Heat shock protein 47 stress responses in Chinese Hamster Ovary Cells exposed to raw and reclaimed wastewater

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As wastewater reclamation and reuse becomes more widespread, risks of exposure to treated wastewater increase. Moreover, an unlimited number of pollutants can be identified in wastewater. Therefore, comprehensive toxicity assessment of treated wastewater is imperative. The objective of this study was to perform a comprehensive toxicity assessment of wastewater treatment systems using stress response bioassays. This powerful tool can comprehensively assess the toxicity of contaminants. In this study, samples from conventional activated sludge treatment, membrane bioreactors (MBRs) with different pore sizes and sludge retention times (SRTs), rapid sand filtration, coagulation, nano-filtration (NF) and reverse osmosis (RO) were investigated. The results of stress response bioassays confirmed that the secondary effluent showed higher stress response than influent indicating that biological treatment generates toxic compounds. The results obtained from molecular weight fractionation of water samples demonstrated that organic matter with a higher molecular weight fraction (>0.1μm) causes toxicity in secondary effluent. Furthermore, supernatant from MBR reactors showed toxicity regardless of SRT. On the other hand, stress response was not detected in MBR permeates except for an MBR equipped with a larger pore size membrane (0.4μm) and with a short SRT (12 days). While rapid sand filtration could not remove the toxic compounds found in secondary effluent, coagulation tests, operated at an appropriate pH, were effective for reducing stress response in the secondary effluent. Experimental findings also showed that stress response was not detected in cases of NF and RO permeate subsequent to MBR treatment.

Introduction

Wastewater reclamation and reuse is gaining widespread acceptance because of increasing water consumption and the scarcity of freshwater resources. Wastewater reclamation and reuse play an important role as an environmentally friendly and cost efficient process in the management of water resources. As early as 1970, only about 216 million m³ of wastewater was reused in California. However, this increased to 648 million m³ in 2001 [1]. In Europe in the early 1990s municipal water reuse was limited to a few cases, but now more than 200 water reuse projects can be identified [2]. With the popularization of wastewater reclamation and reuse, the risks associated with exposure to treated water increase. In Japan for example, reclaimed wastewater standards cover only a portion of the total water quality index such as E. coli bacteria, turbidity, chromaticity and odour [3]. Therefore a more comprehensive toxicity assessment of reclaimed wastewater is crucial. Since many harmful compounds are contained in wastewater, several methods can be used to assess wastewater toxicity. Bioassay is a powerful and effective tool that can assess the toxicity of harmful matter comprehensively as compared to instrumental analysis [4]. Several researchers have used bioassays to assess water toxicity. Ono et al. [5] evaluated the toxicity of concentrated effluent from a wastewater treatment plant using the Umu test. Farré et al. [6] applied Toxalert and Microtox bioluminescence inhibition assays to assess the toxicity of influent and effluent from several wastewater treatment plants. Jimmiano et al. [7] assessed toxicity of raw wastewater and secondary effluent including several advanced treatment systems such as membrane filtration by bioassay using Oryzias latipes. Kontana et al. [8] evaluated ecotoxicological characteristics of reclaimed wastewater using Vibrio fischeri, Daphnia magna and Tetrahymena thermophila assays. In addition to these methods, it is also reported that bioassay methods using cultured human cell and mammalian cell lines are effective to assess toxicity on human systems [9], [10] and [11]. As potable reuse is the trend of today’s water management, the aforementioned methods are recommended.

In this study, we applied stress response bioassays using Chinese hamster ovary (CHO) cells stably transfected with Heat Shock Protein (HSP) 47 promoter. HSPs are members of a distinctive class of molecules that protect cells against a wide range of injuries. They assist in cellular recovery from stress either by repairing damaged proteins or by removing them to restore protein homeostasis [12]. In this respect, it is known that the stress response system, in particular HSP, functions in all mammalian tissues and cells. Therefore, in bioassay systems utilizing this stress response, it is not necessary to take into account
consideration the basic problems regarding cell specificity [13].
HSP47 molecules are 47-kDa collagen-specific molecular
chaperones that are required for molecular maturation of various
target proteins, collagen is the only substrate protein for HSP47.
On the other hand, collagen is the major component of the
extracellular matrix and the most abundant protein in mammals,
making up 25% of the total protein to support and maintain cell
and tissue structures. It is a main component of cartilage,
ligaments, bone and teeth, and is responsible for maintaining the
strength and elasticity of many soft tissues, including skin and
blood vessels [14].

