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Organic matter concentration and characteristic dynamics in surface waters post-bushfires and cyclones: fDOM sensors for environmental monitoring and control†

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This study presents the findings of an investigation on the dynamics of dissolved organic matter (DOM) concentration and characteristics of four Australian rivers and reservoirs after their catchments had been severely burned by bushfires (wildfires) or impacted by a tropical cyclone. Dissolved organic carbon (DOC) increased immediately following the events, and subsequently decreased. The findings indicate rapid stabilisation of water quality, based on the measured parameters, following the commencement of the first winter after the events (which occurred in mid/end summer). In the fire-affected Middle River catchment, DOC decreased from 30.7 mg L⁻¹ to 10.2 mg L⁻¹ over approximately seven months. In the case of the Herring Lagoon catchment, which was affected by cyclone Uesi, DOC decreased from 15.6 mg L⁻¹ to 1.2 mg L⁻¹ over approximately ten months. However, the DOM present in the surface water exposed to the cyclone showed higher molecular weight, coagulability and UV-vis absorbance than the DOM present in the surface water of fire-affected catchments. The observed rapid increase and then reduction in DOM concentrations after extreme climate events indicates the need for short-term and rapid responses for drinking water treatment. The fluorescence signal of a field-deployable fluorescent DOM (fDOM) sensor showed potential as an online monitoring tool for assessing DOM concentration in surface waters, including under extreme conditions. The rapid identification of high DOM loadings in surface waters following extreme climate events (e.g. using a field deployed fDOM sensor) along with its coagulability characteristics could assist in catchment management and drinking water treatment by enabling timely control decisions in response to the impacts of such events.

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Environmental significance

Understanding the impact of natural disasters on water sources is vital for resilience and resource management. This study reveals the response of dissolved organic matter (DOM) concentration and characterisation in surface water sources to severe bushfires and a tropical cyclone. Post-event, DOM concentrations initially surged, followed by a rapid decline, indicating water quality stabilisation within the first winter. Despite similar concentration trends, DOM characteristics differed between fire-affected and cyclone-impacted waters, suggesting varied impacts on treatment processes and ecological health. The research emphasises the need for swift drinking water treatment responses to extreme climate events. The deployment of a field fluorometer proves promising for real-time DOM monitoring, aiding catchment management and enabling timely decision-making for water treatment in the aftermath of environmental crises.

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1 Introduction

Extreme climate events can cause the quality of surface drinking water sources to deteriorate through the degradation of catchments, water flow pathways, and storage reservoirs. These events can impact drinking water treatment processes and infrastructure, and the quality of distributed water.^{1,2} Heavy rainfall and floods (including those associated with cyclones) can significantly increase turbidity and dissolved organic carbon (DOC) loads in surface waters from terrestrial (allochthonous) sources.³ This can change dissolved organic matter (DOM) pool characteristics and treatability, impacting the disinfection by-product formation potential in treated water.⁴

Bushfires are a feature of the Australian landscape, that occur following extended periods of dry weather conditions.^{5,6} Extreme events such as the 2019–2020 Australian bushfires have been attributed to the effects of anthropogenic climate change, which has driven increased temperature extremes and long-term warming.^{6,7} Large-scale bushfires can be detrimental to surface water catchments and associated water stream quality, which is an ongoing concern. This can occur through an increase in catchment runoff following such events, which can transport sediments, ash, soluble organic materials, heavy metals and nutrients into waterways.^{8–10} Bushfires can therefore elevate risk of algal/cyanobacterial blooms in surface waters due to elevated nutrient inflows.^{4,11} Extremely high levels of turbidity (up to 4200 NTU), appearing as pitch-black water, were reported for the Murray River after the 2019–2020 Australian bushfires.⁹ Bushfires can lead to elevated concentrations of organic compounds, including dioxins and fire pyrolysis products such as polycyclic aromatic hydrocarbons, polychlorinated dibenzo- ρ -dioxins and dibenzofurans and polychlorinated biphenyls, which could pose challenges for conventional water treatment processes.^{12–16} Exposure of these organic compounds to chlorine during the disinfection process may lead to the formation of halogenated disinfection by-products, which can pose health concerns^{12–14} or provide a substrate for microbial regrowth in drinking water distribution systems.¹⁷

High DOM concentrations can have a significant impact on freshwater ecosystems. Dissolved organic matter is a substrate for microbial growth that fuels microbial-based food webs^{18,19} and transport metal ions including heavy metals.²⁰ Moreover, high DOM concentrations can attenuate light penetration, which consequently limits photosynthetic activity in the water body.²¹ Conversely, DOM induced attenuation of the ionising irradiation portion of light (*e.g.* ultraviolet light) can protect aquatic organisms from its hazardous impacts.²¹ High DOM concentrations can occur due to transport of terrestrial organic matter when catchments become inundated with surface water flows following heavy rainfall.²² This can elevate heterotrophic bacterial activity, deplete oxygen, and kill aquatic organisms.²³ Nonetheless, extreme events such as bushfires and cyclones can also be beneficial by supplying food and nutrients, which increases river productivity and benefits waterbirds, native fish and other aquatic organisms.²³

Few studies have examined the concentration and characteristics of DOM in waterways after fires. Johnston and Maher (2022)²⁴ studied flow-stratified water quality data from a large coastal catchment (Macleay River, Australia) spanning severe drought and extensive fires followed by flooding. They reported that the levels of several water constituents including suspended sediment, PO₄, NO₃, DOC, K, Ca, SO₄, HCO₃, Mg, Cl, Na, and NH₄ were highly elevated (*i.e.* >100, 22, 21, 7.8, 7.4, 5.7, 4.7, 3.9, 3.6, 2.1, 1.6 and >104 times, respectively) during the initial post-fire period (comparison of the maximum post-fire concentrations with pre-fire median concentrations). The observed ion concentration order was reported to reflect the composition of the ash materials. They reported a fast biogeochemical cycling of dissolved inorganic nitrogen species during initial flow events following the fire, with NH₄-N initially dominant (>80% of dissolved inorganic nitrogen [DIN]) and exceeding ecosystem guideline threshold values (>100 μ M NH₄-N), followed by rapid (approximately one week) nitrification. Sherson *et al.* (2015)²⁵ and Johnston and Maher (2022)²⁴ emphasised the necessity for high-frequency sampling to adequately capture the impacts of drought, fires, and floods on aquatic systems due to the high temporal variability of water quality parameters. Another study by Basso *et al.* (2021)²⁶ on boreal sub-arctic lake water chemistry throughout a summer in northern Alberta, where almost 30% of the area that flows into the reservoir was affected by a bushfire, showed substantial increases in dissolved nitrogen (1.2 times), dissolved phosphorus (2 times) and DOC (1.5 times) compared to reference lakes. These increases were accompanied by a 9% reduction in mean pH. Similarly, Sherson *et al.* (2015)²⁵ studied nutrient dynamics in a stream following a bushfire (the Las Conchas bushfire in June 2011 that burned approximately 36% of the contributing watershed) and during several precipitation events using water quality sensors. They found significant increases in dissolved NO₃-N (6 times the background levels), dissolved phosphate (100 times the background levels), specific conductance (5 times the background levels) and turbidity (>100 times the background levels). These increases were noted to be accompanied by reductions in dissolved oxygen and pH, as reported by Sherson *et al.* (2015)²⁵. The study highlighted the rapid rates of change in water quality during flow events of short duration following bushfires. White *et al.* (2006)²⁷ reported increases (up to thirty times) in turbidity, iron and manganese in the upper catchment storages following a bushfire. These were problematic to water supply, resulting in the construction of a drinking water treatment plant (DWTP) for turbidity removal. White *et al.* (2006) also reported the impact of natural revegetation in the catchment on the improvements in water quality.

