Spatial and temporal variability in *Phormidium* mats and associated anatoxin-a and homoanatoxin-a in two New Zealand rivers

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**ABSTRACT:** Toxic benthic cyanobacterial proliferations in freshwater are becoming increasingly prevalent, and associated animal poisonings are being reported with greater regularity. Despite this, few studies have investigated spatial and temporal variations in freshwater mat-forming *Cyanobacteria* and their associated toxin production. Some sections of the Hutt and Wainuiomata rivers (lower North Island, New Zealand) contain extensive *Phormidium* sp. proliferations that sometimes produce anatoxin-a (ATX) and homoanatoxin-a (HTX). The percentage coverage of *Phormidium* sp. was greater in the summer months and was correlated with warmer water temperatures and stable river flows. Flows in excess of 3 times the mean resulted in the removal of *Phormidium* mats. There was no correlation between the presence/absence of *Phormidium* mats and water-soluble nutrients. The presence and concentration of ATX and/or HTX and their degradation products, dihydroanatoxin-a and dihydrohomoanatoxin-a, was highly variable across all sites and over time. Anatoxin-a and HTX occurrence was restricted to periods of warm water temperatures (above 13.4°C) and below average river flows.

**KEY WORDS:** Anatoxin · Benthic *Cyanobacteria* · Nitrogen · *Phormidium* · Phosphorus · River flow · Temperature

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**INTRODUCTION**

Toxic cyanobacterial proliferations causing human and animal poisonings and fatalities have been documented in fresh and brackish waters worldwide. These events have commonly been linked to ingestion of toxins produced by planktonic *Cyanobacteria* (Negri et al. 1995, Kuiper-Goodman et al. 1999, Saker et al. 1999, Azevedo et al. 2002). However, animal toxicosis linked to benthic *Cyanobacteria* has increased recently (Mez et al. 1997, Hamill 2001, Gugger et al. 2005, Cadel-Six et al. 2007, Wood et al. 2010). The physical, chemical, and biological parameters leading to planktonic cyanobacterial blooms have been extensively studied (e.g. Oliver & Gnaf 2000), as have variables regulating toxin production by planktonic species (see Sivonen & Jones 1999 for review). In contrast, research on benthic *Cyanobacteria* has been initiated largely in response to animal poisonings and has focused on characterising the causative cyanobacterium and toxin and the symptoms in affected organisms (e.g. Cadel-Six et al. 2007, Wood et al. 2010). There is a limited understanding of mechanisms leading to benthic cyanobacterial proliferations and the influences of environmental variables on regulating toxin production of benthic species.

Benthic *Cyanobacteria* produce most of the known cyanotoxins, e.g. microcystins (Mez et al. 1997, Wood et al. 2010), saxitoxins (Carmichael et al. 1997), and cylindrospermopsins (Seifert et al. 2007). In New
Zealand, anatoxin-a (ATX) and homoanatoxin-a (HTX) are the most commonly detected cyanotoxins produced by benthic species (Wood et al. 2007, Heath et al. 2010). These are powerful neuromuscular blocking agents acting through the nicotinic acetylcholine receptor. In affected animals they can cause convulsions, coma, rigors, cyanosis, limb twitching, hypersalivation, and/or death. Research on the regulation of ATX in planktonic species has revealed that toxin production varies among species and with different physicochemical factors, e.g. temperature, light, and phosphorus (Rapala et al. 1993, Rapala & Sivonen 1998). Few studies have investigated ATX regulation and production in benthic mat-forming Cyanobacteria, despite the evident risk posed to human and animal health.

In November 2005, at least 5 dogs died rapidly after contact with water from the Hutt River (lower North Island, New Zealand). Dense mats of Phormidium autumnale were found in the river, and ATX and HTX were identified in the mats and dog stomach contents (Wood et al. 2007). Increased monitoring of Cyanobacteria in subsequent summers identified extensive coverage of Phormidium throughout the middle and lower reaches of the river, and further Cyanobacteria related dog deaths were reported. Phormidium mats were also detected in other rivers in the region (Milne & Watts 2006, Wood et al. 2007), including the Wainuiomata River (Fig. 1).