In previous studies, Narita et al. [15] and [16] performed the
basal cyto-toxicity test using cultured human cell lines, NB-1
assay, to samples of water obtained from wastewater and sludge
treatment plants. The tests revealed that the toxicity of the
effluent was more intensive than the influent and return flows
from sludge treatment facilities and the organic matter released
from activated sludge bacteria during their decay process
contributed to the increase in toxicity in the secondary effluent.
Funamizu et al. [17] found that secondary effluent contains some
organic matter that causes stress in cells by applying stress
response bioassays using CHO cells transfected with HSP47
promoter to samples obtained from a wastewater treatment plant.
These studies showed that secondary effluent contains toxic
compounds, but toxicity characteristics of secondary effluent are
poorly investigated. Investigation of the characteristics of
toxicants in secondary effluents is helpful in selecting advanced
treatment alternatives. Furthermore, toxicity characterization of
samples obtained from MBR as an alternative to activated sludge
treatment processes may provide a new standpoint in terms of
SRT and solid-liquid separation by membrane filtration. A
comparison between the toxicity of MBR sludge and activated
sludge was performed by Cicek et al. [18]. They reported that
overall activity was consistently higher in the MBR, and the
biomass in the MBR had a higher viable fraction than the
activated sludge. Kimura et al. [19] and Miyoshi et al. [20]
reported that accumulation of soluble microbial products (SMP)
dissolved organic matter in the MBRs was significant in the
case of shorter SRT. Alfieri et al. suggested that MBR could be
operated at higher SRT without drawbacks in terms of biological
activity [21]. Moreover, sand filtration and coagulation as typical
tertiary treatment processes; along with NF and RO subsequent to
MBR as advanced treatment processes were also investigated.
Furthermore, HSP 47 has proven its effectiveness as a highly
sensitive system that can be used for studying the effect of trace
contaminants such as heavy metals, organic pollutants and
biotoxins. Gupta et al 2010 [22] summarized current development
in the application of stress genes and their products HSPs in
toxicology. We used in this study the HSP 47. Although
evaluating one particular HSP does not provide adequate
information on the toxicity of chemicals, this work is a first
attempt to investigate the effect of bacterial by-products (e.g.;
LPS endotoxin) on water toxicity using bioassays. This research
needs to be followed up by other investigation to study the
interaction of several compounds and the contribution of each in
the context of overall toxicity and synergetic, antagonistic and
additive effects that may occur.

The objectives of this study are: 1) to characterize the toxicity
of effluent from an activated sludge treatment process by
molecular weight distribution and HSP bioassay; 2) to assess the
effectiveness of toxicity treatability of pilot-scale MBRs with
different SRTs in attenuating water toxicity based on HSP
bioassays for MBR permeates and supernatant of mixed liquor in
MBRs; and 3) to assess the toxicity treatability of sand filtration,
batch-scale coagulation tests, pilot-scale NF and RO as advanced
treatment processes.

Materials and methods

Chemicals and reagents
The following reagents were purchased from several
manufacturers and were used to prepare the culture medium and
the required solutions: NaCl, KH₂PO₄, KCl, NaH₂PO₄·2H₂O,
bovine serum albumin, NaN₃, dimethyl sulfoxide, glycline, NaOH,
and sodium dodecyl sulfate (all from Wako), Na₂HPO₄·7H₂O
(from MP Biomedicals), fetal bovine serum (from Biowest),
Geneticin (G418) (from Invitrogen), kanamycin solution, trypsin
(ethylenediaminetetra-acetic acid [EDTA]), MgCl₂·6H₂O, and 4-
methylumbelliferyl-β-galactose (MUG) (all from Sigma-Aldrich),
and lysis buffer (from Promega).

Samples
As illustrated in Figure 1, wastewater samples were collected
from an existing municipal wastewater treatment plant in Sapporo,
Japan operated with a conventional activated sludge process and
a rapid sand filtration process as advanced treatment. As the
overall goal is to promote potable reuse, it is fully understandable
that the type of raw water used for the investigation will affect the
results. Nevertheless, in this attempt comparison between levels of
stress responses before and after treatment process will be interpreted with consideration of the bacterial products. These
products are released during biological reaction and their impacts
on cells are independent of the type of the water.