The environmental risks associated with extreme climate events demonstrate the need for high-frequency water quality monitoring of vulnerable drinking water sources.^{4,28} Online, real-time monitoring systems may assist in providing treatment and disinfection for impacted influent surface water in water treatment plants (WTPs).²⁹ Hurst *et al.* (2004)²⁹ proposed the integration of an online DOC analyser with a model-based coagulation control system to determine the required coagulant



dose for waters varying in DOC variation circumstances during extreme events. Moreover, fluorescent DOM (fDOM) sensors can detect fluorescence signals from the fluorophore portion of the DOM pool. The fDOM sensor signal can be used as either semi-quantitative (as a surrogate parameter to estimate DOC concentration) or semi-qualitative (DOM pool optical and coagulability characteristics) information.^{30,31} Field fDOM measurements are prone to interference from signal attenuation/quenching factors, such as temperature, turbidity, pH, and the inner filter. Interference with the fDOM signal can be exacerbated in water sources impacted by extreme climate events, though this aspect requires further investigation.

The impacts of bushfires and cyclones on water quality have been widely investigated. However, investigations have focused predominantly on the discharge of sediments, ash, turbidity, dissolved metals, DOM concentration, and nutrients (N and P) discharged from catchments into waterways.^{2,9,10,25,32,33} This study used multi-analytical techniques to investigate DOM concentration and characteristics, Chlorophyll-a (Chla) and turbidity following bushfires and a cyclone. This is reported for (1) river waters of two Australian catchments following the extreme bushfire events in the summer of 2019/2020 over a period of approximately two years, and (2) two reservoirs impacted by Tropical Cyclone Uesi in 2020 over a period of approximately one year. We also investigated the application of the EXO fDOM sensor signal for monitoring DOM concentration and characteristics, in comparison with the DOC and A_{254} indices.

2 Materials and methods

2.1 Study area

Water samples ($n = 51$) were collected between September 2019 and November 2021 from four water sources in Australia: 25 from the Orroral River (P01) site located in the Australian Capital Territory (ACT), 6 from the Leslie Harrison Dam (P03) and Herring Lagoon (P05) sites located in Queensland (QLD) and 20 from the Middle River (P11) site located in South Australia (SA) – see Fig. 1 and Table A1, ESI.† The water sampling points were coded, as detailed above and as previously

reported.^{34,35} Of the catchments investigated, only the Orroral River (P01) is not used for drinking water supplies.

The P03 and P05 sites experienced Tropical Cyclone Uesi on February 4–14, 2020, QLD^{36,37} – see Table S1, ESI.† The P01 and P11 sites experienced severe bushfires from December 2019 to February 2020 in Canberra and SA.³⁸ Liu *et al.* (2022)³⁹ reported that more than 30% of Kangaroo Island (mainly the western region) burned during a bushfire that occurred between October 2019 and February 2020. Another severe bushfire burned 80% of the Namadgi National Park (Canberra, Australia) area.⁴⁰ Cyclone Uesi caused flooding due to storms with up to 220 mm of heavy rain in south-east Queensland.^{37,41}

2.2 Water samples and water quality analyses

Water samples were collected soon after bushfires (coded as P01_01 and P11_01), and for almost two years thereafter (coded as P01_02 to P01_25 and P11_02 to P11_20) – see Fig. 2 and Table S1, ESI.† For the cyclone – a water sample was collected before the event at site P03 (coded as P03_01), two samples were collected soon after the event (coded as P03_02 and P05_01), and three follow up samples were collected for almost one year thereafter (coded as P03_03, P05_02 and P05_03) – see Fig. 2 and Table S1, ESI.†

Multi-analytical techniques – including commonly used and advanced analytical techniques – were used to investigate a wide range of water quality parameters in water samples. Parameters included DOC, A_{254} , colour, EXO fDOM_s (fluorescence excitation/emission [ex/em] wavelengths of $[365 \pm 5]/[480 \pm 40]$ nm, YSI, Yellow Springs, OH, USA), turbidity, and Chla (using a fluorescence sensor with an ex/em wavelength of $[470 \pm 15]/[685 \pm 20]$ nm, Xylem Analytics, USA⁴²) calibrated using 625 μg per L rhodamine WT dye (66 μg per L Chla), high pressure size exclusion chromatography (HPSEC) with a UVA₂₆₀ detector (HPSEC-UVA₂₆₀) and HPSEC with a fluorescence-humic detector at an ex/em wavelength of 320/430 (HPSEC-Fl_Hu). The details of the acquisition of some of these water quality parameters (including DOC, A_{254} , colour, EXO fDOM_s, and turbidity) have been previously reported.⁴³ EXO fDOM_s were corrected for turbidity, temperature and pH impacts (referred to as EXO fDOM_{s,III}), and EXO fDOM_{s,III} were corrected for the inner filter effect (IFE) impact (referred to as EXO fDOM_{s,IV}) following the correction approach presented in Daraei *et al.* (2023).³⁴ An alkalinity test was conducted for the samples using the titration method.⁴⁴

Specific optical characteristics of DOM were calculated for each sample, namely specific UVA₂₅₄ ($\text{SUVA}_{254} = \text{UVA}_{254}/\text{DOC}$), specific colour (Sp. Colour = Colour/DOC), and specific fDOM_s (Sp. fDOM = fDOM_s/DOC). The UV-vis absorbance spectra were collected and analysed as described in Daraei *et al.* (2023).⁴⁵

For a limited number of samples (mostly collected immediately after the events), several additional water quality parameters were acquired. These included total nitrogen (TN), total nitrate plus nitrite, total Kjeldahl nitrogen (TKN), total phosphorus (TP), total organic carbon (TOC), heavy metals, and alkalinity. These data were obtained through a NATA accredited (<https://nata.com.au/>) laboratory (AWQC, SA Water).

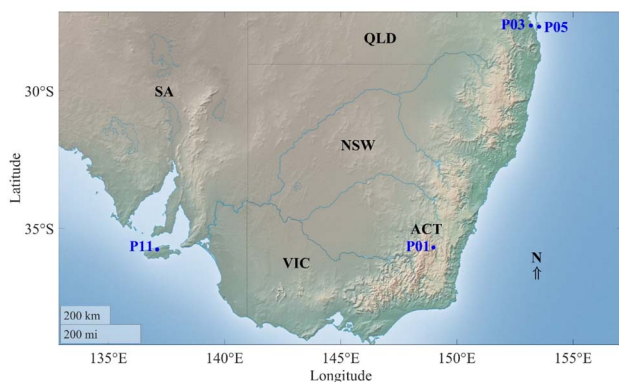


Fig. 1 Water sampling locations – Orroral River (P01), Leslie Harrison Dam (P03), Herring Lagoon (P05) and Middle River (P11).



