

# **Assessment of Acid Sulfate Soil Materials (Phase 2): Glen Esk-Rusty Waterhole**

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**FINAL REPORT** 



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#### **Cover Photograph:**

*Photograph of site 28200\_1 in the Glen Esk-Rusty Waterhole. Photographer: Mitch Tulau*

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## **EXECUTIVE SUMMARY**

Detailed assessments of acid sulfate soils within the Murray-Darling Basin (MDB) are conducted as a two-phase process under the MDB Acid Sulfate Soils Risk Assessment Project (ASSRAP). An initial Phase 1 acid sulfate soil investigation of the Glen Esk-Rusty Waterhole in March 2010 showed acid sulfate soils to be a priority concern within this wetland (Ward *et al.* 2010). Based on Phase 1 recommendations, a Phase 2 investigation was undertaken for the Glen Esk-Rusty Waterhole to determine the nature, severity and the specific risks associated with acid sulfate soil materials. Phase 2 activities included soil laboratory analysis, a risk assessment, and interpretation and reporting, including discussion on broad acid sulfate soil management options.

The Phase 2 assessment of the Glen Esk-Rusty Waterhole examined both the contaminant and metalloid dynamics and reactive metals associated with surface layers from two sites within the wetland. These results were used to determine the risks associated with contaminant mobilisation. The risks associated with both acidification and de-oxygenation were determined primarily using data from the Phase 1 assessment.

The contaminant and metalloid dynamics tests were undertaken to assess the release of metals during a water extraction, and to assess changes with time as saturated soils by incubating soil materials for periods of 1, 7, 14 and 35 days. The degree to which metal and metalloid concentrations exceed ANZECC/ARMCANZ water quality guideline values for environmental protection was used to characterise the degree of hazard. For Glen Esk-Rusty Waterhole, the contaminant and metalloid dynamics test over 35 days showed that under the experimental conditions eight of the metals examined (including cobalt (Co), chromium (Cr), copper (Cu), iron (Fe), manganese (Mn), nickel (Ni), silver (Ag) and zinc (Zn)) were found to exceed the water quality guidelines (ANZECC/ARMCANZ 2000). The guidelines for iron (Fe) and silver (Ag) were exceeded by more than 10 times at both sites. The guideline for iron (Fe) was exceeded by more than 100 times in a sulfuric soil material.

The contaminant and metalloid behaviour often varied between the metals/metalloids examined during the inundation experiments. The reductive dissolution of iron minerals seemed to have partially controlled the release of sorbed arsenic, although the controls on the release of other metals were not clear. The trend with some metals/metalloids indicates further potential for release had the incubation experiments been allowed to proceed for a longer timeframe; it is also likely that many of the metals/metalloids will become incorporated into sulfide minerals following further reduction.

As shown in the table below, the metals found to exceed the ANZECC water quality guidelines during the inundation experiments represent a low to high hazard. The reactive metal concentrations were also found to be sufficiently high to be a potential hazard if released into surrounding waters. In natural systems the dynamics of metal release will be governed by the upward chemical flux, which is a function of soil type, water flow, diffusion and chemistry of the soils near the sediment-water interface (MDBA 2011).

A risk assessment framework was applied to determine the specific risks associated with acidification, contaminant mobilisation and de-oxygenation (MDBA 2011). The Phase 2 assessment identified the following risks associated with the presence of acid sulfate soils in the Glen Esk-Rusty Waterhole:

- high acidification risk.
- medium contaminant mobilisation risk, and
- medium de-oxygenation risk.



These findings indicate that, if not managed appropriately, the acid sulfate soil materials identified in the Glen Esk-Rusty Waterhole have the potential to present a serious risk to the environmental values of both the wetland and adjacent waters. This report outlines the variety of management options available to manage acid sulfate soils in inland aquatic ecosystems. The most appropriate management strategy for the Glen Esk-Rusty Waterhole would be to prevent oxidation of the acid sulfate soil materials identified. Neutralisation may be the best management strategy in the event of disturbance. However, in designing a management strategy for dealing with acid sulfate soils in affected inland wetlands, other values and uses of a wetland need to be taken into account to ensure that any intervention is compatible with other management plans and objectives for the wetland.

It is important to note that the soil materials collected in March 2010 as part of the Phase 1 assessment only provided a snapshot of the acid sulfate soil materials present and the conditions at selected locations in the wetland. While recent inundation within the wetland may have minimised the risks identified in the short-term, it is also likely that this inundation will lead to further formation of acid sulfate soil materials.

It is recommended that, within the context of other management objectives for the wetland, consideration be given to undertaking water quality monitoring to identify potential contamination as a result of the disturbance of acid sulfate soils within the wetland. The presence of some high risks identified in this Phase 2 assessment indicates that senior management attention is probably needed (MDBA 2011).

## **1. INTRODUCTION**

At its March 2008 meeting, the Murray–Darling Basin Ministerial Council discussed the emerging issue of inland acid sulfate soils and the associated risks to Murray–Darling Basin waterways and agreed that the extent of the threat posed by this issue required assessment. The purpose of the Murray–Darling Basin Acid Sulfate Soils Risk Assessment Project was to determine the spatial occurrence of, and risk posed by, acid sulfate soils at priority wetlands in the River Murray system, wetlands listed under the Ramsar Convention on Wetlands of International Importance and other key environmental sites in the Murray–Darling Basin. The project involved the selection of wetlands of environmental significance, as well as those that may pose a risk to surrounding waters. These wetlands were then subjected to a tiered assessment program, whereby wetlands were screened through a desktop assessment stage, followed by a rapid on-ground appraisal, and then detailed on-ground assessment if results of previous stages indicated an increased likelihood of occurrence of acid sulfate soils.

Detailed assessments of acid sulfate soils within the Murray-Darling Basin (MDB) are conducted as a two-phase process under the MDB Acid Sulfate Soils Risk Assessment Project (ASSRAP). Detailed Phase 1 acid sulfate soil assessments have been undertaken in both wetlands and channel systems throughout the MDB as part of the MDB ASSRAP. Phase 1 investigations are initially undertaken to determine whether acid sulfate soil materials are present (or absent) in the study area, and provide characterisation of the properties and types of acid sulfate soils. Phase 2 investigations are only conducted if the acid sulfate soil materials from Phase 1 are determined to be a priority concern for the study area and, based on Phase 1 recommendations, selected samples undergo further investigations to determine the nature, severity and the specific risks associated with the acid sulfate soil materials. Phase 2 activities include: (i) soil laboratory analysis to confirm and refine the hazards associated with contaminant mobilisation and/or deoxygenation, (ii) a risk assessment, and (iii) interpretation and reporting, including discussion on broad acid sulfate soil management options.

