Investigating the fate of natural organic matter at a drinking water treatment plant in South Africa using optical spectroscopy and chemometric analysis

Welldone Moyo¹, Nhamo Chaukura¹, Machawe M Motsa¹, Titus AM Msagati¹, Bhekie B Mamba¹, Sebastiaan GJ Heijman², and Thabo TI Nkambule¹

¹Nanotechnology and Water Sustainability (NanoWS) Research Unit, University of South Africa (UNISA), Johannesburg, South Africa
²Department of Civil Engineering and Geosciences, Technical University of Delft, Delft, The Netherlands

The removal dynamics of biodegradable dissolved organic carbon (BDOC) and natural material (NOM) polarity fractions at a water treatment plant (WTP) in South Africa was studied using UV-Vis absorbance, fluorescence excitation-emission matrix, and two-dimensional synchronous fluorescence spectroscopy (SFS). This study gave insights into the transformation of NOM due to treatment processes. The objectives of the study were: (i) to use chemometric analysis and two-dimensional SFS correlations to investigate the evolution of NOM arising from treatment processes, and (ii) to access the chemical profile dynamics of polarity and BDOC fractions throughout the treatment train. The UV_fra absorbance, which indicates aromaticity, reduced by 45% along the WTP. Gaussian fitting of UV-Vis data showed a decreasing trend in intensity and number of bands along the treatment process. The removal efficiency of NOM components followed the order: humic-like (HL) > tyrosine-like (TYL) > fulvic-like (FL) > tryptophan-like (TPL) > microbial-like (MBL). At the source, the relative distribution of the hydrophobic (HPO), hydrophilic (HPI), and transphilic (TPI) fractions was 45%, 31%, and 24%, respectively. The HPI was recalcitrant to treatment, and the TYL component of the HPI fraction was conjectured to be a disinfection byproduct limiting reagent. The HL and FL components of the BDOC fraction were the major substrates for bacterial growth. According to two-dimensional-SFS correlation, TYL, TPL, and MBL varied concurrently across the treatment stages. Used for the first time in South Africa, the robustness of a multi-dimensional approach of optical methods coupled with chemometric tools for the assessment of the fate of NOM along the treatment processes was revealed by this study.

INTRODUCTION

The fate of natural organic matter (NOM) in drinking water treatment processes remains contentious due to the heterogeneous nature and structural complexity of NOM. Compounding this is its temporal and spatial variability, making efforts to remove it uncertain (Nkambule et al., 2012). A single set of results is therefore not enough to either project these to a new treatment system or predict future trends for an existing system. Detrimental to water treatment and distribution, NOM is the major contributor to the fouling of membranes and activated carbon, is a precursor to the formation of disinfection by-products (DBPs), impacts on the organoleptic properties of water, and certain fractions of NOM promote biological growth in distribution networks (Lobanga et al., 2013; Lyon et al., 2014; Hua et al., 2015; Park et al., 2016). Therefore, closely monitoring the transformation of NOM along the treatment train is essential for the management of NOM-related problems. Current research characterizes NOM with emphasis on properties such as biodegradable dissolved organic carbon (BDOC), polarity, UV absorption, fluorescence intensity, and molecular weight fractions (He et al., 2013; Chen et al., 2017; Li et al., 2017a).

Biodegradable dissolved organic carbon is the fraction of NOM that is responsible for providing heterotrophic bacteria with energy and nutrition (Li et al., 2017b). Its measurement entails quantification of the reduction of dissolved organic carbon (DOC) after a predefined timeframe, ranging from a few days to several weeks and depending on, inter alia: (i) incubation timeframe, vessel type, size, and type (batch culture, flow-through bioreactor); (ii) initial DOC concentration; (iii) supplementary nutrient additions and/or additives; (iv) type of bacterial strain, amount of inoculums, and monitored biodegradation product (change in DOC over time, evolution of carbon dioxide); and (v) temperature and frequency of measurements (Mcdowell et al., 2006; Lohwacharin et al., 2014; Marais et al., 2016; Prest et al., 2016). However, these measurement approaches are lacking in that they measure the reduction in bulk NOM over time, disregarding the fact that NOM comprises of a wide continuum of carbon (e.g fluorescent – aromatic and conjugated and non-fluorescent – aliphatic and saturated). Therefore, a measure of DOC reduction cannot adequately identify the specific DOC fraction available for mineralization by heterotrophic bacteria. Knowledge of the character of the labile fraction available for assimilation by heterotrophic bacteria is necessary for adjusting the treatment protocol to target the problematic NOM fraction and hence starve the bacteria.

