PASSIVE AND SEMI-ACTIVE TREATMENT OF ACID ROCK DRAINAGE FROM METAL MINES - STATE OF THE PRACTICE

Prepared for
U.S. Army Corps of Engineers
696 Virginia Road
Concord, Massachusetts  01742

April 2, 2003

URS

URS Corporation
477 Congress Street Annex, Suite 3A
Portland, Maine  04101
# TABLE OF CONTENTS

1. **Section 1 Introduction** ................................................................................................... 1-1
   1.1 Limitations of Available Information ........................................................................... 1-1

2. **Section 2 State of the Practice** .................................................................................. 2-1
   2.1 Passive Treatment History .......................................................................................... 2-1
   2.2 Currently Available Passive Treatment Components ............................................... 2-2
      2.2.1 Aerobic Wetlands ................................................................................................. 2-3
      2.2.2 Anoxic Limestone Drains ..................................................................................... 2-4
      2.2.3 Open Limestone Channels ................................................................................... 2-5
      2.2.4 Settling Ponds ...................................................................................................... 2-5
      2.2.5 Successive Alkalinity Producing System (SAPS) .................................................. 2-5
      2.2.6 Solid-Reactant Anaerobic or Sulfate Reducing Bacteria (SRB) Bioreactors ........... 2-6
      2.2.7 Liquid-Reactant (Semi-active) Bioreactors .......................................................... 2-9

3. **Section 3 Case Studies** .................................................................................................. 3-1
   3.1 Burleigh Tunnel Pilot Cells .......................................................................................... 3-1
   3.2 Brewer Pad 5 Pilot Cell .............................................................................................. 3-1
   3.3 Ferris-Haggerty Mine/Osceola Tunnel ....................................................................... 3-2
   3.4 Doe Run Mine ............................................................................................................ 3-2
   3.5 Leviathan Mine ........................................................................................................... 3-3

4. **Section 4 Acid Rock Drainages at the Elizabeth Mine** ................................................. 4-1
   4.1 GEOLOGY AND MINE FEATURES ......................................................................... 4-1
   4.2 GEOCHEMISTRY OF ACID ROCK DRAINAGE ....................................................... 4-1

5. **Section 5 Conceptual Design Approach for the Elizabeth Mine** ................................. 5-1
   5.1 Conceptual Design ...................................................................................................... 5-1
   5.2 Feasibility Level Cost Estimate ................................................................................... 5-2

6. **Section 6 References** .................................................................................................... 6-1
TABLE OF CONTENTS

Tables

Table 1  Feasibility Level Capital Cost Estimate
Table 2  Feasibility Level Operation and Maintenance Cost Estimate

Figures

Figure 1  Flowchart for Selecting a Passive Acid Rock Drainage Treatment System Based on Water Chemistry and Flow
Figure 2  Anoxic Limestone Drain Plan Profile and Section
Figure 3  Typical Cross-Section View of a Successive Alkalinity Producing System (SAPS) Treatment Component
Figure 4  Schematic Cross Section of Upflow and Downflow Sulfate Reducing Bacteria Anaerobic Bioreactor

Appendices

Appendix A  Bibliography
Appendix B  Overview Papers
Appendix C  Manganese Removal Papers
Appendix D  Papers by Jim Gusek et al. on Solid-Reactant SRB Bioreactors
Appendix E  Papers by Tim Tsukamoto et al. on Liquid-Reactant SRB Bioreactors
SECTION ONE

Introduction

This document presents an overview of current state-of-the-practice applications for passive and semi-active treatment systems to treat acid rock drainage (ARD) associated with metal mines. Acid rock drainage is a common effluent of metal mining and one of the major environmental impacts resulting from mining activities. It is caused by the natural weathering of pyrite and other metal sulfides in the mineral deposit, or through the accelerated weathering of waste products generated by the mining process. The sulfide minerals react with oxygen in air or pore water and produce sulfuric acid. Acid rock drainage is thus low-pH water with elevated concentrations of iron, sulfate, and heavy metals and metalloids of varying composition dependant upon the originating mineral deposit type.

The purpose of this document is to identify passive and semi-active treatment technologies that could potentially be used at the Elizabeth Mine Site (herein after referred to as the Site), present general capital, operation and maintenance costs for specific treatment technologies, and discuss the relative success of these treatment technologies at other sites. Technologies reviewed are inclusive of the passive treatment methods of aerobic wetlands, anoxic limestone drains (ALD), open limestone channels (OLC), settling ponds, successive alkalinity-producing systems (SAPS), and solid-reactant anaerobic sulfate-reducing bacteria (SRB) bioreactors, and the semi-active method of liquid-reactant SRB bioreactors.

