

## Article

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# Regulated Inductively Coupled Plasma–Optical Emission Spectrometry Detectable Elements in Utah Lake: Characterization and Discussion

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# Regulated Inductively Coupled Plasma–Optical Emission Spectrometry Detectable Elements in Utah Lake: Characterization and Discussion

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**Abstract:** During the 2021 ( $n = 15$ ) and 2022 ( $n = 13$ ) summers, we measured the total and dissolved ( $<0.45 \mu\text{m}$ ) concentration of 25 elements in Utah Lake using Inductively Coupled Plasma–Optical Emission Spectrometry (ICP–OES) with detection limits in the order of a few parts-per-billion (ppb). This resulted in 1400 measurements, which is a unique dataset in terms of sensitivity and temporal resolution. Regulated elements are not commonly measured at the ppb-level; thus, these data provide insight into both the behavior and existence of these elements in an aquatic environment and have implications for both the management and regulation of the lake. Utah regulates twelve of these elements. While ICP–OES has ppb-level sensitivity, it is not the approved regulatory analysis method for these elements. All regulations are for dissolved concentrations, except aluminum (Al) and phosphorus (P), which are for total recovery. We found total Al above the allowable concentration, but dissolved concentrations were well below allowable concentrations. We attribute high total concentrations to suspended clays. This suggests that regulatory methods should be reviewed for lakes with a high suspended-solid content. Dissolved copper (Cu) concentrations were below regulatory levels in 2021, but some samples were above regulatory levels in 2022. This could be related to the use of Cu-based algaecide treatments, or from other sources. Lead (Pb) data were inconclusive; dissolved Pb concentrations were well below the acute (1 h average) limit, but the chronic concentration limit (4 h average) was below the ICP–OES minimal detection limit. Arsenic (As) concentrations exhibited a seasonal trend that we attribute to groundwater inflows—they were below regulatory levels for aquatic environments but around the levels for drinking water. This ppb-level study with high temporal resolution provides insight into regulated elements in Utah Lake previously not available due to the high sensitivity of the method and measurements of both total and dissolved concentrations.

**Keywords:** Utah Lake; inductively coupled plasma–optical emission spectrometry; water quality; micronutrient; lake management; trace elements; state regulations



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## 1. Introduction

### 1.1. Study Overview

Utah Lake (UL) is approximately 40 km by 21 km, with a surface area of 390 km<sup>2</sup> at maximum fill, and is in north-central Utah in the United States. UL waters are characteristically turbid and unstratified because of wave action and bioturbation and are near the solubility limit of calcite, with calcite precipitation commonly occurring [1].

There is a significant body of published research on UL. Early published UL research from 1931 [2] addresses algae, with additional work on phytoplankton communities published from the 1960s to the present [3–12]. Numerous studies on nutrients have been published [13–20], with studies on the local geology that contains formations with elevated phosphorus (P) [20–22]. Early remote-sensing studies from the 1970s used UL in some of the first papers from NASA’s Landsat satellite [23,24]. Other early remote-sensing studies from the 1980s demonstrated the heat capacity mapping mission (HCMM) [25] and AVHRR sensor [26] on UL. Recent studies using remote-sensing data address long-term trends and variability in algal blooms [27–37].

With the exception of P, there has been little published work on regulated elements in UL water [38]. A 2014 study focused on compliance with EPA standards and identifying pollution sources for arsenic (As) and other heavy metals in Utah Lake and its tributaries [39]. Despite the lack of studies on regulated elements in UL, their concentrations are a concern as excessive or deficient amounts of these elements can have damaging effects on aquatic life [40–42] and affect biogeochemical processes [43]. The impacts of trace elements in aquatic systems have been studied worldwide [44,45], demonstrating that in most ecosystems, nutrients are the limiting factor in algal growth, though sometimes other trace elements limit growth. Bayer, et al. [46] studied Lake Hayes in New Zealand in regard to the importance of macro- and micronutrients. They reported on nitrogen (N), P, copper (Cu), iron (Fe), boron (B), molybdenum (Mo), zinc (Zn), and silicon (Si) and found that the limiting factors for phytoplankton growth in Lake Hayes were N, Zn, and B. P. Dengg, et al. [47] evaluated the trace elements of manganese (Mn), cobalt (Co), Fe, Zn, and Mo in the waters of the Taupo Volcanic Zone (TVZ) in New Zealand and found that Fe was a colimiting micronutrient for cyanobacteria. Studies in southeastern Australia showed that trace micronutrients are “an important regulator of the severity of cyanobacterial blooms” [48,49]. This research shows that understanding trace micronutrients is required to effectively manage algal blooms, as these trace elements may control ecological processes. Our study starts to provide these data for UL.

We provide data on regulated elements that can be detected using Inductively Coupled Plasma–Optical Emission Spectrometry (ICP-OES). We selected ICP-OES for this study for two reasons: (1) it measures 25 elements per sample, providing a comprehensive dataset, and (2) it has very low parts-per-billion (ppb) detection limits for these elements. These regulated detectable elements (DEs) include heavy metal and nonmetal elements in UL. Over two summers, we measured both dissolved and suspended concentrations of DEs in UL approximately weekly, with 15 samples in 2021 and 13 in 2022. We analyzed both total and dissolved concentrations of 25 elements for each sample. This resulted in 1400 data points over the critical warm growing season.

For this manuscript, we evaluated the 12 regulated elements that are detected by ICP-OES. For five of these elements, which are well under regulatory levels of concern, we provide only a minimal discussion. We provide a more detailed discussion of the seven elements more likely to be of concern, either from toxic excess or deficiency. We did not follow regulatory methods for the elements presented in this analysis as we used a standard ICP-OES method—ICP-OES is not generally stated for most standard methods as it is complex and expensive, and for most elements, the low detection levels are not required. Also, ICP-OES measures elemental concentrations, while many regulations are based on ionic or molecular forms. Our analysis of this large range of elemental concentrations (25) at the very low ppb detection level for both total and dissolved concentrations provides insight into UL conditions and potential elements of concern, at both toxic excess and toxic deficiency levels. It characterizes if the elements are in the dissolved state, and thus more biologically available, or associated with suspended solids. This analysis is not meant to determine if UL waters are out of compliance with Utah regulatory standards, but rather to provide important information about the condition of the lake and facilitate future research.

## 1.2. ICP-OES Detectable Elements in Aquatic Systems

Trace metals and other elements are natural and present in all ecosystems, and water concentrations are influenced by geology, land use, erosion, ecology, and geochemistry. Ecological processes also affect concentrations as aquatic biota ingest and absorb metals. These organisms are affected in both beneficial and adverse ways by the concentrations of these elements. These trace elements typically have no visible indicators in surface waters, although at toxic or deficient levels, they can have visual impacts on the ecology [50]. The toxicity of trace elements is a particular concern for UL as it hosts a large migratory bird population along with the endemic and threatened June Sucker fish species. All metals are toxic at certain thresholds [51] and impair the survival, reproduction, and behavior of aquatic life.

Some common toxic DEs in lakes and reservoirs, including metals, found at concentrations that can cause detrimental impacts include arsenic (As), cadmium (Cd), chromium (Cr), lead (Pb), nickel (Ni), selenium (Se), copper (Cu), and zinc (Zn) [50]. Even at non-toxic concentrations, As, Cd, and Pb, can affect the mobility, feeding, and navigation behaviors of invertebrates [52–54]. These effects cascade through aquatic ecosystems [55,56]. In larger organisms such as fish, Pb, Cd, Ni, and Cr can impact growth rates, biological processes, and reproductive health [57,58]. The impacts of aluminum (Al), Cd, Cu, Zn, Pb, Ni, As, and Se on birds include behavioral impairments and reduced reproductive success [59–61].