Besides BDOC, the other most studied parameter of NOM is polarity. The polarity of organic matter influences its palatability by heterotrophic bacteria and is used as a measure of chemical changes caused by environmental and treatment processes (Liao et al., 2015a). Low molecular weight hydrophilic fractions of NOM are more palatable to bacteria than high molecular weight

Correspondence

Thabo TI Nkambule

Email
nkambtt@unisa.ac.za

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hydrophobic fractions (Chen et al., 2014). However, the available analytical techniques are not capable of concurrently characterizing NOM polarity fractions under prevailing environmental settings.

The polarity rapid assessment method (PRAM) partitions NOM based on adsorption preference onto solid-phase extraction (SPE) sorbents (Rosario-Ortiz et al., 2004a). Studies on PRAM characterization applied to precursors of nitrogenous DBPs reported higher selectivity than the conventional XAD resin approach (Chen et al., 2016). Another study modified the PRAM method by eluting the fractions sequentially instead of the parallel elutions of fractions producing three fractions, namely, the hydrophobic, hydrophilic and transphilic fractions (Nkambule., 2012a). PRAM fractionation has been widely applied to the bulk NOM without further analysis of the produced fractions (Rosario-Ortiz et al., 2007b; Nkambule, 2012a; Chen et al., 2014; Liao et al., 2015b). Therefore, additional analysis of the fractions can be performed to reveal more information on the characteristics of the fraction. To this end, this study used fluorescence excitation-emission matrix (FEEM) spectroscopy for an in-depth investigation of the polarity and BDOC fractions character.

Fluorescence spectroscopy is a robust tool for characterizing fluorescent dissolved organic matter (FDOM) in various natural and engineered aquatic systems (Baghth et al., 2010; Sanchez et al., 2013; Tijani et al., 2014; Zhang et al., 2015). Owning to its simplicity in application, FEEM spectroscopy has promise in predicting the removal of NOM during both conventional and advanced drinking water treatment processes (Kastl et al., 2016; Wang and Benjamin, 2016; Li and Hur, 2017). Coupled with multivariate statistical methods such as parallel factor analysis (PARAFAC), self-organizing maps (SOM), principal filter analysis (PFA), and fluorescence regional integration (FRI), EEMs have been used to identify NOM components and provide further insight into the environmental dynamics of NOM in diverse aquatic ecosystems (Henderson et al., 2009; Bieroza et al., 2011; Korak et al., 2013; Sanchez et al., 2013).

Complementary to the FEEM spectra, synchronous fluorescence spectroscopy (SFS) shows advantages of better spectral deconvolution, light dispersion reduction, and well-resolved peaks and shoulders (Guo et al., 2013; Costa et al., 2016). Coupling two-dimensional correlations to SFS (2D-SFS) produces a powerful chemometric tool that can resolve superimposed peaks by extrapolating them to the second dimension. This can also be used to determine the order of spectral changes in response to external perturbations (Chen et al., 2017). The combination of these techniques in the assessment of NOM fractions provides data on the character of NOM at each treatment stage.

Herein, we propose the assessment of NOM and its BDOC and polarity fraction dynamics with insights from UV-Vis absorbance with Gaussian fitting, FEEM, and 2D-SFS for tracking NOM transformation throughout the treatment train. The majority of treatment plants in South Africa follow the conventional treatment processes, with flocculation-coagulation as the major organics removal stage, and chlorination at the disinfection stage (Nkambule et al., 2012; Marais et al., 2016; Chaukura et al., 2018). To the best of our knowledge, this is the first study conducted in South Africa to investigate the occurrence, fate, and removal efficiencies of specific NOM fractions including in-depth transformations of BDOC and polarity fractions brought about by the treatment processes. The objectives of the study were to: (i) investigate the fate of NOM and subsequent transformations brought about by treatment processes; (ii) evaluate the chemical profile dynamics of polarity and BDOC fractions throughout the treatment train; and (iii) apply 2D-SFS correlation for the assessment of dynamic spectral properties of NOM due to water treatment processes.

**MATERIALS AND METHODS**

**Bulk sample analysis**

**Sampling**

Sampling was conducted at a water treatment plant in the Gauteng Province of South Africa. The plant has a maximum treatment output of 42 ML·day⁻¹. Triplicate samples were collected after each treatment stage. The treatment processes involve coagulation/flocculation using a combination of ferric sulphate and a commercial organic flocculant with pH correction using lime, followed by dissolved air floatation filtration (DAFF) through sand beds, and finally chlorination. Portable multimeters were used to measure turbidity, pH, temperature, and conductivity on site. Samples were transported to the laboratory in ice boxes and then filtered using a 0.45 μm GF/F filters (Sigma Aldrich, Germany) prior to storage at 4°C and analysed within 48 h.

