Development and Validation of an ICP-MS Method for the Detection of Platinum in the Lebanese Aquatic Environment

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Abstract: Lebanon is a Middle Eastern country located on the eastern Mediterranean coast. Compared to other countries in the MENA (Middle East North Africa) region, Lebanon is considered the richest in water resources. However, due to inadequate water management, Lebanese water resources are under stress. Water pollution is one of the main problems causing major concerns. The contamination of Lebanese surface water originates predominantly from the discharge of untreated municipal and industrial wastewater. With only a few studies investigating the level of water contamination in Lebanon, this research is the first to report the level of platinum in numerous rivers, three wastewater treatment plants, and two untreated sewage outfalls in coastal areas across Lebanon. To determine the platinum levels, an Inductively Coupled Plasma-Mass Spectrometry (ICP-MS) method was developed and validated in compliance with the ICH guidelines. The method demonstrated acceptable sensitivity with LOQ and LOD equal to 2.35 and 0.56 ng L⁻¹, respectively. The level of total platinum in the Lebanese water samples ranged between 22.44-53.32 ng L⁻¹. These concentrations were all above the baseline concentration of platinum in tap water indicated by WHO and aligned with previous studies in other countries. Although the baseline concentration of platinum in Lebanese water resources is unknown and the source of the total platinum detected in this study could not be identified, these preliminary findings could serve as a foundation for future research.

Keywords: water contamination; platinum; Lebanon; ICP-MS

1. Introduction

Lebanon is a Middle Eastern country bounded by the Mediterranean Sea with a coastline length of about 225 km. Despite its small area (10,452 km²), Lebanon is privileged with plentiful water resources [1]. These consist of rivers, springs, water storage and groundwater with a flow ranging between 2000 and 2700 million m³ per year.

With no clear and updated reports on the state and use of water resources, estimations and projections for water consumption and demand concluded that Lebanon would encounter chronic water shortages due to the poor management of the seasonal water imbalance. The factors promoting this imbalance are (1) minimal water storage capacity; (2) water loss into the sea; (3) increased water demand; and (4) insufficient water networks [1–3].

Another environmental stress affecting the Lebanese water environment is its deteriorated quality. This originates from anthropogenic sources and has previously resulted in numerous water-borne disease outbreaks such as cholera, typhoid fever, shigella infection and hepatitis A & E [4]. The water resources are mainly contaminated by untreated sewage from households and industries, agricultural runoff, dumpsite leachate and recreational activities [5].

Few academic studies and reports have assessed the contamination levels in the water, with only microbiological and heavy metals evaluations [6–8]. Despite examining...
the occurrence of several heavy metals in the Lebanese water resources and sediments, including Cu, Cd, Co, Pb, As, Ag and Hg [9,10], platinum has never been assessed.

Platinum (Pt) is a noble, scarce, and precious metal. Consequently, the presence of platinum in soil, water and air is at very low concentrations, often ng L\(^{-1}\). Naturally, platinum occurs in its native form (chemically uncombined) in alluvial deposits and rare ores such as Cooperite, mainly in South Africa, the Soviet Union, and the United States. Anthropogenically, platinum may arise from sources such as chemotherapeutic drugs, catalytic converters or industrial usage, and it is a potential contaminant in water discharges [11,12].

Platinum has been widely used since the 1970s in the mining and automobile industry. The most common oxidation states are II and IV [13]. Gastrointestinal absorption of platinum salts in rats is extremely low (<1% of oral dose) but much higher in humans (42–60%) [14]. The acute toxicity of platinum salts depends on their water solubility and speciation. Pt(II) forms a tetra-coordinate aqua ion [Pt(H\(_2\)O)\(_4\)]\(^{2+}\), but PtOH\(^+\), Pt(OH)\(_2\), or [Pt(OH)\(_5\)(H\(_2\)O)]\(^{-}\) in river water and [PtCl\(_5\)(OH)]\(^{-}\) in seawater. These have been proposed to be the dominant inorganic species of platinum. A range of Pt(IV) salts has been reported to have rodent oral LD\(_{50}\) (lethal concentration 50) values of 10–1100 mg kg\(^{-1}\). The mean UK human dietary intake was found to be ca. 0.2 µg/day [13], and a human permitted daily exposure (PDE) of 108 µg/day (orally) has been advised [15].