Some metals are toxic at excess levels and also harmful when deficient. Zn, Cu, and Ni negatively impact biological activity if they are present in excess levels as well as if they are missing or below toxic deficit concentrations [40–42,55,56,62]. Some elements, such as boron (B), iron (Fe), manganese (Mn), and molybdenum (Mo), are beneficial at certain concentrations and act as micronutrients that are essential for aquatic life. Micronutrients have been found to be limiting or colimiting factors for algal growth [46,63–66]. In 2008, Downs, et al. [42] conducted a review of micronutrients in 56 freshwater lakes and found that “the proportion of the lakes analyzed in which micronutrient limitation was found was 76% for molybdenum; 74% for iron; 67% for boron, 67% for cobalt, and 20% for copper.” Similar to the effects of toxicity, nutrient deficiencies for algae and phytoplankton affect the entire biota of lake systems, including invertebrates, migratory birds, fish, and other organisms [55,56].

The US Environmental Protection Agency (EPA) states that sources and activities “such as mines, smelters, firing ranges, municipal wastewater treatment outfalls, industrial point sources, urban runoff, landfills, and junkyards are potential sources for heavy metals and other harmful elements that can cause impairment” [50]. Other sources include the combustion of fossil fuels, phosphate fertilizers, metallo-pesticides, and road salts [51,67]. All of these sources are currently or have historically been present within the UL watershed. The Geneva Steel plant operated from 1944 to 2002 on the northeastern shore of UL. This plant produced steel using “coal-derived coke,” with effluent drains near the lake that served as a substantial source of metals to UL [38,68,69]. Anthropogenic nutrient sources include seven wastewater treatment plants (WWTPs) that discharge into the lake, while the southern end of UL is surrounded by agricultural land, and the western side has active gravel mines. The population in the last 40 years has nearly tripled, increasing from ~220,000 in 1980 to ~640,000 in 2020 [70].

In addition to the potentially toxic DEs, ICP-OES measures several other elements that have important functions or impacts on aquatic ecosystems, including barium (Ba), calcium (Ca), potassium (K), sodium (Na), sulfur (S), B, silicon (Si) and P.

B and S are naturally present in aquatic environments and serve as essential nutrients. However, human activities can lead to elevated concentrations, potentially reaching toxic levels [71–73]. Toxic S concentrations are typically rare in surface waters unless there is a direct source of pollution. B is also influenced by human activities, and the range between toxic deficit and excess B concentrations is relatively small compared to other nutrients [74].

Ba occurs naturally and is also released by industrial processes primarily in the form of airborne particles that can eventually settle in nearby surface waters. UL is particularly

susceptible to capturing atmospherically deposited dust due to its location and large surface area [16,17,19,20]. Ba usually exists in a precipitated form in aqueous systems, with alkaline environments, such as UL, further limiting its solubility [75].

K and P are essential nutrients for aquatic primary production. Since plants require less K than P, and K is not usually a limiting nutrient. Neither element becomes toxic at the concentrations typically found in surface waters, although high concentrations of P can lead to eutrophication, which negatively impacts water quality. Natural P and K [76] sources include geologic formations after weathering. In the UL watershed, the Delle Phosphatic and Meade Peak geologic formations contribute large amounts of P to the lake [1,18].

Na is a key component of salinity and can severely degrade freshwater systems if present in high concentrations. It enters surface waters through natural weathering, but also from road salts [77].

The impacts of Se are not well understood in freshwater systems due to the complexity of its geochemical interactions and cycling. However, it is toxic at very low concentrations and can degrade aquatic systems when it enters surface waters through anthropogenic activity [78]. While Se is necessary for aquatic biota, there is a very narrow concentration window in which Se turns from beneficial to toxic [79].

These discussions of the origins and impacts of DEs on the UL ecosystem are not comprehensive. We included this review to demonstrate the potential impacts and functions of these DEs in the UL ecosystem and illustrate the importance of understanding their ambient concentrations and behaviors.

### 1.3. Regulated DE Water Quality Standards Applicable to UL

We analyzed 25 elements detectable by ICP-OES, 12 of which are regulated by water quality standards promulgated by the Division of Water Quality of the Utah Department of Environmental Quality in compliance with the Clean Water Act [80–82]. To contextualize our discussion of water column concentrations, we outline the relevant standards.

The publication “Standards of Quality for Waters of the State” [80–82] categorizes waterbodies according to their use and significance and prescribes different standards for each category. UL is classified as follows:

- Class 2A: “Protected for frequent primary contact recreation where there is a high likelihood of ingestion of water or a high degree of bodily contact with the water” (swimming, kayaking, diving, water skiing, etc.);
- Class 2B: “Protected for infrequent primary contact recreation. Also protected for secondary contact recreation where there is a low likelihood of ingestion of water or a low degree of bodily contact with water” (e.g., boating, wading, etc.) [82];
- Class 3B: “Protected for warm-water species of game fish and other warm water aquatic life, including the necessary aquatic organisms in their food chain”;
- Class 3D: “Protected for waterfowl, shore birds, and other water-oriented wildlife not included in Classes 3A, 3B, or 3C, including the necessary aquatic organisms in their food chain”;
- Class 4: “Protected for agricultural uses including irrigation of crops and stock watering” [80–82].

For the Class 3B and 3D standards, the regulations are for dissolved elements in units of  $\mu\text{g}/\text{L}$ . For Class 4 standards, regulations are for dissolved elements in units of  $\text{mg}/\text{L}$ . We converted the Class 3B and 3D standards to units of  $\text{mg}/\text{L}$  to match our data units. Table 1 lists the most stringent acute (1 h average) and chronic (4-day average) standards for the 12 regulated DE analytes [80].

The standards for Al, Cd, Cr, Cu, Ni, Pb, and Zn depend on water hardness. Typically, the adjustment factor for these “hardness dependencies” would be determined using the hardness level detected in each sample; however, our samples were not tested for hardness, so we used a representative average hardness level for the lake. Using data from the Utah Ambient Water Quality Management System (AWQMS) collected from 1978 to 2015, we found the mean hardness, based on  $\text{CaCO}_3$ , of UL water to be 197.4  $\text{mg}/\text{L}$ ,

with a range from 123 to 291 mg/L. We used the mean value to calculate a reference concentration for elements with hardness dependencies for discussions. For Cd, Cr, Pb, and Zn, the conversion is a natural log equation based on the hardness of the water with a specified conversion factor [80]. Cr standards are for hexavalent chromium—a toxic ion. We measured elemental Cr, which provides an upper bound, but does not measure hexavalent Cr. Again, we want to stress that our analytical methods do not follow required regulatory methods. Instead, we determined elemental concentrations using ICP-OES, which has sensitivity in the few parts-per-billion range, depending on the element, and uses either filtration for dissolved concentrations or digestion for total concentration.

**Table 1.** Most stringent applicable standard for each regulated element in UL.

DE	Acute Standard (mg/L)	Chronic Standard (mg/L)	Designated Use
Aluminum (Al) *	0.75	0.75	3B, 3D
Arsenic (As) **	0.10	0.10	4
Boron (B) **	0.75	0.75	4
Cadmium (Cd)	0.0018	0.00072	3B, 3D
Chromium (Cr) (Hexavalent) ***	0.016	0.011	3B, 3D
Copper (Cu)	0.013	0.009	3B, 3D
Iron (Fe) **	1	1	3B, 3D
Nickel (Ni)	0.468	0.052	3B, 3D
Phosphorus (P) *	0.025	0.025	3B
Lead (Pb)	0.065	0.0025	3B, 3D
Selenium (Se)	0.0184	0.0046	3B, 3D
Zinc (Zn)	0.12	0.12	3B, 3D

Notes: \* Concentration based on total recovery criteria. \*\* Measured as maximum not acute and chronic. \*\*\* We measured elemental Cr, not hexavalent.

The standards for Al depend on pH and hardness. Since the pH of UL water is nearly always greater than 7.0 and the hardness, based on CaCO<sub>3</sub>, is greater than 50 mg/L, we use the acute Al criterion of 750 µg/L (expressed as total recoverable Al) for our discussion. The Utah Department of Water Quality (UDWQ) is currently in the process of adopting the EPA's recommended Al criteria, which depends on dissolved organic carbon (DOC), hardness, and pH. We were told in conversations with representatives of the UDWQ that the current value of 750 µg/L is acceptable as a reference point. Al and P are the only regulated elements with a standard for total recoverable concentration—all other elements require the analysis of dissolved concentrations. As discussed below, we performed a complete digestion of unfiltered water samples for our total concentration measurements.