**Dissolved organic carbon, fluorescence and UV absorbance analysis**

The filtered samples were analysed for DOC using a total organic carbon analyser (Teledyne Tekmar, TOC fusion). Fluorescence EEMs and absorbance spectra were measured in the wavelength range 200–800 nm at Δλ = 2 nm excitation intervals, and 248.58–830.59 nm at Δλ = 3.28 nm emission intervals using a fluorescence spectrometer (Aqualog, HORIBA, Jobin Yvon). To determine the overall removal of NOM and efficiency of the treatment steps, the UV absorbances of water samples were measured at a wavelength, \( \lambda = 254 \text{ nm (UV}_{254} \). The character of the three fractions (hydrophobic, hydrophilic, transphilic) obtained through the polarity rapid assessment method (PRAM) was also analysed using UV_{254} measurement. Further, simulated synchronous fluorescence spectroscopy at a wavelength offset of \( \Delta \lambda = 18 \text{ nm} \) was used to study the fluorescence characteristics of water samples collected after each treatment stage at the same wavelength range as the EEMs (Guo et al., 2013).

**In-depth NOM fraction analysis**

**Polarity rapid assessment method**

The modified-polarity rapid assessment method (m-PRAM) was used to partition NOM into three fractions, namely: the hydrophobic (HPI), hydrophilic (HPO) and transphilic (TPI) fractions (Nkambule et al., 2012). In this method, a 24-position vacuum manifold (Sigma Aldrich, Germany) was connected to a vacuum pump operated at 5 inches Hg, 10 kPa, and 20 mL glass test tubes were used for collecting the SPE filtrate. The C18 (non-polar), CN (polar) and NH₂ (weak anionic) cartridges, each of size 100 mL aliquots was added to each cartridge to wash off the residual methanol and loose organic matter until a steady-state UV 254 was obtained. Thereafter, 20 mL of sample was introduced into the cartridges sequentially starting with C18 cartridge; the permeate thereof was introduced into the CN cartridge, then ultimately to the NH₂ cartridge. About 10 mL 0.1 M NaOH was eluted through the C18 and CN cartridges to elute the HPO and HPI fractions, respectively. The TPI was the fraction which was not retained by the NH₂ cartridge.
Biodegradable dissolved organic matter
An inoculum of biologically active sand (BAS) was collected from the DAFF beds at the WTP. Excess carbon in the sand was removed by washing with 0.1 M sodium thiosulphate solution until a steady UV\textsubscript{254} absorbance close to zero was obtained. Thereafter, the sand was rinsed in deionized water and the supernatant analysed for DOC, UV\textsubscript{254}, and fluorescence, and used as the baseline. The washed sand (100 g) was placed in 500-mL Erlenmeyer flasks covered with aluminum foil, and the water sample (300 mL) was added. Three solution concentrations of glucose (5, 8 and 10 mg\textcdot L\textsuperscript{-1}) were used as control. The flasks were incubated at 22°C in a water bath for 5 days, and daily measurements of DOC, UV\textsubscript{254}, and fluorescence were carried out.

Chemometric analysis
Fluorescence regional integration of removed fractions
The EEM spectra were divided into 5 wavelength-dependent regions representative of specific components of NOM (TYL, TPL, FL, MBL, and HL) (Ejarque et al., 2017). The equations for the calculations of the quantity of respective components under the EEM curve is shown in Box 1 (He et al., 2013).

Box 1. Equations used to determine the quantitative contribution of each region

\[
\theta_i = \sum_{\text{ex}} \int (\lambda_{\text{ex}}, \lambda_{\text{em}}) \Delta \lambda_{\text{ex}} \Delta \lambda_{\text{em}} \text{d} \theta_i \\
\theta_{\text{Tot,5}} = \theta_{\text{Tot,5,ex}} = \sum_{\text{ex}} \theta_i \\
P_i = \frac{\theta_i}{\theta_{\text{Tot,5}}} \times 100\% 
\]

where \(\theta_i\) is the volume bound by region \(i\) and \(\theta_{\text{Tot,5,i}}\) is the total volume of all five regions; the specific region is denoted by number \(n\); and \(P_i\) is the percentage volume of specific region \(i\).