With the discovery of its chemotherapeutic properties, platinum-based drugs have been developed and administered to treat various types of cancer, such as testicular, ovarian and lung cancer [16,17]. The most commonly used drugs, cisplatin, carboplatin and oxaliplatin, are eliminated within 24 h via urine with excretion rates of 28 ± 4%, 77 ± 5% and 36 ± 8%, respectively [18].

Improvements in analytical techniques have facilitated the detection of trace elements in various environmental samples. Nevertheless, the ICP-MS technique is currently the most preferred method for this type of analysis due to its higher sensitivity and selectivity [19], especially for not regularly monitored elements such as thallium and platinum [20,21]. Besides environmental analysis, ICP-MS has been applied in several fields, including clinical, pharmaceutical, food, fertiliser and geological analysis [22,23].

Based on the data gathered from the literature, the table below represents the occurrence of dissolved platinum in the aquatic environment (Table 1).

| Sample Source           | Dissolved Pt (ng L\(^{-1}\)) | Ref. |
|-------------------------|------------------------------|------|
| East Asia—Rivers        | 0.035–0.123                  | [24] |
| France—Gironde estuary  | 0.068–0.164                  | [25] |
| Japan—Estuary           | 0.936–6.867                  | [26] |
| Japan—Seawater          | 0.080–1.990                  | [27] |
| Japan—Seawater          | 0.037–0.049                  | [28] |
| Japan—Seawater          | 0.078–0.388                  | [29] |
| Japan—Seawater          | 0.039–0.293                  | [30] |
| Spain—Estuary           | 0.004–0.121                  | [31] |
| Spain—Wastewater influent | 3.95–38.68                  |      |
| Spain—Wastewater effluent | 56.08–75.79                 | [17] |
| Spain—Hospital wastewater | 81.94–13,913                |      |
Table 1. Cont.

| Sample Source                          | Dissolved Pt (ng L\(^{-1}\)) | Ref.   |
|----------------------------------------|------------------------------|--------|
| Spain—Hospital wastewater              | 144                          | [32]   |
| Spain—Wastewater treatment plant influent | 7.9                         |        |
| Spain—Wastewater treatment plant effluent | 5.9                         |        |
| Slovenia—Hospital wastewater           | 352                          |        |
| Slovenia—Wastewater treatment plant influent | 23.3                        | [32]   |
| Slovenia—Wastewater treatment plant effluent | 12.8                        |        |
| Slovenia—Hospital wastewater           | 226–352                      | [33]   |
| Slovenia—Wastewater treatment plant effluent | 23–27                       |        |
| Austria—Hospital wastewater            | 4700–145,000                 | [18]   |
| Austria—Hospital wastewater influent from in-patient ward | 3240–266,000               | [34]   |
| Austria—Hospital wastewater effluent   | 1770–144,000                 |        |
| France—Hospital wastewater             | 80–610                       | [35]   |
| France—Wastewater influent             | 10–20                        |        |
| France—Wastewater effluent             | 10                           |        |
| UK—Hospital wastewater                 | 20–140,000                   | [36]   |
| UK—Main drain                          | 30–100,000                   |        |

With the continuous discharge of platinum into the environment from hospital/household wastewater and road runoff, it has become essential to assess the environmental impact of these contaminants on the aquatic biota. A small number of studies have evaluated the effect of the element platinum and the anticancer drug cisplatin. However, these revealed that platinum has an impact on the mobility and survival of *Daphnia magna* with EC\(_{50}\) (48h) = 110 µg L\(^{-1}\) (half maximal effective concentration) and LC\(_{50}\) (48h) = 157 µg L\(^{-1}\) [37]. Cisplatin has also affected the reproduction of crustaceans by reducing the offspring percentage [38], the number of eggs and the population growth rate [39]. Moreover, it has caused DNA damage at concentrations starting from 300 ng L\(^{-1}\) [40] and modified the activity of several enzymes at 100 ng L\(^{-1}\) [41].

Considering the potential occurrence of platinum-based anticancer drugs in the aquatic environment, even at very low concentrations and the lack of data on the occurrence of platinum in the Lebanese water sources, this research aims to give a comprehensive assessment of multiple sites and sources and offer preliminary findings for future research, with the purpose of highlighting the state of waste management in Lebanon. This will be achieved by determining the level of total platinum in Lebanese rivers, wastewater treatment plants and sewage outfalls by ICP-MS.