Total elemental Al concentrations can be misleading, as clay minerals, which make up a significant fraction of suspended solids, have high Al concentrations and are captured by a total digestion approach. However, the Al found in clay minerals is not bioavailable or reactive and is of less concern than dissolved Al. For this study, we analyzed filtered and unfiltered samples to show the impact of suspended solids on elemental concentrations. We performed a complete digestion of the unfiltered samples, which includes the suspended clay minerals in the sample. The filtered samples only measured dissolved elements. For Al concentrations, we did not attempt to follow the UDWQ analysis methods but assumed that “total recoverable Al” required unfiltered lake water to be analyzed after aggressive acid digestion, which would include the nonreactive Al bound up in clay particles. The regulatory methods under consideration use less-aggressive digestion.

#### 1.4. Study Goals and Objectives

This study presents research based on a unique high-sensitivity (ppb-level) weekly dataset collected over two years, measuring both dissolved and total concentrations of 25 elements. This dataset includes 1400 measurements that characterize the weekly variation and temporal behavior of these elements over the study period. These data and

our analysis provide insights into lake geochemistry and potential anthropogenic impacts that are not found in the literature, for either Utah Lake or other water bodies, because of the number of elements analyzed for both total and dissolved concentrations, the high sensitivity of the analytical method, and the temporal resolution of the dataset—all of these factors are unique in studies of aquatic environments (to our knowledge). Regulated elements are not commonly measured with detection limits in the order of a few ppb, except for lead (Pb).

This dataset helps characterize both the behavior and presence of these elements in an aquatic environment. For example, the analysis of what percentage of the 1400 samples were above detection limits for the various elements could only be conducted on this one-of-a-kind dataset and helps in understanding both the variability and behaviors of these 25 elements in both total and dissolved phases. These measurements start to characterize how these elements are present in the water column—whether they are dissolved and readily bioavailable or associated with suspended solids. We provide an in-depth analysis of the regulated elements that have concentrations near regulatory limits or have other interesting behaviors. The goal is not to evaluate regulatory compliance, as our methods are not the approved analysis methods, but to provide data for managers and regulators that can be used to design sampling schemes, evaluate management practices, and review regulatory analysis methods. We show interesting seasonal variations along with differences between total and dissolved concentrations that can help inform decisions.

Our goal in this manuscript is to present and describe a unique, ppb-level dataset collected weekly in Utah Lake. These data, consisting of 1400 measurements, provide managers and regulators with information that can be used to develop management strategies that encourage beneficial use and mitigate potential harmful conditions. Our research goals present our characterization of this dataset to include insights such as which elements are commonly present in Utah Lake, in what phase, and their seasonal variation. For regulated elements, we compare our data to regulatory limits and discuss the implications. Unique aspects of this research include the analysis of the percent of measurements below the ppb-level detection limits and how often this occurred, and data for elements with concentrations at or near regulatory limits that characterize their behavior and variability. One distinctive insight includes a discussion of current aluminum (Al) regulations, which are based on total recovery standards but can easily be exceeded if there are suspended clay-minerals that contain Al but are not bioavailable, and another is the presence of Cu above regulatory limits and potential Cu sources. Both provide managers with information to help them make decisions on the monitoring and management of Utah Lake. While these findings and data are specific to Utah Lake, our data collection and analysis methods provide a case study for other researchers. This approach can be used in other water bodies to better understand complex geochemistry with relatively limited study requirements.

## 2. Materials and Methods

### 2.1. Field Study

We collected UL water samples approximately once per week over two sampling seasons: June–October 2021 and May–August 2022. We collected lake background samples roughly 400 m offshore (Figure 1).

At the beginning of the 2021 and 2022 sampling seasons, the depth of the water column was 2.5 m at the sample location, which is a typical depth for the lake at that distance from shore. By the end of the season, the depth was 1.5 m as the lake levels decreased due to evaporation and outflow through the Jordan River. The seasonal pattern of drawdown and refill is also typical for the lake. We analyzed the samples using ICP-OES analysis on a Thermo-Scientific iCAP7400 (ThermoFisher Scientific, Waltham, MA, USA) for both dissolved (filtered on a 0.45  $\mu\text{m}$  filter) and total (unfiltered and digested) samples.



**Figure 1.** Study locations: 0.4 km (0.25 mi) offshore, northwest of Lindon marina.

This is a unique and comprehensive longitudinal dataset for UL, with near-weekly samples collected on 28 days over two summer sampling campaigns. While spatially limited to the northeastern corner of the lake, we assume these data characterize general conditions in UL, though it is an area likely to be impacted by human activities.

The purpose of our study is to begin to characterize and understand the presence and behaviors of ICP-OES regulated DEs in UL. The sample locations are near the largest WWTP outfall and the former site of the Geneva Steel Plant, and they consequently represent an area more likely to have higher DE concentrations than other areas of the lake.

## 2.2. Sample Collection

Over the two sampling seasons, weather permitting, we collected water samples approximately weekly from June to October in 2021 ( $n = 15$ ) and from May to August in 2022 ( $n = 13$ ) near the northeast corner of Utah Lake (Figure 1). We followed sampling procedures from the “Standard Operating Procedure for Lake Water Sampling and Data Collection” from the UDEQ [37]. We used 1 L dip samplers that were triple-rinsed in lake water at the sample location prior to collecting the sample. After the triple-rinse, we inverted the dip sampler and submerged it to approximately elbow depth. We then inverted the sample cup to fill it with water. This reduced potential contamination from surface particles. We used the collected water to triple-rinse a new, pre-labeled, opaque 250 mL plastic sample bottle, then filled the bottles, leaving no headspace, and immediately placed the sample on ice in a cooler. Samples were either analyzed or frozen within a few hours of collection.

Coincident with water samples, we used YSI ProDSS (YSI, Yellow Springs, OH, USA) water quality sondes (probes) to collect data on chlorophyll-a and phycocyanin. For these measurements, we fully submerged the probes and logged the sensor data at a depth of ~30 cm.

### 2.3. Analysis Methods

#### 2.3.1. Laboratory Analysis

We split each water sample for ICP-OES analysis for dissolved (particles < 0.45 µm) and total concentrations. For dissolved concentrations, we filtered each sample using a new 0.45 µm membrane filter. For total concentrations, we performed microwave-assisted acid digestion using the EPA 3015A method. Table 2 summarizes the methods used for the laboratory analysis of the water samples.

**Table 2.** Lab analytes and methods.

Analyte	Model/Method	Equipment
DE Total (digested)	EPA 3015A	Thermo Scientific™ 7400 ICP-OES
DE Dissolved (filtered)	0.45 µ filter	Thermo Scientific™ 7400 ICP-OES (ThermoFisher Scientific, Waltham, MA, USA)

Table 3 lists the minimum detection limits (MDLs) for our ICP-OES method. MDLs vary by element but are approximately in the range of a few µg/L or at the part-per-billion (ppb) level. Se has the highest MDL of 7.36 ppb, while Ca has the lowest MDL at 0.02 ppb or 20.0 ppt. We selected ICP-OES for sample analysis because it is highly sensitive and allows us to easily analyze multiple trace elements in the UL water column.

**Table 3.** ICP-OES minimum detection limits (MDLs).

Analyte	Detection Limit (µg/L)	Analyte	Detection Limit (µg/L)
Aluminum (Al)	1.51	Molybdenum (Mo)	1.11
Arsenic (As)	4.74	Sodium (Na)	1.80
Boron (B)	1.26	Nickel (Ni)	2.29
Barium (Ba)	0.17	Phosphorus (P)	5.66
Calcium (Ca)	0.02	Lead (Pb)	4.50
Cadmium (Cd)	0.19	Sulfur (S)	2.22
Cobalt (Co)	1.16	Selenium (Se)	7.36
Chromium (Cr)	0.85	Silicon (Si)	7.20
Copper (Cu)	2.36	Strontium (Sr)	0.04
Iron (Fe)	0.80	Titanium (Ti)	0.58
Potassium (K)	5.10	Vanadium (V)	0.80
Magnesium (Mg)	0.04	Zinc (Zn)	0.60
Manganese (Mn)	0.21		

Additionally, water samples were tested for volatile suspended solids and total suspended solids. These results were correlated with ICP-OES measurements to further contextualize if total DE concentrations are associated with suspended sediments.