FRI was applied in the quantification of fractions making up the BDOC fraction:

\[
\text{BDOC}_{\text{EEM}} = \text{EEM}_{\text{day1}} - \text{EEM}_{\text{day5}} 
\]

Parallel factor analysis
PARAFAC model was generated using the SOLO software (EigenVector Inc, USA) which is inbuilt in the fluorescence spectrometer (Aqualog, HORIBA, Jobin Yvon). Disproportionate leverage that affects the validation of the model was corrected by disregarding fluorescence intensity below 240 nm. Non-negativity constraints were applied for all models, and outlier tests and leverage plots were used to exclude outlier samples from the dataset. The PARAFAC modeling was validated using a series of predetermined criteria according to Murphy et al. (2013): (i) core consistency examination, (ii) spectral loading shape evaluation, (iii) contribution of the influence of specific sample or wavelengths on the leverage, (iv) analysis of the residual, and (v) the split half criterion. Following validation, the quantitative distribution of the components after each treatment stage was quantified using their maximum fluorescence intensities (\(F_{\text{max}}\)). In-depth information on the PARAFAC model can be accessed in Murphy and Stedmon (2017).

Gaussian fitting
The PeakFit v4.12 curve-fitting software was employed to deconvolute the UV-Vis spectrum and fit the Gaussian peaks.

The location of the maxima, intensity, and energy of each Gaussian peak characterized the extent and efficiency of each treatment process. Prior research (e.g., Dryer et al., 2008a; Dryer et al., 2008b) revealed that these bands have a Gaussian nature when expressed in photon energy (Eq. 2):

\[
E(\text{eV}) = \frac{1240}{\lambda(\text{nm})} 
\]

Two-dimensional correlation spectroscopy
Two-dimensional correlation synchronous fluorescence spectroscopy was conducted to determine the rate of compositional variation of NOM due to the impact of treatment using the ‘2Dshige software’ (Kwansei-Gakuin University, Japan). The order of treatment was the external perturbation. Nodea and Ozaki provide a detailed description of the algorithms used in the 2Dshige software (Chen et al., 2017). In brief, 2D-SFS produces a synchronous and asynchronous map, auto-peaks and cross-peaks characterize a synchronous map and the peaks are located along the 1:1 slope line and off the diagonal line, respectively, while cross peaks characterize asynchronous map. These peaks are either positive or negative to show the sequential order of dynamic spectral variations.

RESULTS AND DISCUSSION
Removal of bulk parameters
There was a 45% reduction in UV\textsubscript{254} absorbance along the WTP (Fig. 1). This indicates the WTP has the capacity to significantly reduce the aromatic content of NOM throughout the treatment stages. Disinfection was the major contributor to UV\textsubscript{254} reduction (27%). This is attributed to the reduction of activated aromatic rings, conjugated double bonds, and long aliphatic chains of NOM to smaller and hydrophilic organic fractions by chlorine (Hur et al., 2014; Golea et al., 2017). Coagulation/flocculation and DAFF contributed 12.9% and 12.7%, respectively, to the elimination of UV\textsubscript{254} absorbance. Although coagulation is usually the major NOM removal process, research has shown that SUVA < 2 L\textcdot mg\textsuperscript{-1}·m\textsuperscript{-1} implies that the major portion of NOM in the water is of non-humic substances and that less than 25% should be removed when SUVA < 2 (Lavonen, 2015). UV\textsubscript{254} is used in the water treatment industry to indicate organic matter containing conjugated functional groups with high aromatic rings and with characteristic π-π* electron transitions, typical for groups containing alkenes and carbonyl bonds, an attribute of humic substances (Matilainen et al., 2002; Li and Hur, 2017). Therefore, UV\textsubscript{254} can be used as a semi-quantitative indicator of the concentration of NOM in natural waters. The results indicate that the water is mainly non-humic with poor removal by coagulation. Additional to UV\textsubscript{254}, several absorbance-related ratios have been proposed as tools for characterizing NOM (Li and Hur, 2017). For instance, the ratio UV\textsubscript{233}/UV\textsubscript{253} has been used as an indicator for the propensity to form DBPs during water treatment. Aromatic rings rich in -OH, -COOH, -RCOR, -COOR groups give a high quotient, whereas a low quotient is expected from unsubstituted aromatic rings (Li et al., 2017a). The average of the UV\textsubscript{253}/UV\textsubscript{233} ratio decreased along the WTP from 1.219 raw to 0.808 final water, indicating the treatment regime progressively destabilized the aromatic structures, consequently lowering the potential to form hazardous DBPs.

