Physico-chemical parameters and culturable yeast diversity in surface water: a consequence of pollution

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Rivers in the North West Province (NWP) of South Africa are polluted by various anthropogenic activities that are associated with agricultural, domestic, industrial, and mining activities. Pollutants and effluent resulting from these activities impact the physico-chemical and biological characteristics of river water. More particularly, domestic and agricultural pollution has been associated with the occurrence of pathogenic yeast species in water. The aim of this study was to determine physico-chemical parameters, yeast levels, and the antifungal susceptibility of pathogenic yeasts in river water. Physico-chemical parameters and yeast levels were determined using standard procedures. Yeasts were identified by biochemical tests and 26S rRNA gene sequencing. Disc diffusion antifungal susceptibility tests were conducted on identified potential pathogenic yeasts. Physico-chemical parameters were within target water quality ranges (TWQR) for livestock farming but were mostly out of range for irrigation. Yeast levels (incubation at 37°C) ranged from 363 to 1 778 CFU/L. There were significant differences (p < 0.05) in the physico-chemical parameters and yeast levels between some seasons and among the river systems under study. A positive association was observed between temperature, COD and yeast levels in all the river systems. Ascomycetes, which were the most prevalent isolates, were identified as Candida spp. (35%), Pichia spp. (13%), Cyberlinera spp. (12%), Meyerozyma spp. (11%), Clavispora spp. (10%), Saccharomyces spp. (6%), Kluyveromyces spp. (5%), Yamadazyma spp. (4%), Trichosporon spp. (3%), and Wickerhamomyces spp. (1%). Antifungal resistance of the potential pathogenic yeasts was as follows: flucytosine and miconazole (100%) > fluconazole (78.5%) > econazole, and miconazole and ketoconazole (49.6%) > nystatin (15.2%). The river water systems explored in the study are used for agricultural, domestic and recreational purposes. Direct contact with the water, especially by immune-compromised people, may pose a health threat and should be further investigated.

INTRODUCTION

Organic and inorganic pollutants, heavy metals and biological materials from industries, mines, wastewater treatment plants (WWTPs) and agricultural settings are known to be indiscriminately discharged into river systems (Gupta et al., 2017). These surface and subsurface water resources are used for drinking water supply, domestic use, agricultural irrigation and aquatic life. Pollutants affect the physico-chemical parameters and microbial density present in water systems (Traoré et al., 2016) and thus water use prospects are affected. The Sustainable Development Goals (SDG) adopted by all United Nations member states in 2015 stipulate that all people should have access to clean water, sanitation, good health, and wellbeing by 2030 (UNDP, 2015). However, water pollution derived from human activities threatens the achievement of the SDG targets.

Current microbial studies use standard faecal indicator bacteria such as *Escherichia coli* and intestinal enterococci to determine water pollution levels (Kirschner et al., 2019). However, some studies have also reported high yeast levels and the presence of certain types of yeasts as an indication of contaminated water environments (Van Wyk et al., 2012; Monapathi et al., 2017). Some studies have also deemed yeasts as a complement to faecal indicator bacteria counts when water quality is monitored (Medeiros et al., 2008; Brandão et al., 2010). A rapid response of yeasts to organic contamination makes some yeasts important indicators of nutrient enrichment because these convert easily accessible carbon sources into energy for reproduction (Brandão et al., 2010).

Pathogenic yeast species have been isolated from freshwater environments (Monapathi et al., 2020). They can cause various diseases – from superficial mucosal infections to life-threatening systemic disorders, especially in immunocompromised people (Ravikumar et al., 2015). These patients are generally associated with cancer, organ transplants, and HIV (Shiels et al., 2017). The situation is exacerbated as pathogenic yeast species have developed resistance to several of the clinically used antifungal agents (Ksiezopolska and Gabaldón, 2018).