1.1 LIMITATIONS OF AVAILABLE INFORMATION

The evaluation of passive treatment systems has involved numerous discussions with system designers and operators. Numerous examples of failed systems have been identified. In the majority of these cases, evaluations to determine how and why systems failed were not generally performed, or this information was not otherwise made available to URS. It was found that the owners were unwilling to invest additional money to evaluate the failure cause without financial benefit. Therefore, detailed information on system failures is somewhat limited.

Additionally, as many treatment technology applications have been research-based, for proprietary reasons developers have been guarded in discussing details of either processes or applications with URS.
This section describes the generalized history of passive treatment systems and currently available passive treatment system components.

The overview is based on application experience combined with information provided by several well-known experts in the field, an extensive literature search, and review of recent technical publications. Appendix A presents a bibliography of reviewed publications, arranged by topic. Appendix B includes several overview papers useful in defining the technologies and applications history. Significant recent publications outlining case studies of individual applications are included as Appendices C, D, and E. Information provided and case studies reviewed include those associated with publicly available records. Information on systems developed and/or operating under confidentiality agreements are not included.

2.1 PASSIVE TREATMENT HISTORY

The cost of potentially perpetual treatment of ARD using traditional active lime treatment technology has caused the mining and regulatory communities to search for less expensive alternative treatment solutions. In the late 1970s and early 1980s, evidence from coal mine drainages suggested that wetlands, with their complex mix of biological, chemical, and physical processes, could potentially treat ARD. In the 1980s, the use of constructed wetlands was attempted to mimic natural wetlands and treat ARD.

Researchers developed hypotheses as to which wetland processes (e.g., plant uptake, adsorption, microbial reactions) were most important in the treatment of ARD. Early research in the field of passive wetland treatment focused on maximizing these processes. In the late 1980s and early 1990s, a practical understanding of the biological and chemical processes was developed and several authors (i.e., Brodie 1992; and Hedin, Nairn, and Kleinmann 1994) published guidelines for designing treatment systems (e.g., aerobic constructed wetlands) based on the influent water chemistry (Figure 1).

In one early investigation of a treatment wetland (Brodie 1992), ARD emanating from the base of a coal debris pond flowed under an old roadbed made of limestone rubble. The limestone from the roadbed was found to add alkalinity to the anoxic subsurface flow, which in turn allowed iron to precipitate when it became oxidized in the adjacent wetland. From this research originated the ALD concept. Open limestone channels were also used to add alkalinity to ARD before it enters a constructed wetland.

Much of the early work focused on passive treatment of ARD at eastern U.S. coal mine sites. These techniques were then applied to hard-rock drainages. Investigators found that typical hard-rock ARD had lower pH (i.e., <4) and higher concentrations of heavy metals (e.g., copper, zinc) that could not be treated as effectively with aerobic surface-flow constructed wetlands (Day, Filipek, and Papp 1986). In response, researchers attempted to employ subsurface processes of natural wetlands, mainly bacterially mediated anaerobic sulfate reduction systems (e.g., SRB systems). The sulfide produced by this process was found to bind with iron and most heavy metals, re-forming the same minerals that had originally been mined. Constructed wetlands designed to form metal sulfides are referred to as SRB constructed wetlands.

To maximize the subsurface processes, the ARD flow was discharged through an organic substrate that typically had relatively low permeability. The treatment processes that remove the
metals tended to further decrease the permeability during operation. Finally, researchers found that anaerobic systems typically function better without vegetation and when they are covered. Current anaerobic system designs typically include pipes, valves, back-flush mechanisms, and other engineered features not found in natural wetlands. Thus, the anaerobic systems have evolved into designed systems that attempt to mimic natural geochemistry. These systems are better termed anaerobic bioreactors than anaerobic constructed wetlands because their design must counter physical and biological processes that occur naturally in wetlands.

The advancement of ARD treatment technologies for hard rock drainage has occurred in conjunction with advances in coal mine drainage treatment methods. For example, the SAPS used to treat components of ARD is a variant of the anaerobic bioreactor developed for coal mine drainage. Kepler and McCleary (1994) first reported using this concept to solve the problem of precipitation of iron hydroxide minerals and concomitant clogging in ALDs whenever the influent ARD was oxygenated and had high concentrations of iron. The goals of a SAPS design are to remove oxygen and reduce Fe$^{3+}$ to Fe$^{2+}$, but not necessarily to precipitate metal sulfides. Thus, there is no attempt to maximize SRB processes.