2. Materials and Methods

2.1. Chemicals and Instrumentation

Ultrapure water (18.2 M\(\Omega\)-cm, 0.22 µm filtered) was provided by an Avidity Science (Aylesbury, UK) water purification system. MF-Millipore membrane filters (0.45 µm pore size) were obtained from Merck Millipore (Cork, Ireland). Platinum and Rhenium standard solutions for ICP were purchased from Sigma-Aldrich (Gillingham, UK) in 2% HNO\(_3\) and at a concentration of 1000 mg L\(^{-1}\). For metal dissolution and stabilisation, the acid HNO\(_3\) (trace analysis grade) was obtained from Fisher Chemical (Loughborough, UK).

The determination and quantification of total platinum were carried out on a 7700 series ICP-MS instrument from Agilent Technologies (Didcot, UK) coupled with an ASX-520 series autosampler. The ICP-MS was controlled by Agilent Masshunter 5.1 software (version D.01.01). The operating parameters of the instrument are presented in Table 2.
Table 2. Operating parameters of the ICP-MS.

| Parameter          | Type/Value                  |
|--------------------|----------------------------|
| Nebuliser          | MicroMist glass concentric  |
| Spray chamber      | Scott                       |
| Nebulizer gas flow | 1.02 L min⁻¹                |
| Auxiliary gas flow | 0.9 L min⁻¹                 |
| Plasma gas flow    | 15 L min⁻¹                  |
| Omega lens voltage | 11.6 V                      |
| ICP RF power       | 1550 W                      |
| Isotopes measured  | ^{195}Pt, ^{185}Re          |
| Gas mode           | No gas, He: 4 mL min⁻¹       |
| Scan mode          | Peak hopping                |
| Dwell time per isotope | 0.1 s               |
| Replicates         | 5                           |
| Sweeps             | 100 sweeps/replicate        |

2.2. Calibration Standards Preparation:

Stock solutions of platinum and rhenium were prepared at concentrations of 1000 ng mL⁻¹ and 500 ng mL⁻¹ (w/w), respectively, in ultrapure water with 2% HNO₃ (v/v%). The calibration curve points ranging from 0.01 to 10 ng mL⁻¹ (w/w) of platinum, including a blank, were made from the stock solution. The internal standard, rhenium, was added to each solution at a fixed and constant concentration, 1 ng mL⁻¹ (w/w). Throughout this manuscript, the concentrations are presented as nominal concentrations.

2.3. Method Optimisation and Validation

For the method optimisation, a single solution of platinum was prepared at a concentration of 2.5 ng mL⁻¹. Different parameters were investigated separately in order to achieve the highest sensitivity for platinum with the lowest background noise. These include He gas flow rate, cell deflect energy, optical bias, energy discrimination, etc.

Moreover, the signal of hafnium was monitored, as this element is expected to have interferences with platinum through the hafnium oxide (HfO⁺) signals generated in the plasma [19,22]. A pilot study was initially conducted with 3 river samples and 3 wastewater samples to evaluate the presence of hafnium and its potential effect on the platinum signal.

Once optimised, the method was then validated according to the International Council for Harmonization (ICH) guidelines for the following parameters: linearity, limit of detection, limit of quantification, precision and accuracy [42]. The linearity was assessed over the working range of 0.01 to 10 ng mL⁻¹. The limit of detection and quantification was calculated based on the standard deviation of the blank. For precision and accuracy, 3 points covering the working range were selected (QCₗow = 0.5, QCₘedium = 2.5 and QCₜig = 7.5 ng mL⁻¹) and evaluated as percent relative standard (%RSD) deviation and recovery percentage, respectively. The recovery percentage of platinum in the samples was also measured by spiking river and wastewater matrices with the QCs concentrations.

Nevertheless, to avoid any instrumental drift, a calibration was run before every sample analysis. In addition, quality control measures were run throughout the analysis (QCₗow = 0.5 ng mL⁻¹ of platinum) at regular intervals, followed by two instrumental 2% HNO₃ washes, to ensure no memory effects related to the QC standard were carried over.