Sub-therapeutic levels of antifungal agents are constantly leaked into the environment by means of wastewater treatment plant (WWTP) discharges and agricultural runoff (Tran et al., 2017). The release of such effluent into aquatic ecosystems escalates antifungal resistance through continuous exposure of pathogenic yeasts. Antifungal resistance has been observed among environmental isolates (Brilhante et al., 2016; Milanezi et al., 2019; Monapathi et al., 2017, 2018). According to a report by the UN ad hoc Interagency Coordinating Group on Antimicrobial Resistance,

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Figure 1. Map showing the selected river systems in the North West Province of South Africa (Mo = Mooi River; Ma = Marico River; Sc = Schoonspriut River; Kr = Crocodile River)

drug-resistant diseases could cause up to 10 million deaths each year by 2050. Furthermore, by 2030, antimicrobial resistance could force up to 24 million people into extreme poverty (WHO, 2016). Antifungal drug resistance by pathogenic yeasts is a pressing public health crisis. The present study thus aimed to determine physico-chemical parameters and yeast levels as well as the antifungal susceptibility of pathogenic yeasts in four selected river systems in the North West Province (NWP) of South Africa.

METHODS

Study area

Water samples were collected from 4 river systems: the Mooi River, the Schoonspruit River, the Crocodile (West) River, and the Marico River, South Africa (Fig. 1). A Garmin Nüvi 1310 (Garmin, US) global positioning system (GPS waypoints) was used to locate the water sampling sites. Water quality of these river systems is affected by surrounding informal settlements, urbanization, agricultural runoff, and industrial and mining activities (DWAF, 2004, 2007, 2009; Winde, 2010; Barnard et al., 2013; Venter et al., 2013).

Sampling

Sampling was conducted in 2015 and 2016 during the wetwarm (August to March) and dry-cold seasons (May to July) in the NWP. An aquatic microbiology research group at the North West University assisted with sampling. Water samples (1 L) were collected using sterile Schott bottles. The direct sampling technique was used close to the riverbank. Dip sampling was done using a rope where direct access to the surface water systems was limited (Van Wyk et al., 2012).

Physico-chemical parameter analyses

Physico-chemical parameter analyses were done in triplicate. Temperature, pH, dissolved oxygen (DO) and total dissolved solids (TDS) were determined in situ using a 350 multi parameter probe (Merck, Germany). Water samples were transported on ice to the laboratory and analysed within 8 h for nitrates (Method 10206), phosphates (Method 8048) and chemical oxygen demand (COD) (Method 8000) using the Hach Lange DR 2800 system and reagents (Hach Company, 2007). Raw data for physico-chemical parameters have been reported by Bezuidenhout et al. (2018).

Yeast isolation and enumeration

Membrane filtration was done to determine the presence of yeasts in the water samples as described by Van Wyk et al. (2012). Water samples (100 mL) were filtered in triplicate through 0.45 μ m HA membrane filters (Whatman). Membranes were placed on yeast-malt-extract (YM) plates, supplemented with 100 ppm chloramphenicol, and incubated at room temperature (incubation in the laboratory under uncontrolled conditions) and at 37°C for 24 h. Colony counts were done on individual plates. Representatives of morphotypes that grew at 37°C with respect to cellular shapes, colonial elevation, colonial surface, and colonial pigmentation were selected and purified by sub-culturing on YM plates (Wickerham, 1951) and subjected to further analysis.

Biochemical characterisation

Diazonium Blue B (DBB) test

Yeast cultures were grown on YM agar plates and incubated at 37°C for 14–21 days (Kurtzman and Fell, 1998). Petri dishes were

flooded with ice-cold DBB reagent (0.1MTris-HCl; Diazonium Blue B Salt 1 mg/mL) (Prillinger et al., 1999). A dark red to violet red colour occurring within 2 min at room temperature indicated a positive reaction for basidiomycetes (Kurtzman and Fell, 1998).

Molecular identification

DNA extraction

Two millilitres overnight YM broth cultures of the yeast isolates were prepared and centrifuged to obtain a pellet of the cells. This was followed by genomic DNA extraction according to the modified method of Hoffman and Winston (1987).