In this study, 6 sites along the Hutt River and its tributaries, and 2 sites on the Wainuiomata River were surveyed and sampled for 1 yr. Surveying and sampling was undertaken fortnightly during periods of flushing flows or weekly when river flow was more stable. Samples were collected for species identification and ATX and HTX analysis. Monthly water samples were taken for nutrient analysis, and river flow and temperature were logged continuously at different locations along each river. Percentage cover and community composition data were used in concert with physicochemical measurements to elucidate parameters correlated with cyanobacterial mat proliferation and ATX and HTX production.

**MATERIALS AND METHODS**

**Site descriptions.** The Hutt River stretches 54 km south through the Hutt Valley and discharges into Wellington Harbour (Fig. 1). It runs through several populated areas and has over 1 million recreational visits each year. In its upper reaches the river is used to provide the wider Wellington metropolitan area with up to 40% of its potable water (Wood et al. 2007).

Six sites were sampled on the Hutt River and its tributaries and 2 on the Wainuiomata River (Fig. 1). These were selected based on historical knowledge of high benthic cyanobacterial abundance at these locations. The sites were ranked based on a Water Quality Index (WQI) that relies on 6 variables: visual clarity (black disc), dissolved oxygen, dissolved reactive phosphorus, ammoniacal nitrogen, nitrate nitrogen, and faecal coliforms (Perrie 2007). The sites ranged in water quality between ‘poor’ and ‘excellent’ (Table 1). Coarse greywacke, a hard sedimentary rock, was the dominant substrate found at all 8 sites. The Sites 2, 3 and 4 were located on feeder tributaries to the Hutt River just above their confluence.

**Data and sample collection.** At each site, cyanobacterial mat and periphyton percentage substrate coverage was measured in three 1 m$^2$ quadrats, randomly positioned within a larger 10 m$^2$ quadrat, in a run (a swiftly flowing region of river with a relatively smooth surface). The data from the three 1 m$^2$ quadrats were averaged to estimate the percentage cover for each 10 m$^2$ quadrat. Samples were collected within each 10 m$^2$ quadrat by scraping mat material...
Heath et al.: Anatoxin spatial and temporal variability from rocks into a sterile plastic screw-cap bottle (50 ml, Biolab, New Zealand). All samples were placed on ice for transport. On arrival at the laboratory, samples were frozen (−20°C) for toxin analysis or preserved using Lugol's Iodine for morphological identification. Continuous river flow and water temperature data were measured at sampling sites using either Campbell C107 or Unidata LM34 sensors. Continuous readings for Site 4 were obtained from a logger 10 km upstream and readings for Sites 5 and 6 from a logger positioned midway between the 2 sites. We calculated the average river flow over the 12 mo period for each river (Fa). This yearly average value was used to standardise the river flow data as F/Fa (times above average), where F is the river flow at the time of sampling. F/Fa was used as a potential predictor for our statistical analysis. No water temperature was recorded for Sites 4, 7 and 8 and no river flow for Site 8.

Water samples were collected monthly at Sites 1 to 7 to determine nitrate-N, nitrite-N, ammonia-N, total nitrogen, total kjedahl nitrogen, dissolved reactive phosphorus, and total phosphorus. No water samples were collected at Site 8. Samples for dissolved nutrients were filtered through 45 µm Whatman GF/C filters and frozen (−20°C) until further analyses.

Species identification. Subsamples of the preserved field samples were homogenized and allowed to settle in a cavity slide (1 ml), and species were identified under an Olympus (CKX41) inverted microscope. Identifications were made primarily by reference to Komárek & Anagnostidis (2005), McGregor (2007), and Biggs & Kilroy (2000).