To overcome the permeability issues typically associated with anaerobic bioreactors, some designers have added liquid organic reactants, such as alcohol to the treatment process, rather than the compost, wood chips, sawdust, or similar solid organic reactants used in most anaerobic bioreactors as documented by Tsukamoto and Miller. Such systems are referred to in this text as semi-active technologies.

The benefit of these systems is they can be relatively low maintenance, perhaps as low as the solid-reactant systems. In a variant of this method, a small plant can be built to co-treat ARD with a local processing waste or the liquid in sewage so that the two waste streams essentially treat each other. To the best of our knowledge, to date only one such co-treatment pilot system has been built (i.e., Jerome, Arizona) as documented by Allbright (2002).

During the past 20 years since research in passive and semi-active treatment methods began, researchers have gained a better understanding and respect for the complex interaction of the myriad of biological, chemical, and physical processes involved in the formation and treatment of ARD. The flow chart presented in Figure 1 shows an example approach to selecting a functional passive system. The concept of passive treatment has been tempered by experience. To date there has been no successful application of a completely passive method that effectively treats the typically less benign ARD from heavy metal mining. This review has found that all constructed systems to date have required some level of ongoing maintenance and periodic reconditioning (reconstruction) and few are currently in operation.

2.2 CURRENTLY AVAILABLE PASSIVE TREATMENT COMPONENTS

The following section summarizes the state of the practice of each of the common methods of passive and semi-active ARD treatment including aerobic wetlands, ALDs, OLCs, solid-reactant SRB anaerobic bioreactors, SAPS, and liquid-reactant SRB anaerobic bioreactors. To address specific issues relevant to Elizabeth Mine, the discussion focuses on solid- and liquid-reactant anaerobic bioreactors used to treat ARD.
2.2.1 Aerobic Wetlands

Aerobic wetlands and subaerobic wetlands are similar to natural wetlands in that the water flows mainly over the substrate surface. This type of wetland is well understood because it has a relatively long application history in municipal sewage treatment systems. Aerobic wetlands are typically shallow excavations filled with one to three feet of soil, gravel, and/or rocks in a hummocky pattern. The designed hummocks allow for variations in water depth of between one inch to approximately one foot to form a diversity of microenvironments. In these microenvironments, consortia of micro- and macro-organisms carry out a wide variety of biogeochemical processes.

The dominant processes treating mine drainage in the deeper portions of aerobic wetlands are oxidation of Fe$^{+2}$ to Fe$^{+3}$ by aeration spillways, precipitation of Fe$^{+3}$ hydroxides, and settling of the hydroxides in settling ponds. Plants are typically used within the system because they help trap suspended matter and provide additional oxygen. Aerobic wetlands require sufficient alkalinity in the water to keep the pH from falling as a result of the hydrolysis of iron, following the reaction:

$$\text{Fe}^{3+} + 3 \text{H}_2\text{O} \rightarrow \text{Fe(OH)}_3 + 3 \text{H}^+$$

Through trial and error, a number of investigators have concluded that a system consisting only of aerobic wetlands is effective only in treating relatively benign ARD (e.g., pH >4.5, significant alkalinity, and relatively low heavy metal content). However, aerobic wetlands are often included as a final process step in a treatment system containing one or more other passive treatment methods, such as ALDs, OLCs, and/or anaerobic bioreactors. When used with ALDs or OLCs, aerobic wetlands act as oxidation and settling ponds. They are also typically used as polishing cells for effluent from SRB bioreactors to remove the nitrogen and organic breakdown products solubilized during SRB treatment. Aerobic wetlands have been used successfully to treat manganese, as discussed below.

**Manganese Treatment by Aerobic Wetlands**

Nearly all manganese will pass through ALDs, OLCs, and SRB bioreactors because manganese does not readily form manganese sulfides and, without catalysts, manganese requires a pH greater than 9 to precipitate. The dominant manganese removal processes are oxidation of Mn$^{+2}$ to Mn$^{+4}$, precipitation of Mn$^{+4}$ oxides, and settling of the oxides in settling ponds as a final polishing step after treatment for removal of acidity and all other metals.