Amplification and sequence validation

The 26S rRNA gene fragments of the extracted DNA were amplified using the polymerase chain reaction (PCR) procedure as described by Monapathi et al. (2017). Forward primers, NL1 (5-GCATATCAATAAGCGGAGGAAAAG-3) and reverse primers, NL4 (5-GGTCCGTGTTTCAAGACGG-3) (O'Donnell, 1993) were used. PCR products were sequenced using the BigDye Terminator version 3.1 Cycle Sequencing kit (Applied Biosystems, UK) as prescribed by the manufacturer.

Clean-up of the sequencing reaction was achieved using the NaOAc/ EDTA/ethanol method (Applied Biosystems, UK) and sequenced using an ABI 3130 Genetic Analyser (Applied Biosystems, Hitachi) (Jordaan and Bezuidenhout, 2013). BLAST (Altschul et al., 1997) searches (http://www.ncbi.nlm.nih.gov/BLAST) were used to determine the identity of the amplified sequences. Sequence reads were submitted to a NCBI GenBank database.

Antifungal susceptibility tests

Antifungal susceptibility testing using the Kirby-Bauer disc diffusion method (Bauer et al., 1966) was done using YM agar. Diameters of zones of inhibition (in mm) of yeast species against commonly used antifungal agents: fluconazole (FCN; 25 μ g),

econazole (ECN: 1 µg), ketoconazole (KCA: 15 µg), miconazole (MCL: 1 µg), metronidazole (MZ: 5 µg), flucytosine (FY: 1 µg), and nystatin (NY: 100) were measured after a 24-h incubation period at 37°C. These were compared to the susceptibility breakpoints for fluconazole and *Candida* spp. (Pfaller et al., 2006). Zone breakpoints and interpretative categories for antifungal agents were classified as resistant (\geq 19 mm), susceptible-dose dependent (15 to 18 mm), or susceptible (\leq 14).

Data and statistical analyses

Microsoft Excel (2016) was used to analyse data. Descriptive and one-way ANOVA followed by Tukey's HSD (Honestly Significant Difference) Test for unequal sample sizes were used for post-hoc comparisons of physico-chemical and microbiological parameters where p < 0.05 was deemed significant. Significant differences were denoted by different alphabetical letters (a, b, c, d, e, etc.) between variables. Redundancy analysis (RDA) was used to test for association between physico-chemical parameters and yeast levels. Data output was provided as correlation biplots. The correlation significance level was set at p < 0.05.

RESULTS AND DISCUSSION

Physico-chemical parameters and yeast levels

The mean values of the physico-chemical parameters were compared to water quality standards defined in Total Water Quality Ranges (TWQR) for agricultural applications (irrigation and livestock farming) (DWAF, 1996a, 1996b). Temperatures for the wet-warm seasons were significantly higher (p < 0.05) than for the dry-cold seasons (Fig. 2 I). During the wet rainy season, organic pollutants from domestic wastewater, effluent from wastewater treatment plants, and agricultural runoff are leached into environmental water which results in increased water temperature (Susilowati et al., 2018). The high temperatures observed in the present study resulted from discharges and run-off from agricultural settings, informal settlements, and urbanization activities in the vicinity of the river systems.



Figure 2. Mean±SD values for physico-chemical and microbiological parameters in selected surface water resources in the NWP; (I) temperature (°C), (II) yeast levels at 37°C (CFU/L), (III) chemical oxygen demand (COD) (mg/L), (IV) dissolved oxygen (DO) (mg/L)



Figure 3. Redundancy analysis (RDA) ordination biplots illustrating correlation between environmental variables and yeast levels (YL) in the river systems under study: (I) Mooi River (II) Schoonspruit River (III) Crocodile (West) River and (IV) Marico River. Temp = temperature; TDS = total dissolved solids; COD = chemical oxygen demand: DO = dissolved oxygen; NO3- = nitrates (NO₃-); PO42- = phosphates (PO₄²⁻)