Detailed Phase 1 acid sulfate soil assessments were undertaken at almost 200 wetlands and river channels throughout the Murray-Darling Basin. In the Edward and Wakool Rivers region, 12 wetlands were investigated by Southern Cross GeoScience (Ward *et al*. 2010). From these Phase 1 investigations, Glen Esk-Rusty Waterhole, Boiling Downs Creek and Wakool River Billabong (Wetland ID 20246) were selected for further investigation. This report outlines the results of Phase 2 activities on selected samples from the Glen Esk-Rusty Waterhole (Figure 1-1).

Following the Glen Esk-Rusty Waterhole Phase 1 assessment (Ward *et al.* 2010) and the priority ranking criteria adopted by the Scientific Reference Panel of the MDB ASSRAP (see Table 1-1), selected sites from within the Glen Esk-Rusty Waterhole were chosen for Phase 2 detailed assessment. The Glen Esk-Rusty Waterhole Phase 1 assessment identified there were three high priority sites based on the presence of sulfuric material, two high priority sites based on hypersulfidic material and two high priority sites based on monosulfidic material in the Glen Esk-Rusty Waterhole (Ward *et al.* 2010). In addition, all four sampling sites examined had a high priority ranking for Phase 2 detailed assessment based on monosulfidic black ooze (MBO) formation hazard (Ward *et al.* 2010). Phase 2 investigations were carried out on selected samples from two of the high priority sites (i.e. Sites 28200\_1 and 28200\_2) identified in the Phase 1 assessment.



**Figure 1-1: Map showing the areas assessed in the Glen Esk-Rusty Waterhole (Wetland ID 28200) during the Phase 1 assessment.** 





A summary of the soil laboratory analyses undertaken as part of the Phase 2 assessment and the sample selection criteria for each analysis are given in Table 1-2. Soil samples identified to undergo Phase 2 laboratory analysis are primarily from the surface layer, as this is the soil most likely to have initial contact with water. A list of the samples selected for Phase 2 analysis for the Glen Esk-Rusty Waterhole is presented in Table 1-3.









*1 n = total number of samples analysed* 

# **2. LABORATORY METHODS**

## *2.1. Laboratory analysis methods*

### **2.1.1. Summary of laboratory methods**

A list of the parameters measured and each of the method objectives for the Phase 2 assessment are summarised below in Table 2-1. All soil samples analysed in this Phase 2 assessment were collected and subsequently stored as part of the Phase 1 field assessment.





Guidelines on the approaches that were followed as part of this Phase 2 assessment for the contaminant and metalloid dynamics (CMD) and monosulfide formation potential (MFP) methods are presented in full in the detailed assessment protocols (see Appendices 7 and 8, MDBA 2010). Any variations to the two methods outlined in the detailed assessment protocols are presented in Sections 2.1.2 and 2.1.3. The reactive metals method has only recently been added to the Phase 2 assessment procedure and is presented in Section 2.1.4.

## **2.1.2. Contaminant and metalloid dynamics method**

The guidelines for the contaminant and metalloid dynamics method are outlined in Appendix 7 of the detailed assessment protocols (MDBA 2010). In this study supernatant was collected and assessed at four intervals including 24 hours, 7 days, 14 days and 35 days. The concentration of 15 metals/metalloids (i.e. aluminium (Al), antimony (Sb), arsenic (As), cadmium (Cd), chromium (Cr). cobalt (Co), copper (Cu), iron (Fe), lead (Pb), manganese (Mn), nickel (Ni), selenium (Se), silver (Ag), vanadium (V) and zinc (Zn)) was determined by ICP-MS (Inductively Coupled Plasma - Mass Spectrometry) (APHA 2005). Redox potential (Eh) and pH were determined at each interval using calibrated electrodes linked to a TPS WP-80 meter; Eh measurements are presented versus the standard hydrogen electrode. Electrical conductivity (EC) was determined using a calibrated electrode linked to a TPS WP-81 meter. All parameters were measured on filtered (0.45 µm) water samples.

## **2.1.3. Monosulfide formation potential method**

The guidelines for the monosulfide formation potential method are outlined in Appendix 8 of the detailed assessment protocols (MDBA 2010). No samples from the Glen Esk-Rusty Waterhole were selected for monosulfide formation potential analysis.

### **2.1.4. Reactive metals method**

In this Phase 2 assessment a reactive metals method was carried out instead of the x-ray fluorescence (XRF) spectrometry method outlined in the detailed assessment protocols (MDBA 2010). While the XRF method provides data on the total elements in the soil, the reactive metals method gives an indication of the potential metal concentrations that may be released into the surrounding waters. In this method samples for analysis were prepared by disaggregation (not grinding) using a 'jaw crusher', and then sieved to include only the <2 mm fine earth fraction. A total of 2.5 g sediment was added to 40 mL of 0.1 M HCl, gently mixed for 1 hour and filtered through a pre-acid washed 0.45 micron nitro-cellulose filter. As with the contaminant and metalloid dynamics method, the metals were determined by ICP-MS and included aluminium (Al), antimony (Sb), arsenic (As), cadmium (Cd), chromium (Cr). cobalt (Co), copper (Cu), iron (Fe), lead (Pb), manganese (Mn), nickel (Ni), selenium (Se), silver (Ag), vanadium (V) and zinc (Zn). The reactive metals test was conducted on all samples that underwent the contaminant and metalloid dynamics test.

### *2.2. Quality assurance and quality control*

For all tests and analyses, the quality assurance and quality control procedures were equivalent to those endorsed by NATA (National Association of Testing Authorities). The standard procedures followed included the monitoring of blanks, duplicate analysis of at least 1 in 10 samples, and the inclusion of standards in each batch. In addition, the contaminant and metalloid dynamics tests were duplicated.

Reagent blanks and method blanks were prepared and analysed for each method. All blanks examined here were either at, or very close to, the limits of detection. On average, the frequencies of quality control samples processed were: 10% blanks, ≥ 10% laboratory duplicates, and 10% laboratory controls. The analytical precision was ±10% for all analyses.

# **3. RESULTS AND DISCUSSION**

### *3.1. Summary of soil laboratory results*

### **3.1.1. Contaminant and metalloid dynamics data**

The contaminant and metalloid dynamics data for the four Glen Esk-Rusty Waterhole soil materials examined (i.e. 28200\_1.1, 28200\_1.2, 28200\_2.1 and 28200\_2.2) are presented in Appendix 1 (Tables 8-1 to 8-4) and summarised in Table 3-1. Table 3-1 also compares the pore-water metal contents to the relevant national water quality guideline for environmental protection (ANZECC/ARMCANZ 2000). Results for all parameters measured are presented in Figures 3-1 to 3-4.