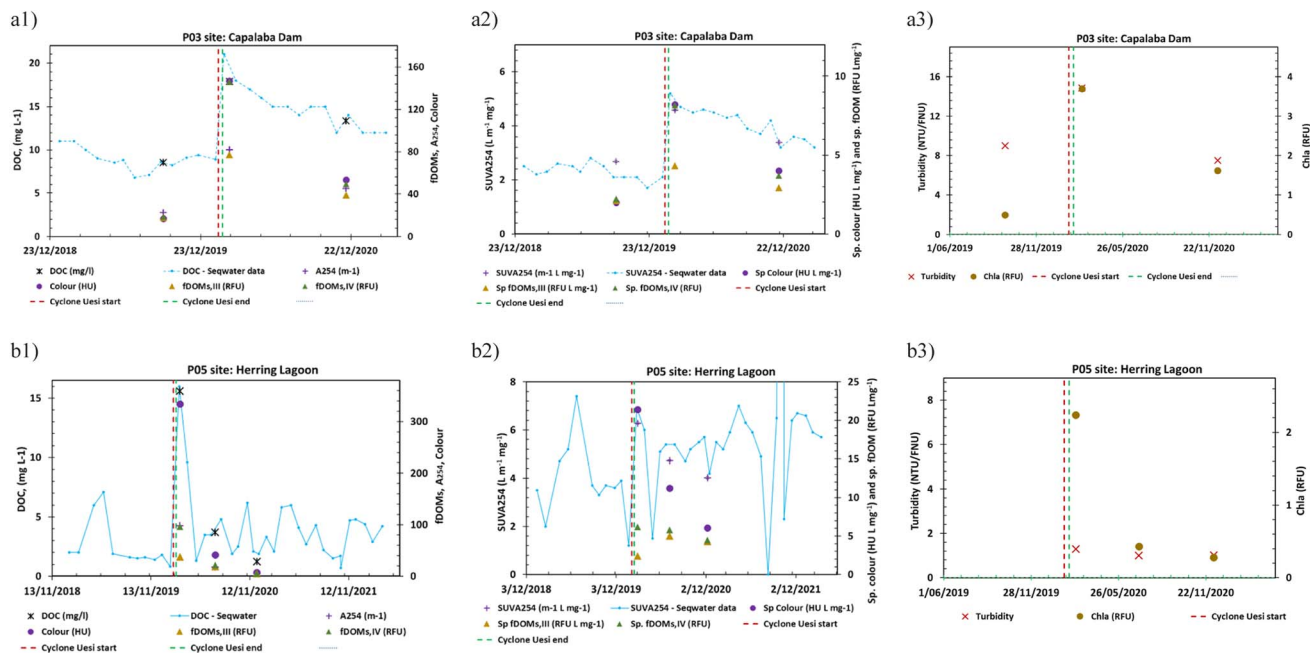


Fig. 2 Chronological variations of (1) DOM concentration (measured as DOC, A₂₅₄, fDOM_{s,III}, fDOM_{s,IV} and colour), (2) DOM optical characteristics (measured as SUVA₂₅₄, Sp. fDOM_{s,III}, Sp. fDOM_{s,IV} and Sp. colour) and (3) turbidity and Chla in (a) Leslie Harrison Dam, P03 and (b) Herring Lagoon, P05 surface water sources. *Seqwater (South-East Queensland water authority, Queensland, Australia) recorded DOC and SUVA₂₅₄ data with higher chronological resolution have been added to the diagrams. †Full-page size graphs are provided in the ESI.†

The DOM coagulability, which is defined as the removable DOC percentage by the enhanced coagulation process, and DOM removal rate, which is defined by Daraei *et al.* (2023)³⁵ as the maximum DOC removal rate per 100 mg alum as Al₂(SO₄)₃ · 18H₂O during the enhanced coagulation process and in the lower range of applied alum doses (the range with the highest efficiency for removing DOM) were attained using jar test data as previously reported in details in ref. 35 and 45 studies.

2.3 Advanced DOM characteristics and data analysis

Fluorescence excitation-emission matrices (fEEMs, *i.e.* 3D spectra) and HPSEC data (*i.e.* 2D chromatograms) were collected and analysed as described in Daraei *et al.* (2023).⁴⁵ These were obtained using an LS55 (PerkinElmer, USA) benchtop fluorometer. fEEMs were corrected before any further data analysis, using a script developed in MATLAB software (according to Murphy *et al.* (2013)⁴⁶ instruction). Corrections included blank correction using a Milli-Q water fEEM, masking and smoothing primary and secondary Rayleigh peaks, IFE correction using UV-vis absorbance data, and converting signal intensity to a standardised Raman unit. Following this, fEEMs were analysed through visualisation of the 3D spectra, followed by determination of the biological fluorescence index (BIX, ranging from 0 to 1)⁴⁷ and humification fluorescence index (HIX, ranging from 0 to 1),⁴⁸ and PARAFAC analysis.⁴⁹ Details of the PARAFAC model development for the fEEMs of the water samples, the selection procedure for the number of model components, and the PARAFAC model statistical validation have been previously reported.⁴³

HPSEC with UV-detection (HPSEC-UVA₂₆₀) and humic fluorescence detection (HPSEC-FlHu, $\lambda_{ex/em} = 280/330$ nm) were performed according to the method of Chow *et al.* (2008)⁵⁰ to determine the apparent molecular weight (AMW) profile and weight-average molecular weight (WAMW). The HPSEC data were also analysed using five previously introduced molecular size fractions (area) according to the method described by de Oliveira *et al.* (2018)³¹ – see Table A2, ESI† for MW areas and corresponding fractions. The HPSEC data were analysed by a peak fitting technique as previously described in Daraei *et al.* (2023).⁴⁵ HPSEC-UVA₂₆₀ chromatograms were partitioned into seven components (peaks, *i.e.* HPSEC-UVA260-PF-P_x peaks) with peak centres corresponding to molecular weights (MWs) of ~457 Da (PF-P₁), 1000 Da (PF-P₂), 1698 Da (PF-P₃), 2454 Da (PF-P₄), 3162 Da (PF-P₅), 4169 Da (PF-P₆) and 36 308 Da (PF-P₇), respectively. The HPSEC-Fl-Hu chromatograms were partitioned into seven components (Peaks, *i.e.* HPSEC-Fl-Hu-PF-P_x peaks) with peak centres (MWs) of ~398 Da (PF-P₁), 776 Da (PF-P₂), 1140 Da (PF-P₃), 1841 Da (PF-P₄), 2690 Da (PF-P₅), 4169 Da (PF-P₆) and 70 280 Da (PF-P₇), respectively. For each sample, the peak centre, peak width and peak height were adjusted using the genetic algorithm (GA) function,³¹ and some differences in peak centres are therefore expected among the different samples.

2.4 Precision estimation

Using two water samples, the coefficient of variation (CV) of several key water quality analyses was less than 4.5% for all parameters (except true colour for one sample: CV = 12.1%), as shown in Table A3, ESI.†



3 Results and discussion

3.1 Chronological water quality index variations

The results of DOM concentration and characteristic (commonly used indices) analyses are presented in Fig. 2a₁ and b₁ (cyclone) and Fig. 3a₁ and b₁ (bushfires). These data were used for the calculation of SUVA₂₅₄ and specific colour (sp. colour) as common DOM characteristic parameters, as well as sp. fDOM_s parameter (*e.g.* YSI EXO fDOM sensor) and online TOC/DOC analysers. The results of the chronological tracking of turbidity and Chla are also presented in Fig. 2a₃ and b₃ (cyclone) and Fig. 3a₃ and b₃ (bushfires).

In most cases, there were similar correlations between DOC and its optical surrogate parameters (*i.e.* A₂₅₄, colour and corrected fDOM_s) as measures for DOM concentration (Table A4, ESI†). This indicates the potential of these optical parameters (signals) for site-specific tracking of DOM concentration, including the sites that experienced the cyclone or bushfires. However, there were different variation rates for these optical parameters (*i.e.* greater changes in colour than A₂₅₄ and fDOM signals; Fig. 2a₁ and b₁ [cyclone] and Fig. 3a₁ and b₁ [bushfires]). This might be due to the different DOM fractions and associated optical characteristics. In the case of Fig. 2a₁, colour changes are approximately two times greater than A₂₅₄, which may indicate a greater variation in the portion of the humic-like DOM fraction (more associated with the colour index) rather than the protein-like fraction in the total DOM concentration variation. In the case of the EXO fDOM signal, interfering factors such as turbidity and IFE may also cause these different

variation rates. Daraei *et al.* (2023)³⁴ previously reported a study on these interfering factors as well as a correction technique for such interferences.