Extraction and detection of anatoxin-a and homoanatoxin-a. Frozen mat material was lyophilized (FreeZone6, Labconco). Lyophilized material (100 mg) was resuspended in 10 ml of double distilled water (DDW) containing 0.1% formic acid and sonicated (Cole Parmer 8890, Biolab) for 15 min. Samples were centrifuged (4000 × g, 10 min). The pellet was re-extracted a second time using 5 ml DDW and supernatants combined.

All samples were analysed for ATX, HTX, and their degradation products: dihydroanatoxin (dhATX), dihydrohomoanatoxin (dhHTX), epoxyanatoxin-a (epATX), and epoxhomoanatoxin-a (epHTX), using liquid chromatography-mass spectrometry (LC-MS). Anatoxins were separated by Acquity uPLC (Waters Corp.) using a 50 × 1.0 mm Acquity BEH-C18 (1.7 µm) column (Waters Corp.). The mobile phase A (0.1% formic acid in water) and mobile phase B (0.1% formic acid in acetonitrile) were used at a flow of 0.3 ml min⁻¹, isocratic for 1 min at 100% A, followed by a gradient to 50% B over 2 min. Injection volume was 5 µl. The Quattro Premier XE mass spectrometer (Waters-Micromass) was operated in ESI+ mode with capillary voltage 0.5 kV, desolvation gas 900 l h⁻¹, 400°C, cone gas 200 l h⁻¹ and
cone voltage 25 V. Quantitative analysis was by multiple reaction monitoring (MRM) using MS-MS channels set up for ATX (166.15 > 149.1; retention time (rt) 1.0 min), HTX (180.2 > 163.15; rt ca. 1.9 min), dhATX (168.1 > 56; rt 0.9 min), and dhHTX (182.1 > 57; rt ca. 1.9 min). The instrument was calibrated with dilutions in 0.1% formic acid of authentic standards of ATX (A.G. Scientific).

Nutrient analyses. Filtered nutrients were analysed on a Lachat QuickChem FLOW Injection Analyser (FIA + 8000 Series, Zellweger Analytics) and a Konelab Aquakem 600 Discrete Analyser (Thermo Scientific). Nitrate-N was analysed using automated AZO dye colorimetry, and total oxidised nitrogen (nitrate-N + nitrite-N) were analysed using automated cadmium reduction following the methods of the American Public Health Association (Eaton et al. 2005). Total kjedahl nitrogen was digested in sulphuric acid and analysed using phenol/hypochlorite colorimetry (Eaton et al. 2005). Nitrate was calculated by subtracting nitrite-N from total oxidised nitrogen (nitrate-N + nitrite-N), and total nitrogen was calculated by the addition of total kjedahl nitrogen, nitrite-N, and nitrate-N. Total phosphorus was digested with acid persulphate before using ascorbic acid colorimetry (Eaton et al. 2005). Dissolved reactive phosphorus was analysed using molybdenum blue colorimetry (Eaton et al. 2005).

Statistics. Stepwise logistic regression was used to establish the best predictors of Phormidium mat coverage and ATX and/or HTX presence across all 8 sites. No physicochemical data were collected for Site 8, and therefore this site was excluded from the analysis. Predictors were considered significant at the 95% (p < 0.05) confidence interval. River flow and water temperature were found to be significant predictors. In all cases, binary predicative’s were improved by altering the default of 0.5 probability to a cut value that maximised the prevalence of the presences in the data. This decreased the probabilities of false negatives (i.e. Cyanobacteria predicted to be absent when in fact it was present).

For water temperature and river flow (m$^3$ s$^{-1}$), a 5-day average prior to sampling was correlated with each sampling point. Standardisation of river flow was achieved by dividing the 5-day average river flow by yearly average river flow. All data were analysed using SPSS Statistical Software, version 16.0.