Samples were also collected from a pilot-scale treatment process
operated in the same plant. It consists of MBRs, fed with the
same raw wastewater, that are operated as an alternative to
conventional activated sludge systems. The MBR permeate was
further treated with a parallel set of nano-filter (NF) and reverse
osmosis (RO) membranes. Additionally, a batch scale coagulation test was conducted on secondary effluent.
Figure 1: Schematic diagram of wastewater treatment plant and pilot-scale treatment system

MBR
Three types of MBRs were operated in parallel in this study. Operational conditions of the pilot-scale MBRs are summarized in Table 1. MBR18 was operated with baffled MBR (BMBR). The BMBR examined in this study was equipped with 6.8m² flat-sheet-type micro-filtration (MF) membranes (Toray, Tokyo, Japan) and operated at 18 days SRT. The membrane was made of polyvinylidenefluoride (PVDF) and had a nominal pore size of 0.1μm. On the other hand, MBR50 and MBR12 are submerged MBRs with SRTs of 50 days and 12 days respectively. Hollow-fiber MF membranes (Mitsubishi Rayon Engineering, Tokyo, Japan) made from PVDF polymer were used in MBR50 and MBR12. The nominal pore size of these membranes is 0.4μm.

Table 1. Details of three MBRs.

<table>
<thead>
<tr>
<th>ID</th>
<th>MBR Type</th>
<th>Volume (m3)</th>
<th>Membrane Type</th>
<th>Nominal Pore Size (μm)</th>
<th>SRT (day)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MBR50</td>
<td>Submerged</td>
<td>0.00255</td>
<td>Hollow fiber</td>
<td>0.4</td>
<td>50</td>
</tr>
<tr>
<td>MBR18</td>
<td>Baffled Submerged</td>
<td>0.712</td>
<td>Flat sheet</td>
<td>0.1</td>
<td>18</td>
</tr>
<tr>
<td>MBR12</td>
<td>Submerged</td>
<td>0.00255</td>
<td>Hollow fiber</td>
<td>0.4</td>
<td>12</td>
</tr>
</tbody>
</table>

NF/RO
Two 2-inch spiral-wound NF/RO membrane modules were used in this study: NF membrane (LES90, Nitto Denko; Tokyo, Japan) and RO membrane (ES10, Nitto Denko; Tokyo, Japan). Membrane characteristics of the NF/RO membranes are given in Table 2. Membrane flux and recovery, in the continuous operation, were fixed at 460Lm⁻²day⁻¹ and 70% respectively. Permeate samples were collected after one day of operation.

Table 2. Characteristics of NF and RO Membranes

<table>
<thead>
<tr>
<th>Membrane ID</th>
<th>Material</th>
<th>Water permeability (Lm⁻²day⁻¹Pa⁻¹)</th>
<th>Zeta potential (mV)</th>
<th>Nominal salt rejection (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>NF (LES90)</td>
<td>Polyamide</td>
<td>1.6</td>
<td>-8.6</td>
<td>90.0</td>
</tr>
<tr>
<td>RO (ES10)</td>
<td>Polyamide</td>
<td>1.2</td>
<td>-15.3</td>
<td>95.5</td>
</tr>
</tbody>
</table>

Coagulation
Secondary effluent from the wastewater treatment plant was used for the batch-scale coagulation test. Coagulation was conducted in 500mL glass beakers with the following sequence: 1minute of rapid mixing at 110 rpm, 30 minutes of flocculation at 25 rpm, 30-min of settling. In this experiment, polyaluminum chloride (PAC) as coagulant was added at a concentration of 10 mg Al/L. Several sets of coagulation tests were conducted at different pH values. The pH levels in the tests were adjusted using HCl (1N) or NaOH (1N) before adding the PAC. Dissolved organic carbon (DOC) and turbidity were measured after the test.

Molecular weight distribution (MWD)
In order to obtain the information about which molecular fractions cause the toxicity, we applied bioassays to samples at different MWDs. Samples were fractionated into different molecular weights using a series of MF membranes (ADVANTEC, Tokyo, Japan) and ultra filter (UF) membranes (Alfa Laval, Lund, Sweden). Pore sizes of the MF membranes were 0.1μm and 0.2μm. The molecular weight cut off of the UF membranes were 100kDa and 25kDa. The molecular weight fractionation was carried out on two types of secondary effluent. The first one was fed with only raw wastewater (Case A: samples taken from plant A), while the other one was fed with a mixture of raw wastewater and rejected water from a sludge treatment facility (Case B: samples taken from plant B). Return flow from the sludge treatment facility was mixed into raw wastewater at a rate of 20%. Supernatant of mixed liquor in MBR was fractionated as well.