The pH, electrical conductivity (EC) and redox potential (Eh) dynamics over 35 days of inundation for the Glen Esk-Rusty Waterhole soil materials are presented in Figure 3-1. The pH was below the ANZECC guideline of 6.5 for all soil materials throughout the timeframe of the experiment. An increase in pH was observed during the incubation with three of soil materials, with one soil (i.e. 28200 2.1) showing minimal pH change. The sulfuric surface soil material at site 28200 1 (i.e. 0-5 cm) had an initial pH of less than 4. All soil materials showed a decrease in Eh during the inundation experiments. The data indicates that the increase in pH observed with the majority of soil materials is a consequence of reduction processes consuming acidity. Previous studies have often found inundation removes the acidity in partially-oxidised sediments as the acidity gets consumed from the reduction of iron (III) oxides, sulfates and other oxidised species by anaerobic bacteria (Dent 1986).

The electrical conductivities for all samples decreased during the experiment (Figure 3-1). Surface soil materials analysed from both sites (i.e. 28200 1.1 and 28200 2.1) had electrical conductivity values that exceeded the ANZECC guideline of 2200 µS/cm for the first three sampling periods, however, by day 35 these had fallen below the upper guideline value (Figure 3-1). The decrease in conductivity with time would suggest the formation of insoluble mineral phases during the inundation experiment.

It is well established that inundating oxic soils can dramatically alter the mobility of metals and metalloids. The contaminant and metalloid dynamics results for the Glen Esk-Rusty Waterhole soil materials are presented in Figures 3-2 to 3-4. Under the experimental conditions eight of the metals examined (i.e. cobalt (Co), chromium (Cr), copper (Cu), iron (Fe), manganese (Mn), nickel (Ni), silver (Ag) and zinc (Zn)) were found to exceed the ANZECC water quality guidelines during the inundation experiments (Table 3-1). Cadmium was below the limit of detection for all soil materials over the 35 day inundation period. Some of the metals (i.e. aluminium (Al), cobalt (Co), copper (Cu), iron (Fe), manganese (Mn) and zinc (Zn)) were above the ANZECC guideline at all sampling intervals with one or more soil materials. Many of the metals showed the greatest exceedance with the sulfuric surface soil material at site 28200 1; copper (Cu) and nickel (Ni) only exceeded the quideline with the sulfuric soil material. The guidelines for iron (Fe) and silver (Ag) were exceeded by more than 10 times at both sites. The guideline for iron (Fe) was exceeded by more than 100 times in the sulfuric soil material.

The metal/metalloid behaviour during the 35 day inundation period often varied between the metals/metalloids examined (Figures 3-2 to 3-4). The magnitude of mobilisation is affected by many factors that include but are not exclusive to: 1) the abundance and form of metal and metalloid contaminants; 2) the abundance and lability of organic matter; 3) the abundance and reactivity of iron minerals; 4) availability of sulfate; 5) acid/alkalinity buffering capacity; 6) pH; 7) EC; 8) clay content; 9) microbial activity; 10) temperature; and 11) porosity (MDBA 2010). It is expected that the increase in the iron (Fe) concentration observed during the incubation is largely a consequence of ferric iron  $(Fe^{3+})$  reduction

releasing ferrous iron (Fe<sup>2+</sup>) into solution (Figure 3-3). Burton *et al.* (2008) found significant mobilisation of arsenic (As) associated with ferric iron reduction following the inundation of acid sulfate soil materials, and this was also observed in this study (Figure 3-2).

The trend with some metals/metalloids indicates there is the potential for further release had the incubation experiments been allowed to proceed for a longer timeframe. However, it is also expected that many of the metals/metalloids will become incorporated into iron sulfide minerals (due to sorption to and/or co-precipitation) or precipitated as non-ferrous sulfides following further reduction.

<b>Parameter</b>	units	<b>ANZECC</b> <b>Guidelines</b>	28200_1.1 $(0-5 cm)$		28200_1.2 $(5-10 \text{ cm})$		28200_2.1 $(0-5 cm)$		28200 2.2 $(5-10 \text{ cm})$	
			Min.	Max.	Min.	Max.	Min.	Max.	Min.	Max.
pH		$6.5 - 8.0$	3.70	4.91	4.77	5.96	5.66	5.83	5.84	6.45
$EC^*$	$\mu$ S cm $^{-1}$	125-2,200	1,495	3,229	744	1,477	1,263	2,592	964	1,883
Eh	mV	$\blacksquare$	299	459	262	415	209	331	176	300
Ag	$\mu$ g l <sup>-1</sup>	0.05	< 0.1	1.1	< 0.1	0.3	< 0.1	1.1	< 0.1	0.4
$AI^A$	mg $\Gamma^1$	0.055	0.12	0.39	0.02	0.05	< 0.01	0.02	< 0.01	0.04
$\mathsf{As}^{\mathsf{B}}$	$\mu$ g l <sup>-1</sup>	13	1.1	7.1	1.0	6.0	1.0	8.4	1.0	5.3
Cd	$\mu$ g l <sup>-1</sup>	0.2	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1
Co	$\mu$ g $\Gamma^1$	2.8	6.0	8.5	1.0	1.5	2.4	2.9	1.6	3.4
$Cr^C$	$\mu$ g l <sup>-1</sup>	$\mathbf{1}$	1.0	2.0	1.0	4.2	1.0	1.5	1.0	1.7
Cu <sup>H</sup>	$\mu$ g l <sup>-1</sup>	1.4	1.4	3.8	1.0	1.0	1.0	1.0	1.0	1.0
Fe	mg $\Gamma^1$	0.30	14.72	52.75	0.33	18.01	11.85	19.23	8.88	22.53
Mn	mg $\Gamma^1$	1.70	2.96	4.01	1.41	1.70	2.17	2.73	1.51	1.73
Ni <sup>H</sup>	$\mu$ g l <sup>-1</sup>	11	6.5	12.3	1.1	1.9	1.1	2.5	1.5	3.2
Pb <sup>H</sup>	$\mu$ g l <sup>-1</sup>	3.4	1.0	1.2	1.0	1.0	1.0	1.0	1.0	1.0
Sb	$\mu$ g l <sup>-1</sup>	9	1.0	1.0	1.0	1.3	1.0	1.0	1.0	1.0
Se	$\mu$ g l <sup>-1</sup>	11	1.0	4.6	1.0	1.5	1.0	1.0	1.0	1.0
$\vee$	$\mu$ g l <sup>-1</sup>	6	1.0	2.1	1.0	2.0	1.0	1.0	1.0	1.2
$Zn^{H}$	$\mu$ g $\Gamma^1$	8	18.0	26.4	$1.2$	6.9	1.5	9.1	3.2	9.3