Seqwater (South-East Queensland water authority, Queensland, Australia) also recorded DOC and SUVA₂₅₄ data which have been added to the data acquired in this study for the P03 and P05 sites (see Fig. 2). This combination of the data, with high temporal resolution and coverage can provide more reliable indicators of bushfire and cyclone effects on the surface water sources than the three-point based datasets we have recorded in our study.

3.1.1 DOM concentration and characteristics (commonly applied indices). High DOM concentrations with high SUVA₂₅₄ occurred immediately after Cyclone Uesi at the P03 and P05 sites. This was followed by a decrease in both DOM concentration and its chromophoric characteristics over time. This may be due to the fresh terrestrial origin of DOM transported to waters by heavy rains from the cyclone.⁵² In contrast, a high DOM concentration (up to ~31 mg L⁻¹) but a medium-low level of SUVA₂₅₄ (<4 L m⁻¹ mg⁻¹) was observed immediately following the bushfires (P01 and P11 sites). This was followed by a decrease in the DOM concentration and a slight increase in the SUVA₂₅₄ level over time. This could be due to the transport of DOM with a lower chromophoric characteristic from fire (combustion) or subsequent microbial growth conditions. Our post-fire DOM results are similar to those of Sherson *et al.* (2015)²⁵, who demonstrated an increase in DOM shortly following bushfire events. Chen *et al.* (2020)⁵³ reported lower coagulability for DOM sourced from the bushfire ash due to the

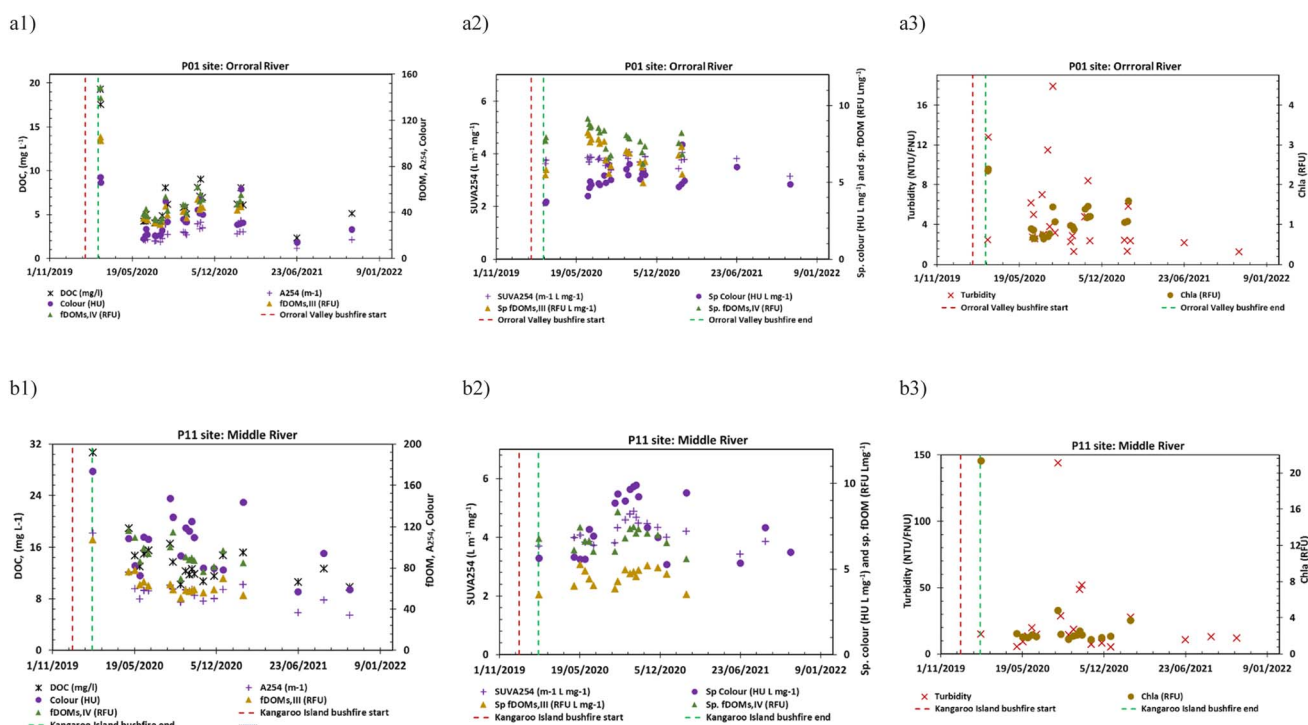


Fig. 3 Chronological variations of (1) DOM concentration (measured as DOC, A₂₅₄, fDOM_{s,III}, fDOM_{s,IV} and colour), (2) DOM optical characteristics (measured as SUVA₂₅₄, Sp. fDOM_{s,III}, Sp. fDOM_{s,IV} and Sp. colour) and (3) turbidity and Chla in (a) Orroral River, P01 and (b) Middle River, P11 surface water sources. *Full-page size graphs are provided in the ESI.†



loss of side chains and the decrease in DOM MW. The discharge of such materials can occur *via* (1) airborne ash and smoke, (2) precipitation events and (3) fire extinguishing activities during the bushfire and soon after the event.^{52,54,55} Aerosol emissions from bushfires can increase the atmospheric transport of macronutrients and bio-essential trace metals such as nitrogen and iron, respectively.⁵⁶ For the Middle River water sample collection site, a layer of ash was present on the surface of the water stream and surrounding soil area.

Xu *et al.* (2024)⁵⁷ also observed the emergence of recalcitrant dissolved black carbon (DBC), dissolved black nitrogen (DBN), and dissolved organic sulphur, alongside elevated levels of condensed aromatics, proteins, and unsaturated hydrocarbons, coupled with reduced lignin content in river waters post-wildfire occurrences. Their findings unveiled that DBC constituted 9.5–19.2% of DOC in these water bodies, potentially attributed to atmospheric deposition facilitated by wind dynamics. The Hu *et al.* (2024)⁵⁸ review also showed that the thermal effects of wildfire on DOM in watersheds likely impacts the DOM coagulability and reactivity to yield DBPs.

3.1.2 EXO fDOM signal. The results presented in Fig. 2 and 3 also indicated the potential of EXO fDOM sensors as field-deployable instruments to capture the timing and magnitude of climate event impacts on surface waters. The EXO fDOM signal results (without filtration for turbidity removal and with potential for online, real-time applications) showed high correlation (correlation coefficient up to 0.99, Table A4, ESI†) with DOC and UVA₂₅₄, which have been recorded after filtration for turbidity removal (see Fig. 2a₁ and b₁ [cyclone] and Fig. 3a₁ and b₁ [bushfires]).

The fDOM data reported here were obtained from samples collected for laboratory tests, ensuring consistency with other benchmark laboratory-based techniques, rather than from field deployment of the EXO fDOM probe and online data collection. Therefore, further assessments of the probe are necessary to evaluate its potential under online operation under field conditions for extended durations. To address this, we conducted a trial of online data collection from the raw water chamber of a water treatment plant operated by Seqwater in Queensland, drawing water from a reservoir. The trial involved a deployment period over four months (and ongoing), utilising a plastic fouling guard-equipped EXO sonde. Data were collected at ten-minute intervals, immediately following operation of an anti-fouling brush system to clean the probes' windows and tips for each reading. The EXO sonde was powered by four D-size batteries, providing approximately three months of continuous operation under these conditions. However, longer-term power supply options, such as a cable plugged into a power outlet, are available for the sonde in powered sites. Our results demonstrate high performance, reliable, and consistent data collection, even during and after a severe cyclonic event affecting the catchment, reservoir, and plant area. Additionally, the data exhibited strong correlation ($r > 0.95$) with discrete daily A₂₅₄ and monthly DOC data from the plant. The trial revealed the effectiveness of the combined anti-fouling system employed. Calibration checks confirmed the adequacy of monthly checks and indicated the need for approximately bi-

monthly recalibration for pH probes, while demonstrating the stability of the fDOM probe over more than four months of operation with less than 5% drift from calibration standards. Detailed quantitative findings from this continuous implementation stage will be provided in a forthcoming publication on the trial.