Yeasts require carbon sources for energy and growth (Tomova et al., 2019); thus a polluted organic state of the river systems will favour the growth of yeasts. Yeast levels enumerated at 37°C were between 363 and 1 778 CFU/L (Fig. 2 II). High yeast levels were recorded for all sampling sites during the wet-warm seasons. At room temperature (RT), the yeast levels followed a similar trend and a positive association is depicted in Fig. 3 (I, II, III, IV). At room temperature, the yeast levels were substantially higher (data not shown). An increase in temperature results in increased concentration of dissolved substances and a decreased concentration of dissolved substances and a decreased concentration of dissolved oxygen (Rajwa-Kuligiewicz et al., 2015; Susilowati et al., 2018). Maximum COD concentrations for wet-warm and dry-cold seasons were 72.5 mg/L and 42.3 mg/L, respectively (Fig. 2 III).

From RDA, a positive association was observed between yeast levels, temperature, and COD in all the river systems (Fig. 3 I, II, III, IV). A negative correlation was observed with DO. This conformed to lower DO levels (\leq 7.8 mg/L) during the wetwarm season (Fig. 2 IV).

The pH values for the wet-warm and dry-cold seasons varied between 7 and 8 but were within the acceptable TWQR for irrigational use (6.5–8.4). The pH values were slightly alkaline. Organic matter from anthropogenic pollutants decreases the pH in water (Schneider and Le Campion-Alsumald, 1999). Maximum TDS values recorded were 720 mg/L and 711 mg/L for the wetwarm and dry-cold seasons (Fig. 2 III). Significantly higher TDS values were observed for the Schoonspruit River samples compared to the other rivers. RDA showed a positive association between TDS, yeast levels, temperature, and COD for the Marico and Schoonspruit River samples (Fig. 3 II, IV). TDS values were within TWQR for livestock farming (<1 000 mg/L) but above the recommended values for irrigation use (>40 mg/L). The high TDS values that were found in the NWP rivers by the current and by previous studies (Van Wyk et al., 2012; Monapathi et al., 2017) could have occurred as a result of dolomitic springs which are water sources for some of the river systems (Van Der Walt, 2002).

Nitrate levels for the wet-warm and the dry-cold seasons were $\leq 1.8 \text{ mg/L}$ and $\leq 1.51 \text{ mg/L}$, respectively, while the phosphate concentrations for the wet-warm and dry-cold seasons were recorded as $\leq 5.4 \text{ mg/L}$ and ≤ 1.8 , respectively. No significant seasonal differences were observed per river system and among the river systems for the latter nutrient. Moreover, no clear association could be established between the nutrients and yeast levels, except for phosphates in the Marico and Schoonspruit Rivers.

The river systems explored in the present study are used for irrigation purposes. However, the mean nitrate levels were above the permissible TWQR levels (<0.5 mg/L) for irrigation use. Dryland farming practices in the NWP use fertilisers and animal manure (Jordaan and Bezuidenhout, 2016). The nutrients observed in the river systems could have resulted from surrounding agricultural activities.

Biochemical and molecular identification of yeasts

DBB staining classified yeasts as ascomycetous yeasts. In the four rivers, a total of 256 isolates were obtained and identified (Mooi River = 99; Schoonspruit River = 84; Marico River = 43; Crocodile (West) River = 30. BLAST searches determined the identity of the amplified 26S rRNA gene sequences. The following genera were identified: predominant *Candida* spp. (35%), *Pichia* spp. (13%), *Cyberlinera* spp. (12%), *Meyerozyma* spp. (11%), *Clavispora* spp. (10%), *Saccharomyces* spp. (6%), *Kluyveromyces* spp. (5%), *Trichosporon* spp. (4%), *Yamadazyma* spp. (3%), and *Wickerhamomyces* spp. (1%).