RESULTS

Nutrients

Total nitrogen (TN) and total phosphorus (TP) were generally found at low levels throughout the course of this investigation (Table 1). The lowest nutrient loadings were observed at the head waters of both the Hutt (Site 1, TN 0.174 g m$^{-3}$, TP 0.009 g m$^{-3}$) and Wairarapa Mata (Site 7, TN 0.074 g m$^{-3}$, TP, 0.072 g m$^{-3}$) rivers, whilst the highest concentration were found in the lower reaches of the Hutt River (Site 6, TN 0.403 g m$^{-3}$, TP 0.018 g m$^{-3}$) and at Site 2, which had increased nitrogen and phosphorus levels (TN 0.802 g m$^{-3}$, TP 0.023 g m$^{-3}$). Site 7 had a low TN:TP ratio (2.6); the remaining sites all had ratios above 15, with Sites 2, 5 and 6 above 25 with ratios of 34.3, 26.9, and 27.4, respectively. The TN:TP ratios increased with distance down the river. A t-test found there was no significant difference between N:P ratios with and without the presence of Phormidium mats (p = 0.3350, df = 89).

Temporal and spatial variability in Phormidium mat abundance

Cyanobacterial mats were recorded at all 8 study sites. The dominant genus in these mats was Phormidium. Although genetic analysis was not conducted as part of this study, polyphasic analyses of samples collected simultaneously identified Phormidium autumnale as the dominant cyanobacterium in the mats (Heath et al. 2010). Species from the Pseudanabaenaceae family such as Leptolyngbya spp. and Pseudanabaena spp. were observed in low concentrations within the mats (Heath et al. 2010). Phormidium mats were generally most abundant in summer and less frequently observed in winter (between May and October 2008). This higher coverage coincided with periods of stable river flow and warmer water temperatures (Fig. 2).

Percentage Phormidium mat cover was often over 50% at 4 of the 6 Hutt River sites (Sites 2, 3, 5, and 6) during summer, with the highest coverage (70%) observed at Sites 2 (6 January 2008) and 5 (27 February 2008) (Fig. 2). At Site 1, Phormidium mats were observed from the initial sampling (6 December 2007) until the 6 January 2008 (Fig. 2a). Mats were not present again until late February 2008, when the maximum percentage cover was less than 10%. Site 2 had extensive Phormidium mat coverage at the time of initial sampling (peaking at 70%, 6 January 2008), this steadily decreased until there were no Phormidium mats observed (26 March 2008; Fig. 2b). Phormidium mat coverage (up to 25%) was recorded in May and June 2008 and then no mats were observed until November 2008. No Phormidium mats were observed at Site 3 in January and only minimal coverage (<15%) in February (Fig. 2c). In March 2008, Phormidium mat coverage peaked at 60%, and then occurred sporadically in April and May 2008 before consistent mats
Fig. 2 (above and following page). Percentage mat coverage (bars) and anatoxin-a, and homoanatoxin-a concentrations of *Phormidium* sp., as well as temperature and river flow (solid lines) between 1 December 2007 and 31 December 2008. Error bars represent standard error. (a) Te Marua (Site 1), (b) Mangaroa (Site 2), (c) Akatarawa (Site 3), (d) Whakatikei (Site 4), (e) Silverstream (Site 5), (f) Boulcott (Site 6), (g) Manuka (Site 7), (h) Richard Prouse Park (Site 8). Water temperature was not measured for Sites 4, 7 and 8; river flow was not measured at Site 8. ATX: anatoxin-a; HTX: homoaatoxin-a; dhATX: dihydroanatoxin-a; dhHTX: dihydrohomoanatoxin-a
were observed between September and December 2008 (Fig. 2c). Site 4 experienced peak Phormidium coverage in December 2007 (30%) and patchy or low (<10%) percentage cover between January and March 2008 (Fig. 2d). Low (<15%) Phormidium mat cover was observed once in April and once in June 2008 with no further mats recorded until December 2008. Consistent Phormidium mat coverage (<40%) was observed at Site 5 from December 2007 to the end of March 2008. After that, Phormidium mats (5%) were observed once in June 2008 and were not recorded again until November 2008 (Fig. 2e). Phormidium mat abundance at Site 6 peaked at 60% in December 2007 and early January 2008. Throughout the rest of the sampling period, abundance was patchy and low (<10%; Fig. 2f).