Treatment of manganese requires a special type of aerobic wetland that is highly oxidizing and contains populations of microorganisms that help catalyze the precipitation reaction by locally raising the pH to between 8 and 9. To achieve these conditions, an aeration spillway typically precedes the wetlands to ensure that the water is highly oxidized.

Manganese wetlands are typically shallow rock wetlands colonized with an algal mat of green algae and cyanobacteria designed to locally raise pH and the oxidation-reduction potential (Eh) (Wildeman et al. 1993). A wetland without the appropriate algae and bacteria must be approximately five times as large as one with such a mat in order to achieve similar levels of effectiveness.
Adsorption and/or coprecipitation of heavy metals onto iron or manganese oxide minerals is a function of pH. The higher the pH, within the range of natural stream environments, the greater the adsorption (Filipek, Nordstrom, and Ficklin 1987). If the wetland is able to precipitate manganese, it will also be capable of significant removal of certain heavy metals, including copper, lead, and zinc (Filipek, Chao, and Carpenter 1981).

Proprietary methods for designing and constructing mixed microbial mats to treat manganese and other heavy metals are available through developers such as Microbial Aquatic Treatment Systems, Inc. Refer to Appendix C for information on this technology. However, applications experience has shown that similar results can typically be obtained by inoculating a wetland with sufficient matting (i.e., 5 to 10 percent of the wetland area) or by inoculating a shallow rock wetland with a naturally occurring consortium of green algae and cyanobacteria.

The process of obtaining a consortium of algae and bacteria acclimated to site conditions can be as simple as placing hay bales in a stream near the site in spring or summer and allowing the growth of green bacteria and algae to develop on the hay, which then can be dispersed into the constructed wetland as inoculum. Inoculating is effective because the algae and bacteria tend to “self-organize” to form and maintain their appropriate micro environments and the process can be achieved at a significantly reduced cost compared to the proprietary mats.

### 2.2.2 Anoxic Limestone Drains

An ALD is a trench filled with crushed high-calcium limestone, sealed with geotextile or plastic, and covered with clay or soil to prevent oxygen inflow (Figure 2). It is typically built into a hillside or tailing pile to capture ARD that has not yet been exposed to oxygen. As the acidic water flows through the ALD, the acid dissolves some of the limestone, which adds alkalinity to the water and raises the pH.

The US Bureau of Mines (Hedin, Nairn, and Kleinmann 1994) has produced guidelines for ALD design. These guidelines include: 1) the limestone should have at least 80-percent CaCO$_3$ content as the calcium-rich limestone is significantly more reactive than magnesium-rich limestone; 2) the limestone should consist of Nos. 3- or 4-sized materials (e.g., baseball-size) to minimize compaction and permeability decrease with time; 3) the limestone should be buried beneath several feet of clay, with heavy plastic separating the clay and limestone to limit oxygen infiltration; 4) the limestone should be inundated at all times; and, 5) to ensure limestone dissolution, the ALD is to be sized to provide a minimum retention time of 14 hours.

According to Hedin et al., a 14-hour retention time requires approximately 3000 kilograms (kg) of calcium carbonate for each liter per minute (l/min) of flow, regardless of water chemistry. These guidelines for designing an ALD have not changed significantly since published in 1994.

A properly designed ALD can raise the alkalinity of water flowing through it to about 200 milligrams per liter (mg/L) CaCO$_3$ equivalent. The guidelines for ALDs (Figure 1) indicate that they are most functional if the influent has the following characteristics:

- Dissolved oxygen (DO) <2 mg/L, oxidized Fe$^{+3}$ <10 percent of the total iron, and
- Aluminum is <25 mg/L in the initial drainage.

Iron (III) and aluminum will precipitate on and coat, or armor, the limestone. The coating and aluminum floc will significantly reduce the limestone’s reactivity and the permeability of the
system, especially in low-flow conditions. Similarly, high concentrations of sulfate (i.e., > 1,500 mg/l) in acid rock drainage can cause precipitation of gypsum as the limestone dissolves, which will also reduce permeability (Nairn, Hedin, and Watzlaf 1991).

### 2.2.3 Open Limestone Channels

The OLC is a variant of the ALD and is used to treat discharges that are oxygenated and contain Fe$^{+3}$ or high aluminum content. The OLC can be effective in adding alkalinity to ARD and raising the pH. However, OLCs require an environment that will self-scour the exposed limestone surface.