Specific UV absorbance values reduced along the WTP from 1.456 ± 0.025 L\textcdot mg\textsuperscript{-1}·m\textsuperscript{-1} to 1.204 ± 0.020 L\textcdot mg\textsuperscript{-1}·m\textsuperscript{-1} at final water (Fig. 1). The hydrophobic character as measured by SUVA decreased down the treatment train, with a significant reduction (\(p < 0.05\)) observed during the disinfection (18%) treatment...
SUVA < 2 L·mg⁻¹·m⁻¹ indicates a hydrophilic character of NOM; this fraction is often recalcitrant to treatment (Marais et al., 2017). Because of this, it was expected that coagulation should not cause a substantial reduction in NOM (11.6%). Another probable contributing factor to the low NOM removal at this stage could be due to the high pH value (7.29), which offsets the optimal function of coagulants such as ferric salts, whose optimum pH range is 4–6 (Swartz et al., 2004). Research has shown that SUVA gives an indication of the composition of NOM so that treatment processes can be tailored for its removal (Roccaro et al., 2015). SUVA is a semi-quantitative measure of the aromatic and hydrophobic character of the organic matter (Aschermann et al., 2016).

**Perturbations of latent features of the UV-Vis absorbance spectra throughout the treatment stages**

Gaussian fitting on DOC-normalized spectra was processed to identify the bands making up the spectra (Fig. 2). A very close fit of the experimental data and the Gaussian distribution bands was observed ($R^2 > 0.995$). The contributing bands were operationally named: A01 (< 240 nm), A02 (4.66 eV or 266 nm), A03 (4.16 eV or 298 nm), A04 (3.84 eV or 323 nm), A05 (3.49 eV or 355 nm), A06 (3.25 eV or 381 nm) and A07 (3.02 eV or 411 nm). Literature reported similarly positioned Gaussian bands obtained after deconvolution of UV-Vis spectra (Dryer et al., 2008; He et al., 2015). Band A01 peaks at wavelength ($\lambda$) less than 240 nm; consequently it could not be effectively resolved by the deconvolution. Additionally, at $\lambda < 240$ nm there are spectral interferences from nitrates, nitrites, and inorganic ions (Roccaro., 2015). Thus, spectra at wavelengths less than 240 nm are not discussed further in this study.

The electron transfer band of benzene at approx. 253 nm is related to band position A2 (254–260 nm) (Korshin et al., 2017). Band A3 (270–280 nm) was ascribed to $\pi$-$\pi^*$ electron transitions of phenolic arenes and aniline derivatives substituted by at least two rings (Sharpless and Blough, 2014). At present, there is not enough evidence to state with certainty that the band A03 can be assigned to specific DBPs’ formation. More research exploring changes in the intensity, position, and shape of this band characteristics is required. However, it is established that connections exist between the intensity of the differential absorbance at 272 nm and individual DBPs (Korshin et al., 2007; Lavonen, 2015). The intensities of all bands decreased at the disinfection stage and, interestingly, bands in the longer wavelength (i.e. A05, A06, and A07) disappeared at this stage. Absorption at long wavelengths is usually due to intramolecular charge transfer brought about by electron donor/acceptor interactions (Sharpless and Blough, 2014). Such interactions are low energy transitions prone to chemical and biological reactions, and are hence sensitive to treatment processes and can thus be easily removed.
It is noteworthy that the intensities of the bands reduced at specific treatment stages. However, the positions of the maxima of the Gaussian bands did not change considerably at specific treatment stages. The reduction of intensity of these bands is a measure of the efficiency of the WTP in removing chromophoric moieties. In previous studies, Gaussian decomposition of absorbance spectra has proved to be a useful tool in probing the changes in the molecular structures of NOM (Yang et al., 2017).

In-depth NOM fraction analysis at each treatment stage

Polarity fractionation dynamics along the treatment train

At the source, the relative distribution of the HPO, HPI, and TPI was 45%, 31%, and 24%, respectively (Fig. 3a). These results are in agreement with previous reports that surface water NOM consists of about 50% of HPO while the hydrophilic fraction contributes from 25–40%, and the transphilic fraction occupies the remainder (Universiti et al., 2016). The WTP reduced the hydrophobic fraction from 45% content in raw water to 36% in final water. Previous studies have indicated that the HPO fraction is most amenable to removal in conventional treatment processes compared to other fractions (Kim and Yu, 2005; Knowles, 2011; Bagloth, 2012; Hesham et al., 2013). However, this was not the case in this study because the water had low SUVA values, and for SUVA < 2 L-mg$^{-1}$ m$^{-1}$ it is expected that less than 25% of HPO fraction should be removed (Lavonen, 2015).