#### 2.3.2. Filtration

Dissolved particles are often defined as smaller than 0.45 µm, while suspended particles are larger than 0.45 µm [50,83]. To separate the dissolved from the suspended samples, we used Nyaflo® Membrane Disc Filters (Cytiva, Marlborough, MA, USA) with 0.45 µm pores. For this study, we used the terms dissolved and filtered interchangeably, and the terms total and unfiltered interchangeably. Total or unfiltered concentrations include both the suspended and dissolved fractions.

We found through our quality assurance analyses that trace elements on the filters contaminated our samples for As, B, K, Mg, Na, S, and Zn. We used the same brand of 0.45 µm filters for all samples, though due to the number of samples we analyzed, using the same filter batch was not possible. Overall, we found that the level of contamination in the filtered samples was small. In this study, we assumed that all samples were equally

impacted by filter contaminants, which allowed us to make comparisons and observe trends based on the sample results, which would still be accurate, even if the specific concentrations for these elements amounts are not.

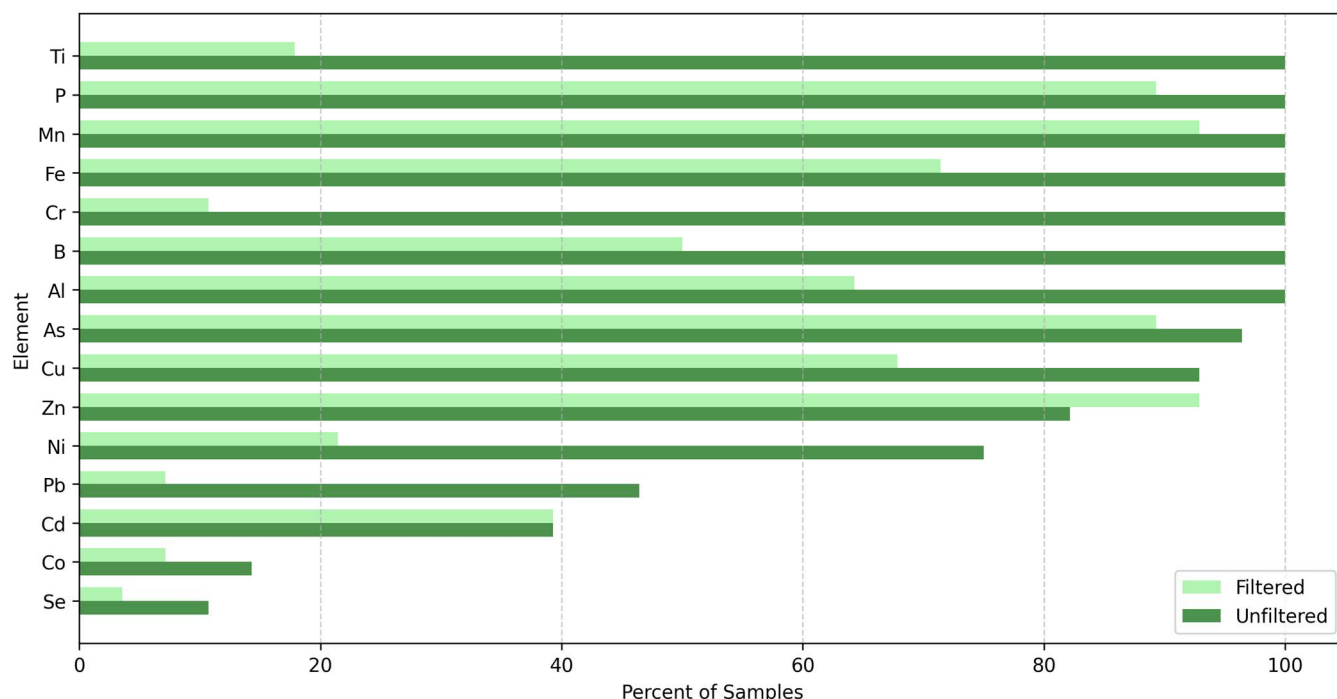
#### 2.4. Data Cleaning

Prior to analysis, we replaced measurements with concentrations below the MDL (Table 3) with a value of half of the MDL to preserve the detection of the element for analysis. The final dataset contained 28 samples collected over the 2021 and 2022 sampling periods (June through October and May through August, respectively), for which we analyzed 50 DE analytes, comprising 25 filtered and 25 unfiltered elements. The 2021 sampling season has a sampling gap from late July through late August due to equipment challenges but resumed in September and October. The 2022 sampling period was more consistent during the sampling season but was shorter, ending in August.

### 3. Results and Discussion: Regulation Levels

#### 3.1. MDL Analysis

We analyzed how often a detection for an element was below the MDL. This analysis showed how often a sample contained the element at or below detection limits. Figure 2 shows the percentage of DE measurements that were above the MDL for both unfiltered and filtered samples by element. If every sample for both the unfiltered and filtered samples was above the MDL, we excluded that DE from the graph for simplification. Forty percent of the DEs—Ba, Ca, K, Mg, Mo, Na, S, Si, Sr, and V—had all measurements above the MDL and are not included in the graph.



**Figure 2.** Percentage of samples for each ICP-OES analyte with measured concentrations above the MDL. Analytes with 100% detection in both filtered and unfiltered samples are excluded from the plot.

Co and Se concentrations fluctuate around the MDL in UL, with less than 20% of filtered or unfiltered samples having Co and Se concentrations above the MDL, and the remainder had concentrations only slightly above the MDL. These elements are present in UL, but in trace amounts.

Ti, Cr, Ni, and Pb were above the MDLs in less than 20% of filtered samples but were over the MDLs in unfiltered samples. These elements are mainly associated with suspended solids, with very low dissolved concentrations. Ti and Cr were above the MDLs in 100% of the unfiltered samples but only 20% of the filtered samples. From this, we can infer that these metals are mainly contained in the suspended solids and sediments of UL and are not present as dissolved particles.

Unfiltered Ni and Pb were above the MDLs in about 75% and 45% of cases, respectively. These elements similarly show that they are prevalent within the water column more in the unfiltered than the filtered state.

One metal, Zn, had more filtered samples over the MDL than unfiltered samples. This was caused by contamination from the filters, which consequently only affected the dissolved samples. We assume that although the dissolved Zn values are not accurate, the changes and trends in concentration are still relatively correct. As Zn is regulated and a vital micronutrient, we included the filtered results in this study. The unfiltered results are not contaminated because they did not come into contact with the filter and are therefore representative of our samples.

### 3.2. Specific Element Selection

We highlight eight ICP-OES elements—Cu, Zn, Ni, Al, Pb, P, Ba, and As—that were close to or over the most stringent applicable standards or exhibited interesting characteristics. The other five regulated DEs—B, Cd, Cr, Fe, and Se—are well below regulatory limits and we do not discuss or present them in this study.

We analyzed the distributions as box-and-whisker plots grouped by year (2021,  $n = 15$ ; 2022,  $n = 13$ ) and phase (filtered and unfiltered). In these plots, the line in the middle of the box is the median concentration, the box ends are the 25th and 75th percentiles, and the whiskers represent 1.5 times the interquartile range ( $1.5 \times \text{IQR}$ ). Outliers, or values outside  $1.5 \times \text{IQR}$ , are shown as dots.

We included lines on each boxplot indicating the most stringent acute (1 h average) and chronic (4-day average) standards (Table 1). All regulations are for dissolved (filtered) concentrations except Al and P, which are based on the total (unfiltered) concentration.

We did not follow state-approved methods of analysis. Consequently, our findings cannot be used to determine or indicate the impairment status of UL; we include the regulatory criteria only in order to provide context.

