Regulated Coal Mine Wastewater Contaminants Accumulating in an Aquatic Predatory Beetle (*Macrogyrus rivularis*): Wollangambe River, Blue Mountains New South Wales Australia

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Abstract: This study investigates contaminants from a single coal mine wastewater discharge released to the Wollangambe River accumulating in an aquatic predatory beetle (*Macrogyrus rivularis*). The study was undertaken within the Wollangambe River and its surrounding tributaries. The coal mine wastewater discharge is regulated by the New South Wales Environment Protection Authority and the regulation of the pollutants only concentrates on water column concentrations. The majority of the Wollangambe River flows within the World Heritage Greater Blue Mountains National Park and is protected through many layers of legislation from state to federal and international (Threatened Species Conservation Act 2005, Environment Protection and Biodiversity Conservation ACT 1999, United Nations Educational, Scientific and Cultural Organization 2000). Results show that many contaminants are at statistically higher concentrations within the water column, stream sediment and beetles sampled when compared between reference and impacted sample locations. Analysis of Similarity (ANOSIM) found significant differences for contaminants in beetles sampled at impacted sites compared to reference sites with no significant difference recorded between reference sites. Biota and/or Environmental matching (Best) found Manganese, Cobalt, Nickel and Zinc as the factors which have the greatest influence in differences. The implications that contaminants from the regulated wastewater being discharged may be accumulating within aquatic biota is of major concern as the regulation of the wastewater only concentrates on water column pollutants and is not taking into account the greater environmental ramifications of the pollution.

Keywords: Water Pollution, Bioaccumulation, Heavy Metals, Macroinvertebrates, Freshwater Ecology, Coal Mine Pollution, Regulated Water Pollution

1. Introduction

In recent years anthropogenic activities and the subsequent contamination from pollutants of global ecosystems has reached unparalleled heights. Exponentially increasing industrial and mining activates have led to further environmental pollution through wastes produced by these activities. A host of pollutants from mining and industrial activities, when released to the environment, have the potential to accumulate within biota at toxic concentrations and cause chronic ecological impacts in ecosystem food chains [1-4]. Metal pollution is a global environmental issue and has been for many decades. In many cases metal pollutants are directly discharged into waterways from anthropogenic activities. Many of these metal pollutants become absorbed to suspended particulates within the water column. Once absorbed to these suspended particulates the metal contaminated suspended particulates eventually deposit
Aquatic ecosystems are often more highly sensitive to contamination from anthropogenic activities, especially as waterways are often used as sources of discharge for many industrial wastes [8]. Heavy metal contamination within aquatic environments can persist much longer than terrestrial organic pollutants. This is due to the lack of a “biodegradation function” of heavy metals in aquatic ecosystems in comparison to a terrestrial ecosystem [9, 3]. Bioaccumulation of contaminants within fish species have been recorded worldwide as a result of metal and coal mining activities [3, 10-13] along with other aquatic fauna and flora [14, 15].

One study found bioaccumulation of arsenic and selenium within caddisflies of coal ash polluted sites [16] whilst increases in the levels of pollutants were also recorded for turtles, crayfish, tadpoles and varying fish species by [17]. Bioaccumulation of wastewater contaminants within macroinvertebrates from metal mining activities have been reported in Canada, Australia and North America [18-23].

The pH of water is identified as an important factor directly linked to the speciation and bioaccumulation of metals and metalloids. pH will affect the bioavailability of metals and metalloids by influencing their solubility and subsequent ability of bioaccumulate to a medium [17, 24]. The individual constituents of coal can also influence metal bioavailability [17]. Atkinson et al. found that lower water pH allowed for greater bioavailability and sequestration of heavy metals in biota. They found that iron and manganese oxidatively precipitated increasingly as pH decreased. This decreasing pH had a secondary effect which allowed a greater sequestration rate of lead and zinc [25].

Coal mining practices are well documented to contribute to an array of differing environmental problems including air pollution, fire hazards, ground subsidence or deformation, surface and or ground water pollution. Surface water pollution is a major environmental problem associated with coal mining and it occurs through the discharge of mine wastewaters that are contaminated by various disturbances associated with mining practices [26-30]. Water pollution from coal mining occurs as large volumes of surface and groundwater are required to be removed from most underground coal mines. This is generally through the pumping of the wastewater to the surface. Without this, groundwater would flood most sections of the underground mining operation [27, 31]. Coal mine wastewater will often be contaminated due to the disturbance of the local geology associated with mining activities. The exact nature of the water contamination will vary depending on local factors such as groundwater geochemistry, hydrology and mineralogy of the local strata. In addition to the physical activity of the mining operation and the removal of the wastewater, other activities will also often contaminate water used throughout a mining plant which can include; coal washing and the inclusion of other wastes generated by the surface operation at the mine such as sewage wastes [31].

Coal mine wastewater discharges in New South Wales, Australia are regulated by the New South Wales Environmental Protection Authority (NSW EPA) and environmental protection of receiving waterways is implemented through Environmental Protection Licenses (EPL’s), under the Protection of the Environment Operations Act 1997 (POEO Act 1997) [34]. EPL’s set discharge limits for water quality and chemical properties in which coal mine wastewaters that are discharged to the environment must adhere to [32, 33]. In many cases the EPL’s for coal mine wastewater discharges are failing to protect the receiving waterways ecosystems by failing to identify ecologically hazardous chemicals in the waste discharges that may accumulate or become persistent in the receiving waterways ecosystem. This non-wholistic regulation is failing to protect the aquatic environment of these receiving waterways by not taking into account legacy pollutants. Studies have been conducted on the impacts of metal bioaccumulation from coal mining activities on terrestrial and aquatic plant and fish tissue with none investigating the bioaccumulation of pollutants from actively licenced and regulated coal mine wastewater discharges.

This study investigates if the contaminants from a single licenced and regulated coal mine wastewater discharge is accumulating in the ambient environment. More specifically within stream sediments and a single species of aquatic predatory beetle (Macrogyrus rivularis) from the family Gyrinidae within the high conservation Wollangambe River. Macroinvertebrates are widely used as ecological indicators of water quality due to well-established methods and ease of sampling [34]. Macroinvertebrates are regarded as being effective indicators as they have relatively long-life cycles and different taxonomic groups have differing sensitivities to disturbance and water pollution [34, 35]. There is a large amount of well-established literature on freshwater macroinvertebrates and their applications to assess various human-induced anthropogenic disturbances including a broad variety of pollution types [35] including coal mine drainage [27, 30, 36, 37]. The beetle species (Macrogyrus rivularis) was used as it was identified as a reliable source of sampling, its predacious habit allowing it to consume other aquatic fauna which may be contaminated and due to its long-life cycle (2 years) allowing for a prolonged period in which it can accumulate contaminants from the wastewater [38]. This stream also allows for a great test case as other than the coal mine wastewater the stream resides within an untouched pristine catchment with the two reference locations having no anthropogenic influence on water quality or chemistry [33, 39].