There was no Phormidium sp. mat coverage at Site 7 (Wainuiomata River) until February 2008. Consistent coverage was then observed until the end of March 2008, where it peaked at 15% (Fig. 2g). Phormidium coverage at Site 8 was low and patchy, with consistent coverage only observed from late January 2008 to early March 2008. No Phormidium mat coverage was observed after June 2008 (Fig. 2h).

**Predicting the presence of Phormidium mats**

Using stepwise logistic regression, river flow and water temperature were identified as the only significant physicochemical parameters (p < 0.05) correlated with the presence of Phormidium mats. Consequently, these were the only variables included in the logistic model (Fig. 3). At a cut value of 0.5, the model successfully predicted 80.9% of the observed results, correctly predicting 82.6% of the times where Phormidium mats were absent and 78.8% of the time they were present. Using this model, the probability of Phormidium mats (P) occurring for any given river flow and water temperature can be calculated (Fig. 3b). The model demonstrates that with a yearly average river flow (1 in the model: refer to Table 1 for yearly average river flows) and water temperatures of 15°C, the probability of Phormidium mats occurring is 31%. With a river flow value of 0.5 and water temperatures of 20°C, the probability is 71% (Fig. 3b).

**Spatial and temporal variation in anatoxin-a and homoanatoxin-a**

ATX and/or HTX were detected at all Hutt River sites, and concentrations were highly variable, both within and between sites (Fig. 2a–f). The highest concentrations of ATX (1.7 mg kg⁻¹), HTX (23.5 mg kg⁻¹), and dhATX (535 mg kg⁻¹) were all observed at Site 2 (13 February 2008), and the highest concentration of dhHTX (95.4 mg kg⁻¹) was detected at Site 6 (6 January 2008). Some temporal similarities were observed, with maximal ATX and/or HTX concentrations occurring at Sites 3 and 4 on 6 December 2007, at Sites 2, 5 and 6 on 6 January 2008 and Sites 1, 2 and 4 on 12 February 2008 (Fig. 2a-f). ATX was only detected at Sites 2 and 6, while HTX was observed at all sites, with the exception of Site 1. The
most common compound, dhATX, was found at all sites, while dhHTX was observed at all sites except Site 1. At all Hutt River sites degradation products (dhATX and dhHTX) were always found in much higher concentration than their parent compounds, and dhATX was consistently found in higher concentrations than dhHTX. ATX and/or HTX were only detected in the summer and spring, i.e. December 2007 to April 2008 (Fig. 2a–i).

At the Wainuiomata River, no anatoxins were detected at Site 8 except on 14 March when dhATX (0.5 mg kg⁻¹) was detected (Fig. 2h). In contrast ATX, HTX, and their degradation products were detected at Site 7 (Fig. 2g). The highest concentrations of ATX (3.4 mg kg⁻¹) and HTX (283 mg kg⁻¹) at this site were both recorded on 4 April 2008 (Fig. 2g). This was the only site where the parent toxins (ATX and HTX) were found in higher concentrations than their degradation products.

Rapid temporal changes in anatoxin concentrations were observed, despite there being little change in Phormidium sp. percentage cover. For example, at Site 2 the combined ATX, HTX, dhATX, and dhHTX concentration increased from 201 mg kg⁻¹ on 6 February 2008 to 620 mg kg⁻¹ on 13 February 2008, without any change in Phormidium mat coverage (Fig. 2b). At Site 7, on 13 February 2008, the combined ATX, HTX, dhATX, and dhHTX concentration was 129.8 mg kg⁻¹, but 1 wk later on 20 February, HTX was the only compound detected (0.6 mg kg⁻¹; Fig. 2g).