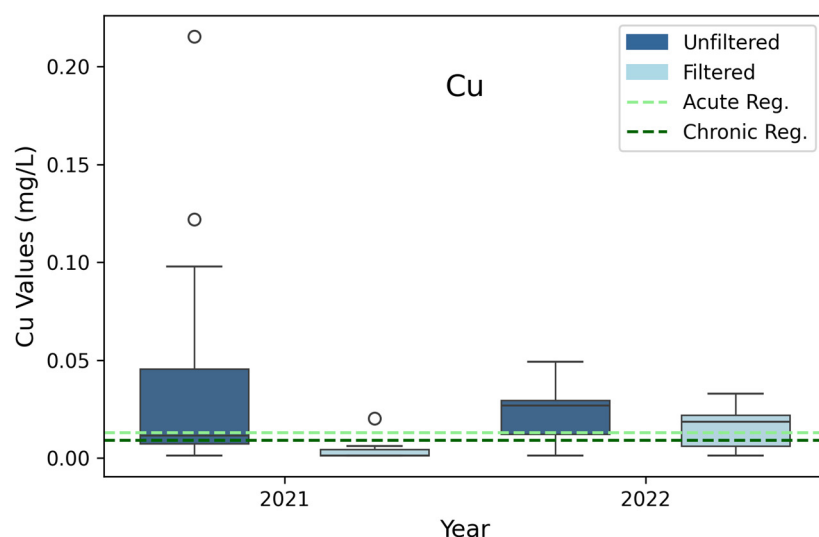
In the following figures, the dark-blue boxplots represent unfiltered data, and the light-blue boxplots represent filtered data. The dark-green dotted horizontal line represents the most stringent chronic standard, and the light-green line represents the most stringent acute standard. In a few cases where the acute and chronic regulations are the same, only the dark-green line is visible.

### 3.3. Copper (Cu)

Cu (Figure 3) has harmful effects at both excess and deficient concentrations. It is a micronutrient vital for a healthy ecosystem. For the unfiltered Cu concentrations, the median concentration for both years is above or close to the regulatory criterion, with concentrations more varied in 2021 ( $n = 15$  in 2021, and  $n = 13$  in 2022). Median filtered Cu concentrations are below the criteria in 2021 but above them in 2022.

For both the total and dissolved measurements, over 70% of the water samples contained concentrations of Cu over the MDL. Only dissolved Cu is regulated. All the filtered Cu samples in the 2021 sampling season, except one, were below both the acute and chronic regulatory criteria (Figure 3). However, in the 2022 sampling season (Figure 3), 7 of the 13 samples were above 0.013 mg/L, which is the acute standard. The highest concentration was 0.0329 mg/L. The State regulations for Cu are only applicable to the filtered samples, but for context, unfiltered Cu was above regulatory levels during both sampling seasons. For reference, the UL sediments on average contain 19.5 mg/kg of Cu. We found that unfiltered Cu concentrations are strongly correlated with unfiltered Ni and Fe, with Pearson

correlation coefficients (PCC) of 0.901 and 0.752, respectively. This might indicate mineral forms in the suspended solids.



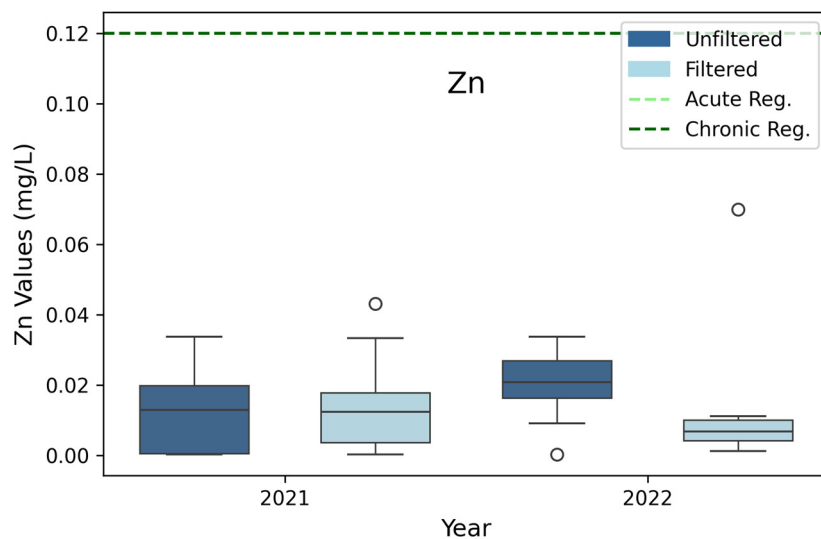
**Figure 3.** Distributions of Cu grouped by year, 2021 ( $n = 15$ ) and 2022 ( $n = 13$ ); and phase, unfiltered (dark blue) and filtered (light blue). The plot includes reference lines for 0.013 mg/L (acute light green) and 0.009 (chronic dark green) regulatory values. The circles are outlier data points.

The elevated levels of Cu may also be related to the use of copper sulfate-based algaecide, which the State began using to treat intense algal blooms in marinas in 2020 [84,85]. Our study location is located less than 1 km (0.6 miles) from Lindon Marina and 2 km (1.25 miles) from American Fork Marina. Lake currents are generally along the north shoreline towards the east, which could carry Cu from the American Fork Marina to our study site. The Utah Department of Environmental Quality (UDEQ) released an interim report on UL marina HAB treatments in which they found that “copper concentration increased considerably” in the days following treatment, but it returned to below toxic levels within a week [84]. The higher concentrations of dissolved Cu in 2022 relative to 2021 may also be due to the lower lake levels during the 2022 sampling season. On average, the lake had about 49 million cubic meters (39,700 ac-ft) less volume in 2022 than in 2021, which may have increased the water column concentrations of several elements [86]. Although the use of algaecide and low lake levels are one potential cause of the elevated levels of Cu observed in both years, they may not be the actual drivers.

Additional research is needed to identify and characterize the source and behavior of these metals. This is important as our results show that levels of dissolved Cu in the lake could be approaching harmful concentrations.

### 3.4. Zinc (Zn)

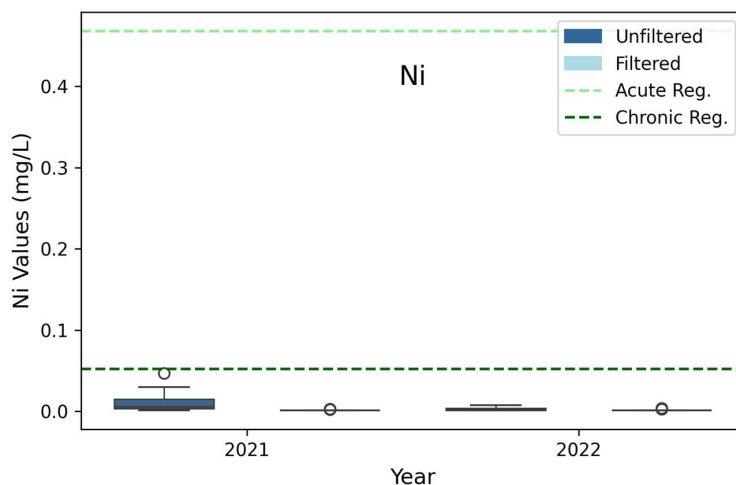
Zn (Figure 4) has both toxic effects at both excess and deficient concentrations. It is a micronutrient vital for a healthy ecosystem. Although filter contamination slightly increased concentrations of our filtered Zn samples, both filtered and unfiltered concentrations of Zn were well below the most stringent regulatory standard. More than 80% of Zn samples, both the filtered and unfiltered, measured above the MDL (Figure 2). In 2022, the difference between the total and the dissolved Zn was larger than in 2021, but it remained low. Filtered Zn concentrations, which are lower than those shown because of contamination from the filter, are low enough that it is more likely to be a limiting or colimiting nutrient for algal growth in UL because of deficient concentrations than a toxin due to high concentrations. For reference, Zn concentrations average 145.5 mg/kg in our UL sediment samples.



**Figure 4.** Distributions of Zn grouped by year, 2021 ( $n = 15$ ) and 2022 ( $n = 13$ ); and phase, unfiltered (dark blue) and filtered (light blue). The plot includes reference lines for 0.12 mg/L (acute light green) and 0.12 mg/L (chronic dark green) regulatory values. The acute and chronic standards for Zn are the same. Circles are outlier data points; the light green and dark green lines are the same value and only the dark green can be seen.

