Mitigating AMD Impacts in New Zealand Using Engineered Wetlands¹

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Abstract. Coal mining in New Zealand has caused perturbation of water resources and biodiversity. Contaminants impairing local waterbodies include acidity, iron, aluminum, arsenic, manganese, nickel, zinc, copper, sulfate and suspended solids. Exposure of sulfur containing rocks, such as pyrite, to atmospheric oxygen during mining operations produces acid mine drainage (AMD). Sulfuric acid and metal acidity are generated and can accentuate metal mobilization and bioavailability. Metals favor the dissolved state in acidic environments but form less toxic precipitates when exposed to adequate alkalinity. Metal toxicity effects are synergistic dependent on metals speciation and their concentrations.

New Zealand is in the initial stages of acid mine drainage mitigation and has yet to develop proven treatment technologies. Implementation of passive treatment methods, such as engineered wetlands, have successfully reduced acid mine drainage impacts worldwide. Design criteria for these systems are improving while their limitations are well documented. We are currently collecting water quality and flow data from selected AMD-impacted sites. We are in the process of designing pilot-scale engineered wetland systems to ameliorate acid mine drainage in New Zealand. Sequential-treatment trains will be constructed and their performance evaluated in order to optimize design effectiveness.

New Zealand acid mine drainage characteristics and complex topography offer unique challenges for implementing treatment systems. The AMD typically contains very high aluminum concentrations (commonly exceeding 50 g/m³) and has an aluminum to iron concentration ratio of three to one. Abundant steep topography can be exploited to create adequate driving head for implementing systems such as SCOOFI reactors while reducing and alkalinity producing systems can also be employed. Precipitation of up to six meters per year contributes to dynamic hydraulic characteristics and will offer unique design and treatment challenges.

Additional Key Words: aluminum, iron, water quality, New Zealand, acid mine drainage, coal mining, steep topography

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Introduction

It is reported that coal mining in New Zealand began in the 1830s with mining districts becoming established in the 1870s for commercial purposes (Pattrick, 2004, Solid Energy, 2005). Solid Energy New Zealand Limited (SENZ) represents a state owned and operated coal company developing from a government trading enterprise established in 1901 and privatized in 1987 as Coal Corporation of New Zealand. Current production exceeds 4 billion kilograms of coal annually from its seven underground and opencast mines in the North and South Islands (Solid Energy, 2005). Most acid mine drainage (AMD) impacts occur on the South Island from estuarine coal formations containing high concentrations of sulfur (Pers. Comm. Paul Weber, Environmental Research Manager SENZ, 2005). North Island coals and a significant amount of South Island coals do not typically generate AMD since they formed from lacustrine accretions containing low sulfur concentrations. Historical coal mining has resulted in contamination of numerous streams with AMD. Solid Energy inherited many of these environmental liabilities and are in the initial stages of addressing AMD impacts from historical and current mining activities. The contents of this report will focus on AMD issues from South Island coal mining as these sites are most impacted by AMD.

Problem Statement

Numerous streams in New Zealand contain high concentrations of acidity and metals from unabated AMD runoff. Carbonaceous mudstones and coal represent the primary sources of acidity by producing sulfuric acid when exposed to oxygen (Trumm et al., 2005). Metal cations also contribute acidity upon hydrolyzing with water. Metals from the parent rock material leach into the aqueous phase when exposed to acidity. Sandstones abutting the mudstones contribute metals leachate, and some sulfur, when exposed to acidity but are not the major acidity contributors.

Pyrite (FeS₂) represents the primary source of iron. The primary sources of aluminum include potassium feldspar or microcline (KAlSi₃O₈), muscovite (KAl₂(AlSi₃O₁₀)(F,OH)₂) and kaolinite (Al₂Si₂O₅(OH)₄). The sequence of reactions for pyrite oxidation is well documented (e.g. Watzlaf et al., 2003). However, it is important to note that following initial pyrite

oxidation, ferric iron acts as an oxidizing catalyst for acidity generation. This reaction is summarized as follows (Equation 1):

$$FeS_2 + 14Fe^{3+} + 8H_2O \rightarrow 15Fe^{2+} + 2SO_4^{2-} + 16H^+$$
Pyrite + Ferric Iron + Water \rightarrow Ferrous Iron + Sulfate + Acidity

When potassium feldspar is exposed to acidity, kaolinite and silicic acid are produced (Equation 2) (Younger et al., 2002; Watzlaf et al., 2003). Kaolinite is further degraded by acidity, resulting in mobile aluminum ions (Equation 3). Aluminum precipitates in the presence of water (measured at a pH = 4.8 from Stockton Mine Water (Pers. Comm. Joseph Holman, Environmental Engineering Lab Technician, University of Canterbury, 2005)) and produces acidity (Equation 4). Thus, aluminum-containing rock which is exposed to proton acidity results in the release of aluminum (Equations 2 and 3), coupled with the production of acidity upon aluminum hydroxide formation (Equation 4), and so the process of aluminum released into solution can be perpetuated. In New Zealand West Coast rock formations, aluminum-containing rocks and low pH (typically near or less than 3) contribute to the relatively high aluminum to iron proportions uncharacteristic of the majority of AMD water elsewhere.