Factors contributing to anatoxin-a and homoanatoxin-a presence/absence

Logistic regression was used to determine the best predictors of ATX and/or HTX presence/absence when Phormidium mats were present. Water temperature and river flow were the only significant (p < 0.05) physicochemical predictors of toxin presence. The logistic model successfully predicted 72.1% of the observed results, correctly predicting 64.7% of the times when ATX and/or HTX were absent and 81.5% of the time when they were present. ATX and/or HTX were observed to occur in water temperatures >13.4°C. Furthermore, ATX and/or HTX were only detected when river flows were low (below half the yearly average) and stable. The logistic model shows that when river flow is average (1 in the model) and water temperature is 15°C, the probability of ATX and/or HTX being detected when mats are present is 28%. At a river flow value of 0.5 and a water temperature of 20°C the probability increases to 72% (Fig. 4).

DISCUSSION

Phormidium mat presence and physicochemical parameters

The data from the Hutt and Wainuiomata rivers demonstrate that Phormidium mat coverage can be
highly variable spatially and temporally within rivers. Despite this variation, a seasonal factor contributing to *Phormidium* mat coverage was apparent. Peaks in *Phormidium* mat coverage for all sites were observed in summer (December to March). Similar trends in total periphyton (including cyanobacterial mats) have been documented previously in New Zealand and have been attributed to a lower frequency of flushing river flows during summer (Biggs & Price 1987, Biggs & Close 1989, Biggs 1990, Clausen & Biggs 1997).

In this study, river flow and water temperature were found to be significant physicochemical parameters predicting *Phormidium* mat coverage. High river flows have the ability to pick up rocks and pebbles, and thereby remove attached mat material from substrates (Biggs & Close 1989). Water turbulence has been previously described as a major controlling factor for benthic *Cyanobacteria* in river systems (Biggs 1990, Milne & Watts 2006, Wood et al. 2007), lake systems (Johnson & Castenholz 2000, Dasey et al. 2005), and marine environments (Thacker & Paul 2001). The ability of river flow to flush *Phormidium* mats has led local government agencies in the Wellington region to adopt this measure as one of the factors used to predict *Phormidium* mat percentage coverage (Milne & Watts 2006). Two weeks without a river flow of 3 times the long term median is used as an early warning indicator of the strong likelihood of benthic *Phormidium* mat proliferation.

Water temperature was shown to have a strong positive relationship with *Phormidium* mat coverage. This was consistent with Biggs (1990) who has previously elucidated temperature as a significant parameter in *Lyngbya* (a cyanobacterium that can also grow on the benthos) proliferations in New Zealand rivers. Furthermore, this is consistent with previous planktonic *Cyanobacteria* research, where optimal temperatures maximise growth (Robarts & Zohary 1987, Song et al. 1998). The model developed in this study successfully predicted 80.9% of our field observations. The probability of *Phormidium* sp. being present dramatically increased when river flow was half the yearly average and water temperature above 14°C. These results support Greater Wellington Regional Council’s use of river flow as a predictor of the presence of mat-forming benthic *Phormidium* sp., and suggest that water temperature should be incorporated into the model for greater predicative power.

No other physicochemical parameters were significant predictors of *Phormidium* mat presence. Research on planktonic and benthic *Cyanobacteria* in marine, lake, and culture environments has shown nutrients such as phosphorus and nitrogen to be key parameters responsible for growth (Paerl 1996, Downing & Watson 2001, Vilalta et al. 2003). In New Zealand, benthic *Cyanobacteria* have previously been identified proliferating in sites with low nutrients (Biggs & Price 1987, Biggs 1990). Some *Cyanobacteria* from the LPP (*Lyngbya, Phormidium, Plectonema*) group possess the ability to fix nitrogen, therefore enhancing their ability to persist in nitrogen poor environments (Bergman et al. 2006, Pankratova et al. 1998). Low levels of DIN observed throughout this investigation may favour algal species with the ability to fix atmospheric nitrogen. At Site 2, expansive proliferations of chain diatom *Melosira varians* were observed; coincidently, this was the only site with elevated DIN levels. Those sites with the highest cyanobacterial coverage (2, 3, 5 and 6) were found to have high TN:TP ratios (above 20:1) providing evidence that nitrogen, rather than phosphorus, may be the nutrient limiting *Phormidium* mat growth (Borchardt 1996). It has been suggested that essential nutrients can be sourced from the geology of the surrounding catchment (Leland & Porter 2000), and therefore it is plausible that nutrients can be dissolved from the substratum and utilised. Biggs (1990) found that *Lyngbya* proliferations were highly associated with hard sedimentary rocks, which provide a potential source of phosphorus.