It is hypothesised that the contaminants from the single coal mine wastewater are accumulating within a single species of predacious aquatic beetle found within the Wollangambe River downstream of the coal mine wastewater discharge.
2. Methods

This study was conducted at two upland streams found within the Blue Mountains area of Sydney, Australia. One stream being the Wollangambe River, the other Bell Creek which is a tributary of the Wollangambe River. Both of which flow mostly within the Blue Mountains World Heritage National Park Estate and the majority of their catchments are naturally vegetated (Figure 1). Four sample sites were used in total for this study, Wollangambe 1 (W1) and Bell Creek (Bell) both as reference sites and Wollangambe 3 (W3) and Wollangambe 5 (W5) as impact sites. The reference site (W1) is located approximately 200m above the discharge point and is a first order stream [40], whilst (W3) is approximately 500m downstream and is a second order stream [40]. Both sample locations share similar altitudes between 1025m and 960m above sea level (Figure 1 and Table 1). Two sample locations were located approximately 18km downstream of the coal mine. One being the reference site (Bell Creek) and is a naturally vegetated tributary of the Wollangambe River and is a first order stream [40]. Bell Creek’s paired impact site (W5) was located on the Wollangambe River and was approximately 200m downstream of the confluence of Bell Creek and the Wollangambe River and is a second order stream [40]. Both lower sample sites share similar altitudes between 760m and 740m above sea level (Figure 1 and Table 1). Stream order level was determined by the Strahler 1952 method [40].

Figure 1. Map of Australia and sample locations and Clarence colliery. Red X depicts approximate location of study area. W1 (reference) sample location is located approximately 200m above the coal mine wastewater discharge and its paired (impact) sample location W3 is located approximately 200m downstream of the wastewater inflow. W5 (impact) sample location is located on the Wollangambe River approximately 18m downstream of the wastewater inflow and its paired (reference) sample location Bell Creek (Bell) is a clean tributary of the Wollangambe River and is also located approximately 18km downstream of the coal mine wastewater discharge just upstream of W5.

Table 1. Sample location name, longitude and latitude, altitude (Metres above sea level) and stream order of the four sample locations used in this study. Stream order is derived from the Strahler method (Strahler 1952).

| Sample location      | longitude    | latitude           | Altitude (ASL) | Stream order |
|----------------------|--------------|--------------------|----------------|--------------|
| W1 (reference)       | 150.249101 E | -33.455964 S       | 1025 m         | 1            |
| Bell Creek (reference)| 150.353770 E| -33.490335 S       | 760 m          | 1            |
| W3 (downstream)      | 150.257359 E | -33.455673 S       | 960 m          | 2            |
| W5 (downstream)      | 150.355208 E | -33.487474 S       | 740 m          | 2            |

Five replicated water samples (grab samples) and insitu field water quality parameters were sampled and measured at the same time as sediment and beetle sampling occurred from the same four sample locations above. Field meters included a TPS WP-82V meter with a YSI dissolved oxygen probe for Dissolved Oxygen, TPS WP-88 Turbidity meter with a TPS turbidity sensor for Turbidity and a TPS WP-81 Conductivity, pH and Temperature meter with TPS Conductivity and Temperature probe and a TPS submersible k407 pH sensor. Five field grab samples were collected in commercial preserved sample containers provided by a commercial laboratory (EnviroLab) (to allow for quality control of sampled water) at each site (20 total) and analysed using standard methods (APHA 1998) by Envirolab (Chatswood, NSW) a National Associations of Testing Authorities accredited laboratory for fifteen metals (Aluminium, Barium, Boron, Cadmium, Chromium, Cobalt, Copper, Iron, Lead, Manganese, Molybdenum, Nickel, Strontium, Uranium and Zinc).

Stream sediments were sampled at the same time as water and beetle samples were collected from the same four sample locations. This was achieved by sampling stream sediments within unused commercial laboratory sample containers provided by (EnviroLab) (to allow for quality control of sampled sediments) in a zone of accumulated sediment, following standard methods recommended by the Victorian Environmental Protection Authority [41]. Samples were placed into sealed glass sampling jars supplied by a commercial laboratory and stored. Five samples were collected at each sample location (20 total) and analysed using standard methods (APHA 1998) by Envirolab (Chatswood, NSW) a National Associations of Testing Authorities accredited laboratory for the same fifteen metals analysed for water samples (Aluminium, Barium, Boron, Cadmium, Chromium, Cobalt, Copper, Iron, Lead, Manganese, Molybdenum, Nickel, Strontium, Uranium and
Six replicated predacious aquatic beetle samples (Macrogyrus rivularis) from the family Gyrinidae were collected on one occasion at all four sample sites (24 total beetles from 4 sites). The beetle species was used as it was identified as a reliable source of sampling, its predacious habit allowing it to consume other aquatic fauna which may be contaminated and due to its long-life cycle (2 years) [38]. Sampling was achieved by identifying and catching beetles from the surface or mid water column as they retreated using a macroinvertebrate sampling ‘kick’ net with a frame of 30 x 30 cm and 250 µm mesh. Beetles were caught and stored in deionised water within individual sample location sample containers. These containers were sealed and labelled on collection and dispatched to the commercial laboratory (EnviroLab) and analysed for 21 metals (Aluminium, Arsenic, Barium, Beryllium, Boron, Cadmium, Chromium, Cobalt, Copper, Iron, Lead, Manganese, Molybdenum, Nickel, Selenium, Silver, Strontium, Titanium, Uranium, Vanadium and Zinc).

In the laboratory Ultra High Purity (UHP) water was used to wash the beetles and remove any potentially contaminated residues from their outer surface (precipitates). They were then kept in individual sealed plastic containers. The samples were dried in a 60-degree Celsius oven for 48 hours prior to being ground using a clean pestle and mortar. Approximately 0.05 grams of the ground samples were weighed into plastic digestion tubes. High purity Nitric and Hydrochloric acid was added to the tubes and the samples were digested at 98 Degrees Celsius for 1.5 hours. The samples were then made up to 20mL with UHP water. These samples were analysed by Inductively Coupled Mass Spectrometry (ICP-MS). Quality control measures were used to ensure the integrity of the results including blanks and spiked laboratory control samples (LCS).