### 3.5. Nickel (Ni)

Ni (Figure 5) has adverse effects at both excess and deficient concentrations. It is a micronutrient vital for a healthy ecosystem. Both total and dissolved Ni concentrations in 2021 and 2022 were well below acute regulations, and just below chronic State regulations. Most of the Ni in UL appears to be associated with suspended solids rather than being dissolved in the water column. This matches Figure 2, which shows that ~70% of the unfiltered Ni measurements were over the MDL, compared to only ~20% of the dissolved measurements being over the MDL. These low Ni concentrations suggest that, like Zn, Ni is more likely to be a limiting or colimiting micronutrient because of deficient concentrations than a toxin in UL. Unfiltered Ni is correlated with unfiltered Cu and Fe, with PCCs of 0.901 and 0.735, respectively. This suggests a mineral phase in the suspended solids. For reference, our UL sediments samples average 11.5 mg/kg of Ni.

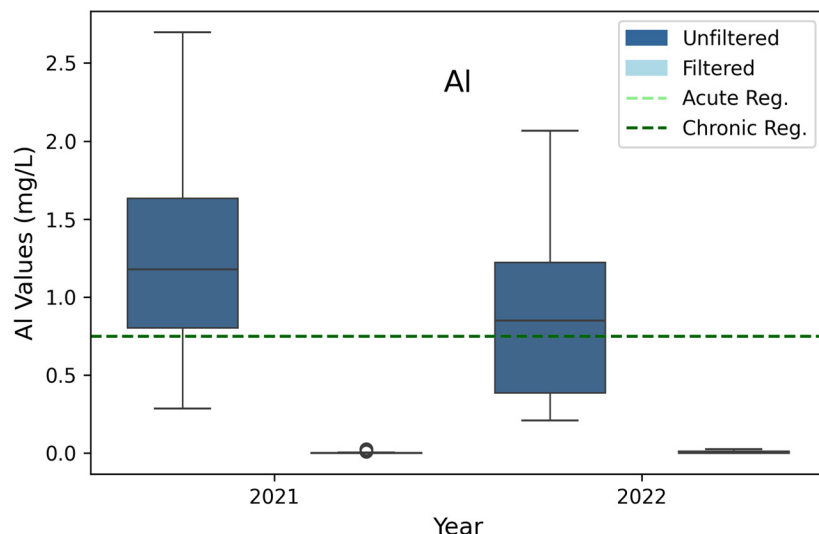


**Figure 5.** Distributions of Ni grouped by year, 2021 ( $n = 15$ ) and 2022 ( $n = 13$ ); and phase, unfiltered (dark blue) and filtered (light blue). The plot includes reference lines for 0.468 mg/L (acute light green) and 0.052 mg/L (chronic dark green) regulatory values. The light blue boxes are the 2nd and 4th from the left, but are so thin they show as black lines. The circles are outliers.

### 3.6. Aluminum (Al)

Al is a nonessential metal that is toxic at high concentrations [59–61,87]. State regulations state that Al concentrations are based on total recoverable Al, not dissolved. We assume that our unfiltered, digested samples represent total recoverable Al in UL water.

Unfiltered (total) Al median measurements are above both the acute and chronic criteria (Figure 6). Dissolved Al concentrations, however, are well below the criteria. Over 100% and 60% of the unfiltered and filtered Al measurements, respectively, were above the Al MDL (Figure 2).



**Figure 6.** Distributions of Al grouped by year, 2021 and 2022; and phase, unfiltered (dark blue) and filtered (light blue) with reference lines for 0.75 mg/L (acute) and 0.75 mg/L (chronic) regulatory levels. The light blue boxes are so thin they show as black lines. Circles are outlier data points; the light green and dark green lines are the same value and only the dark green can be seen.

The large difference between filtered and unfiltered concentrations indicates that Al is associated with suspended solids in the UL water column. The unfiltered concentrations represent a total digestion of all suspended solids, including clay minerals. Al and Si ratios in the samples are consistent with Al and Mg clay minerals, providing evidence that high Al concentrations are related to suspended clays. We attribute the high total Al concentrations to suspended clay particles in the water column.

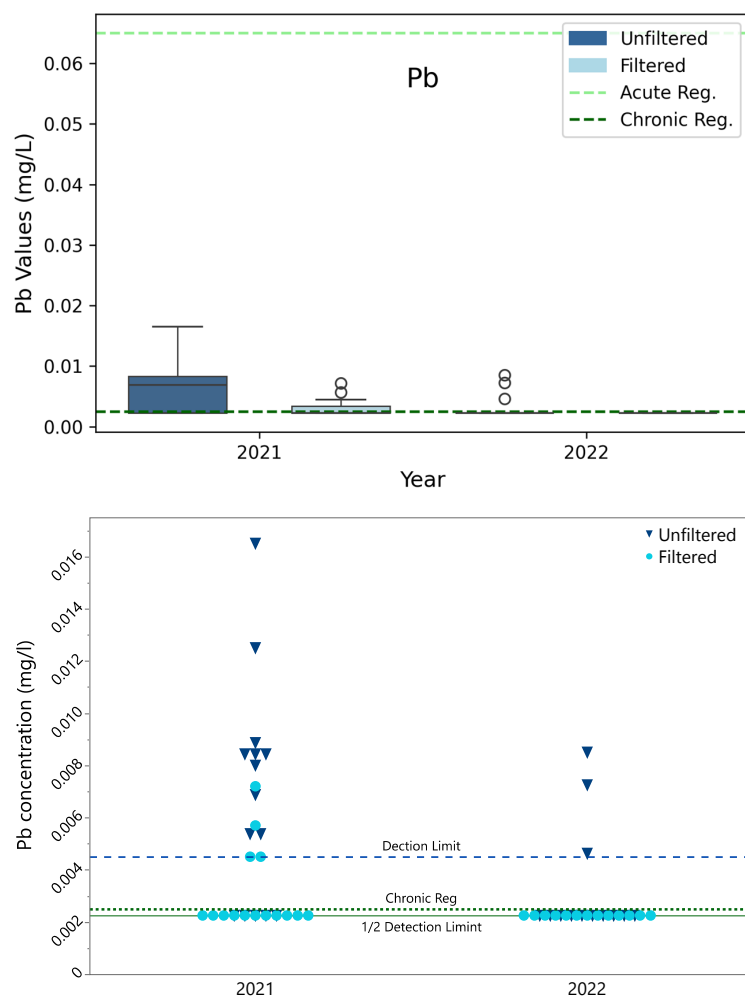
As noted earlier, we did not follow regulatory analysis methods. State water quality regulations require measurements of the dissolved (filtered) concentrations of elements, except for Al and P. For Al, the State regulations require “total recoverable” concentrations, which we took to mean unfiltered samples, but we did not use the State-approved method for Al analysis. Figure 6 shows that the unfiltered Al samples, when measured using full digestion and ICP-OES, exceed regulatory criteria in both the 2021 and 2022 seasons for both the acute and chronic regulations. The filtered (dissolved) samples, however, are well below the State regulations for both sampling periods. Our results show that dissolved Al concentrations—the phase likely to be bioavailable—are well below levels of concern. While unfiltered concentrations are above levels of concern, they are likely caused by suspended clays.

For reference, the average concentration of Al in the sediment is 8975.8 mg/kg, or almost 0.9% by mass. This concentration of Al is an order of magnitude higher than the amount of Zn and two orders of magnitude greater than the concentrations of Cu and Ni in the sediments.

Unfiltered Al was strongly correlated with unfiltered Ti, Mn, volatile suspended solids, and total suspended solids, with PCCs of 0.916, 0.750, 0.745, and 0.737, respectively.