There was no significant difference between the quantity of the HPI fraction in the raw and final water ($p < 0.01$). However, there was a significant drop in the hydrophilic fraction at the disinfection stage from 41% to 33%. The HPI has high THM and HAA formation potential for waters with low humic content characterized by low SUVA compared to the HPO fraction (Karapinar et al., 2014). Additionally, the HPI fraction is characterized by a low C/O ratio and SUVA < 2 is indicative of less aromatic carbon content (Marais et al., 2017). Therefore, it can be conjectured that the drop in the HPI fraction was due to the formation of DBPs. Treatment options such as coagulation are not effective for removing this type of fraction (Bond et al., 2010; Matilainen et al., 2010; Sarathy et al., 2011). There was a 2% drop in quantity of HPO and a 6% increase in quantity of HPI accompanied by a 12.7% decrease of UV-Vis absorbance at the DAFF stage. The drop in the HPO fraction at this stage is attributable to microbial degradation of high molecular conjugated aromatic structures to simpler aliphatic compounds in the sand filter beds. Heterotrophic bacteria transform NOM by degrading organic matter rich in carbonyl and hydroxyl groups, which elevates the HPI content at the expense of HPO concentrations (Universiti et al., 2016). There was no significant reduction ($p < 0.01$) in the transphilic fraction throughout the treatment stages.

The impact of the treatment stages on the EEM spectra of the PRAM fractions was quantitatively analysed using the FRI technique (Fig. 3b). At all treatment stages, the intensity of HPO in Region V was higher than in other regions. This was expected because this is the fingerprint region for the HPO fraction. Region V is due to the HL fluorescence, while Region IV describes the microbial by-products’ fluorescence, and Region III is for FL material. Regions I and II are due to protein-like fluorescence (TYL and TPL, respectively) (Chen et al., 2003). It was of interest to carry out an in-depth study of the compositional variation of HPI because it is the main predecessor to THM and HAA formation in low-humic water such as in this case (Universiti et al., 2016).

The evolution of the HPI component throughout the treatment train showed the TYL component had an upward trend (20–50%) while the HL component declined (51–18%) from raw to DAFF. This suggests the treatment processes were able to break down the humic fraction into simpler fractions which mimicked a proteinaceous-like spectral signature. Notably, there was a 14% decline in the TYL component and an 11% increase of the HL component at the disinfection stage. The reason could be the formation of DBPs whose spectral signature was similar to the HL component. These findings suggest the TYL component of the HPI fraction is the DBP-limiting reagent. Therefore, sequestrating proteinaceous material before the disinfection stage should subsequently reduce DBP quantity.

The dynamics of biodegradable dissolved organic carbon along the treatment train

The BDOC for raw water was 5.07 ± 0.99 mgC·L$^{-1}$ measured as DOC (Fig. 4a), which was higher than reported values for surface waters (Charnock and Kjonnø, 2000, 0.21–2.79 mgC·L$^{-1}$; Najm et al., 2000, 0.05–0.85 mgC·L$^{-1}$; Volk and LeChevallier., 2002, 0.03–1.03 mgC·L$^{-1}$). This is probably because the DOC of this study area was more amenable to bacterial attack than that reported in literature. BDOC is traditionally measured as a change in DOC over a timeframe with no further regard given to DOC composition (Nkambule et al., 2012a). The BDOC fraction progressively decreased from raw, flocculation, DAFF, and disinfection (47%, 39%, 38% and 11%, respectively), as the water was losing its hydrophobic character ($R^2 = 0.87$). It
has been reported that water with high humic content is more susceptible to biodegradation than low humic waters (Volk et al., 2000). Samples at disinfection showed the lowest BDOC fraction value because it had the lowest DOC content; therefore the change in DOC was minimal compared to other treatment stages. This result is contrary to previous studies which reported an increase in the BDOC fraction at the disinfection stage (Terry and Summers, 2018). Overall, the quantity of the BDOC fraction upon disinfection depends on the quantity and quality of NOM, chlorine dosage, and contact time (Dong et al., 2014).