3.1.3 Chlorophyll-a (Chla). Middle River had the highest post-fire Chla of $\sim 85 \mu\text{g L}^{-1}$ (21.4 RFU), which was approximately four times greater than the average level of Chla concentrations in subsequent samples. This level of Chla can be associated with a 50% probability of exceeding microcystin health recommended concentrations.⁵⁹ The level of $\sim 85 \mu\text{g L}^{-1}$ of Chla can potentially be an indicator for exceeding the US-EPA recommendation based maximum allowable concentration (MAC) for microcystin in drinking water for children (*i.e.* 23 $\mu\text{g per L Chla}$ potentially indicative of 0.3 $\mu\text{g per L microcystin}$) and in drinking water (*i.e.* 68 $\mu\text{g per L Chla}$ potentially indicative of 1 $\mu\text{g per L microcystin}$), and US-EPA recommendation based MAC for microcystin in adult's drinking water (*i.e.* 84 $\mu\text{g per L Chla}$ potentially indicative of 1.6 $\mu\text{g per L microcystin}$). This highlights the necessity for specific water treatments at DWTPs for Chla and microcystin removals where these occur in source waters. Nonetheless, the maximum recorded Chla for the other sites and samples were $< \sim 23 \mu\text{g L}^{-1}$, which is below the guideline MAC levels.

The increase in Chla levels following cyclones has been described as relating to the increase in discharges of run-off water (containing nutrients) into surface waters, which can lead to the growth of phytoplankton populations.^{32,60} Growth of phytoplankton populations, which consume nutrients, acts as a natural stabilising mechanism and regulates nutrient levels in water sources.^{32,61} Peat *et al.* (2005)⁶² reported increased Chla levels in catchment stream flows following a bushfire.

3.1.4 Turbidity. Turbidity and Chla demonstrated similar trends (correlations) to each other over time at the sites studied (Fig. 2 and 3). Several studies previously reported an increase in turbidity and sediments in surface waters associated with either burned catchment areas following bushfires^{9,25,27,60} or extreme discharge events (*e.g.* following cyclones, typhoons, storms and extreme rainfalls).^{29,63–65} Rasul *et al.* (2013)³² reported increased Chla in a catchment affected by a cyclone.

3.1.5 Nutrients and alkalinity. For Orroral River and Middle River samples collected immediately after bushfires, the TN levels were 1.55 and 5.88 mg L^{-1} , total nitrate and nitrite were 0.207 and 0.055 mg L^{-1} N, TKN was 1.34 and 4.82 mg L^{-1} , TP was 0.18 and 0.31 mg L^{-1} , TOC was 18.2 and 32.6 mg L^{-1} and alkalinity was 144 and 173 mg L^{-1} , respectively. These values are in the range of previously reported studies of catchments impacted by bushfires.¹⁶ The high alkalinity values of samples collected immediately following bushfires may be due to the alkalinity of wood combustion ash.⁶⁶

3.1.6 Heavy metals. Post-bushfire and cyclone concentrations of heavy metals and other elements (*i.e.* Ag, As, B, Ba, Be, Ca, Cd, Co, Cr, Cu, Fe, K, Mg, Mn, Na, Ni, P, Pb, S, Sb, Se, Si, Sr, Ti, V, Zn and U) were all below Australian Drinking Water Guidelines (Table A5, ESI†). Minor effects of fire on streamflow ionic composition have previously been reported.^{67,68} This was



explained by (1) the ash element adsorption on the humic and clay complexes in the catchment area prior to discharge into the streams, and (2) the adsorbed elements in the catchment area soil being consumed by the vigorously regrowing vegetation.⁶⁷ Johnston and Maher (2022)²⁴ reported general increases in TSS, DOC, electrical conductivity and several ions (*i.e.* HCO_3^- , SO_4^{2-} , Ca^{2+} , K^+ , PO_4^{3-} and NO_3^-) immediately after a severe drought and extensive fires followed by flooding in a large coastal catchment (Macleay River, Australia).

Uzun *et al.* (2020)⁶⁹ compared the water quality impacts of two California bushfires (Rocky and Wragg Fires, 2015) to an unburned reference watershed. They reported that the concentrations of DOC, DON, ammonium ($\text{NH}_4^+/\text{NH}_3$), and SUVA₂₅₄ levels were significantly higher during the first year ($67 \pm 40\%$, $418 \pm 125\%$, $192 \pm 120\%$, and $31 \pm 17\%$, respectively).⁶⁹ Similarly, Chen *et al.* (2018)⁶⁵ reported major increases in both dissolved nutrients and suspended particulates during storms in the Jurong River Estuary, southeast China. These increases were related to river inflows, surrounding runoff, pore water supply, and nutrients desorbed from scoured sediment.⁶⁵

3.2 fEEM data

Both HIX and BIX data demonstrated potential for capturing the variations in DOM characteristics following the cyclone (Fig. 4). HIX values in the samples collected immediately

following the cyclone event were high for both the P03 and P05 sites (0.97 and 0.98, respectively), and slightly decreased in subsequent samples (0.94 and 0.85, respectively). In contrast, BIX was low at these sites immediately after the cyclone event (0.41 and 0.30, respectively), and increased in the subsequent samples (0.53 and 0.48). Heavy rainfall and subsequent surface water flows associated with the cyclone event may increase the terrestrial fraction of humic substances in the DOM pool immediately after the cyclone, resulting in observed increases in HIX and SUVA as well as the observed decreases in BIX.

Xu *et al.* (2016)⁷⁰ reported similar increases in HIX and SUVA₂₅₄ after a typhoon in the South Tiaoxi River, China. They reported increases in both HIX (0.75 to 0.79) and DOC (3.17 to 3.64 mg L^{-1}), though a decrease in BIX (0.83 to 0.75), which is consistent with the results of the current study. Similar increases in HIX (0.74 to 0.91) and DOC (2.0 to 5.1 mg L^{-1}) and decreases in BIX (0.81 to 0.79) were reported by Begum *et al.* (2023)⁷¹ following a storm affecting an agricultural watershed in South Korea.

As detailed in Daraei *et al.* (2023),⁴⁵ the humic-like components PAR-C₁, PAR-C₂ and PAR-C₄ are among the most common fluorophores found in freshwaters, with a large number of matches in the OpenFluor database library.⁷² These are associated with high molecular weight and aromatic terrestrial compounds.⁷² PAR-C₃ and PAR-C₅ components are