### 3.7. Lead (Pb)

Pb is a nonessential metal that is toxic at high concentrations [59–61,87]. In our samples, dissolved Pb distributions had median values well below the acute standard for both filtered and unfiltered samples. However, the median values were often near or above their chronic State-regulation level (Figure 6). These results are misleading, however, as the acute Pb standard is below the ICP-OES method MDL for Pb. The bottom panel of Figure 7 shows that four dissolved Pb samples were above or at the acute standard, with two samples having values of 0.0072 and 0.0057 and two at the MDL of 0.0045. The remaining data were below the MDL and were set to one-half of the MDL to generate the plots and statistics. All the samples were well below the chronic standard. In 2021 and 2022, ten and three of the unfiltered samples were above the acute standards, respectively. Of the remaining samples, five and ten samples for 2021 and 2022, respectively, were below the MDL. Actual dissolved Pb concentrations could be lower (or higher) than this value. This is a limitation of our study caused by the lack of sensitivity of the ICP-OES compared to the regulatory standard. The EPA standard for Pb analysis uses a more sensitive ICP-MS method, which we did not use.



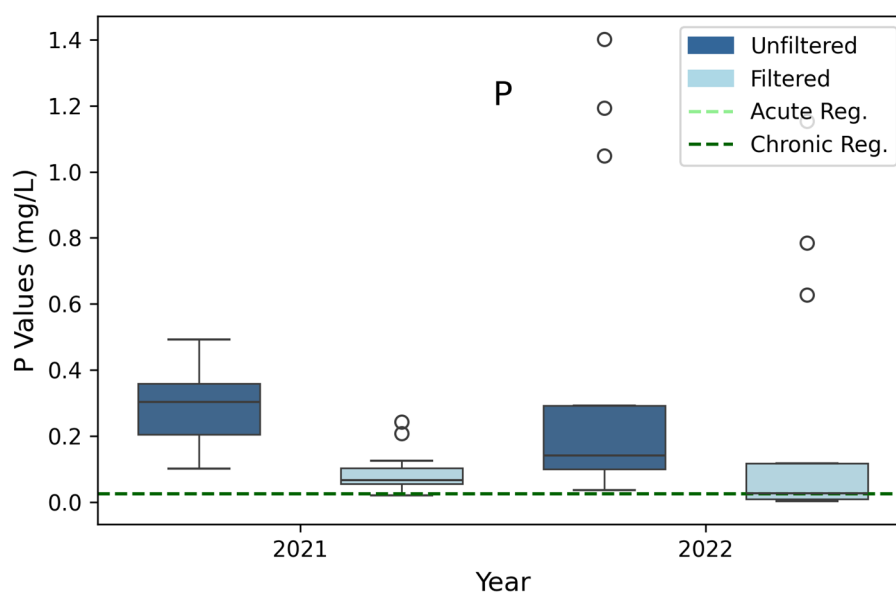
**Figure 7.** Distributions of Pb grouped by year (**Top Panel**), 2021 ( $n = 15$ ) and 2022 ( $n = 13$ ); and phase, unfiltered (dark blue) and filtered (light blue) with reference lines for 0.065 mg/L (acute) and 0.0025 mg/L (chronic) regulatory levels; Pb data, grouped by year (**Bottom Panel**) with lines for the detection limit, the chronic regulatory level, and  $\frac{1}{2}$  the detection limit. There are only two data points above detection limits, with two at detection limits. In the top panel, the boxes are so thin they show as black lines. Circles are outlier data points.

The Pb chronic regulation concentration is half of the ICP-OES MDL for Pb. Since we replaced 90% of measurements that were below the MDL with half of the MDL, most of the dissolved Pb measurements are exactly at the chronic regulation. Our data cannot be used to evaluate Pb concentrations relative to the chronic standard.

Figure 2 shows that except for two samples, all the samples with Pb concentrations over the MDL were for unfiltered samples, indicating that the Pb is associated with suspended solids. Both the filtered and unfiltered Pb distributions exhibit large differences between 2021 and 2022, with 2022 exhibiting significantly lower concentrations. We are unaware of any specific cause for this, although it is possible that higher levels of spring runoff in 2021 in comparison to 2022 led to greater inputs of Pb into the lake in 2021.

### 3.8. Phosphorus (P)

P is a limiting macronutrient in many freshwater lakes [88], but if sufficient P is available, other factors, such as micronutrients or light, can become limiting factors. Both total and dissolved P distributions are above the acute and chronic State regulations at the 25th percentile, with the exception of filtered samples in 2022, where the median is above the limits, but the 25th percentile is below (Figure 8). UL is listed as impaired due to high P concentrations, and the State has begun the process of developing a total maximum daily load (TMDL) for the lake [89]. Our data agree and show that P concentrations in both the filtered and unfiltered samples are above the regulatory levels. The MDL analysis further emphasizes the presence of P, with over 80% of the filtered and all the unfiltered samples over the MDL (Figure 2).



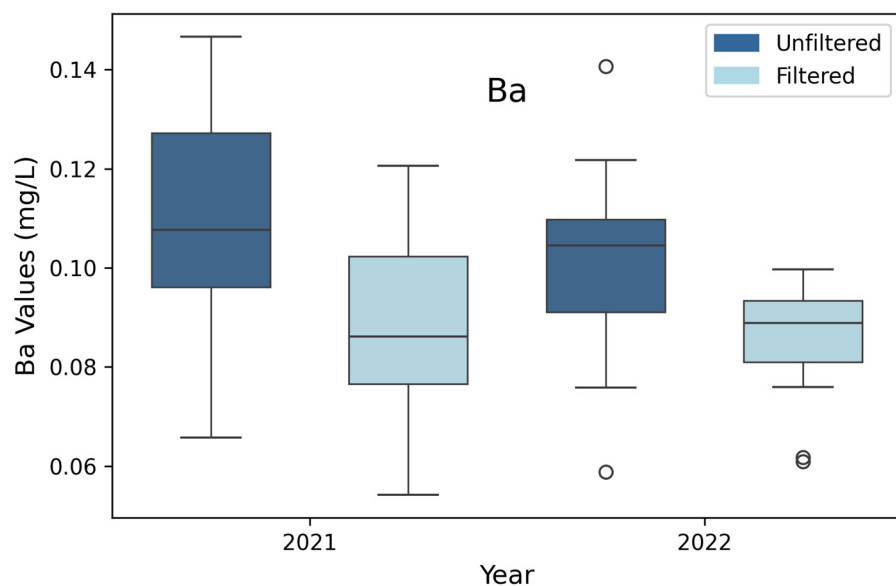
**Figure 8.** Distributions of P grouped by year, 2021 and 2022; and phase, unfiltered (dark blue) and filtered (light blue) with reference lines for 0.025 mg/L (acute) and 0.025 mg/L (chronic) regulatory levels. Circles are outlier data points; the light green and dark green lines are the same value and only the dark green can be seen.

Filtered and unfiltered P concentrations are strongly correlated, with a PCC of 0.949, but neither filtered nor unfiltered P correlates strongly with any other parameters we measured. The strong correlation between unfiltered and filtered P concentrations indicates that they are dependent on each other. That is, when unfiltered concentrations increase, due to higher turbidity or sediment resuspension, filtered or dissolved concentrations increase as well. These data support the idea that dissolved P concentrations in UL are governed by a sorption process that keeps sorbed (unfiltered) and dissolved (filtered) P in a constant ratio [1]. Because of the large reservoir of P in the UL sediments, water column concentrations are relatively stable, as was shown by Taggart, et al. [1]

Dissolved P concentrations were not correlated with probe measurements of either chlorophyll or phycocyanin, both of which indicate phytoplankton growth. This suggests that phytoplankton growth is not responding directly to bioavailable P concentrations, meaning dissolved P is not a limiting factor for algal growth in the UL water column. This analysis indicated that P concentrations, both filtered and unfiltered, are independent of other processes, which is again consistent with P in UL acting within a sorption-based system, where water column concentrations are in equilibrium with P-rich sediments of geologic origin [1].

### 3.9. Barium (Ba)

Ba is a potentially toxic metal that is not currently regulated in Utah [80]. Ba occurs naturally and is released by industrial processes, where it becomes mobilized as airborne particles that could enter UL through atmospheric deposition [75]. UL is susceptible to atmospheric deposition, as indicated by several studies [16,17,19]. Based on this, we have chosen to include a discussion on Ba. Figure 9 shows the Ba distributions for both filtered and unfiltered measurements. The Ba distributions for 2021 and 2022 sampling seasons are consistent and emphasize the presence of Ba within the UL ecosystem. There is a difference, but not a large one, between the unfiltered and filtered distributions. This indicates that a significant portion of Ba is dissolved, with some portions associated with suspended solids. About half of the Ba in UL is present in the dissolved phase.