By definition, BDOC is the fraction of DOC taken up and mineralized by heterotrophic microorganisms within a predefined timeframe (Terry and Summers, 2018). It was of interest to follow BDOC dynamics throughout the creation stages conceptually depicted here by the raw water sample so that the specific components making up the BDOC fraction can be tracked for the purposes of identifying the most amenable component for uptake by heterotrophic bacteria (Fig. 4b). It was observed that the FL fraction increased (19%) while the proteinaceous fractions increase was 5% and 23% for TPL and TYL, respectively, and the HL fraction declined (4%) from Day 1 to Day 2, corresponding to a 44% decline in DOC in the same period (Fig. 4b insert). The increase in proteinaceous material was perhaps due to some species of heterotrophic bacteria secreting extracellular polymeric enzymes to break down the substrate for ease of assimilation whilst in the process consuming HL material (Universiti et al., 2016). The disproportionate decrease in DOC was probably due to other non-fluorescent sources of carbon contained in the DOC available for uptake by the heterotrophic bacteria. From Day 2 to Day 3 all fractions showed a decreasing trend, with FL and HL having the greatest decline (28 and 17%, respectively), while proteinaceous matter showed the least decrease (each less than 8%). At this stage, the implications are that the heterotrophic bacteria have acclimatized to the available substrate (mainly FL and HL) (Filip et al., 2000). The downward trend of FL and HL persisted throughout the study period. Previous reports indicate that humic matter is the preferred material for uptake by heterotrophic bacteria (Prest et al., 2016). From Day 3 to Day 4 there was a spike in TPL (26%) and TYL (20%) with a corresponding 40% rise in DOC. This is the point of cell death where the bacteria had outgrown the available nutrients (Terry and Summers, 2018). The rise in DOC was thus attributed to dead bacteria plus the non-fluorescent and/or non-biodegradable organic matter (Terry and Summers, 2018). Day 4 to Day 5 saw a decline in all fractions with TPL and TYL, giving a 25% reduction each with a corresponding 27% decline in DOC. At this point, the dead bacteria served as substrates for higher bacteria or other organisms in the food chain. According to the FRI results, the contributions of HL, FL, MBL, TYL and TPL were 27%, 19%, 5%, 6% and 5%, respectively, to the total BDOC fraction (Fig. 4c). These results suggest HL and FL are the major substrates for bacterial assimilation. Control and monitoring of these fractions is thus necessary to curb the concentrations of BDOC.

The management of the BDOC fraction is crucial because, if left unchecked, it is this fraction of DOC that is responsible for bacterial regrowth in the water distribution system. The BDOC fraction is also responsible for the alteration of the physicochemical properties of the treated water, impacting on taste and odour, elevation of turbidity, and loss of residual chlorine, subsequently increasing the formation potential of DBPs (Vital et al., 2010; Li et al., 2017b).

Fluorescent NOM fraction dynamics at each treatment stage

NOM characterization using synchronous fluorescence spectra

Representative synchronous fluorescence spectra of the raw water and after various treatment stages at an interval of Δλ = 18 nm are illustrated in Fig. 5a and quantitative distributions of fluorescent fractions are shown in Fig. 5b. PARAFAC analysis was carried out

![Figure 4](image-url)

**Figure 4.** (a) BDOC fraction dynamics throughout the treatment train; (b) BDOC compositional dynamics throughout the formation stages and (c) BDOC fraction EEM spectra
to complement the SFS data (Fig. 5c, 5d and 5e). Previous reports suggest that the fluorescence peaks centered in the ranges of 265–295, 295–360, 360–420, 420–460, and 460–520 nm are assigned to TYL, TPL, MBL, FL, and HL matter, respectively (Pan et al. 2017; Su et al., 2016). Area integration of the raw water sample revealed the TPL component contributed 51.75% abundance followed by MBL (27.24%), TYL (9.74%), FL (7.22%), then HL (4.06%). There was a significant change in the relative component abundance exhibited by the HL and TPL from raw to disinfection, which was a 16 and 14 times reduction, respectively, whereas there was a 3-, 2- and 3-fold change in TPL, MBL, and FL, respectively ($p > 0.05$). The total removal efficiency of the HL component (93.68%) was the highest, followed by TYL (92.83%), FL (71.23%), TPL (60.58%), then MBL (58.52%) (Fig. 5b). Coagulation removed 88.87% TYL, followed by HL (54.44%), FL (36.55%), TPL (30.07%) then MBL (26.22%), whereas disinfection removed 67.38% HL, followed by FL (44.92%), TPL (41.40%), MBL (40.13%), then TYL (29.53%). Thus the main component removal stages were coagulation and disinfection.