Toxicity test (HSP assay)
Chinese hamster ovary (CHO) cells stably transfected with (+) or without (-) a HSP 47 promoter were used for this experiment. It has already been revealed that the production of stress proteins HSP 47 is induced as a result of the reaction of CHO cell with a stressor such as wastewater contaminants [13]. The use of mammalian cells such as CHO cells to assess the stress response pathway related to HSP 47 has high relevance to human health. However, as potable reuse is the target, further investigations on ingestion of contaminated water have to be carried on using cells relevant to digestive tract such as hepG2 or Caco 2 cells. Furthermore, although being with high relevance to human health, using CHO cells to detect only one particular HSP is too limited to be correlated with in vivo tests. This work is a first attempt to study about stress response induced by bacterial by-products and further investigations are needed.
In this article, heat shock protein 47-promoter-transfected cells will be abbreviated into HSP(+). The CHO cells were provided by S. Yokota (Kaneka, Osaka, Japan) and were grown as adherent monolayer in 75-cm² tissue culture flasks using F12 Medium (Invitrogen, Carlsbad, CA, U.S.A.), supplemented with 10% fetal bovine serum, 200 μg/mL of G418, and 0.1 g/L kanamycin solution. The cultures were maintained in a 5% CO₂ incubator at 37 °C. Cell passage was carried out at 80% confluence at a 1:2 ratio using 0.25% trypsin (1 mM EDTA).

HSP47 activity can be determined using HSP47-transformed CHO cells by incubating the latter with the test samples and measuring the enzymatic activity of β-galactosidase. When introduced into a chromosome, the HSP47 plasmid can express β-galactosidase efficiently during stress induction. Experimental CHO cells were transformed by inserting the β-galactosidase gene downstream of the HSP47 promoter while control CHO cells had the β-galactosidase gene under the control of the SV40 pA promoter [23]. Isoda et al. [13] developed this system for detecting trace amounts of environmental pollutants and natural toxins and we followed this protocol in this study. Cells were exposed to the sample for three hours and the amount of HSP was measured using a multi-detection micro plate reader in terms of fluorescence (Multi-Detection Micro-plate Reader POWERSCAN HT). In the evaluation, the relative value of HSP production was calculated by the following equation:

\[
\text{The relative stress response} = 100 \times \frac{\text{sample HSP(+)} - \text{sample HSP(-)}}{\text{control HSP(+)} - \text{control HSP(-)}}
\]

HSP assay is not very accurate and requires a statistical analysis. Student’s t test was adopted to statistically assess the differences between the assay results of samples and negative controls. The error bar used in this article represent the 95% confidence interval, which is wider than standard error bar. This fact must be taken into account when analysing statistical significance. Using standard deviation quantifies variability, but does not consider sample size. Hence, it tells nothing about whether the difference is, or is not statistically significant. It should be noticed that, even when the 95% confidence intervals overlap, the difference could be statistically significant.

**RESULTS AND DISCUSSIONS**

**HSP assay of raw wastewater, secondary effluent and effluent from sand filtration**

Figure 2 shows the results of HSP assays of raw wastewater, secondary effluent and effluent from rapid sand filtration. In the figure, the vertical axis shows the relative stress response of each sample compared to the control. Control value is fluorescence from the cells exposed to the medium, which did not contain any samples. When relative stress response exceeds 100% and it is significantly different from the control (p<0.05), we define it as toxic to the cell and significance is denoted with asterisks in the figure. The error bar shows the range of observed fluorescence. In the assay, samples were diluted with medium several times to obtain the dose and response relationship such as the low response at low dose range, and the low response at high dose range because of the damage to the cell itself caused by too high a dose.