Ziemkiewicz et al. (1997) found that armored limestone in a turbulent stream was 50- to 90-percent as effective as unarmored limestone in neutralizing acid. The highest neutralization rates were associated with channels having slopes of between 45- and 60-percent, but some neutralization occurred on slopes as gentle as 8-percent. Accordingly, if sufficient topographic relief is present, an OLC is significantly more cost effective, is more easily monitored, and is more likely to be effective than an ALD for oxidized, aluminum-rich waters.

Design of an OLC must have a significant vertical gradient to allow for turbulent flow to strip off precipitates and must contain a number of small ponding areas between turbulent points to collect the resultant precipitates. Sizing considerations are similar to those for an ALD, except that typically a 50-percent efficiency should be assumed for yearly dissolution tonnages to account for potential armoring.

### 2.2.4 Settling Ponds

Settling ponds are used to collect treated or partially treated waters discharging from an ALD or OLC. These ponds allow iron and other precipitates to settle and are useful in providing a more constant flow rate into a downgradient treatment cell (e.g., SRB bioreactor). Settling ponds should be sized to allow a retention time of approximately 14 days.

Upon emerging from an ALD, water with reduced Fe$^{+2}$ must be oxygenated in the atmosphere to allow oxidation of the iron and precipitation of iron oxyhydroxides such as ferrihydrite. Oxygenation of the water can be most simply accomplished by having it flow over a series of riffles before it enters a pond. At common temperatures, oxygen is soluble in water to approximately 8 mg/L. Stoichiometrically, 8 mg/L of oxygen can oxidize approximately 50 mg/L of Fe$^{+2}$. Therefore, each settling pond can only remove approximately 50 mg/L of initially reduced iron unless it is equipped with an active aeration device to add oxygen directly into the settling pond. To achieve this aeration by passive means, the site must have sufficient topographic relief and area to allow for a number of small settling ponds in series, each at a lower elevation than the previous. Passive oxygenating structures such as riffles are then placed in between each pond.

### 2.2.5 Successive Alkalinity Producing System (SAPS)

The goal of a SAPS is to add alkalinity to ARD and then precipitate iron hydroxides upon subsequent oxygenation using two separate steps to limit iron hydroxides from armoring the limestone. A SAPS is a variant of the anaerobic systems used mainly to treat coal mine drainage. Successive Alkalinity Producing Systems can be designed specifically for those
instances that are not appropriate for ALDs (i.e., waters with DO concentrations greater than 5 mg/L and high concentrations of oxidized Fe$^{+3}$).

Zipper and Jagie (2001) developed guidelines for the design and construction of SAPS (Appendix B). The SAPS uses ponded water above a 2-layer solid system of organic material overlying limestone (Figure 3). The hydraulic head differential achieved by the ponded water forces flow downward through the organic material to remove oxygen from the water and reduce iron. The reduced water then flows through the limestone and receives alkalinity without immediate precipitation of iron hydroxides. Once the water has obtained alkalinity from the SAPS, it flows into a settling pond where it becomes re-oxidized and iron hydroxides precipitate and settle out.

This system uses a much thinner layer of organic substrate than the SRB anaerobic bioreactor, but still suffers from progressive reduction in permeability and short-circuiting. Based on Demchak, Skousen, and Morrow (undated), the SAPS designed to date have been unable to eliminate the short-circuiting that lower their effectiveness in removing copper and zinc as sulfides. Demchak et al. recommend that several treatment cells be developed in series to combat this problem.

### 2.2.6 Solid-Reactant Anaerobic or Sulfate Reducing Bacteria (SRB) Bioreactors

The chemical processes in anaerobic bioreactors are bacterial oxidation of organic matter with concomitant reduction of DO, ferric iron (Fe$^{+3}$), and sulfate. Because sulfate-reducing bacteria (SRB) play a major role in this type of bioreactor, the anaerobic bioreactor is often called an SRB bioreactor. As sulfate reduction occurs, the produced sulfide then reacts with iron, copper, zinc, and cadmium to form metal sulfides. Metal sulfide production is the only effective passive means of reducing copper, zinc, and cadmium concentrations in ARD.

Reduction occurs in the absence of oxygen, which requires that flow be uni-directional and preferably vertical throughout the organic bioreactor material (i.e., substrate) within the subsurface. Uniform flow rates and even flow distribution through the substrate are critical in effective SRB bioreactor treatment. Accordingly, the bioreactor must be appropriately engineered to maximize vertical flow and, as with SAPS, to minimize short-circuiting.