Multivariate analysis was performed using Single Factor ANOVA and the software package Primer 6 for analysis of similarity (ANOSIM) to test for differences between combined reference sites (W1 and Bell) and both individual impact sites (W3 and W5). Water quality and chemistry, stream sediment chemistry and aquatic predacious beetle chemistry individually. Primer 6 was also used to identify which water quality and chemistry parameters influenced the dissimilarity in beetle contaminants through a BIOENV test. Percentage increases from water column concentrations (µg/L) to aquatic beetle concentrations (µg/kg) were calculated to assess potential uptake differences between reference water chemistry to impacted water chemistry.

3. Results

Mean water column concentrations of Barium, Boron, Cobalt, Iron, Manganese, Nickel, Strontium and Zinc were found to be statistically different from reference to impacted sample sites (Table 2). Reference sites recorded mean barium of 9.5 and 8.6 µg/L whilst in comparison downstream sites recorded mean barium of 20.8 and 16.0 µg/L. Reference sites recorded mean boron of 8.0 µg/L in comparison downstream sites had mean boron of 9.1 and 9.5 µg/L. Reference site Cobalt concentrations were below laboratory detectable limits. Whilst in comparison downstream sites recorded mean cobalt of 26.5 and 1.7 µg/L. Mean iron concentrations at reference sites was 300 and 381.7 µg/L whilst in comparison downstream iron was of 134.3 and 66.8 µg/L. Reference sites recorded mean manganese concentrations of 22.8 and 35.2 increasing downstream to a mean of 178.5 and 29.2 µg/L. Reference sites recorded mean nickel of 0.5 µg/L just above the laboratory threshold. Whilst in comparison downstream mean nickel was 76.7 and 17.7 µg/L. Reference strontium concentrations were recorded a mean of 3.6 and 5.2 µg/increasing downstream to 59 and 36.2 µg/L. Reference sites recorded mean zinc of 2.4 and 3.7 µg/L and downstream sites a mean of 105 and 20.8 µg/L. Cadmium, chromium, copper, lead, molybdenum and uranium were all at concentrations within water that were below laboratory detectable limits (Table 2).

Mean stream sediment concentrations of Aluminium, Barium, Cadmium, Chromium, Cobalt, Copper, Iron, Manganese, Molybdenum, Nickel, Strontium, Uranium and Zinc were found to be statistically different from reference to impacted sample sites (Table 3). Bell Creek (reference) recorded the lowest mean aluminium in stream sediment of 233.3 mg/kg, whilst in comparison W1 (reference site), W3 and W5 (impacted sites) had mean aluminium of 1248, 5083 and 1034 mg/kg respectively. Reference sites recorded mean barium of 4.75 and 18.0 mg/kg whilst downstream recorded mean barium of 101.6 and 7.5 mg/kg. Both reference sites Bell Creek and W1 as well as the downstream site W5 all recorded cadmium concentrations lower than laboratory detectable limits. W3 recorded mean cadmium of 0.47 mg/kg. Bell Creek, W1 and W5 all recorded replicates which were below laboratory detectable concentrations for chromium. W1 and W5 both recorded mean chromium of 1 mg/kg whilst Bell Creek recorded a mean of 0.25 mg/kg. In comparison W3 recorded chromium concentrations between 2-8 mg/kg and a mean of 4 mg/kg. The reference site Bell Creek recorded below detectable laboratory concentrations of cobalt for all replicates sampled. W1 ranged between below laboratory concentrations and 5 mg/kg, whilst in comparison W5 was found to have mean cobalt of 6 mg/kg and W3 recorded mean concentrations of 552 mg/kg. The reference sites Bell Creek recorded below detectable laboratory concentrations of copper for all replicates sampled and W1 recorded a mean of 3.5 mg/kg. Whilst the downstream sites W3 and W5 recorded mean copper of 17.0 and 2.4 mg/kg respectively (Table 3).

Bell Creek and W1 recorded mean iron of 1443 and 4000 mg/kg whilst in comparison W3 and W5 recorded mean iron of 9950 and 1982 mg/kg. Manganese recorded at the reference sites (Bell and W1) was mean 13.3 and 127.5 mg/kg in comparison W3 and W5 recorded mean manganese of 5474 and 76 mg/kg respectively. Bell Creek, W1 (reference sites) and W5 (impacted site) samples recorded concentrations of molybdenum within sediment below laboratory detectable limits. Whilst in comparison W3 (downstream site) had mean molybdenum of 4
mg/kg. W3 recorded one replicate below laboratory detectable limits for molybdenum. Bell Creek recorded mean nickel below laboratory detectable limits whilst W1 (reference sites) recorded mean nickel of 2 mg/kg. In comparison W3 and W5 (downstream sites) recorded mean nickel of 606.7 and 19 mg/kg with all samples recording detectable concentrations. Reference site mean strontium was found to be 1.25 and 5 mg/kg and ranged from below laboratory detectable limits and 9 mg/kg across both sites. Whilst in comparison W3 and W5 (downstream sites) recorded mean strontium of 12.3 and 2.8 mg/kg respectively. All sample locations recorded low concentrations of uranium in sediment with the reference stream Bell Creek recording all replicates at concentrations below laboratory detectable limits. W1 was recorded between below laboratory limits and 0.2 mg/kg with a mean of 0.06 mg/kg. W5 was similar to both reference uranium concentrations measuring between below laboratory detectable limits and 0.3 mg/kg with a mean of 0.08 mg/kg. Whilst in comparison W3 mean uranium of 0.53 mg/kg and ranged between 0.1-1.4 mg/kg. The reference site Bell Creek recorded all replicates at concentrations below laboratory detectable limits for Zinc. W1 was found to have a mean of 1.75 mg/kg. Whilst in comparison W3 and W5 (downstream sites) had mean zinc of 734 and 25 mg/kg respectively (Table 3).

| Water Parameter | Statistics | Reference Sites |
|-----------------|------------|-----------------|
| pH              | F value    | p value         | df  |
|                 |            |                 | 2   |
|                 | 1544.1     | 4.8E-40         | 2   |
| Electrical Conductivity | 11232.8  | 4.9E-58         | 2   |
| Dissolved Oxygen | 0.28      | 0.75            | 2   |
| Turbidity       | 6.29       | 0.004           | 2   |
| Water Temperature| 10.7      | 0.0002          | 2   |
| Aluminium       | 4.05       | 0.03            | 2   |
| Barium          | 65.3       | 3.4E-12         | 2   |
| Boron           | 31.6       | 1.09E-07        | 2   |
| Cadmium         | n/a        | n/a             | 2   |
| Chromium        | n/a        | n/a             | 2   |
| Cobalt          | 5.57       | 0.008           | 2   |
| Copper          | n/a        | n/a             | 2   |
| Iron            | 18.7       | 3.75E-06        | 2   |
| Lead            | n/a        | n/a             | 2   |
| Manganese       | 4.24       | 0.02            | 2   |
| Molybdenum      | n/a        | n/a             | 2   |
| Nickel          | 11.3       | 0.0002          | 2   |
| Strontium       | 88.6       | 5.38E-14        | 2   |
| Uranium         | n/a        | n/a             | 2   |
| Zinc            | 7.50       | 0.002           | 2   |