**Table 3-1. Summary of contaminant and metalloid dynamics data** 

*Exceeded ANZECC Guideline (x1) Exceeded ANZECC Guideline (x10) Exceeded ANZECC Guideline (x100)* 

*Notes accompanying this table can be found on the following page.* 

*The ANZECC guideline values for toxicants refer to the Ecosystem Protection – Freshwater Guideline for protection of 95% of biota in 'slightly-moderately disturbed' systems, as outlined in the Australian Water Quality Guidelines for Fresh and Marine Water Quality (ANZECC/ARMCANZ 2000).* 

*\* ANZECC water quality guidelines for lowland rivers in South-east Australia are provided for salinity (there are currently no trigger values defined for 'Wetlands').* 

*Values outside the ranges defined in the ANZECC guidelines are indicated with yellow, orange and red background colours.* 

- 
- 
- A Guideline is for Aluminium in freshwater where  $pH > 6.5$ .<br>
<sup>B</sup> Guideline assumes As in solution as Arsenic (AsV).<br>
<sup>C</sup> Guideline for Chromium is applicable to Chromium (CrVI) only.<br>
<sup>H</sup> Hardness affected (refer to Guide
- 



**Figure 3-1: pH, EC and Eh dynamics over 35 days for the Glen Esk-Rusty Waterhole sites (28200\_1 and 28200\_2).** 



**Figure 3-2: Contaminant and metalloid dynamics (Ag, Al and As) over 35 days for the Glen Esk-Rusty Waterhole sites (28200\_1 and 28200\_2).** 



**Figure 3-3: Contaminant and metalloid dynamics (Cd, Co, Cr, Cu, Fe and Mn) over 35 days for the Glen Esk-Rusty Waterhole sites (28200\_1 and 28200\_2).** 



**Figure 3-4: Contaminant and metalloid dynamics (Ni, Pb, Sb, Se, V and Zn) over 35 days for the Glen Esk-Rusty Waterhole sites (28200\_1 and 28200\_2).** 

### **3.1.2. Reactive metals data**

The determination of the reactive metal fraction provides an indication of the total potential metal release from the sediment into surrounding waters. The reactive metals test used in this study gives an indication of the metals and metalloids that are more strongly bound to minerals (or weakly soluble with an acid extraction) than would be soluble with a water extraction. The moderately strong acid used (i.e. 0.1 M HCl) indicates the "stored metals" and metalloids associated with iron and manganese oxides, organic materials as well as acid soluble minerals. While the ANZECC sediment quality guidelines (ANZECC/ARMCANZ 2000) are based on total metal/metalloid concentrations, a reactive metal concentration near to or above guideline values indicate an elevated hazard.

The reactive metals data for the Glen Esk-Rusty Waterhole soil materials showed all metals/metalloids were  $\leq$  17% of the ANZECC trigger value for the total metal/metalloid concentration (see Table 8-5, Appendix 1). While all reactive metal concentrations were below the ANZECC trigger value, the metal concentrations measured were sufficiently high to be a potential hazard if the total reactive fractions were to be released into a surrounding water body (i.e. above ANZECC water quality quidelines).

## *3.2. Interpretation and discussion of results*

The contaminant and metalloid dynamics test undertaken as part of this Phase 2 assessment assists in determining the impacts on water quality by simulating the release of metal and metalloid concentrations that may occur under anaerobic conditions over a several week period. The contaminant and metalloid behaviour of the four Glen Esk-Rusty Waterhole soil materials during the 35 day inundation often varied between the metals/metalloids examined (see Figures 3-2 to 3-4). The reductive dissolution of iron (Fe) minerals seemed to have partially controlled the release of sorbed arsenic (As), although the controls on the release of other metals were not clear.

The contaminant and metalloid dynamics data for the four soils from Glen Esk-Rusty Waterhole showed eight of the metals examined (i.e. cobalt (Co), chromium (Cr), copper (Cu), iron (Fe), manganese (Mn), nickel (Ni), silver (Ag) and zinc (Zn)) exceeded the ANZECC water quality guidelines during the inundation experiments (Table 3-1). The guidelines for iron (Fe) and silver (Ag) were exceeded by more than 10 times at both sites, with the quideline for iron (Fe) being exceeded by more than 100 times in the sulfuric surface soil material (i.e. 0-5 cm) at site 28200\_1. Cadmium (Cd) was below the limit of detection for all soil materials over the 35 day inundation period.

While the contaminant and metalloid dynamics test gives an indication of the metal/metalloid content of the soil, the overlying water will rarely have the concentration measured in solution during this test due to dilution in the receiving waters. It can therefore be assumed that if a metal/metalloid concentration did not exceed the ANZECC water quality guideline during the test it does not represent an environmental hazard. Thresholds for the degree of hazard associated with the contaminant and metalloid concentrations were developed with respect to the ANZECC water quality guidelines, and a summary of the degree of hazard each of the metals/metalloids pose at the sites examined in the Glen Esk-Rusty Waterhole is given in Table 3-2. Note the background colours presented in Table 3-1 also correspond to the degree of hazard (i.e. no colour (no hazard), yellow (low hazard), orange (moderate hazard) and red (high hazard)).





The eight metals found to exceed the ANZECC water quality guidelines during the inundation experiments represent a low to high hazard (Table 3-2). Iron (Fe) was the only metal/metalloid observed to have a high hazard, and only with the surface soil material at site 28200 1 (Table 3-1). Only iron (Fe) and silver  $(Ag)$  were found at a concentration that represents a moderate hazard (Table 3-1). All metals with a low hazard had this degree of hazard at both sites, except for copper (Cu) and nickel (Ni) which were only a hazard with the

sulfuric surface soil material from site 28200 1. Many of the metals posed a greater degree of hazard with the sulfuric soil material due to their greater solubilities at lower pH values.

The reactive metal concentrations were significantly less than the ANZECC trigger value for total metals, although they were sufficiently high to be a potential hazard if released into a surrounding water body. In natural systems the dynamics of metal release will be governed by the upward chemical flux, which is a function of soil type, water flow, diffusion and chemistry of the soils near the sediment-water interface (MDBA 2011).