2.4. Sample Collection and Preparation

Grab samples were collected from the 5–10 January 2022 from different rivers and wastewater treatment plants (WWTPs) covering all the Lebanese regions, as illustrated
in Figure 1. In total, 22 samples were collected, including 14 from rivers, three from wastewater treatment plants (influent and effluent) and two from sewage outfalls to the sea. Sample bottles were formerly soaked and washed with 5% HNO₃ and triple rinsed with the sample water before collection.

Upon reaching the laboratory, samples were filtered through 0.45 µm syringe filters and acidified with HNO₃ at a concentration of 2%. The effect of filtration was assessed over three concentrations covering the working range (0.1, 1 and 10 ng mL⁻¹) to determine if the platinum concentration is reduced by the retention on the filter material. All samples were stored in a fridge at 3 °C until analysis. Before analysis, the internal standard, rhenium, was added at a concentration of 1 ng mL⁻¹.

Figure 1. Map of Lebanon with the sample collection sites (green: rivers; red: WWTPs; orange: sewage outfalls).

3. Results and Discussion
3.1. Method Development and Validation

The platinum signal was initially monitored in a real-time scan to detect which parameter was enhancing or suppressing the signal of ¹⁹⁵Pt. All the parameters were assessed individually and were found to not cause any significant effect, except for the Helium gas flow rate. For this reason, this parameter was examined further by analysing a solution of platinum at variable flow rates (0 to 12 mL min⁻¹). The figure of merit obtained showed that the platinum signal was the highest at “no gas mode” and at 4 mL of He min⁻¹. All the other parameters were operated as default, as shown in Table 3.
Table 3. ICP-MS operating parameters for the detection of platinum.

| Parameter | No Gas Mode | Helium Mode |
|-----------|-------------|-------------|
| Plasma    |             |             |
| Extract 1 | 0 V         |             |
| Extract 2 | -200 V      |             |
| Omega bias| -90 V       |             |
| Omega lens| 10.8 V      |             |
| Lenses    |             |             |
| Cell entrance | -30 V      | -40 V       |
| Cell exit  | -50 V       | -60 V       |
| Deflect   | 14.4 V      | 1.2 V       |
| Plate bias| -40 V       | -55 V       |
| Cell      |             |             |
| He flow   | 0 mL min⁻¹   | 4 mL min⁻¹  |
| OctP bias | -8 V        | -18 V       |
| OctR RF   | 180 V       |             |
| Energy discrimination | 5 V | 3 V |

The signal of the element hafnium was monitored for both isotopes $^{178}$Hf and $^{179}$Hf in 6 environmental samples. Hafnium could be detected in the environment from its adoption in autocatalysts with zirconium or from mineral dust particles [43]. Nevertheless, the counts of the interferent were less than five times the counts of platinum measured in the samples, which suggests that the formation rate of hafnium oxide was negligible. Hence, mathematical correction and further analysis of hafnium were not required [24,44].

In the analysis of the environmental samples, it was noticed that under “Helium mode”, the counts of hafnium were reduced by 60% (±6%). Helium is known to be more effective against all polyatomic interferences, including $^{178}$Hf and $^{179}$Hf [19,43], especially in complex matrices, in addition to reducing the responses of several elements such as Na, K, Ca, Mg, etc., which are expected to be abundant in environmental samples [45]. Thus, “Helium mode” concentrations could represent a more accurate indication of platinum levels in real samples where the levels of interferent ions are variable. For this reason, “Helium mode” was adopted for the validation and sample analysis procedures.

For the filtration procedure, it was experimentally found that platinum at low, medium and high concentrations did not vary by more than ± 9% in filtered and unfiltered standard solutions. Hence, the quantification of total platinum in real samples was performed in filtered samples. The same results were obtained in a study by Vidmar et al., where the variation was not more than ± 5% after filtration with 0.45 µm cellulose nitrate membrane filters [32].

The method was validated when the operating parameters for “Helium mode”, as stated in Table 3, were performing over a period of two days. All the platinum signals were corrected by the response of the internal standard. The calibration curve obtained demonstrated linearity over the working range with a regression coefficient greater than 0.999. The instrumental limit of detection and limit of quantification corresponding to $3.3 \times SD_{blank}$ and $10 \times SD_{blank}$ demonstrated high sensitivity for platinum with values in the range of several ng L⁻¹. The precision was assessed within one day (intraday) and between days (interday), which are equivalent to the method repeatability and intermediate precision, respectively. The precision met the acceptance criteria with a % RSD lower than 5%. Similarly, the accuracy met the acceptance range of 80–120%. The detailed results are presented in Table 4.