Table 2. Water quality and water chemistry parameters, sample locations, Single Factor ANOVA. F value, p value and degrees of freedom (df) between groups indicates statistical difference between merged upstream (reference) and individual downstream (W3 and W5) sample locations, range and mean for water chemistry for all four sample locations. All water chemistry data is measured in µg/L, pH in pH units, Electrical Conductivity in µs/cm, Dissolved Oxygen in % saturation, Turbidity in Nephelometric Turbidity Units and Temperature in degrees Celsius. BD = below laboratory detectable limits, n/a = not any.

| Water Parameter | W3 Range | W3 Mean | W5 Range | W5 Mean |
|-----------------|----------|---------|----------|---------|
| pH              | 7.27-7.37| 7.29    | 7.35     | 6.97-6.98|
| Electrical Conductivity | 316-316   | 316     | 316      | 217.8-217.9|
| Dissolved Oxygen | 92.4-92.5 | 92.4   | 91.2     | 89.6-90.1 |
| Turbidity       | 0.5-0.7  | 0.62    | 3.4      | 0.4-1.2  |
| Water Temperature | 12.4-12.4 | 12.4  | 12.4     | 10.8-10.8|
| Aluminium       | BD-210   | 71.7    | 50       | BD-40   |
| Barium          | 19-23    | 20.8    | 20.5     | 14-19   |
| Boron           | 1-9      | 9.5     | 9.5      | 9-10    |
| Cadmium         | BD       | BD      | BD       | BD      |
| Chromium        | BD       | BD      | BD       | BD      |
| Cobalt          | 3-47     | 26.5    | 28       | BD-3    |
| Copper          | BD       | BD      | BD       | BD      |
| Iron            | 38-420   | 134.3   | 87.5     | 42-82   |
| Lead            | BD       | BD      | BD       | BD      |
| Manganese       | 43-430   | 178.5   | 200      | 22-34   |
| Molybdenum      | BD       | BD      | BD       | BD      |
| Nickel          | 24-130   | 76.7    | 70       | 8-25    |
| Strontium       | 45-68    | 59      | 59       | 28-47   |
| Uranium         | BD       | BD      | BD       | BD      |
| Zinc            | 21-190   | 105     | 99       | 8-31    |

Table 2. Continued.
### Table 3

| Sediment/Parameter | Statistics | Reference Sites |
|--------------------|------------|-----------------|
|                   | F value    | p value | df | W1 Range | W1 Mean | W1 Median | Bell Range | Bell Mean | Bell Median |
| **Aluminium** | 9.52      | 0.003 | 2 | 790-2200 | 1247.5 | 1000 | 190-280 | 233.3 | 230 |
| **Barium**     | 16.1      | 0.0002 | 2 | 10-33 | 18 | 14.5 | 3-8 | 4.75 | 4 |
| **Boron**      | 103.9     | 6.03E-07 | 2 | BD | BD | BD | BD | BD | BD |
| **Cadmium**    | 10.2      | 0.003 | 2 | BD | BD | BD | BD | BD | BD |
| **Chromium**   | 9.99      | 0.003 | 2 | BD-2 | 1.0 | 1 | BD-1 | 0.25 | BD |
| **Cobalt**     | 18.4      | 0.0002 | 2 | BD-5 | 2.0 | 1.5 | BD | BD | BD |
| **Copper**     | 9.59      | 0.003 | 2 | 2-6 | 3.5 | 3 | BD | BD | BD |
| **Iron**       | 8.39      | 0.005 | 2 | 2300-7800 | 4000 | 2950 | 670-2500 | 1442.5 | 1300 |
| **Lead**       | 3.02      | 0.09 | 2 | 3-8 | 4.5 | 3.5 | BD-2 | 0.5 | BD |
| **Manganese**  | 36.4      | 4.7E-06 | 2 | 38-300 | 127.5 | 86 | 7-22 | 13.3 | 12 |
| **Molybdenum** | 10.6      | 0.002 | 2 | BD | BD | BD | BD | BD | BD |
| **Nickel**     | 20.8      | 0.0002 | 2 | 1-4 | 2 | 1.5 | BD | BD | BD |
| **Strontium**  | 1.19      | 0.3 | 2 | 3-9 | 5.0 | 4 | BD-4 | 1.25 | 0.5 |
| **Uranium**    | 19.7      | 0.002 | 2 | BD-0.2 | 0.06 | BD | BD | BD | BD |
| **Zinc**       | 27.5      | 3.32E-05 | 2 | 1-3 | 1.75 | 1.5 | BD | BD | BD |

### Table 3. Continued.

| Sediment/Parameter | Downstream Mine Sites | | |
|--------------------|------------------------|---|---|---|---|---|---|---|---|
|                   | W3 Range | W3 Mean | W3 Median | W5 Range | W5 Mean | W5 Median |
| **Aluminium** | 1500-8700 | 5083.3 | 5200 | 230-2200 | 1034 | 310 |
| **Barium**     | 37-170 | 101.6 | 88 | 2-22 | 7.5 | 3 |
| **Boron**      | BD-6 | 2 | BD | BD | BD | BD |
| **Cadmium**    | BD-1 | 0.47 | 0.5 | BD | BD | BD |
| **Chromium**   | 2-8 | 4.0 | 2 | BD-3 | 1.0 | BD |
| **Cobalt**     | 61-1000 | 551.8 | 635 | BD-23 | 6.0 | 0.5 |
| **Copper**     | 8-29 | 17.0 | 15.5 | BD-7 | 2.4 | BD |
| **Iron**       | 2400-19000 | 9950 | 10350 | 490-3900 | 1982 | 1100 |
| **Lead**       | 5-17 | 8.4 | 6 | BD-6 | 2.0 | BD |
| **Manganese**  | 270-9300 | 5474 | 6900 | 3-220 | 76.0 | 11 |
| **Molybdenum** | BD-9 | 4 | 4 | BD | BD | BD |
| **Nickel**     | 160-1100 | 6067 | 630 | 2-67 | 19.0 | 2 |
| **Strontium**  | 3-27 | 12.3 | 9 | BD-8 | 2.8 | 0.5 |
| **Uranium**    | 0.1-1.4 | 0.53 | 0.4 | BD-0.3 | 0.08 | BD |
| **Zinc**       | 300-1100 | 734 | 760 | 2-97 | 25.0 | 2 |