![Figure 2: HSP assay results of raw wastewater, secondary effluents and effluents from rapid sand filtration. Asterisk in the figure shows the sample with statistically significant difference with control (t-test, p<0.05).](image)

The data in the figure show that the stress response was detected in secondary effluent and effluent from rapid sand filtration; by contrast, raw wastewater showed less stress than that of the influent. This result indicates the possibility that toxic organic matter was generated during the activated sludge treatment process and was not removed by rapid sand filtration. Eckenfelder reported that effluent toxicity might be created in the biological treatment process itself [24]. Rappaport et al., using the Ames test, showed a greater mutagenic response in secondary effluent than that of the influent [25]. Narita et al. have found that the organic matter released from activated sludge bacteria during their decay process contributed to the increase in toxicity in the secondary effluent ([15] and [16]). Guizani et al. studied the endotoxic activity of samples from a wastewater treatment plant. They reported that the ratio of endotoxic material concentration to chemical oxygen demand (COD) of the treated effluent was higher than that of the influent [26].

Heavy metals are hardly removed by conventional wastewater treatment facility (operated using activated sludge process). According to the annual report of Sapporo sewage works (2009) [30], the investigated treatment plant (raw water) has trace heavy metals below detection limit, except for arsenic and zinc that account for 4μg/l and 1.6 μg/l, respectively. These levels of heavy metals remained constant after treatment. In addition, Guizani and al. 2009 [26] found that the organic matter initially found in wastewater has decreased after treatment. However, HSP response of the effluent was found to be higher that that of the influent. This increase in the stress response is likely to be associated with the release of bacterial by-products (LPS endotoxin) as reported by Guizani et al 2009 [26]. Ben Fredj et al. 2010 [31] reported that, in certain conditions, presence of organic matter (bacterial by-products in this case) masks the heavy metal effect.
HSP assay of fractionated secondary effluent samples

Figure 3 illustrates the HSP assay result of two types of fractionated secondary effluent. Figure 3(a) corresponds to samples from plant A (case A) and Figure 3(b) corresponds to samples from Plant B (case B). Both cases showed significant stress response in samples with relatively high MW fractions. By contrast, MW fractions less than 0.1μm did not show significant stress response, therefore, the higher MW fraction (>0.1μm) causes toxicity to cells. Furthermore, the stress response of case B was more intensive than case A. This result agrees with our previous research. Funamizu et al. demonstrated that the rejected water from a sludge treatment plant such as overflow from thickener and rejected water from dewatering processes are some of the origins of the toxicity of effluent from wastewater treatment plants [17]. Guizani et al. also found that high endotoxic activity was confirmed in the rejected waters from sludge treatment facilities [27].

![HSP assay of fractionated secondary effluent samples](image)

**Figure 3(a):** HSP assay of fractionated secondary effluent; (a) HSP assay results of biologically treated sewage (Case A). Asterisks in the figure show the sample with statistically significant difference with control (t-test, p<0.05).

**Figure 3(b):** HSP assay of fractionated secondary effluent; (b) HSP assay results of Biologically treated mixture of sewage and rejected water from sludge treatment facilities (Case B). Asterisks in the figure show the sample with statistically significant difference with control (t-test, p<0.05).

HSP assay of the effluent from three MBRs and fractionated samples from supernatant from MBR reactors

Because secondary effluent showed toxicity, we focused on the treatment capacity of MBR as an alternative to the activated sludge process. Membranes at longer SRTs are expected to have better biodegradation of organic matter and removal of toxic compounds as compared to the activated sludge process. In this study, three types of MBRs with different SRTs were examined. Figure 4 shows the HSP result of effluent from the three MBRs. No significant stress response was detected in the MBR permeates except MBR12, which was operated under a shorter SRT (12 days) and equipped with a large pore size (0.4μm). DOC concentration of effluent from MBR12 was also high as shown in Table 3.

![HSP assay of effluent from three MBRs](image)

**Figure 4:** HSP assay of effluent from three MBRs. Asterisks in the figure shows the sample with statistically significant difference with control (t-test, p<0.05).