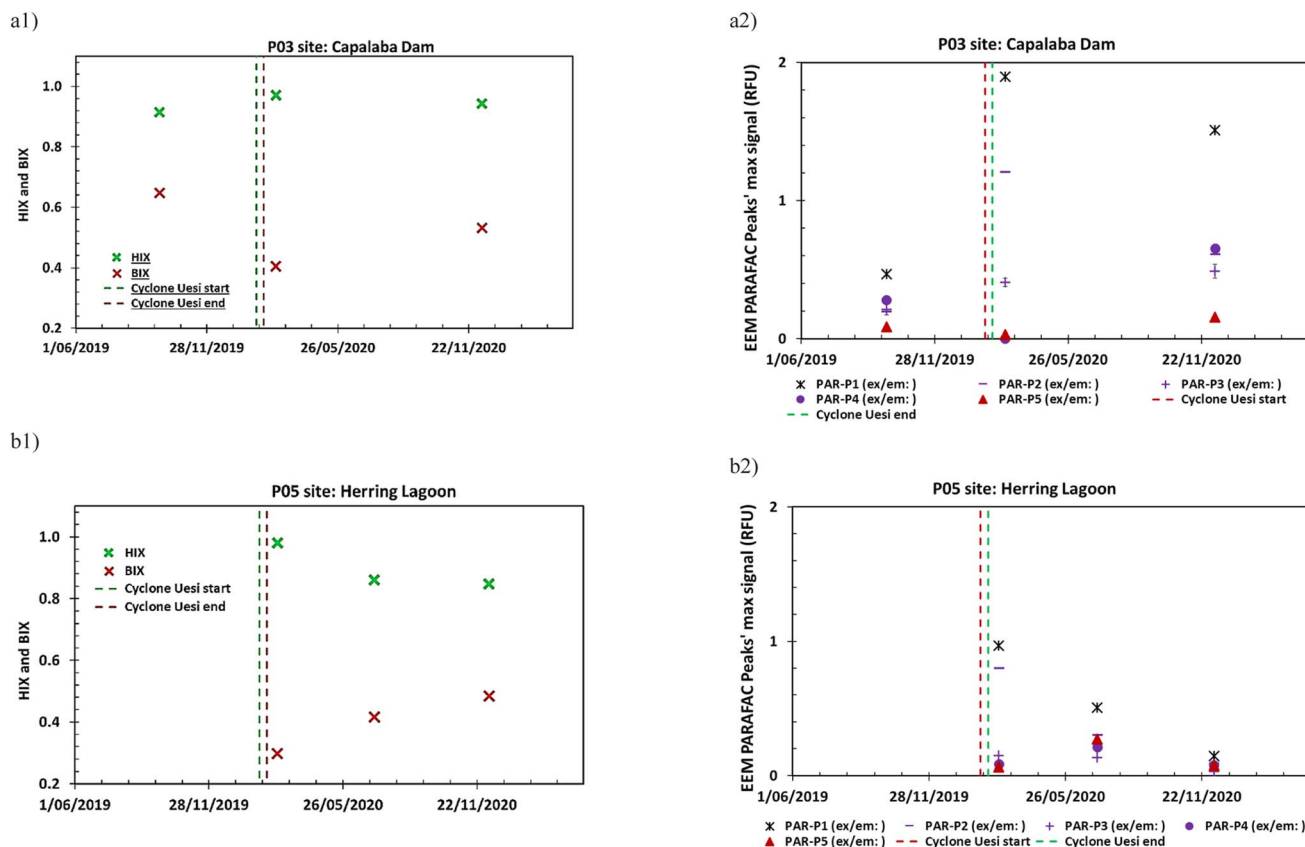


Fig. 4 Chronological variations of (1) FRI based technique indices (HIX and BIX) and (2) PARAFAC based indices (PAR-P1 to PAR-P5) extracted from fEEM data of (a) Leslie Harrison Dam, P03 and (b) Herring Lagoon, P05 surface water sources. *Full-page size graphs are provided in the ESI.†



classified as microbial protein-like components related to the production of DOM within aquatic ecosystems.⁴³ The PARAFAC results for the P03 and P05 sites were used to provide a new ratio index (sum of protein-like peaks to sum of humic like peaks, *i.e.* $[(\text{PAR-C}_3 + \text{PAR-C}_5)]/[\text{PAR-C}_1 + \text{PAR-C}_2 + \text{PAR-C}_4]$) which can be similar in nature to the BIX index and likely more reliable due to the robustness of the PARAFAC technique. For sites that experienced the cyclone, the variations of this new ratio index are comparable to BIX index variations. The new index varied from 0.14 (soon after the event) to 0.23 (in the subsequent samples) at site P03 (and from 0.11 to >0.35 at site P05). However, the new index would benefit from further validation, although this is beyond the scope of this study. The PARAFAC components may provide more information on DOM characteristics by being linked with DOM fractions and molecular level characteristic data, though further investigation is required. Nonetheless, the results presented here can provide a basis for future research.

Xu *et al.* (2016)⁷⁰ reported a four-component PARAFAC model, including two protein-like components and two humic-like components in samples collected before and after a typhoon-induced heavy rainfall event in the South Tiaoxi River, China. Begum *et al.* (2023)⁷¹ reported a three-component PARAFAC model for DOM in the samples they studied. They reported an increase in the mean protein-like portion of the DOM following the storm, which is not consistent with our results, as well as a decrease in BIX. This could be due to the statistical uncertainty associated with their mean data, as mentioned in their study.

For sites that experienced bushfires, the BIX index demonstrates greater potential than HIX for capturing variation in

DOM characteristics (see Fig. 5). In contrast to the sites that experienced the cyclone, high BIX values were found in samples collected immediately after the bushfires, which subsequently decreased over time, with fluctuations likely associated with seasonal effects. This could be due to the discharge of DOM with chromophoric characteristics following bushfires (combustion products and fire extinguishing agents), which differ from those following cyclone events. The Hu *et al.* (2024)⁵⁸ review study shows that generally, the proportion of tyrosine-like and humic acid-like FRI showed a decreasing trend after wildfires. The HIX values are generally consistent, with small variations over time. Uzun *et al.* (2020)⁶⁹ and Johnston and Maher (2022)²⁴ reported an opposite HIX trend to what we observed in the current study (*i.e.* increasing soon after the bushfire and then decreasing over time). This might be due to the application of a different formula for the calculation of HIX,⁴⁸ which may be another reason for higher reliability of the BIX index for comparisons. However, as a PARAFAC-based alternative index to the BIX, the newly suggested index in our study (*i.e.* $[\text{PAR-C}_3 + \text{PAR-C}_5]/[\text{PAR-C}_1 + \text{PAR-C}_2 + \text{PAR-C}_4]$) varied from 0.34 (soon after the event) to <0.20 (in the subsequent samples) at site P01 and from 0.42 to <0.20 at site P11.

3.3 HPSEC data

Weight-averaged apparent molecular weights (WAMWs) were generally higher in the samples collected soon after the cyclone (within approximately two to three weeks, see Fig. 6) than subsequent samples (collected after approximately 10 months, see Fig. 6). This might be due to the presence of a larger amount