### Table 4

| Beetle/Parameter | F value | p value | df | W1 Range | W1 Mean | W1 Median | Bell Range | Bell Mean | Bell Median |
|-----------------|---------|---------|---|----------|---------|-----------|------------|-----------|-------------|
| **Aluminium**  | 3.04 | 0.07 | 2 | 3713-8184 | 6023.3 | 6496 | 3963.4-7394.8 | 4990.8 | 4578 |
| **Arsenic**   | 1.56 | 0.23 | 2 | 177-496.2 | 501.2 | 514 | 260-745.9 | 318.8 | 326.1 |
| **Barium**    | 0.99 | 0.38 | 2 | 177.8-908.2 | 479.9 | 394.4 | 205.8-724.2 | 405.9 | 339.7 |
| **Beryllium** | 9.87 | 0.0009 | 2 | 0-3.2-74 | 1.3 | 1.36 | 0.65-2.83 | 1.4 | 1.21 |
| **Boron**     | 1.23 | 0.31 | 2 | 947-6-1573 | 1250.9 | 1309 | 1039-4-4739 | 1869.2 | 1494 |
| **Cadmium**   | 1.50 | 0.24 | 2 | 6-62.1 | 19.3 | 12.7 | 13.0-142.8 | 59.5 | 33.7 |
| **Chromium**  | 0.16 | 0.85 | 2 | 31.7-258.7 | 118.1 | 88.9 | 42.8-171.4 | 94.9 | 70.5 |
| **Cobalt**    | 4.31 | 7.44E-10 | 2 | 49.45-176.9 | 87.2 | 75.2 | 83.88-492.7 | 206.9 | 147.1 |
| **Copper**    | 0.66 | 0.53 | 2 | 13791-29869 | 23279.8 | 25755 | 12563-10288 | 17147 | 17810 |
| **Iron**      | 5.21 | 0.01 | 2 | 32925-59215 | 45530.3 | 47658 | 34133-71457 | 43733.4 | 37886 |
| **Lead**      | 1.16 | 0.33 | 2 | 12.9-48.5 | 30.8 | 30.1 | 18.3-43.4 | 32.5 | 34.2 |
| **Manganese** | 19.5 | 0.0001 | 2 | 17998-35040 | 26850.7 | 27718 | 27758-55660 | 43243.1 | 43229 |
| **Molybdenum** | 10.1 | 0.0008 | 2 | 46-64.66 | 53.4 | 49.4 | 40-69-61.75 | 49.8 | 48.5 |
| **Nickel**    | 30.2 | 4.97E-07 | 2 | 16-45-97.61 | 62.1 | 73.4 | 12-27-106.1 | 40.9 | 27.5 |
| **Selenium**  | 14.8 | 8.33E-05 | 2 | 918-1855 | 1376.1 | 1274 | 1339-2019 | 1726.1 | 1732 |
| **Silver**    | 1.78 | 0.19 | 2 | 21.5-70.4 | 47.7 | 52.3 | 15.9-50.8 | 31.0 | 31.7 |
| **Strontium** | 6.17 | 0.007 | 2 | 2285-4543 | 3445.5 | 3376 | 3260-6176 | 4730.3 | 4642 |
| **Titanium**  | 0.48 | 0.62 | 2 | 109.9-270.3 | 189.7 | 189.6 | 178.8-203.8 | 189.0 | 189.1 |
Mean aquatic beetle concentrations of Arsenic, Beryllium, Cobalt, Iron, Manganese, Molybdenum, Nickel, Selenium, Strontium, Uranium and Zinc were found to be significantly different between reference and coal mine impacted samples (Table 4). When analysed by ANOSIM, significant differences were found between W1 and W3 (R statistic 0.963), W1 and W5 (R statistic 0.639), Bell and W3 (R statistic 0.924) and Bell and W5 (R statistic 0.523) whilst no significant difference was recorded for W1 and Bell (R statistic 0.639). Biota and/or Environmental matching (BEST) results show that Manganese, Cobalt, Nickel and Zinc have the strongest influence in the differences recorded across beetle sample locations (best results, Corr = 0.944), (method, BIOENV).

Reference site beetles recorded mean arsenic concentrations of 318.8 and 501.2 µg/kg increasing slightly to 320.3 and 454.5 µg/kg at both downstream sites W3 and W5 (mean 474 and 541 µg/kg). Mean Beryllium at reference sites was recorded at 1.42 and 1.3 µg/kg. The downstream sites W3 and W5 recorded mean Beryllium results of 10.3 and 2.7 µg/kg. Reference sites recorded mean cobalt concentrations of 209.6 and 87.2 µg/kg, in comparison W3 and W5 results showed mean cobalt of 6270.5 and 1089.8 µg/kg. Iron at reference sites was recorded at 43373 and 45530 µg/kg. Whilst in comparison W3 and W5 recorded mean iron results of 72586 and 68016 µg/kg. Reference site results show mean manganese concentrations of 34243 and 45530 µg/kg. The downstream sites W3 and W5 recorded mean manganese of 117148 and 70644 µg/kg. Beetles sampled at both reference sites recorded mean molybdenum of 49.8 and 53.4 µg/kg. In comparison W3 and W5 (downstream sites) recorded mean molybdenum of 100.7 and 86.2 µg/kg. Nickel concentrations from both reference sites was 40.9 and 62.1 µg/kg (mean), whilst in comparison mean nickel at the downstream sites (W3 and W5) was recorded at 4491.1 and 39571 µg/kg respectively. In comparison W3 and W5 recorded mean selenium results was 2767 and 2106 µg/kg. Reference site selenium concentrations were recorded at 39751-132119 µg/kg. Mean strontium concentrations were recorded at reference sites as 4730.3 and 3445.5 µg/kg with the downstream sites (W3 and W5) recording mean strontium of 1726 and 1376 µg/kg respectively. In comparison W3 and W5 recorded mean strontium results was 2013.7 and 1909.6 µg/kg respectively. Mean uranium concentrations were recorded at reference sites as 4.7 and 86.2 µg/kg. In comparison W3 and W5 (downstream sites) recorded mean uranium results of 4.7 and 86.2 µg/kg. Mean zinc results for both reference sites recorded concentrations of 75490 (W1) µg/kg and 3502.3 µg/kg respectively. Mean iron results at reference sites were 68016 µg/kg and at downstream sites were 56530 µg/kg.
Figure 2. Mean beryllium with standard error bars for aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

Figure 3. Mean Cobalt with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

Figure 4. Mean Iron with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

Figure 5. Mean Manganese with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

Figure 6. Mean Molybdenum with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

Figure 7. Mean Nickel with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.
Figure 8. Mean Selenium with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

Figure 9. Mean Strontium with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

Figure 10. Mean Uranium with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

Figure 11. Mean Zinc with standard error bars in aquatic predacious beetles (Gyrinidae) measured in µg/Kg. Bell and W1 are reference samples whilst W3 and W5 are coal mine impacted samples.