$$KAlSi_3O_8 + H^+ + 9/2H2O \rightarrow 2H_4SiO_4 + 1/2Al_2Si_2O_5(OH_4)$$
 (2)

Potassium Feldspar + Acidity + Water → Silicic Acid + Kaolinite

$$Al_2Si_2O_5(OH_4) + 6H^+ \rightarrow 2Al^{3+} + 2H_4SiO_4 + H_2O$$
 (3)

Kaolinite + Acidity → Aluminum Ions + Silicic Acid + Water

$$Al^{3+} + 3H_2O \rightarrow Al(OH)_3 + 3H^+$$
 (4)

Aluminum Ions + Water → Aluminum Hydroxide (ppt) + Acidity

Other metals such as arsenic, manganese, nickel, zinc and copper can also leach from source rocks under acidic conditions and increase AMD toxicity to aquatic species.

Proposed Methodology

Passive treatment systems are proven treatment technologies if designed and implemented properly (Younger et al., 2002). Possible systems may include inorganic media passive systems (IMPs). Candidate IMPs include surface catalyzing oxidation of ferrous iron reactors (SCOOFIs) to exploit the New Zealand steep topography and oxic limestone drains and open-channel limestone reactors to generate alkalinity. Engineered wetland types appropriate for New Zealand AMD include sedimentation ponds, reducing and alkalinity producing systems (RAPS), sulfate-reducing bioreactors (such as compost wetlands) and aerobic wetlands, amongst others.

Our research consists of designing and evaluating the performance of multiple pilot-scale engineered wetlands to determine which systems offer the most cost effective treatment options for ameliorating AMD. The initial six-month period will consist of characterisation of AMD at the research field site. Characterisation of samples will quantify contaminant concentrations and loading variability during periods of high and low flows. Basic water quality parameters including pH, turbidity, conductivity and temperature will be continuously recorded using data loggers. Flow ranges will also be recorded using data loggers. Multiple sampling trips will aim to comprehensively characterise the variability in seasonal loadings of metal contaminants to the impacted watershed.

Design criteria developed overseas will be adopted for choosing and sizing system designs. Systems will be designed and constructed to include adequate replication and variation in design parameters. Different cell sequences will be considered to evaluate their treatment effectiveness. Rates of sedimentation and precipitation of metal-hydroxides, their rate-limiting steps and characterising water throughout the treatment process, in addition to materials used and cost implications, will be considered in the final designs.

Water and soil substrate samples will be collected and analysed periodically to determine chemical dynamics and metal speciation within the systems. This data will provide information on the chemical transformations and consequent treatment efficiency of the systems. Chemical modelling will be performed to evaluate treatment limits based on treatment performance

measured from flow and chemical data. An evaluation of the long term suitability of these systems and the potential for metal precipitates to remobilize will be considered. Results will also be used to develop design criteria for passive treatment systems at other sites Because every site is heterogeneous and offers unique challenges, system designs will be modified based on site characteristics, constraints, flow and contaminant concentrations. Since this project officially commences January 2006, we have not yet ascertained the most appropriate designs. These will depend on contaminant loading regimes deciphered by May 2006.

Site Selection

The site where the trialed engineered wetlands will be constructed is under final consideration. Efforts will focus on AMD on the West Coast of the South Island where unique challenges such as high aluminum concentrations and six meters of rainfall annually predominate. A summary of candidate sites including advantages and disadvantages of site selection is given in Table 1. At present, Sullivan Mine is the preferred site for rehabilitating since it offers the best rehabilitation return for the efforts expended (see advantages).

Current Data

Sporadic water quality and flow data exists for numerous AMD sources and streams impacted by AMD on the West Coast. However, data collection has not been consistent, quality assured or considered analytes necessary to obtain adequate design criteria at most sites. The pH in many West Coast South Island streams is naturally acidic fluctuating between 4.0 and 5.0 (Harding, 2005). The pH of AMD is commonly less than 3.0 at source and throughout some impacted streams (Trumm et al., 2005). Background aluminum concentrations in some non-AMD impacted streams can also exceed permitted trigger values (Pers. Comm. Phil Lindsay, Environmental Operations, SENZ, 2005). At the Stockton site, acidity can range between 200-800 g CaCO₃/m³, dissolved ferrous iron of 15-50 g/m³, dissolved aluminum of 50-100 g/m³ and sulfates from 200-800 g/m³ (1 g/m³ = 1 mg/L). Data collected from Sullivan Mine AMD between February 2001 and June 2002 contained an average pH of 2.8, 192 g acidity/m³ as CaCO₃, 366 g sulfate/m³, 13.75 g dissolved aluminum/m³, 0.012 g dissolved arsenic/m³, 41.58 g

dissolved iron/m³, 0.5 g dissolved manganese/m³, 0.12 g dissolved nickel/m³ and 0.7 g dissolved zinc/m³ (Trumm et al., 2005). These concentrations for Al, Fe, Ni and Zn were all above the acceptable trigger values stipulated in New Zealand legislation (ANZECC, 2000).