Additionally, the coefficient of variation of the QC runs throughout the analysis was consistently lower than 5%, and the instrumental blanks did not show detectable traces of platinum, which validated the results obtained for the environmental samples.
Table 4. Validation parameters of the ICP-MS method.

| Parameters                        | Value                          |
|-----------------------------------|--------------------------------|
| Range (ng L\(^{-1}\))            | 10–10,000                     |
| Regression coefficient (R\(^2\)) | 1                              |
| Regression equation               | \(y = 0.5022x + 0.001\)       |
| Accuracy (%) (±SD)                | 99.13 (± 4.35)                |
| Interday precision (%) (±SD)      | 0.54 (± 0.24)                 |
| Intraday precision (%) (±SD)      | 2.08 (± 0.91)                 |
| LOD (ng L\(^{-1}\))              | 0.56                           |
| LOQ (ng L\(^{-1}\))              | 2.35                           |
| % Recovery in spiked river and wastewater samples (n = 6) (±SD) | 106.39 (± 6.92) |

3.2. Application to Lebanese Surface Water and Wastewater Samples

Total platinum concentrations in Lebanese rivers and wastewater samples were all above the instrumental limit of quantification, as shown in Table 5.

Table 5. Concentrations of total platinum in Lebanese water samples in ng L\(^{-1}\) (± SD of the concentrations measured).

| Sample Location       | Geographic Coordinates | Total Dissolved Platinum Concentration (ng L\(^{-1}\)) (± SD) |
|-----------------------|------------------------|------------------------------------------------------------|
| Antelias River        | Lat 33.914801 Long 35.594127 | 38.42 ± 2.88                                               |
| Al Kalb River         | Lat 33.95461 Long 35.60112 | 37.88 ± 4.21                                               |
| Zahrani River         | Lat 33.49531 Long 35.334795 | 37.36 ± 3.15                                               |
| Hasbani River         | Lat 33.40703 Long 35.66732 | 32.61 ± 3.11                                               |
| Litani River          | Lat 33.338889 Long 35.245278 | 34.45 ± 0.94                                              |
| Wazzani River         | Lat 33.274254 Long 35.618137 | 34.59 ± 2.32                                              |
| Awali River           | Lat 33.5888 Long 35.386    | 39.93 ± 3.88                                              |
| Damour River          | Lat 33.705833 Long 35.439444 | 31.63 ± 2.04                                              |
| Orontes River         | Lat 36.045278 Long 35.963611 | 31.04 ± 4.79                                              |
| Al Bared River        | Lat 34.51241 Long 35.95452  | 33.34 ± 2.15                                              |
| Ibrahim River         | Lat 34.062222 Long 35.642222 | 28.74 ± 2.21                                              |
| Al Jawz River         | Lat 34.273804 Long 35.690072 | 30.62 ± 2.70                                              |
| Ostouene River        | Lat 34.58189 Long 36.10626  | 36.38 ± 2.88                                              |
| Abou Ali River        | Lat 34.4579 Long 35.8415    | 39.41 ± 2.14                                              |
| WWTP Joub Jannine Influent | Lat 33.637569 Long 35.776514 | 24.91 ± 1.77                                              |
| WWTP Joub Jannine Effluent | Lat 33.637569 Long 35.776514 | 27.21 ± 2.20                                              |
| WWTP Al Ghadir Influent | Lat 33.81311 Long 35.48340  | 25.35 ± 2.29                                              |
| WWTP Al Ghadir Effluent | Lat 33.81311 Long 35.48340  | 31.79 ± 5.97                                              |
| WWTP Saida Influent   | Lat 33.64271 Long 35.39983  | 24.76 ± 1.36                                              |
| WWTP Saida Effluent   | Lat 33.64271 Long 35.39983  | 22.44 ± 2.19                                              |
| Ramlet al. Baida Outfall | Lat 33.895944 Long 35.467917 | 24.57 ± 2.88                                              |
| Jiyeh Outfall         | Lat 33.661225 Long 35.417343 | 53.32 ± 1.64                                              |
During the monitored period, the dissolved platinum concentrations ranged between 22.44 and 53.32 ng L$^{-1}$. In comparison with previous studies conducted in different countries, the results obtained were at the same level in terms of concentration range, especially for the wastewater samples. For example, total platinum in Slovenian wastewater samples ranged between 12.8 and 27 ng L$^{-1}$; in Spanish wastewater samples, it ranged between 3.97 and 75.79 ng L$^{-1}$ [17,32,33]. In France, platinum concentrations ranged between 10 and 20 ng L$^{-1}$ during working days and 10 ng L$^{-1}$ during non-working days [35].