4. Discussion

This study may be the first to investigate the bioaccumulation of contaminants from a licenced and regulated coal mine wastewater discharge on an aquatic predacious beetle (*Macrogyrus rivularis*).

Results show that the one species of aquatic predacious beetle has increased concentrations of contaminants at impacted sample locations when compared to non-impacted reference sites. Many of the contaminants can be directly linked to the coal mine wastewater discharge as shown by water column and stream sediment results. In its crux, this one aquatic predacious beetle is at the lower trophic level of the food chain and is a food source for many other aquatic and terrestrial species. The implications of concentrating large amounts of contaminants within lower trophic order species, whom are predated on by aquatic and terrestrial species may allow for a link to continue the biomagnification of the contaminants found within the impacted sampled beetles. The ability for these aquatic beetles to be prey for terrestrial species is also of major concern as this may also allow for the mobilisation of these contaminants from the aquatic ecosystem to the terrestrial environment.

There are many studies investigating water column pollution from metal contamination from a broad range of mining activities. Many investigate the links between metal mining impacts on water chemistry, stream sediments, aquatic flora, aquatic fauna, including; fish, turtles to an array of macroinvertebrates [1, 3, 5, 6, 14, 15]. Only a few studies have investigated the bioaccumulation of metals from coal mining activities within macroinvertebrates. Most often these impacts are from waste coal ash dams or coal fine spills but not discharged, licenced and regulated coal mining wastewaters.

Miller et al. 2013 studied the bioaccumulation of the metal selenium from abandoned coal mine pit lakes within two fish species. It was reported that selenium was bioaccumulating
within the fish tissue to levels above USA EPA tissue guidelines and concluded that the current reclamation practices implemented in the abandoned coal mine pit lakes were failing and that significant risk to wildlife and human health was of concern [11]. Within Australia, Telford et al. 2009 and Jasonsmith et al. 2008 have conducted studies assessing the bioaccumulation of metals from mining activities. Telford et al. 2009 investigated bioaccumulation of metals in aquatic gastropods from antimony and arsenic mines and found that concentrations of arsenic were statistically higher in mining impacted samples [21]. Jasonsmith et al. 2008 studied the bioaccumulation of selenium in water, sediment, zooplankton, benthic material, benthic algae, oligochaetes, gastropods, crustaceans, insects and fish residing in a coal power stations cooling reservoir in Lithgow, NSW close (within 15km) to the vicinity of this study. It was found that selenium was found in low concentrations within oligochaetes, gastropods, bivalves and crustaceans sampled and in contrast insects sampled recorded concentrations some 1000 times higher [23]. Jasonsmith et al. 2008 concluded that the detritus invertebrates bioaccumulated selenium in much greater concentrations than oligochaetes, gastropods, bivalves and crustaceans they sampled [23].

Swansburg et al. 2002 found many metal concentrations below laboratory detectable limits within water samples in their study whilst recording detectable (in some cases statistically different) limits of the same heavy metals in chironomid tissue. Their study investigated metal mining impacts on clean dilute (EC < 100 µS/cm), circumneutral (pH 6.5-7.5) streams in New Brunswick, Canada. Cadmium, Chromium, Cobalt, Copper, Lead and Nickel all recording detectable limits within chironomid tissue whilst only Molybdenum still recorded below detectable limits. Significant differences between reference and mine impacted chironomid samples were recorded for copper, cadmium and zinc only [22]. This is similar to this study with many contaminants within the water column recording below laboratory detectable limits at both reference and impacted sample locations, becoming more detectable within stream sediment and in turn sampled beetles at impacted sites.

Swansburg et al. 2002 found increases between 3 and 16 times were recorded across the five streams for zinc in chironomid tissue. The greatest increase being the stream impacted by the metal mine facility “Caribou” was reported to increase some 30 times from a reference mean of 113000 mg/kg and an impacted mean of 1813000 mg/kg [22]. This current study found zinc in beetles to be statistically significant recording increases at lower concentrations close to the vicinity of 1.5 times greater at impacted sites and increasing from water column concentrations to beetle concentrations in the magnitude of 77000 times. Cadmium was found by Swansburg et al. 2002 to be nearly 65 times higher in impacted chironomid tissue samples at the same mining site increasing from 600 to 372000 mg/kg [22]. This study found cadmium to increase two-fold between the reference site W1 and the impacted site W3 but was not statistically significant. Copper was recorded by Swansburg et al. 2002 to increase 11 and 12 times at two impacted sites from 13mg/kg upstream to 153mg/kg downstream and 10mg/kg upstream to 115 mg/kg downstream. Copper in this current study was not statistically significantly different with W1, W3 and W5 all recording similar copper concentrations with Bell Creek recording copper concentrations approximately 0.15 times less.

Swansburg et al. 2002 found mine effected chironomids to have deformed mentums. It was also reported increases in cobalt for four of their five sample streams [22]. The highest increase from reference to mine effected cobalt in chironomid tissue was 600 (reference) to 3200 (mine impacted) µg/kg just over five times higher. This study found cobalt increased from 87.2 µg/kg at the reference site W1 to 6271 µg/kg directly below the coal mine wastewater inflow at site W3 which is over 70 times higher than W1. Water column concentrations of cobalt in this current study increased some 50000 times from water column concentrations to beetle concentrations.

Otter et al. 2012 found Selenium and Arsenic bioaccumulated at differing concentrations within two fish species they studied impacted by a wet coal ash spill [17]. Statistical differences were recorded for both metal concentrations when compared between reference and impacted samples. Otter et al. 2012 concluded that the difference in bioaccumulated selenium and arsenic across species in impacted sites was due to the differing stomach pH of the species [17]. In this current study, the reference sites used may have a greater functional ability to sequester and bioaccumulate heavy metals due to their naturally mildly acid pH in comparison to the impacted sites whom have a “treated” alkaline pH [17, 24, 25]. This is of major concern as impacted sites beetles in this study have significantly higher concentrations of heavy metals whilst having a much lower ability to bioaccumulate these contaminants.