**Spatial and temporal variation in anatoxin-a and homoanatoxin-a production**

The presence/absence and concentrations of ATX and/or HTX varied between sites and over time. In New Zealand, some water managers use set thresholds of *Phormidium* sp. percentage coverage to issue health warnings (Wood et al. 2009). In this study, we found no correlation between the coverage of *Phormidium* mats and ATX and/or HTX concentrations. Previous investigation have shown that cyanobacterial mats consist of mixed toxic and non-toxic strains of *Phormidium autumnale* (Cadel-Six et al. 2007, Heath et al. 2010). It is possible that the variations in ATX and/or HTX concentrations observed in this study were due to changes in the relative abundances of toxic and non-toxic *Phormidium* strains in a mat, rather than changes in the amount of ATX and/or HTX produced. Previous studies have indicated that ATX quota’s only vary 1 to 7 fold as environmental factors (i.e., temperature) are manipulated (Sivonen & Jones 1999). Up and down-regulations in ATX production are therefore unlikely to explain the variability observed among sampling sites in this study.

River flow and water temperature were the only significant parameter in the logistic model used to predict ATX and/or HTX presence. The model successfully predicted 72.1% of our field observations, successfully predicting 81.5% of the times ATX and/or HTX was
present. Toxins were only detected in mats when temperatures were above 13.4°C and river flows were below half the yearly average. Research on some planktonic cyanobacteria has shown changes in ATX production with temperature; however, even at low temperatures, toxic genotypes still produce toxins (Rapala et al. 1993, Rapala & Sivonen 1998). Toxin production at all temperatures, coupled with the presence of non-toxin producing mats at low river flows and warm water temperatures, suggests that these variables do not have a significant effect on the regulation of ATX and/or HTX, rather that these conditions encourage the growth of ATX and/or HTX producing strains. There is currently no method to assess the proportion of ATX and/or HTX producing strains within mats. The recent discovery of a putative gene involved in ATX and/or HTX production (Cadel-Six et al. 2009, Méjean et al. 2010) provides the opportunity to use quantitative molecular techniques to monitor levels and expressions of this gene and will enable the quantification of toxic strains within a sample. This ultimately will assist in the development of predictive models aimed at providing early warning of cyanobacterial mat proliferation and toxin production.

Dihydro- degradation product concentrations were considerably higher than their parent compounds. Anatoxin is unstable especially in sunlight and at high pH, whereas the dihydro- degradation products are more stable (Smith & Lewis 1987). On some sampling occasions, degradation products were the only compounds detected. The detection of these compounds may provide information of previous or nearby toxic mats and these compounds should be monitored in routine cyanotoxin analysis. At Site 7, the concentrations of degradation products were considerably lower than that of their parent compounds. This site had extensive shading provided by overhanging vegetation. In the reduced light environment there may have been less degradation of ATX and/or HTX.

CONCLUSIONS

The results of this study revealed that percentage coverage and presence of Phormidium mats are spatially and temporally variable in the Hutt and Wainuiomata rivers. ATX and HTX concentrations were also found to vary among sampling times and sites. In contrast to previous reports, ATX and/or HTX concentrations did not correlate with Phormidium mat coverage. River flow was shown to control Phormidium mat coverage, and water temperature was identified as important for Phormidium mat growth and ATX and/or HTX production. Higher temperatures are likely to result in faster growth rates and these conditions may be more favourable for ATX and/or HTX producing strains. The ability of Phormidium to obtain essential nutrients for growth from sedimentary rock substrate and via nitrogen fixation may give it a competitive advantage in the mostly oligotrophic conditions in these rivers.

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