). Due to this deprivation, microbiological quality of the RHRW is compromised and treatment prior to use is
thus necessary. The absence of these first flush diverters contributes to increased chances of contamination of RHRW, hence faecal material and other debris on the roof and/or gutter washes into rainwater tanks. Chidamba & Korsten (2018a) established a link between indicator bacteria and pathogens isolated from rooftop surfaces and RHRW. The presence of such pathogens could be transmitted to leafy green vegetables when RHRW is used for irrigation purposes. Several studies have linked irrigation water with pathogens isolated from fresh produce (Allende & Monghan 2015; Gelting et al. 2015), therefore, substandard irrigation water is a contamination risk factor of fresh green leafy vegetables. The presence of human pathogens on irrigated crops due to the use of contaminated irrigation water is well described (Pachepsky et al. 2011).

In this study, heterotrophs were isolated in RHRW at all sampling sites in concentrations higher than recommended for domestic and irrigation purposes according to the Department of Water Affairs Standard, South Africa (DWAF 1996). Concentrations higher than 1,000 CFU/mL in water used for drinking indicate contamination or aftergrowth in the water and pose an increased risk of infectious disease transmission if consumed (DWAF 1996). Even though heterotrophic bacterial counts indicate the general microbial quality of water, they do not indicate possible faecal pollution (DWAF 1996). However, DWAF (1996) recommends that water with high concentrations of heterotrophs requires treatment prior to use. The prevalence of Enterococci was higher than E. coli in this study. This is consistent with other reports (Spinks et al. 2006; Ahmed et al. 2011). These results suggest that RHRW from these sampling sites is not fit-for-irrigation of fresh products consumed raw unless treated prior to use according to the South African irrigation water standards (DWAF 1996).

The reliance of total coliforms and indicator bacteria for water quality assessment has come under scrutiny in the recent past (Chidamba & Korsten 2018a), as findings emerged that pathogenic E. coli could not be isolated from water samples positive for indicator bacteria of faecal origin. However, the use of total coliforms and indicator bacteria to evaluate water quality is by far the most widely used method in laboratories. The simultaneous use of E. coli and Enterococci for potential faecal contamination assessment is recommended (Savichtcheva & Okabe 2007; Ahmed et al. 2011; Chidamba & Korsten 2015a), and our study also supports this approach.

Although the presence of human bacterial pathogens has been widely reported in RHRW (Ahmed et al. 2011), the same cannot be said about human fungal pathogens. Gikas & Tsihrintzis (2012) mentioned that water harvesting systems could transmit microorganisms able to affect human health. Conventional culture techniques
underestimate microbial populations (Ahmed et al. 2011), hence utilization of molecular methods is increasingly used to provide a better understanding of spp. present. Pyrosequencing of the ITS 1 and ITS 2 hypervariable regions of the 18S rRNA was used to investigate the fungal communities in RHRW. Results of this study indicated a diverse fungal community in RHRW.

Pencillium and Cryptococcus spp. were mostly detected in RHRW. Some Penicillium spp. are pathogenic to plants and others can affect human health, such as P. digitatum (Pers.) Sacc. (Hageskal et al. 2009; Oshikata et al. 2013). Further, mycotoxins and other metabolites that are produced by these organisms have been detected in storage water tanks (Paterson et al. 1997). Prolonged water storage may therefore lead to increased mycotoxin concentration, and could pose a potential risk to consumers with persistent consumption (Hageskal et al. 2009). In such a case, immunocompromised individuals are at a higher risk due to their diminished immune system. HIV prevalence in rural communities in South Africa has been reported (Shisana et al. 2014) and is considered high (39.4%) by world standards.

Human pathogenic yeasts, Cryptococcus spp., belonging to a diverse filamentous fungal division Basidiomycota, were also detected in RHRW. Cryptococcus spp. is an opportunistic human pathogen of global importance (Loftus et al. 2005), which has been isolated in bird faeces (Soltani et al. 2013). Effects of continuous exposure to Cryptococcus spp. include fever, coughing and shortness of breath.

Monchy et al. (2011) highlighted the significance of further characterization of human pathogenic microbes and understanding their effects on human health due to persistent exposure. The risk posed by the presence of sequences belonging to phylotypes with a known pathogenic history cannot be underestimated. The presence of unidentified sequences signify their absence in cluster analysis databases, hence the need for research to identify more environmental fungal populations and collection of curated sequences for fungal identification (Liu et al. 2015). The use of RHRW with human pathogenic isolates for domestic use and irrigation purposes without any prior treatment could therefore be a cause of concern if pathogens persist up to the point of consumption. Future studies should focus on the link between fungal spp. in RHRW and crop disease or decay.

**CONCLUSION**

This study concluded that RHRW collected in three rural villages was of poor microbiological quality, therefore not fit for domestic use or irrigation purposes unless treated prior to use as per national guidelines (DWAF 1996). The presence of known human pathogenic microorganisms and faecal indicators in RHRW indicate a potential risk if the crops are also irrigated with this water and the product is consumed raw. Better maintenance of RHRW systems are required to reduce contamination levels. Development of multiple detection systems using faecal indicators, E. coli, Enterococci and next generation sequencing techniques could holistically improve water quality assessment.

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