# **4. RISK ASSESSMENT**

### *4.1. Risk assessment framework*

Risk is a measure of both the consequences of a hazard occurring, and the likelihood of its occurrence (MDBA 2011). According to the National Environment Protection Measures (NEPM), risk is defined as "*the probability in a certain timeframe that an adverse outcome will occur in a person, a group of people, plants, animals and/or the ecology of a specified area that is exposed to a particular dose or concentration of a hazardous agent, i.e. it depends on both the level of toxicity of hazardous agent and the level of exposure*" (NEPC 1999).

In this study a risk assessment framework has been applied to determine the specific risks associated with acidification, contaminant mobilisation and de-oxygenation. In this risk assessment framework a series of standardised tables are used to define and assess risk (MDBA 2011). The tables determine the consequence of a hazard occurring (Table 4-1), and a likelihood rating for the disturbance scenario for each hazard (Table 4-2). These two factors are then combined in a risk assessment matrix to determine the level of risk (Table 4- 3).

Table 4-1 determines the level of consequence of a hazard occurring, ranging from insignificant to extreme, and primarily takes account of the environmental and water quality impacts, to the wetland values and/or adjacent waters.





Table 4-2 determines the likelihood (i.e. probability) of disturbance for each hazard, ranging from rare to almost certain. This requires an understanding of the nature and severity of the materials (including the extent and acid generating potential of acid sulfate soil materials, and the buffering capacity of wetland soil materials) as well as contributing factors influencing the risk (MDBA 2011). Examples of disturbance include: (i) rewetting of acid sulfate soil materials after oxidation, (ii) acid sulfate soil materials that are currently inundated and may be oxidised, or (iii) acid sulfate soil materials that are currently inundated and may be

dispersed by flushing (e.g. scouring flows) (MDBA 2011). As mentioned previously, the consequence of a hazard occurring and the likelihood rating for the disturbance scenario for each hazard are then ranked using a standardised risk assessment matrix (Table 4-3).

<b>Descriptor</b>	<b>Definition</b>
Almost certain	Disturbance is expected to occur in most circumstances
Likely	Disturbance will probably occur in most circumstances
Possible	Disturbance might occur at some time
Unlikely	Disturbance could occur at some time
Rare	Disturbance may occur only in exceptional circumstances

**Table 4-2: Likelihood ratings for the disturbance scenario (from MDBA 2011).** 





It is suggested that:

- For *very high* risk immediate action is recommended.
- For *high* risk senior management attention is probably needed.
- Where a *medium* risk is identified management action may be recommended.
- Where the risk is *low or very low*, routine condition monitoring is suggested.

These categories of management responses have been kept quite broad to acknowledge that jurisdictional authorities and wetland managers may choose to adopt different approaches in dealing with acid sulfate soils. The imprecise nature of these management responses is intended to provide flexibility in jurisdictional and wetland manager responses to the risk ratings associated with the acid sulfate soil hazards (MDBA 2011).

## *4.2. Assessment of risks*

The following sub-sections discuss the risks associated with acidification (Section 4.2.1), contaminant mobilisation (Section 4.2.2) and de-oxygenation (Section 4.2.3) in the Glen Esk-Rusty Waterhole. The risks associated with these hazards are dependent on a variety of factors including the scenario, wetland management regime and the species of aquatic organisms present. While likelihood of a disturbance scenario is taken into account in this risk assessment (see Table 4-2), the sensitivities and tolerances of different species of organism to each hazard has not been included. This risk assessment has primarily used the data obtained from both the Phase 1 and 2 acid sulfate soil assessments to give an overall assessment of each risk to the Glen Esk-Rusty Waterhole and adjacent waters.

### **4.2.1. Risks associated with acidification**

The Phase 1 assessment of acid sulfate soil materials in the Glen Esk-Rusty Waterhole indicated the overall degree of acidification hazard was high (Ward *et al.* 2010). While the Phase 1 assessment found moderate net acidities were dominant within this wetland, two sulfuric and three hypermonosulfidic materials had high net acidities. In addition, the water soluble sulfate content of surface soil materials at all four sites examined were over the trigger value for potential monosulfidic black ooze (MBO) formation.

Sulfuric, hypersulfidic and hypermonosulfidic soil materials which all represent an acidification hazard were identified within the Glen Esk-Rusty Waterhole. Soil profiles at three of the four sites examined (i.e. sites 28200\_1, 28200\_3 and 28200\_4) contained sulfuric soil materials with moderate-high net acidities. The surface soils (i.e. 0-10 cm) contained sulfuric soil materials at sites 28200\_1 and 28200\_3, whereas the sulfuric soil material was observed at depth (i.e. 40-90 cm) at site 28200\_4. Hypermonosulfidic soil materials with moderate-high net acidities were observed at the two sites examined at low elevation (i.e. sites 28200\_1 and 28200\_2). A hypersulfidic soil material with a moderate net acidity was observed below a depth of 40 cm at site 28200\_1.

It is expected that the consequence of an acidification hazard occurring would be *moderate* (i.e. short-term damage to wetland environmental values and/or adjacent waters; short-term impact on species and/or drinking water (including stock and domestic) supplies). The likelihood of these disturbance scenarios would be *almost certain*, and therefore there is a *high* risk associated with acidification in the Glen Esk-Rusty Waterhole.

### **4.2.2. Risks associated with contaminant mobilisation**

The high acidification hazard identified in the Glen Esk-Rusty Waterhole Phase 1 assessment indicated that soil acidification may increase the solubility of metals and soil acidity may be sufficient for the mobilisation of aluminium (Al). In addition, the presence of monosulfidic materials in some surface soils and the potential for monosulfidic black ooze (MBO) formation identified in this wetland may also result in an appreciable metal release hazard. The contaminant and metalloid dynamics data showed many of the metals examined (i.e. cobalt (Co), chromium (Cr), copper (Cu), iron (Fe), manganese (Mn), nickel (Ni), silver (Ag) and zinc (Zn)) exceeded the ANZECC water quality guidelines. The metal concentrations that exceeded the guidelines during the contaminant and metalloid dynamics test represented a low to high hazard, with only two metals (i.e. iron (Fe) and silver (Ag)) having a moderate/high hazard (see Table 3-2). Many metals posed a greater degree of hazard with the sulfuric soil material at site 28200 1 due to their greater solubilities at lower pH values.

If insufficient dilution of the contaminants was to occur in the receiving waters, there is a *moderate* consequence of a contaminant mobilisation hazard occurring (i.e. short-term damage to wetland environmental values and/or adjacent waters; short-term impact on species and/or drinking water (including stock and domestic) supplies). This disturbance scenario would be considered *likely*, and therefore there is a *medium* risk associated with contaminant mobilisation in the Glen Esk-Rusty Waterhole.