Table 1. Site selection potential and priorities.

Site	Advantages	Disadvantages	Additional Comments
Echo Mine/	1) Likely improvement of water	1) Active mining from other	1) Multiple potential sites.
Island Block	quality and biodiversity.	companies on site.	2) Three separate streams
	2) Good location for assessing	2) High level of uncertainty of	receiving AMD.
	effectiveness of SCOOFIs	controlling system inputs	3) Most logical trial system
	because of steep topography.	because of active mining.	site based on available land
	3) Closest site to offices in	3) Adequate land area not	area and current mining
	Christchurch.	available at all potential sites.	operations would only
		4) Construction equipment and	protect a few hundred
		materials would have to be	meters of stream.
		brought in from offsite.	
Stockton	1) Likely improvement of water	1) Active mine with	1) Multiple potential sites.
	quality and biodiversity.	uncertainties regarding	2) High political pressure to
	2) Actively mined by SENZ so	controlling system inputs such	mitigate AMD issues here.
	construction equipment and	as sediment loading.	3) Receives the highest
	technical/logistical support	2) Furthest site from offices in	rainfall of all potential sites
	available on site.	Christchurch.	and also has the highest
	3) Adequate land area.		aluminum concentrations.
Sullivan	1) Likely improvement of water	1) Construction equipment and	1) Two adits contribute
	quality and biodiversity.	materials would have to be	AMD to Rapid Stream with
	2) Adequate land area.	brought in from offsite.	most of the contaminant
	3) Underground abandoned	2) Minor road repairs need on	contribution (99 percent or
	mine with consistent flow from	access road.	more) from the lower adit.
	adit prior to discharging into		
	impacted stream.		
	4) Most data available of all the		
	site options.		
	5) Located in a highly valued		
	recreational area with		
	numerous hiking trails.		
	6) Treating the major		
	contaminant source would		
	reduce most AMD inputs into		
	impacted stream and therefore		
	allow for feasible stream		
	restoration.		

Regulatory Criteria

The lack of historical background water quality data and dominance of endemic aquatic organisms in New Zealand make determining acceptable regulatory criteria challenging for freshwater environments (Harding et al., 2005). Overseas toxicity data must be used with

caution in the New Zealand context. New Zealand environmental law is mandated in the Australian and New Zealand Environment and Conservation Council (ANZECC) Water Quality Guidelines (ANZECC, 2000). Applicable toxicity data for freshwater ecosystems is represented as trigger values with four levels of protection based on percent of species not affected as shown in Table 2. The 80-precent level of protection trigger values are typically mandated by local regional councils for streams impacted by AMD.

Table 2. ANZECC trigger values (g/m³) for freshwater toxicants (ANZECC, 2000)

Element	Level of Protection (% Species)				
	99%	95%	90%	80%	
Al (pH<6.5)	ID*	ID*	ID*	ID*	
Al (pH>6.5)	0.027	0.055	0.080	0.015	
As^{3+}	0.001	0.024	0.094	0.360	
As ⁵⁺	0.0008	0.013	0.042	0.140	
Fe	ID*	ID*	ID*	ID^*	
Mn	1.20	1.90	2.50	3.60	
Ni	0.008	0.011	0.013	0.017	
Zn**	0.0024	0.0080	0.015	0.031	
Cu**	0.0010	0.0014	0.0018	0.0025	

^{*} ID indicates insufficient data.

Conclusions

Passive treatment systems including engineered wetlands have been implemented worldwide for successfully ameliorating AMD. It is believed that engineered wetlands can be successfully utilized in New Zealand once adequate water quality and flow data is collected and analyzed (O'Sullivan, 2005). Before appropriate designs can be developed, research needs to be conducted to optimize system designs and sizing. Cost-analysis needs to be performed to determine benefits and feasibility of engineered wetlands versus adopting more active (traditional) chemical dosing treatment approaches. Long-term monitoring should be conducted

^{**} The values listed assume a hardness value of 30 g/m³ as CaCO₃.

to verify system function and determine long-term applicability. Maintenance such as solids removal must be performed to ensure long term operation and prevent the systems from clogging with sediment, ferric and aluminium hydroxides.

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