Figures 4 I and II show a neighbour-joining tree demonstrating a phylogenetic analysis between yeast species. Medium to

high bootstrap confidence of 66 to 100% supported the relationship between the environmental isolates (bolded) and the representative 26S rRNA gene sequences obtained from GenBank database. A phylogenetic tree constructed from sequences of *Candida* species (Fig. 4 I) showed the formation of distinct clusters for each species. Each cluster indicated high sequence similarity between the groups (Ragonnet-Cronin et al., 2013). Clustering of similar species was also observed in non-*Candida* species (Fig. 4 II). Clusters H and K were outgroups for *Candida* and non-*Candida* groups, respectively. Pathogenic species did not form a monophyletic group and were scattered all over the trees. This indicated that pathogenicity amongst these species evolved independently on multiple occasions (Diezmann et al., 2004).



Figure 4. A neighbour-joining tree showing the phylogenetic relationship between environmental *Candida* species (I) and non-*Candida* species (II) (bold) and representative species from GenBank database. A bootstrap test (1 000 replicates) was conducted and the cluster percentages of trees supporting the cluster are provided.



Figure 4 Continued. A neighbour-joining tree showing the phylogenetic relationship between environmental *Candida* species (I) and non-*Candida* species (II) (bold) and representative species from GenBank database. A bootstrap test (1 000 replicates) was conducted and the cluster percentages of trees supporting the cluster are provided.

Table 1. Isolated yeast species' resistance (%) to commonly used antifungal agents. (FCN = fluconazole; ECN = ezonazole; KCA = ketoconazole MCL = miconazole MZ = metronidazole; FY = fluctyosine; NY = nystatin)

Yeast species	Antifungal agents (% resistance)						
	FCN	ECN	KCA	MCL	MZ	FY	NY
Candida albicans, C. glabrata, C. haemulonis, C. orthopsilosis, C. sake, C. tropicalis, Clavispora lusitaniae, Trichosporon ovoides	100.0	100.0	100.0	100.0	100.0	100.0	0.0
Candida bracarensis, Pichia kudriavzevii, Saccharomyces cerevisiae	100.0	0.0	0.0	0.0	100.0	100.0	0.0
Candida parapsilosis	0.0	0.0	0.0	0.0	100.0	100.0	100.0
Cyberlinera fabianii, Cyberlindnera jadinii, Kluyveromyces marxianus, Wickerhamomyces anomalus, Yamadazyma mexicana	0.0	0.0	0.0	0.0	100.0	100.0	0.0
Meyerozyma caribbica, M. guilliermondii	100.0	0.0	0.0	0.0	100.0	100.0	100.0

Table 2. Number of yeast isolates (n (%)) resistant to various commonly used antifungals

Sampling site	Antifungal agents									
_	FCN	ECN	KCA	MCL	MZ	FY	NY			
Mooi River (<i>n</i> = 99)	78(78.8)	58(58.6)	58(58.6)	58(58.6)	99(100)	99(100)	6(6.1)			
Schoonspruit River ($n = 84$)	59(70.2)	35(41.7)	35(41.7)	35(41.7)	84(100)	84(100)	11(13.1)			
Marico River ($n = 43$)	38(88)	21(48.8)	21(48.8)	21(48.8)	43(100)	43(100)	9(20.9)			
Crocodile River ($n = 30$)	26(86.7)	13(43.3)	13(43.3)	13(43.3)	30(100)	30(100)	8(12.5)			
Total (<i>n</i> = 256)	201(78.5)	127(49.6)	127(49.6)	127(49.6)	256(100)	256(100)	39(15.2)			

Pathogenic yeasts

Yeast species that grow at 37°C are opportunistic pathogens (Maciel et al., 2019). The predominantly isolated species in the current study, *Candida* spp., are widely recognized as major causes of morbidity and mortality in the healthcare environment. Some pathogenic *Candida* species are *C. albicans*, *C. glabrata*, *C. tropicalis*, *C. parapsilosis*, and *C. krusei* (Zupančič et al., 2018, Friedman and Schwartz, 2019). Opportunistic non-*Candida* yeasts spp. isolated in the present study were *Clavispora lusitaniae*, *Cyberlinera fabianii*, *Meyerozyma caribbica*, *M. guilliermondii*, *S. cerevisiae*, *T. ovoides*, and *W. anomalus*. These organisms have been implicated in causing cutaneous and/or systemic bloodstream infections such as vaginitis in healthy patients, endophthalmitis, candidemia, trichosporonosis, and catheter-related, neonatal and ocular infections in immune-compromised people (Gabaldón et al., 2016; Maciel et al., 2019; Consortium Opathy and Gabaldón, 2019).