We also applied HSP assays to the supernatant of mixed liquor of two MBRs with different SRTs (50 days and 12 days) to investigate how SRT affects stress response. Samples were centrifuged and then filtered through a 0.45μm membrane filter. Figure 5 shows the results of the HSP assays of these samples. The supernatants showed significant stress response, and in both cases, the stress was significant in samples including higher MW fractions (>0.1μm) described in Figure 5(a). As shown in Figure 5(b), in the 12 days-SRT cases, stress response was significant at all MW ranges. In comparison the DOC concentration (<0.45μm) of MBR12 was higher than MBR50. Holakoo et al. (2006) concluded that longer SRT might lead to the accumulation of higher MW fractions (>100kDa) of biomass-decay-associated SMP [28]. Using sequencing batch MBR, Bin et al. reported that

![HSP assay of effluent from three MBRs](image)

**Figure 5:** HSP assay of effluent from three MBRs. Asterisks in the figure shows the sample with statistically significant difference with control (t-test, p<0.05).
high MW components become more evident at long SRT [29]. MBR operation at shorter SRT induced inadequate biodegradation of toxic organic matter and resulted in toxicity of all MW fractions. Therefore, in the case of MBR12 the MBR permeate showed a stress response because smaller MW fractions carrying toxicants pass through the membrane. The toxic compounds in a supernatant of mixed liquor cannot be removed at longer SRT, but, toxicity was not detected in permeates. This is probably caused by the fact that small molecules aggregated together in larger molecules and were then removed by MBR. Hence, selection of the pore size of the membrane will be important because the size (MW) of organic matter, which has toxicity, is strongly dependent on SRT.

Figure 3(a) and 3(b) show the stress response was detected only in secondary effluent samples larger than 0.1 μm (Figure 3(a) and 3(b)). Appropriate pH causes the settlement of large molecules and thus no significant stress response was observed in the supernatant.

In this study, batch scale coagulation tests of secondary effluent were performed using PAC as coagulant at 10 mg·A/L dosages. In these batch tests, pH was adjusted to several ranges, using HCl(1 N) or NaOH(1 N) before adding the PAC, to assess the pH influence on the removal of toxicants and thus on the stress response. Turbidity and DOC removal were described in Figure 6, and as shown in this figure, the best pH's for removal of both turbidity and DOC were different. While the best pH was in range of 6.7-7.1 in terms of turbidity removal, the best pH for DOC removal was in a lower range (5.0-5.2). Figure 7 illustrates the results of the HSP 47 assay of samples from coagulation tests at different pH values. At a pH ranging from 5.7 to 6.0, 6.7 and 6.9, no stress response was observed. This result indicated that there was no clear relationship between HSP stress response and turbidity or DOC removal. However, pH levels where no stress response had been observed showed relatively high removal in terms of both indexes. It is thought that higher MW components in the secondary effluent cause toxicity, especially MWs of more than 0.1 μm, and they can be removed by coagulation with the appropriate pH adjustment (6-6.7). Indeed, significant stress response was detected only in secondary effluent samples larger than 0.1 μm. Figure 7 shows the result of the HSP assay of permeate from NF and RO subsequent to MBR18. As shown in this Figure, a significant stress response was not detected in any of the samples. Because effluent from MBR18 did not show toxicity, it seems that prior to NF and RO, toxic compounds were removed by MBR18. The MBR-NF/RO treatment system is effective for the removal of toxic compounds, and this system can be multi-barrier to achieve stable, high quality reclaimed water. However, it remains possible that stress response changes depend on combinations of treatments. Looking ahead, further investigations are needed to study the effect of membrane fouling (due to long term operation) on stress response and release of HSP 47 from CHO cells exposed to membrane permeates.

Its well known from the literature that NF and RO cannot remove
Conclusions

Toxicity assessment using stress response bioassays can be an effective tool for selection of wastewater treatment processes and consideration of the operational conditions of each process. In this study, we applied bioassays using CHO cells with HSP47 promoter to wastewater samples including reclaimed wastewater.

The following conclusions were suggested in terms of HSP assays. The HSP assay could detect the toxicity related to organic matter derived from microorganisms in activated sludge. Large molecules (>0.1μm) are the main cause of significant stress response.

While rapid sand filtration cannot remove the toxic compounds detected in secondary effluent, coagulation with an appropriate pH (6–6.7) was effective for reducing stress response. Stress response can be removed by MBR treatment with the appropriate operational conditions such as SRT and pore size of the membrane. The results obtained from this study suggest that it is better to have longer SRTs and membranes with smaller pore sizes.

No significant stress response is detected in permeates from NF and RO subsequent to MBR treatment, and the MBR-NFRO system can be multi-barrier to achieve stable and high quality reclaimed water. Although HSP 47 is used to assess the toxicity induced by heavy metals and hydro-soluble organic pollutants, based on this investigation it can be concluded that bacterial by-products contribute significantly to the release of stress response HSP 47.

Notes and references

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Notes and references

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