### **4.2.3. Risks associated with de-oxygenation**

Monosulfidic soil materials pose a de-oxygenation hazard if disturbed. Whilst the monosulfide formation potential test was not undertaken as part of the Phase 2 assessment for the Glen Esk-Rusty Waterhole, the presence of monosulfidic soil materials was identified within the wetland (Ward *et al.* 2010). Monosulfide concentrations  $(S_{AV}$  up to 0.12% S) were only observed at two sites at a depth of greater than 10 cm and therefore represent a low deoxygenation hazard. However, the water soluble sulfate contents of surface soil materials at all four sites were also over the trigger value for potential monosulfidic black ooze (MBO) formation. The sulfate data indicates the possible development of an appreciable deoxygenation hazard after prolonged wet conditions.

These findings therefore indicate that the de-oxygenation hazard would represent short-term damage to wetland environmental values and/or adjacent waters and short-term impacts on species and/or drinking water (including stock and domestic) supplies (i.e. *moderate* consequence of a hazard occurring). Disturbance will probably occur in most circumstances (i.e. *likely*) and therefore there is a *medium* de-oxygenation risk in the Glen Esk-Rusty Waterhole.

A summary of the risks associated with the presence of acid sulfate soils in the Glen Esk-Rusty Waterhole is presented below in Table 4-4.

#### **Table 4-4: Summary of the risks associated with acid sulfate soils in Glen Esk-Rusty Waterhole.**



# **5. BROAD ACID SULFATE SOIL MANAGEMENT OPTIONS**

This Phase 2 assessment identified the following risks associated with the presence of acid sulfate soils in the Glen Esk-Rusty Waterhole:

- high acidification risk,
- medium contaminant mobilisation risk, and
- medium de-oxygenation risk.

The acid sulfate soil materials identified in the Glen Esk-Rusty Waterhole have the potential to present a serious risk to the environmental values of both the wetland and adjacent waters if not managed appropriately. A variety of options are available to manage landscapes where acid sulfate soil materials are observed. A national guidance document on the management of inland acid sulfate soil landscapes titled "*National guidance for the management of acid sulfate soils in inland aquatic ecosystems"* has recently been released (EPHC & NRMMC 2011). The national guidance document provides a hierarchy of management options for managing acid sulfate soils in inland aquatic ecosystems including:

- *1. Minimising the formation of acid sulfate soils in inland aquatic ecosystems.*
- *2. Preventing oxidation of acid sulfate soils, if they are already present in quantities of concern or controlled oxidation to remove acid sulfate soils if levels are a concern but the water and soil has adequate neutralising capacity.*
- *3. Controlling or treating acidification if oxidation of acid sulfate soils does occur.*
- *4. Protecting connected aquatic ecosystems/other parts of the environment if treatment of the directly affected aquatic ecosystem is not feasible.*

*In some instances it may not be practical or even sensible to undertake any active intervention (for example in a pond used as part of a salt interception scheme), in which case the management objective is:* 

*5. Limited further intervention.* 

In designing a management strategy for dealing with acid sulfate soils in affected inland wetlands, other values and uses of a wetland need to be taken into account to ensure that any intervention is compatible with other management plans and objectives for the wetland.

The possible activities associated with each management objective are summarised in Table 5-1. Further information on each management option is provided in detail in the national guidance document (EPHC & NRMMC 2011).

The presence of acid sulfate soil materials with high acidification risk and medium contaminant mobilisation and de-oxygenation risks would suggest that the most appropriate management strategy for the Glen Esk-Rusty Waterhole would be to prevent oxidation of the identified acid sulfate soil materials. As outlined in Table 5-1, in order to prevent oxidation it is necessary to keep the acid sulfate soils inundated, and if possible avoid flow regimes that could re-suspend these sediments. In the event of disturbance chemical ameliorants such as lime can be added to neutralise the water column and/or sediments. Details on the ameliorants available including their advantages and disadvantages are provided in the national guidance document (EPHC & NRMMC 2011). Controlled oxidation would not be a recommended management strategy in the Glen Esk-Rusty Waterhole due to the lack of neutralising capacity within the sediments examined and the medium risk of contaminant release.





The Phase 1 acid sulfate soil assessment of the Glen Esk-Rusty Waterhole (Ward *et al*. 2010) only provided a snapshot of the acid sulfate soil materials present and the conditions at selected locations in the wetland in March 2010. Since sampling the prolonged drought in the Murray-Darling Basin has come to an end and many regions have experienced major flooding. While flooding was probably not strong enough to scour the acid sulfate soil materials from the Glen Esk-Rusty Waterhole, inundation of this wetland may have minimised the risks identified in this study in the short-term. However, it is also likely that the recent inundation will lead to further formation of acid sulfate soil materials, particularly highly reactive monosulfidic soil materials, within the Glen Esk-Rusty Waterhole.

It should be noted that further understanding of the complex interactions between surface water flow, groundwater processes, biogeochemistry and the different pathways for the development of acid sulfate soils in inland aquatic ecosystems is required for satisfactory management and preventative strategies. A more robust understanding of these complex interactions is needed before implementing any new strategies for multiple benefits.

## **6. CONCLUSIONS AND RECOMMENDATIONS**

This report provides the results of a Phase 2 investigation that was undertaken for the Glen Esk-Rusty Waterhole to determine the nature, severity and the specific risks associated with acid sulfate soil materials. The Phase 2 assessment of the Glen Esk-Rusty Waterhole examined both the contaminant and metalloid dynamics and reactive metals associated with surface layers from two sites within Glen Esk-Rusty Waterhole to identify the risk associated with contaminant mobilisation. The risks associated with both acidification and deoxygenation were primarily determined using data from the Phase 1 assessment (Ward *et al.* 2010).

The contaminant and metalloid dynamics over 35 days of inundation showed under the experimental conditions eight of the metals examined (including (cobalt (Co), chromium (Cr), copper (Cu), iron (Fe), manganese (Mn), nickel (Ni), silver (Ag) and zinc (Zn)) were found to exceed the ANZECC water quality guidelines. The guidelines for iron (Fe) and silver (Ag) were exceeded by more than 10 times at both sites. The guideline for iron (Fe) was exceeded by more than 100 times in a sulfuric soil material.

The contaminant and metalloid behaviour often varied between the metals/metalloids examined during the inundation experiments. The reductive dissolution of iron minerals seemed to have partially controlled the release of sorbed arsenic, although the controls on the release of other metals were not clear. The trend with some metals/metalloids indicates further potential for release had the incubation experiments been allowed to proceed for a longer timeframe; it is also likely that many of the metals/metalloids will become incorporated into sulfide minerals following further reduction.