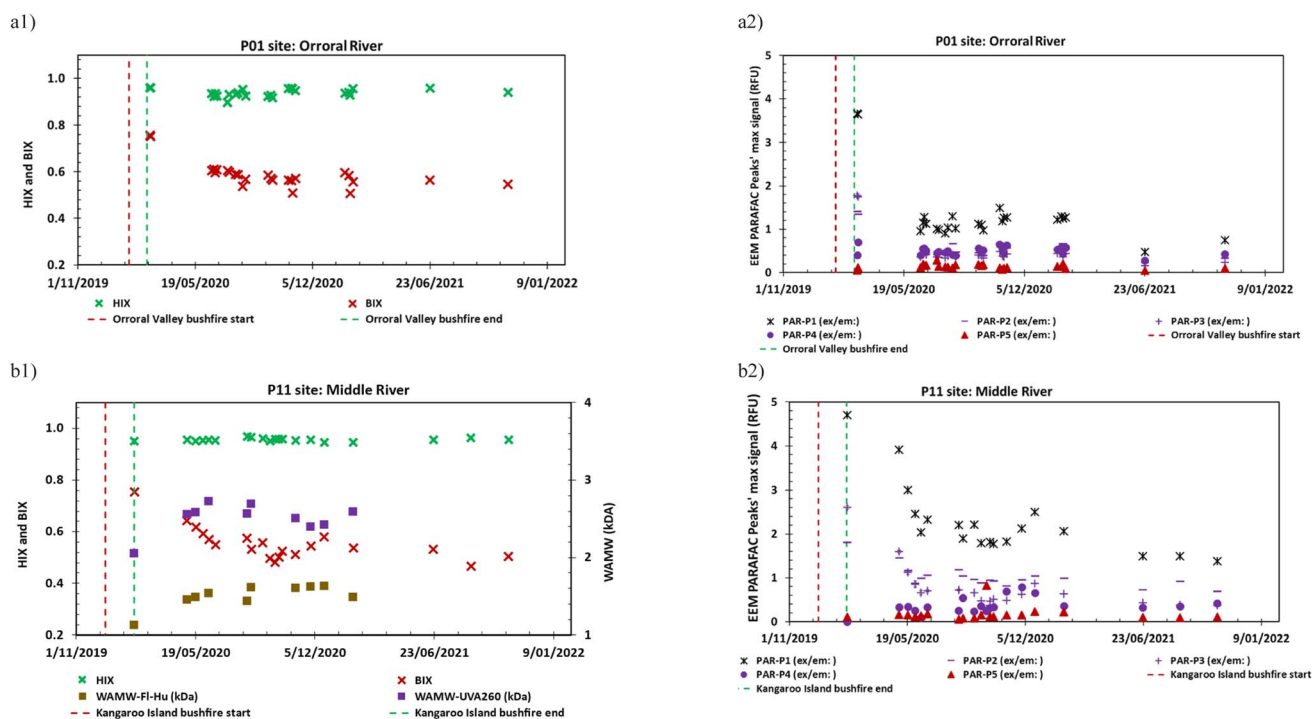


Fig. 5 Chronological variations of the (1) FRI based technique indices (HIX and BIX) and (2) PARAFAC based indices (PAR-P1 to PAR-P5) extracted from fEEM data of (a) Orroral River, P01 and (b) Middle River, P11 surface water sources. *Full-page size graphs are provided in the ESI.†



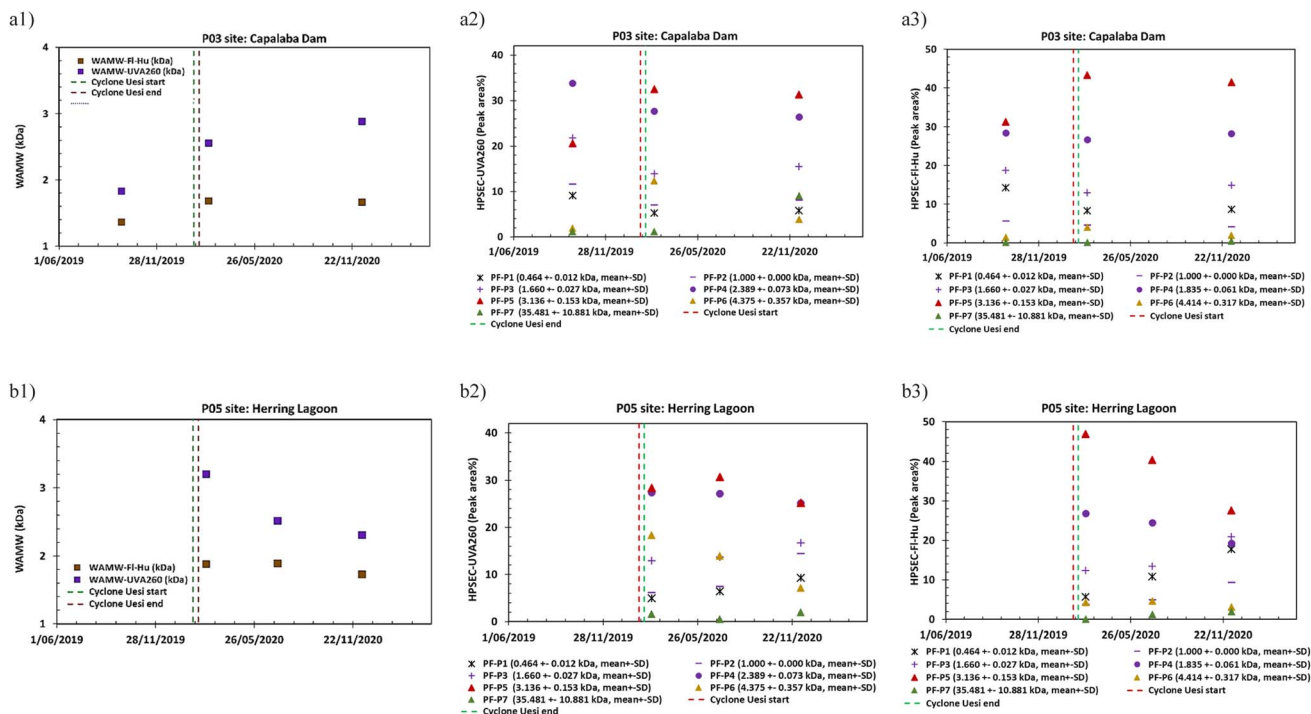


Fig. 6 Chronological variations of the (1) FRI based indices (HIX and BIX), (2) HPSEC-UVA₂₆₀ and (3) HPSEC-FI-Hu extracted from HPSEC data of (a) Leslie Harrison Dam, P03 and (b) Herring Lagoon, P05 surface water sources. *Full-page size graphs are provided in the ESI.†

of fresh terrestrial DOM with a greater molecular weight discharged into the surface waters during heavy precipitations during the tropical cyclone event.

In contrast, WAMWs were lower soon after bushfires (within approximately two to three weeks, see Fig. 7) than in the

subsequent samples (collected after approximately six months). This might be due to the presence of a larger amount of combustion products – including through the airborne smoke and particles,⁷³ with a lower molecular weight in the samples collected immediately after the bushfire.⁷⁴ In Japanese forested