This raises questions as to what will occur after mining activities are no longer undertaken at this colliery as when mining ceases so does the water treatment process. At present pH is increased during the treatment process and water currently being discharged may have a much lower rate of bioavailability of some contaminants for aquatic biota as described by Atkinson et al. 2007 [25]. Once treatment ceases and the Wollangambe Rivers pH will naturally reduce to a background pH which is mildly acidic. This decrease in pH may open a new avenue for increased bioaccumulation from the legacy pollutants within the impacted streams sediments. The findings of this research evoke concerns over the validity of water column pollutant limits if they are in fact allowing for legacy pollutants to bioaccumulate and or magnify within the receiving waterways aquatic ecosystem.

5. Conclusions

Long term legacy pollutants are of great concern worldwide. Sericano et al. 1995 and Ashraf 2011 suggest heavy metals have the potential to accumulate within biota at
toxic concentrations and have chronic ecological impacts within ecosystem food chains [1, 3]. Wang and Rainbow 2008 raised concerns over the longevity of heavy metal pollutants in aquatic environments due to their ability to deposit into waterway sediments which can potentially remain indefinitely [6]. Kolaříková et al. 2012 found that bioaccumulation of pollutants within four macroinvertebrate species within the Elbe and Vltava Rivers in the Czech Republic were still persistent after water quality improvements were implemented. It was reported that these heavy metals concentrations were consistent over their 12-year study and they recorded no significant reduction in heavy metal concentrations other than Hg [7].

The implications that contaminants from the licenced and regulated wastewater being discharged may be accumulating within aquatic biota is of major concern. The implications that this regulated water column pollution is accumulating at magnified rates shows a major floor in water column pollution licensing. It is recommended that further research should be undertaken by the New South Wales Environmental Protection Authority to better assess these broader implications of legacy contaminants from licenced and regulated coal mine wastewater discharges are having on the aquatic ecosystem of EPL protected waterways.

6. Recommendations

It is recommended that further research should be undertaken by the New South Wales Environmental Protection Authority to better assess the implications of coal mine wastewater contaminants bioaccumulation and or biomagnification in EPL protected waterways stream sediments and their biota. If in fact the contaminants are leaving the water column and bioaccumulating within the aquatic biota of their receiving waterways, serious long-term legacy pollutant impacts may persist. Of equal concern is if in fact these pollutants are biomagnifying within the aquatic biota there is feasibility that this may transpose to the terrestrial environment and the extent of the contamination may be more far spreading than this study has found.

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References

[1] Sericano, J. L., Wade, T. L., and Jackson, T. J., 1995. Trace organic contamination in the Americas: An overview of the US national status and trends and the international mussel watch programmes. Mar. Pollution Bulletin. Vol 31. pp 214-225.

[2] Wislocka, M., Krawczyk, J., Klink, A., and Morrison, L., 2004. Bioaccumulation of Heavy Metals by Selected Plant Species from Uranium Mining Dumps in the Sudety Mts., Poland. Polish Journal of Environmental Studies, vol 15. 5. pp 811-818.

[3] Ashraf, M. A., Maah, M. J., and Yusoff, I., 2011. Bioaccumulation of Heavy Metals in Fish Species Collected from Former Tin Mining Catchment. International Journal of Environmental Research. vol 6. 1. pp 209-218.

[4] Nawab, J., Khan, M. T., Khan, K., Huang, Q., and Ali, R., 2015. Quantification of heavy Metals in Mining Affected Soil and their Bioaccumulation in Native Plants Species. International Journal of Phytoremediation. vol 17. Pp 801-813.

[5] Neff, J. M., 1984. Bioaccumulation of organic micro pollutants from sediments and suspended particulates by aquatic animals. Fresenius’ Zeitschrift für analytische Chemie, Vol 319, 2, pp 132-136.

[6] Wang, X. W., and Rainbow, P. S., 2008. Comparative approaches to understand metal bioaccumulation in aquatic animals. Comparative Biochemistry and Physiology Part C: Toxicology & Pharmacology, vol 148, 4, pp 315-323.

[7] Kolaříková, K., Stuchlík, E., Liška, M., Horecký, J., Tátosová, J., Hardekopf, D., Lapašanská, N., Hofická, Z., Hovorka, J., Mihaljevič, M., Fuksa, J. K., and Von Tümpelingm, W., 2012. Long-Term Changes in the Bioaccumulation of As, Cd, Pb, and Hg in Macroinvertebrates from the Elbe River (Czech Republic). Water Air Soil Pollution. vol 223. pp 3511-3526. DOI 10.1007/s11270-012-1129-1.

[8] Allen, H. E., Perdue, E. M., and Brown, D. S., 1993. Metals in Groundwater. Lewis Publishers. pp 437.

[9] Twining, J., Creighton, N., Hollins, S., and Szymczak, R., 2008. ‘Probabilistic Risk Assessment and Risk Mapping of Sediment Metals in Sydney Harbour Embayments’, Human and Ecological Risk Assessment: An International Journal, vol. 14, no. 6, pp. 1202-1225.

[10] Bharti, S., and Banerjee, T. K., 2011. Bioaccumulation of Metals in the Edible Catfish Heteropneustes fossilis (Bloch) Exposed to Coal Mine Effluent Generated at Northern Coalfield Limited, Singrauli, India. Bulletin of Environmental Contamination and Toxicology. vol 87. pp 393-398. DOI 10.1007/s00128-011-0371-3.

[11] Miller, L. L., Rasmussen, J. B., Palace, V. P., Sterling, G., and Hontela, A., 2013. Selenium Bioaccumulation in Stacked Fish as an Indicator of Fishery Potential in Pit Lakes on Reclaimed Coal Mines in Alberta, Canada. Environmental Management. vol 52. pp 72-84. DOI 10.1007/s00267-013-0038-4.

[12] Ajima, M. N. O., Ndodi, P. C., Ogo, O. A., Adaka, G. S., Osuigwe, D. I., and Njoku, D. C., 2015. Bioaccumulation of heavy metals in Mbaa River and the impact on aquatic ecosystem. Environmental Monitoring and Assessment. vol 187. 1-9. DOI 10.1007/s10661-015-4937-0.

[13] Jordanova, A., Strrezov, A., Ayravan, M., Petkovic, N., and Stoilova, T., 1999. Heavy metal assessment in sediments and water from the Bulgarian Black Sea Coast. Water Sci Tech, vol 39. Pp 207-212.