The metals found to exceed the ANZECC water quality guidelines during the inundation experiments represent a low to high hazard (see Table 3-2). The reactive metal concentrations were also found to be sufficiently high to be a potential hazard if released into surrounding waters. In natural systems the dynamics of metal release will be governed by the upward chemical flux, which is a function of soil type, water flow, diffusion and chemistry of the soils near the sediment-water interface (MDBA 2011).

A risk assessment framework was applied to determine the specific risks associated with acidification, contaminant mobilisation and de-oxygenation (MDBA 2011). The Phase 2 assessment identified the following risks associated with the presence of acid sulfate soils in the Glen Esk-Rusty Waterhole:

- high acidification risk,
- medium contaminant mobilisation risk, and
- medium de-oxygenation risk.

These findings indicate that, if not managed appropriately, the acid sulfate soil materials identified in the Glen Esk-Rusty Waterhole have the potential to present a serious risk to the environmental values of both the wetland and adjacent waters. This report outlines the variety of management options available to manage acid sulfate soils in inland aquatic ecosystems. The most appropriate management strategy for the Glen Esk-Rusty Waterhole would be to prevent oxidation of the acid sulfate soil materials identified. Neutralisation may be the best management strategy in the event of disturbance. However, in designing a management strategy for dealing with acid sulfate soils in affected inland wetlands, other values and uses of a wetland need to be taken into account to ensure that any intervention is compatible with other management plans and objectives for the wetland.

It is important to note that the soil materials collected in March 2010 as part of the Phase 1 assessment only provided a snapshot of the acid sulfate soil materials present and the conditions at selected locations in the wetland. While recent inundation within the wetland

may have minimised the risks identified in the short-term, it is also likely that this inundation will lead to further formation of acid sulfate soil materials.

It is recommended that, within the context of other management objectives for the wetland, consideration be given to undertaking water quality monitoring to identify potential contamination as a result of the disturbance of acid sulfate soils within the wetland. The presence of some high risks identified in this Phase 2 assessment indicates that senior management attention is probably needed (MDBA 2011).

## **7. REFERENCES**

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# **8. APPENDICES**

## *APPENDIX 1. SOIL ANALYTICAL DATA*



#### **Table 8-1. Sample 28200\_1.1 contaminant and metalloid dynamics data.**

#### *Notes.*

*The ANZECC guideline values for toxicants refer to the Ecosystem Protection – Freshwater Guideline for protection of 95% of biota in 'slightly-moderately disturbed' systems, as outlined in the Australian Water Quality Guidelines for Fresh and Marine Water Quality (ANZECC/ARMCANZ 2000).* 

*\* ANZECC water quality guidelines for lowland rivers in South-east Australia are provided for salinity (there are currently no trigger values defined for 'Wetlands').* 

*Values outside the ranges defined in the ANZECC guidelines are indicated with red text.* 

*The deviation from the mean is represented by '±'.* 

B<br>
Guideline assumes As in solution as Arsenic (AsV).<br>
C<br>
Guideline for Chromium is applicable to Chromium (CrVI) only.<br>
H Hardness affected (refer to Guidelines).

<b>Parameter</b>	units	<b>ANZECC</b> <b>Guidelines</b>	24 hours		7 days		14 days		35 days	
			Av.	士	Av.	±.	Av.	±.	Av.	±.
pH		$6.5 - 8.0$	4.77	0.00	5.51	0.44	5.96	0.33	5.66	0.85
$EC^*$	$\mu$ S cm <sup>-1</sup>	125-2200	1477	100	1266	41	1307	143	744	5
Eh	mV		415	16	333	24	262	37	279	119
Ag	$\mu g \Gamma^1$	0.05	< 0.10	< 0.10	0.23	0.20	< 0.10	< 0.10	0.25	0.11
$AI^A$	$mgI^{-1}$	0.055	0.02	< 0.01	0.02	< 0.01	0.05	0.02	0.03	0.02
$\mathsf{As}^{\mathsf{B}}$	$\mu g \mid^{-1}$	13	1.0	1.0	1.5	1.0	4.2	1.0	6.0	4.5
Cd	$\mu g \Gamma^1$	0.2	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	0.1	< 0.1	0.1
Co	$\mu g \mid^{-1}$	2.8	1.2	1.0	1.5	1.0	1.1	1.0	1.0	1.0
$Cr^C$	$\mu g \mid^{-1}$	$\mathbf{1}$	1.3	1.0	1.0	1.0	1.1	1.0	4.2	1.9
Cu <sup>H</sup>	$\mu g \Gamma^1$	1.4	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Fe	$mgI^{-1}$	0.30	0.33	0.09	9.86	0.65	18.01	0.32	10.93	9.31
Mn	$mgI-1$	1.70	1.70	0.08	1.52	0.02	1.41	0.12	1.49	0.15
Ni <sup>H</sup>	$\mu g \mid^{-1}$	11	1.1	1.0	1.5	1.0	1.9	1.0	1.2	1.0
Pb <sup>H</sup>	$\mu g \mid^{-1}$	3.4	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Sb	$\mu g \mid^{-1}$	$\boldsymbol{9}$	1.0	1.0	1.3	1.1	1.0	1.0	1.0	1.0
Se	$\mu g \Gamma^1$	11	1.0	1.0	1.0	1.0	1.0	1.0	1.5	1.0
V	$\mu g \, \Gamma^1$	6	1.0	1.0	1.0	1.0	2.0	1.0	1.1	1.1
$Zn^{H}$	$\mu g \Gamma^1$	8	4.8	1.1	6.9	3.3	1.2	$\overline{\phantom{a}}$	2.8	1.0

**Table 8-2. Sample 28200\_1.2 contaminant and metalloid dynamics data.** 

*The ANZECC guideline values for toxicants refer to the Ecosystem Protection – Freshwater Guideline for protection of 95% of biota in 'slightly-moderately disturbed' systems, as outlined in the Australian Water Quality Guidelines for Fresh and Marine Water Quality (ANZECC/ARMCANZ 2000).* 

*\* ANZECC water quality guidelines for lowland rivers in South-east Australia are provided for salinity (there are currently no trigger values defined for 'Wetlands').* 

*Values outside the ranges defined in the ANZECC guidelines are indicated with red text.* 

*The deviation from the mean is represented by '±'.* 

B Guideline assumes As in solution as Arsenic (AsV).<br>
C Guideline for Chromium is applicable to Chromium (CrVI) only.<br>
H Hardness affected (refer to Guidelines).