Antifungal susceptibility pattern

Yeast species showed antifungal resistance to various commonly used antifungals. Table 1 illustrates that pathogenic yeasts *Candida albicans, C. glabrata, C. haemulonis, C. orthopsilosis, C. sake, C. tropicalis, Clavispora lusitaniae* and *Trichosporon ovoides* showed 100% resistance to azoles. Opportunistic non-*Candida* yeasts were also resistant to azoles. In the entire study, all the isolated yeast species were resistant to metronidazole and flucytosine (Table 2), followed by fluconazole (78.5%), econazole, miconazole and ketoconazole (49.6%) and nystatin (15.2%). Predominant resistance to fluconazole was observed in all the river systems. The high resistance to azoles that was observed is worrisome. Due to their broad spectrum of activity and safety profiles, azoles are the most commonly used antifungal drugs (Li et al., 2014).

Several studies have reported on environmental yeast resistance to antifungal drugs. A study conducted by Brilhante et al. (2016) at Catú Lake, Ceará, in Brazil, detected 13 azole-resistant *Candida* isolates. Antifungal resistance was also observed to amphotericin B, itraconazole, and fluconazole. Milanezi et al. (2019) conducted a study in the Dilúvio Stream in Porto Alegre, Rio Grande do Sul, Brazil. The isolated *Candida* spp., *Debaryomyces* spp. and *Rhodotorula* spp. were resistant to itraconazole (57.9%), fluconazole (44.2%), voriconazole (24.2%), amphotericin B 16 (16.8%), and ketoconazole (5.3%).

Monapathi et al. (2017) analysed antifungal susceptibility in the Mooi River and the Harts River in South Africa. All tested yeast isolates (100%) were resistant to miconazole and flucytosine. These were followed by fluconazole (88.5 and 92.5%) in the Mooi and Harts Rivers, respectively). In the Mooi River, 62.8% of the isolates were resistant to econazole and miconazole and 64.1% were resistant to ketoconazole. In the Harts River, 12.5% of the isolates were resistant to the three azole antifungal drugs. These results are similar to the findings of the present study. Azole resistance was common amongst the isolates from the four river systems and thus aquatic environments hosting these pathogenic antifungal resistant yeasts might pose a public health threat. With the high HIV infection rate in South Africa (7.2 million) (UNAIDS, 2018), resistance of pathogenic yeasts to antifungal agents places people at risk who use surface water, such as the communities in the vicinity and visitors to the river areas for recreation or other purposes.

CONCLUSIONS

Physico-chemical parameters and yeast levels were determined in four rivers system in the NWP. The river systems are used for agricultural, domestic, industrial and mining purposes. Physico-chemical parameters were within TWQR for livestock farming but were out of range for irrigation. Higher yeast levels observed during the wet-warm seasons correlated with elevated temperatures and COD levels. Further studies are needed to increase the datasets for yeasts used in conjunction with physicochemical parameters. Isolated yeasts were ascomycetes and were dominated by *Candida* spp., many of which are pathogens/ opportunistic pathogens. Of concern is that these *Candida* spp. that were prevalent are resistant to antifungal agents commonly used in human and animal therapeutic settings. The observed resistance to azole, especially fluconazole, is also a concern. There is high HIV prevalence in South Africa and fluconazole is part of antiretroviral therapy to prophylactically manage potential *Candida* spp. infections. The results demonstrate the attractiveness of using yeasts as a microbiological indicator of organic pollution of aquatic ecosystems. The results also demonstrate that veterinary and agricultural use of antifungal drugs should be regulated and minimised to prevent antifungal resistance of potential pathogenic yeasts in aquatic systems. Further studies on antifungal susceptibility and mechanisms should be conducted to find solutions for resistance and to validate the development of new antifungal drugs.

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