[14] Hill, B. H., Willingham, W. T., Parrish, L. P., and McFarland, B., H., 2008. Periphyton community responses to elevated metal concentrations in a Rocky Mountain stream. Hydrobiologia, vol 428, 161-9.
Amisah, S., and Cowx, I. G., 2008. Impacts of abandoned mine and industrial discharges on fish abundance and macroinvertebrate diversity of the upper River Don in South Yorkshire, UK. J Freshwater Ecology, 15, 237-49.

Reash, R. J., Lohner, T. W., Wood, K. V., 2006. Selenium and other trace metal in fish inhabiting a fly ash stream: implications for regulatory tissue thresholds. Environmental Pollution. Vol 142, 397-408.

Otter, R. R., Bailey, F. C., Fortner, A. M., and Adams, S. M., 2012. Trophic status and metal bioaccumulation differences in multiple fish species exposed to coal ash-associated metals. Ecotoxicology and Environmental Safety, vol 85, pp 30-36.

Peterson, M. J., Smith, J. G., Southworth, G. R., Ryon, M. G., and Eddlemon, G. K., 2002. Trace element contamination in benthic macroinvertebrates from a small stream near a uranium mill tailings site. Environmental Monitoring and Assessment. vol 74. pp 193–208.

Maret, T. R., Cain, D. J., MacCoy, D. E., and Short, T. M., 2003. Response of benthic invertebrate assemblages to metal exposure and bioaccumulation associated with hard-rock mining in northwestern streams, USA. The North American Benthological Society. vol 22. 4 pp 598-620.

Cain, D. J., Luoma, S. N., and Wallace, W. G., 2004. Linking metal bioaccumulation of aquatic insects to their distribution patterns in a mining-impacted river. Environmental Toxicology and Chemistry. vol. 23. 6. Pp 1463–1473.

Telford, T., Maher, W., Krikowa, F., Foster, S., Ellwood, M. J., Ashley, P. M., Lockwood, P. V., and Wilson, S. C., 2009. Bioaccumulation of antimony and arsenic in a highly contaminated stream adjacent to the Hillgrove Mine, NSW, Australia. Environmental Chemistry. vol 6. Pp 133–143. doi: 10.1071/EN08097.

Swansburg, E. O., Fairchild, W. L., Frey, B. J., and Ciborowski, J. H., 2002. Environmental Toxicology and Chemistry. vol. 21. 12. pp 2675–2684.

Jonsomith, J. F., Maher, W., Roach, A. C., and Krikowa, F., 2008. Selenium bioaccumulation and biomagnification in Lake Wallace, New South Wales, Australia. Marine and Freshwater Research. vol 59. pp 1048-1060.

Durães, N., Bobos, I., Ferreira da Silva, E., and Dekayir, A., 2014. Copper, zinc and lead biogeochemistry in aquatic and land plants from the Iberian Pyrite Belt (Portugal) and north of Morocco mining areas. Environmental Science Pollution Research. vol 22 pp 2087–2105. DOI 10.1007/s11356-014-3394-6.

Atkinson, C., Jolley, D. F., and Simpson, S. L., 2007. Effect of overlying water pH, dissolved oxygen, salinity and sediment disturbances on metal release and sequestration from metal contaminates marine sediments, Chemosphere, 69 (9), 1428-1437.

Banks, D., Younger, P. L., Arnesen, R. T., Iversen, E. R., and Banks, S. B., 1997. Mine-water chemistry: the good, the bad and the ugly. Environmental Geology, 32, 157-174.

Jarvis, A. P., and Younger, P. L., 1997. Dominating chemical factors in mine water induced impoverishment of the invertebrate fauna of two streams in the Durham Coalfield, UK. Chemistry and Ecology, vol 13. pp 249-270.

Brake, S. S., Connors, K. A., and Romberger, S. B., 2001. A river runs through it: impact of acid mine drainage on the geochemistry of West Little Sugar Creek pre- and post-reclamation at the Green Valley coal mine, Indiana, USA. Environmental Geology, 1471-1481.

Johnson, D. B., 2003. Chemical and microbiological characteristics of mineral spoils and drainage waters at abandoned coal and metal mines. Water, Air, and Soil Pollution. vol 3. pp 47-66.

Pond, G. J., Passmore, M. E., Borsuk, F. A., Reynolds, L., and Rose, C. K., 2008. Downstream effects of mountaintop coal mining: comparing biological conditions Using family-and genus-level macroinvertebrate bioassessment tools. Journal of the North American Benthological Society, 27: 717-737.

Younger, P. L., 2004. Environmental impacts of coal mining and associated wastes: a geochemical perspective. Geological Society, London, Special Publication 236: 169-209.

Wright, I. A., 2012. Coal mine ‘dewatering’ of saline wastewater into NSW streams and rivers: a growing headache for water pollution regulators. In Grove, J. R. and Rutherford, I. D (eds). Proceedings of the 6th Australian Stream Management Conference, managing for Extremes, 6-8 February 2012 Canberra, Australia, published by the River Basin Management Society pp. 206-213.

Belmer, N., Tippler, C., Davies, P. J., and Wright, I. A., 2014. Impact of a coal mine waste discharge on water quality and aquatic ecosystems in the Blue Mountains World Heritage Area, in Viets, G, Rutherford, I, D, and Hughes, R, (editors), Proceedings of the 7th Australian Stream Management Conference, Townsville, Queensland, Pages 385-391.

Hellawell, J. M., 1986. Biological Indicators of Freshwater Pollution and Environmental Management. London: Elsevier.

Rosenberg, D. M., and Resh, V. H., 1993. Freshwater biomonitoring and benthic macroinvertebrates. Chapman & Hall, New York, London.

Battaglia, M., Hose, G. C., Turak, E., & Warden, B. (2005). Depauperate macroinvertebrates in a mine affected stream: Clean water may be the key to recovery. Environmental Pollution, 138, 132-141.

Wright, I. A. & Burgin, S. 2009a. Comparison of sewage and coal-mine wastes on stream macroinvertebrates within an otherwise clean upland catchment, south-eastern Australia. Water, Air and Soil Pollution, 204, 227-241.

Watts, C., and Hamon, H., 2010. Pictorial Guide to the Australian Whirligig Beetle, Entomology Department, South Australian Museum, Adelaide, South Australia.

Wright, I, A., Belmer, N, and Davies, P, J 2017, Coal Mine Water Pollution and Ecological Impairment of One of Australia’s Most ‘Protected’ High Conservation-Value rivers Water Air Soil Pollution.

Strahler, A., 1952. Dynamic Basis of Geomorphology. Geological Society of America Bulletin. vol. 63. pp. 923-938. doi.org/10.1130/0016-7606 (1952) 63.