<b>Parameter</b>	units	<b>ANZECC</b> <b>Guidelines</b>	24 hours		7 days		14 days		35 days	
			Av.	士	Av.	士	Av.	士	Av.	±.
pH		$6.5 - 8.0$	5.79	0.23	5.77	0.11	5.66	0.74	5.83	1.00
$EC^*$	$\mu$ S cm <sup>-1</sup>	125-2200	2484	$\overline{7}$	2355	5	2592	20	1263	47
Eh	mV		331	24	318	$\mathbf{1}$	209	4	271	134
Ag	$\mu$ g $\Gamma^1$	0.05	< 0.10	0.10	0.12	0.10	< 0.10	< 0.10	1.13	0.98
$Al^A$	$mgI^{-1}$	0.055	< 0.01	0.01	0.01	0.01	0.02	< 0.01	0.01	0.01
$\mathsf{As}^{\mathsf{B}}$	$\mu g \mid^{-1}$	13	1.0	1.0	1.9	1.0	1.6	1.1	8.4	7.1
Cd	$\mu g \Gamma^1$	0.2	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1
Co	$\mu g \Gamma^1$	2.8	2.5	1.0	2.7	1.0	2.9	1.0	2.4	1.7
$Cr^C$	$\mu g \Gamma^1$	$\mathbf{1}$	1.5	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Cu <sup>H</sup>	$\mu g \Gamma^1$	1.4	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Fe	$mgI^{-1}$	0.30	11.85	0.19	17.17	1.18	16.39	10.62	19.23	15.06
Mn	mg $\mathsf{I}^{\text{-1}}$	1.70	2.34	0.01	2.17	< 0.01	2.73	0.19	2.31	0.24
Ni <sup>H</sup>	$\mu g \mid^{-1}$	11	1.1	1.0	1.1	1.0	2.5	1.0	2.0	1.1
$Pb^{H}$	$\mu g \mid^{-1}$	3.4	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Sb	$\mu g \Gamma^1$	9	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Se	$\mu$ g l <sup>-1</sup>	11	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
V	$\mu g \mid^{-1}$	6	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
$Zn^{H}$	$\mu g \, \Gamma^1$	8	1.5	1.0	5.2	1.0	9.1	$\overline{\phantom{a}}$	3.2	2.4

**Table 8-3. Sample 28200\_2.1 contaminant and metalloid dynamics data.** 

*The ANZECC guideline values for toxicants refer to the Ecosystem Protection – Freshwater Guideline for protection of 95% of biota in 'slightly-moderately disturbed' systems, as outlined in the Australian Water Quality Guidelines for Fresh and Marine Water Quality (ANZECC/ARMCANZ 2000).* 

*\* ANZECC water quality guidelines for lowland rivers in South-east Australia are provided for salinity (there are currently no trigger values defined for 'Wetlands').* 

*Values outside the ranges defined in the ANZECC guidelines are indicated with red text.* 

*The deviation from the mean is represented by '±'.* 

B Guideline assumes As in solution as Arsenic (AsV).<br>
C Guideline for Chromium is applicable to Chromium (CrVI) only.<br>
H Hardness affected (refer to Guidelines).

<b>Parameter</b>	units	<b>ANZECC</b> <b>Guidelines</b>	24 hours			7 days	14 days		35 days	
			Av.	士	Av.	±.	Av.	士	Av.	±.
pH		$6.5 - 8.0$	5.92	0.16	6.06	0.04	6.45	0.17	5.84	1.00
$EC^*$	$\mu$ S cm <sup>-1</sup>	125-2200	1883	46	1786	42	1778	71	964	3
Eh	mV		285	10	300	5	176	31	277	130
Ag	$\mu g \Gamma^1$	0.05	< 0.10	0.10	< 0.10	< 0.10	0.21	0.21	0.35	0.28
$AI^A$	$mgI^{-1}$	0.055	< 0.01	0.01	0.03	< 0.01	0.04	0.02	0.02	0.01
$\mathsf{As}^{\mathsf{B}}$	$\mu$ g l <sup>-1</sup>	13	1.0	1.0	2.3	1.0	5.3	1.0	4.4	4.2
Cd	$\mu g \Gamma^1$	0.2	< 0.1	0.1	< 0.1	< 0.1	< 0.1	< 0.1	< 0.1	0.1
Co	$\mu g \Gamma^1$	2.8	2.0	1.0	1.9	1.0	1.6	1.0	3.4	2.2
$Cr^C$	$\mu g \mid^{-1}$	$\mathbf{1}$	1.7	1.0	1.0	1.0	1.0	1.0	1.1	1.0
Cu <sup>H</sup>	$\mu g \Gamma^1$	1.4	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Fe	$mgI^{-1}$	0.30	9.93	2.93	16.01	0.78	22.53	0.94	8.88	7.98
Mn	$mg \, \Gamma^1$	1.70	1.65	0.03	1.51	0.02	1.73	0.04	1.71	0.15
Ni <sup>H</sup>	$\mu$ g l <sup>-1</sup>	11	1.5	1.0	1.6	1.0	2.5	1.0	3.2	2.1
Pb <sup>H</sup>	$\mu g \mid^{-1}$	3.4	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Sb	$\mu g \Gamma^1$	9	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
Se	$\mu g \, \Gamma^1$	11	1.0	1.0	1.0	1.0	1.0	1.0	1.0	1.0
$\vee$	$\mu g \Gamma^1$	6	1.0	1.0	1.0	1.0	1.2	1.0	1.0	1.0
$Zn^{H}$	$\mu g \mid^{-1}$	8	4.4	1.5	3.2	-	7.1	7.1	9.3	8.8

**Table 8-4. Sample 28200\_2.2 contaminant and metalloid dynamics data.** 

*The ANZECC guideline values for toxicants refer to the Ecosystem Protection – Freshwater Guideline for protection of 95% of biota in 'slightly-moderately disturbed' systems, as outlined in the Australian Water Quality Guidelines for Fresh and Marine Water Quality (ANZECC/ARMCANZ 2000).* 

*\* ANZECC water quality guidelines for lowland rivers in South-east Australia are provided for salinity (there are currently no trigger values defined for 'Wetlands').* 

*Values outside the ranges defined in the ANZECC guidelines are indicated with red text.* 

*The deviation from the mean is represented by '±'.* 

B Guideline assumes As in solution as Arsenic (AsV).<br>
C Guideline for Chromium is applicable to Chromium (CrVI) only.<br>
H Hardness affected (refer to Guidelines).





*\* The ANZECC sediment quality guidelines (SQG) are for total metal concentrations (ANZECC/ARMCANZ 2000).* 

*The deviation from the mean is represented by '±'.* 