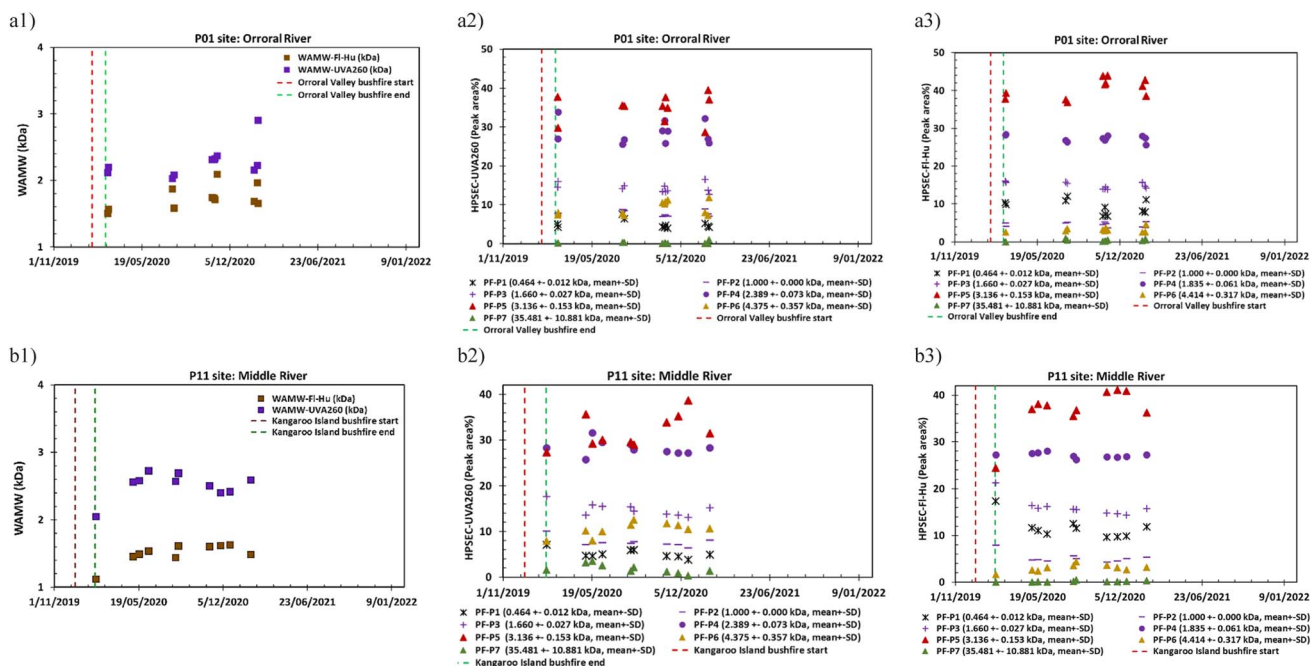


Fig. 7 Chronological variations of the (1) FRI based indices (HIX and BIX), (2) HPSEC-UVA₂₆₀ and (3) HPSEC-FI-Hu extracted from HPSEC data of (a) Orroral River, P01 and (b) Middle River, P11 surface water sources. *Full-page size graphs are provided in the ESI.†



catchments, Sazawa *et al.* (2018)⁷⁵ found that bushfires significantly changed the properties of fulvic acids and increased the concentration of DOC with low molecular weight in surface soil. Similarly, Muri *et al.* (2003)⁷⁶ identified the footprint of a historical bushfire (which had occurred in 1948) in northwest Slovenia through the polycyclic aromatic hydrocarbons and black carbon loads, low SUVA₂₅₄ and molecular weight,⁷⁷ in discharged sediments.

Seven MW-based components or peaks, called PF-P1 to PF-P7, were identified for the samples studied. The results are presented in Fig. 6a₂ and b₂ (cyclone) and Fig. 7a₂ and b₂ (bushfires; for HPSEC-UVA₂₆₀ data) and in Fig. 6a₃ and b₃ (cyclone) and Fig. 7a₃ and b₃ (bushfires; for HPSEC-Fl-Hu data) and Table A2, ESI.†

For sites that had experienced the cyclone, concentration of the coloured (chromophoric) dissolved organic matter (CDOM) and fDOM fractions with higher molecular weight (>~3 kDa) indicated a slight decrease after approximately 10 months (Fig. 6). This aligns with WAMW and Sp. colour trends. However, fractions with lower MW (<~3 kDa) increased slightly, which aligns with the BIX trend (Fig. 4). Li *et al.* (2003)⁷⁸ reported an increase in WAMW of Nagara River water samples from ~3.0 kDa (a few hours prior to the heavy rainfall starting) to 3.5 kDa (a few days after the heavy rainfall) during a typhoon event in Honsyu, Japan. This aligns with our results (Fig. 6a₁) whereby WAMW increased from ~1.8 kDa to ~2.6 kDa. Huang *et al.* (2016) reported higher DOC, A₂₅₄, SUVA₂₅₄, colour, specific colour and WAMW for the DOM pool in urban stormwater, which can occur following cyclones. Moreover, they suggested a likely need for a specific treatment strategy under such conditions.⁷⁹ Furthermore, as previously reported,^{35,45} up to 73% and 90% of the DOC in P03 and P05 samples, respectively, are removable (coagulable) by an enhanced coagulation process. Also, a high DOC removal rate of 12.6 and 18.5 mg DOC per 100 mg alum were reported for P03 and P05 samples, respectively.³⁵

For sites that experienced bushfires, the CDOM and fDOM fractions with higher molecular weight (>~3 kDa) indicated a slight increase with increasing time since fire, which aligns with the WAMW and Sp. colour trends. In contrast, the fractions with lower MW showed a slight decrease with time since fire, which aligns with the BIX trend. This general increase in the humic substance indices could be due to increased discharge of soil leachate biomass into streams. This can happen under particular conditions of rapid revegetation of burned catchment areas following bushfires²⁷ and using minerals available in the ash trapped in the soil of the burned catchment area,^{24,67} increased catchment water yield²⁴ and soil erosion.⁸⁰ This can increase colour, humic index, and WAMW in corresponding streams and water sources.

Similar observations have been previously reported in several studies. Begum *et al.* (2022)⁸¹ studied four DOM sources (*i.e.* field soil, litter, reed and manure), while Begum *et al.* (2023)⁷¹ studied five further DOM sources (*i.e.* soil, leaves, compost, a sewage plant effluent and manure). In both studies, they reported that the highest HIX level was from soil-sourced DOM. Shams (2018)⁸² conducted a comprehensive two year

study on DOM concentration and characteristics and their relationships to trihalomethane formation potential (THMFP) and haloacetic acid formation potential (HAAFP) following a severe bushfire on the eastern slopes of the Rocky Mountains in southwestern Alberta, Canada. Shams (2018)⁸² reported increases in DOC concentration (from <2 mg L⁻¹ to >6 mg L⁻¹), DOM hydrophobicity and disinfection by-product formation potential (DBPFP) following bushfires (and even more following salvage-logging following the fire) and during high discharge events in headwater streams. Shams (2018)⁸² suggested challenges associated with DOM treatability and several detrimental consequences (*e.g.* carbonaceous DBPFP, coagulant demand and membrane fouling) for DWTP intake waters severely impacted by bushfires. In our study, only 62% and 52% of the DOC concentrations in samples collected from sites P01 and P11, respectively, soon after the bushfire events could be removed through coagulation processes, suggesting lower coagulability than DOC content in sites impacted by the cyclone.^{35,45} Furthermore, a lower DOC removal rate of 9.0 and 8.4 mg DOC per 100 mg alum were observed, as reported in Daraei *et al.* (2023).

Our study was conceived after the extreme events of the 2019/2020 bushfires and the 2020 Tropical Cyclone Uesi had occurred, initially in the context of acquiring representative samples of surface waters following such events for a better understanding of changes in the concentrations and characteristics of DOM. This will contribute towards an improved understanding of the impacts of such extreme climate events on the ecological health of fresh surface waters and water resources used for drinking and other purposes. Subsequently, improved predictive capability for the treatment of surface waters following such extreme events for securing drinking water supply was investigated, as reported by Daraei *et al.* (2021).⁸³ Hohner *et al.* (2017)⁸⁴ concluded that the greatest treatment challenges were likely to be immediately after a bushfire and subsequent flow events. However, they suggested a longer-term investigation of the duration of the possible post-fire effects on water treatment process efficiency.

4 Conclusion

This study monitored water quality parameters following bushfires and a cyclone by applying both routine and advanced analytical techniques. The study generated novel insights into potential impacts of these events on surface water quality and the timeframe and processes of subsequent water quality stabilisation. Observed high DOM concentrations (measured as DOC and A₂₅₄) soon after the extreme events (up to >30 mg L⁻¹) can influence the treatment process efficiency and costs at DWTPs. High DOM concentrations, colour and turbidity can also influence aquatic ecological health. A useful feature of this study was chronological tracking of DOC concentration over a relatively long period of 1–2 years, which revealed a general decreasing trend of DOC concentration, likely due to dilution following precipitation events. However, the DOM present in the surface water associated with the cyclone event had different characteristics (SUVA₂₅₄, BIX, WAMW, DOC



coagulability and removal rate) than the DOM present in the surface water associated with bushfire events. The mostly aligned trends exhibited by EXO fDOM_s with DOC and A₂₅₄ indicate the potential benefit of using fDOM field-deployable sensors for continuous/real-time DOM monitoring and water quality determination for freshwater at sites that have experienced extreme climate events.

Conflicts of interest

The authors declared that they had no conflicts of interest with respect to their authorship or the publication of this article.

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