Depending on the nature of NOM, the removal efficiencies at the coagulation stage are influenced by the NOM-coagulant properties such as the electrostatic charge interaction, chemical species, and structures, molecular weight distribution, polarity and hydrophobicity (Jarvis, 2004; Su et al., 2017). These results suggest that the TYL component was more amenable to coagulation than the HL component because it has more charged moieties than the HL component. On the other hand, chlorination results in chemical modifications of NOM fractions by selectively targeting double bonds and conjugated functional groups abundant in the HL fraction compared to the TYL fraction (Golea et al., 2017). Depending on the number and distribution of these moieties, there are differential removal rates with the HL fraction compared to TYL component.

Characterising NOM dynamics using 2D-SFS correlation analysis

Two-dimensional correlation analysis was conducted on the SFS dataset to determine any compositional changes of the fluorescent components due to the treatment processes. Positive auto peaks were observed at the wavelength pairs of 279/279 nm and 320/320 nm on the synchronous map (Fig. 6a). According to Noda’s rule, for a wavelength pair $\lambda_1/\lambda_2$ on the synchronous map, an auto-peak means the transformations at the wavelengths $\lambda_1$ and $\lambda_2$ occur concurrently (Noda and Owasaki, 2004), thus corroborating SFS data which showed a progressive decline in the NOM fractions down the treatment train. The peaks at the excitation wavelengths of 279 and 320 nm were ascribed to TYL and TPL, respectively (Pan et al., 2017). Asynchronous maps indicate only cross peaks (Fig. 6b). For a wavelength pair $\lambda_1/\lambda_2$ in the asynchronous map, a positive cross peak indicates that the rate of change happening at the wavelength $\lambda_1$ is fast paced compared to that taking place at the wavelength $\lambda_2$, while a negative cross-peak indicates the inverse, i.e., the rate of transformation happening at the wavelength $\lambda_1$ is retarded when compared to that taking place at the wavelength $\lambda_2$ (Hur et

Figure 5. Fluorescence characteristics of water at different treatment stages (a) Synchronous scan of water samples after each treatment stage, (b) Removal efficiency of NOM fractions in the water samples after each treatment stage. Validated fluorescent components derived from the Parafac model on drinking water sources showing (c) Component 1 (C1), (d) Component 2 (C2) and (e) Component 3 (C3)
Therefore, these results indicate that changes in TYL, TPL, and MBL varied concurrently as water passed through the different treatment stages.

In the asynchronous map (Fig. 6b), two negative cross peaks occurred at 330/279 and 365/279 nm, while weakly positive cross peaks were observed at 410/330 and 480/330 nm. Spectral changes at the excitation wavelength of 279 nm are faster than at the excitation wavelength of 330 and 365 nm, and spectral changes observed at excitation wavelength 330 nm are faster than at excitation wavelengths of 410 and 480 nm (Noda and Ozaki, 2004). The peak at 279 nm was ascribed to TYL matter, while those at 330, 365, 410 and 480 nm are related to TPL, MBL, FL, and HL matter, respectively. Therefore, based on Noda’s rule, the sequence of changing band order was 279 → 480 → 410 → 330 → 360 nm. This implies the TYL component was more susceptible to treatment, followed by HL, FL, TPL, and MBL, and is consistent with data on the removal efficiency of each component at this WTP.

CONCLUSION

The assessment of the transformation of NOM due to treatment processes in a WTP was followed using BDOC and polarity fraction dynamics with insights from UV-Vis absorbance with Gaussian fitting, FEEM and SFS spectroscopy combined with two-dimensional correlations. The main findings were: (i) there was a reduction in the intensity and number of Gaussian bands down the treatment train implying the treatment plant was efficient in removing chromophoric moieties; (ii) the HPI fraction was the DRP-limiting reagent; (iii) the HL and FL of the BDOC were the major substrates for bacterial assimilation; and (iv) TYL, TPL and MBL varied concurrently as water passed through the different treatment stages. This study has revealed the robustness of a multi-dimensional approach of optical methods coupled with chemometric tools, applied for the first time in South Africa, for the assessment of the fate of NOM throughout the treatment processes.

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Figure 6. (a) Synchronous 2D correlation map; (b) asynchronous 2D correlation map originating from 260 to 550 nm region of SFS of NOM in the DWTP. Red indicates a positive correlation, and blue indicates negative correlation; a darker shade indicates a stronger positive or negative correlation.
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