**Figure 9.** Distributions of Ba grouped by year, 2021 and 2022; and phase, unfiltered (dark blue) and filtered (light blue). Circles are outlier data points.

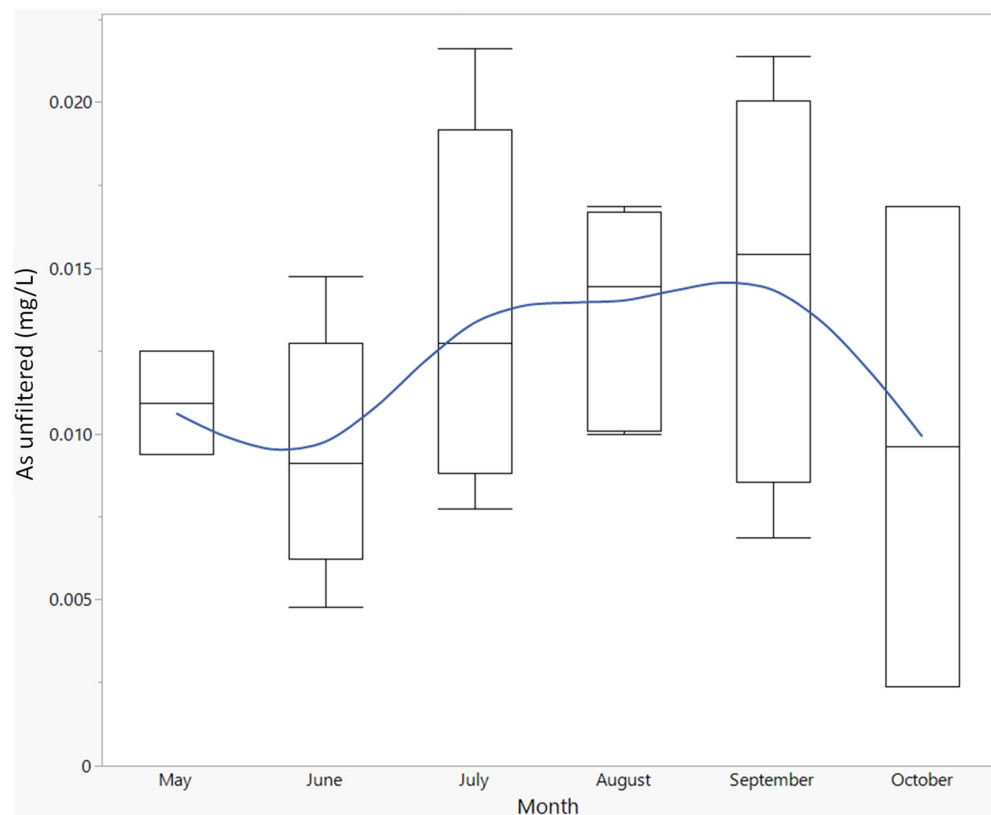
The U.S. EPA released a factsheet on Ba [90] in which they estimate that between the years of 1987 and 1993, 680 kg (1500 pounds) of Ba was released into the waters of Utah, including UL. The MDL analysis (Section 3.1 and Figure 2) revealed that Ba measurements for both unfiltered and filtered samples were over the MDL 100% of the time. Sediment samples show that there is an average of 179.5 mg/kg in the UL sediments. The Ba boxplot (Figure 9) shows that the median unfiltered and filtered samples for Ba are about the same in both the 2021 and 2022 sampling seasons.

Short-term health effects of Ba include gastrointestinal disturbances and muscular weakness, with long-term effects including high blood pressure for levels above the maximum contaminant limit (MCL) [90]. The U.S. EPA MCL for Ba is 2 mg/L for drinking water—nearly two orders of magnitude higher than any of our measured values in UL [90,91]. Within the State of Utah, Ba is only regulated for the designated use of 1C or “Domestic Source” (which does not apply to Utah Lake) at a maximum dissolved limit of

1 mg/L [80]. The Ba distribution plot (Figure 9) shows that both the unfiltered and filtered measurements are well below the EPA MCL of 2 mg/L for drinking water and the 1 mg/L Utah 1C regulation for domestic source water.

### 3.10. Arsenic (As)

Both filtered and unfiltered As concentrations were consistently below the most restrictive aquatic use, which in this case was for agriculture such as irrigation, which has values of 0.10 mg/L for both acute and chronic standards (Table 1). Unfiltered concentrations are near the drinking water standards which are 0.01 mg/L, however, Utah Lake water is not current use for domestic supply. As exhibited a seasonal trend in both years of our data (Figure 10). We postulate that As increases in late summer because groundwater inflows with higher concentrations of As become a larger portion of lake inflows at that time of year [92,93]. As is positively correlated with phycocyanin probe measurements and inversely correlated with filtered Al concentrations. We believe these correlations are spurious and are only present because all three variables have seasonal variation. These variations, however, are due to different processes related to seasonal changes, not chemical or ecological processes involving all three parameters. We could not identify any likely physical process that would drive this correlation. While unfiltered As concentrations are well below the most stringent beneficial use requirements for agricultural and aquatic wildlife, they are, on average, slightly above domestic use standards.



**Figure 10.** Annual trend in unfiltered (total) arsenic (As). The blue line is a smoothed trend through the mean values.

## 4. Conclusions

In this study, we used ICP-OES to examine elements in UL water at a ppb level. We found that total recoverable concentrations of Al in UL exceeded regulatory criteria when using our analytical methods, although we attribute these elevated levels to nonreactive, clay-dominated suspended sediment. Concentrations of dissolved Al, which would be bioavailable and have the potential to be toxic, were well below regulatory criteria. We

consistently measured concentrations of Cu and P above or very close to regulatory limits. The high concentrations of P in UL are well known, but the elevated levels of Cu are not widely recognized, and as these elements have the potential to impair the beneficial uses of UL, further studies should be conducted. Cu especially should be considered because of an ongoing program using Cu-based algaecide to treat intense algal blooms in UL.

Concentrations of other regulated elements remained well below regulatory criteria, which was interesting because anthropogenic inputs of these elements to UL have been and may still be large—the low concentrations we observed demonstrate the lake’s high capacity for absorbing, degrading, and removing pollutants. Additionally, we found no regulations regarding Ba for UL’s designated beneficial uses; however, concentrations of Ba are below drinking water limits, despite nearby anthropogenic sources.

We found a strong correlation between unfiltered (total) and filtered (dissolved) P, indicating that the constant resuspension of P-rich lakebed sediments acts as a source of bioavailable P in the water column. This relationship indicates that dissolved P in the water column is in equilibrium with P sorbed onto lakebed sediments and suspended sediments in a sorption-dominated system.

UL, except for Al and P, is well within compliance for all regulated DE elements associated with its designated uses. It may be beneficial for the State to create a separate designated use category for UL, as has been done with the Great Salt Lake. The DE standards drive many regulatory decisions for point sources around UL, and as such, they could be further considered with the uniqueness of UL.

**Author Contributions:** Conceptualization, R.A.V., K.B.T. and G.P.W.; methodology, R.A.V. and G.P.W.; field data collection, R.A.V., K.B.T., J.B.T., R.L.R., A.C.C., L.M.W. and M.J.O.; software, R.A.V.; validation, J.B.T., R.L.R., A.C.C., A.W.M. and R.B.S.; resources, A.W.M. and G.P.W.; data curation, R.A.V., K.B.T. and A.C.C.; writing—original draft preparation, R.A.V.; writing—review and editing, R.A.V., K.B.T., J.B.T., R.L.R., A.C.C., G.P.W., A.W.M. and R.B.S.; visualization, R.A.V. and G.P.W.; supervision, G.P.W.; project administration, R.A.V., K.B.T. and G.P.W.; funding acquisition, A.W.M. and G.P.W. All authors have read and agreed to the published version of the manuscript.

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**Data Availability Statement:** The data presented in this study are available on request from the corresponding author due to contractual restrictions on data distribution.

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