Nevertheless, the concentrations detected in wastewater from hospitals and surface water were significantly different from those obtained in this present study. As expected, samples collected directly from hospitals showed higher levels of platinum. In Spain, platinum levels ranged between 81.94 and 13,913 ng L$^{-1}$ [17,32], reaching 140,000 ng L$^{-1}$ in the United Kingdom and 266,000 ng L$^{-1}$ in Austria [18,34,36]. In surface water, the concentrations were much lower, with reported concentrations ranging from 0.004 to 0.164 ng L$^{-1}$ in Spain and France [25,31] and from 0.037 to 6.867 ng L$^{-1}$ in Japan [26–30].

From the data collected in this study, it was evident that the concentration of total platinum was of the same order in all the analysed samples. This could be interpreted in three ways: (1) due to the continuous release of untreated or poorly treated waste in surface water, the concentration of platinum has stabilised in the water resources and hence, the concentrations in river samples are similar to the wastewater samples; (2) the detected concentrations could represent the baseline concentration of total platinum in the aquatic environment in Lebanon and/or (3) the platinum could be majorly deposited and accumulated in the sediment/sludge phase.

Since no previous studies have been conducted in Lebanon, the baseline concentration of platinum in the aquatic environment is unknown. Nevertheless, based on reports and previously published studies, the level of platinum estimated in global drinking water was 0.1 ng L$^{-1}$ [46] and not detected when assessed in tap water samples from France and the UK, respectively [35,36]. Consequently, the measured platinum in this study is presumed to be derived from anthropogenic emissions.

The primary source of platinum contamination is from catalytic converters in motor vehicles. The industrial contribution may come from glassmaking, jewellery production, or dentistry and medicine [16]. In medicine, the increased use of platinum-based anticancer drugs has been recorded, with a 50 to 70% administration rate in cancer treatment regimens [47]. With no guaranteed access to Lebanese hospital wastewater, it was not possible to determine the contribution of hospitals regarding platinum contamination. In a study conducted by Goullé et al., it was determined that the hospital contribution was 9.4% of the total platinum in the environment [35]. This estimation depends on several parameters such as sample collection day (working/non-working days), type of sampling, hospital wastewater flow and amount of platinum administered within a selected hospital [48]. Although these results cannot be compared, it could be assumed that platinum-based drugs contribute to the total amount of platinum discharged in the aquatic environment and that current treatment processes are not efficient for the removal of platinum from these wastewater streams.

4. Conclusions

In the present study, an ICP-MS method was developed, validated and applied for the quantitative determination of platinum in the aquatic environment using an Agilent 7700 ICP-MS. As previously stated, this is the first research to report the concentrations of total platinum in the Lebanese aquatic environment. The concentrations detected in rivers and wastewater samples ranged between 22.44 and 53.32 ng L$^{-1}$ with no significant differences between the various sampling points, especially between WWTP influents and effluents. This could indicate that the operating treatment plants are not efficiently eliminating these trace concentrations of platinum.
The levels of platinum obtained were estimated to emerge from anthropogenic sources such as catalytic converters in vehicles and drug excretion of cancer patients as medical discharge. The results obtained were in good agreement with similar studies presented for the aquatic environment, including France, Slovenia and Spain.

Despite the inability to specify the source of the platinum detected within this study, its presence in the environment should be a matter of concern. Considering the PDE of 108 µg/day and neglecting other parenteral sources of platinum, humans could be exposed daily from gastrointestinal ingestion alone to approximately 34 ng L⁻¹, which is way below the permissible limit. However, since platinum is commonly used and increasingly in the medical field and with the continuous discharge of untreated sewage into the Lebanese surface water, more attention and further research should be carried out on the potential long-term effect on aquatic biota and human health. Therefore, the development of specific treatments for its reduction and removal should be investigated and implemented.

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