The organic material substrates typically used in a solid-reactant SRB bioreactor contain a complex mixture of organic compounds, each serving as an energy source for a particular group of bacteria or fungus. Thus, an effective SRB bioreactor must contain microenvironments that allow an entire consortium of microorganisms to prosper (i.e., provide a good energy source for the SRBs and provide for sufficient alkalinity). For SRB bioreactors, a pH of about 5.5 or higher is preferred.

First-generation bioreactors typically used composted cow manure or mushroom compost because they provide significant alkalinity (Filipek et al. 1992; Schafer and Filipek 1995). More recent bioreactors use a combination of limestone, sawdust and alfalfa hay in place of manure because this combination 1) provides alkalinity; 2) provides significantly greater permeability; and, 3) appears to be a better energy source for the bacterial community. Cow manure is typically added only in small quantities as an inoculum to supply an initial source of active SRBs.
The design of a solid-reactant SRB anaerobic bioreactor consists of a lined basin containing the solid organic substrate. In the most effective designs, the substrate is typically sandwiched between pipes set in inert gravel near the top and bottom of the basin (Figure 4). The ARD flows into the bioreactor through one set of pipes and the treated water flows out through the other.

When inflow is through the reactor top, the system is in “downflow” mode. When inflow is through the reactor bottom, the system is in “upflow” mode. The more successful designs allow the flow to be reversed between modes.

Plugging of valves can occur due to the formation of precipitates if the chemical environment changes in the vicinity of the pipes. Precipitation of iron and aluminum hydroxides was a major problem in early SRB bioreactors because the pipes were sometimes laid in a bed of limestone. The limestone raised the pH and caused the same clogging as occurs in ALDs treating water with high concentrations of Fe$^{+3}$ and/or aluminum. Recent systems incorporate pipe valves that allow them to be flushed out periodically to remove precipitates.

On start-up of the SRB bioreactor, the soluble organic compounds are especially prevalent, usually giving the effluent a brownish color. Also, the bacterial population needs time (i.e., two or three months) to become established in the bioreactor. Ideally, only a portion of the total effluent load is allowed to enter the bioreactor during this time period. Recirculating some of the treated effluent back into the reactor makes up the remainder of the flow. Due to the reactions in SRB bioreactors, soluble organic and nitrogen compounds, as well as sulfides, are produced. These compounds must be removed from the effluent before release off-site. Typically, this is accomplished by placing a small aerobic polishing cell in line after the SRB bioreactor.

The bioreactors were originally considered passive because they were believed to be able to function on the order of 20 or more years before requiring major maintenance (i.e., replacement of the treatment media). In the past 10 years a number of pilot SRB bioreactors have been used successfully over short periods to reduce heavy metals such as copper and zinc from sulfate-laden hard-rock drainage. However, none have remained operational for more than a few years without significant overhaul or modification. Between 3 and 4 years is the maximum reported operational period of an SRB bioreactor. Most systems were terminated for various reasons after a year or less. Identified causes of termination have included:

- Systems being overtaken by volunteer plants that added oxygen to the subsurface through their roots;
- Short-circuiting due to compaction and low permeability;
- Systems being overwhelmed by storm surges with acidity and metal loading significantly greater than design loads; and
- System desaturation.

Sizing of the organic portion of constructed SRB bioreactors is based on four primary criteria:

- Sulfate loading;
- Heavy metal loading;
- Residence time required for the sequence of bacterial reactions to occur; and
- Minimum vertical transport time allowed by the permeability of the substrate.
Sizing calculations must be conducted for each criteria to determine which is the limiting factor.

**Sulfate Loading**

Under optimum field conditions, sulfate reduction occurs at the rate of about 0.3 moles sulfate per cubic meter per day (mol/m$^3$ day). Thus, the flux of sulfate into the system must be less than this rate. Sulfate loading is a very conservative factor because it assumes that the bacteria reduce all sulfate. Typically, sulfate concentrations do not drop below a few hundred milligrams per liter. The calculation is useful, however, because in order to function properly, the system requires that sulfate be present at levels in excess of metals.

**Heavy Metal Loading**

The flux of heavy metals, including iron, copper, lead, zinc, and cadmium, into the system must be less than the rate of sulfate reduction. Typically, a flux value of 0.15 mol/m$^3$ day heavy metal is used.

**Residence Time**

Metal sulfide precipitation requires at least 3 to 5 days to occur. Therefore, the bioreactor must be sized to allow a minimum of 3 days residence time within the unit. The porosity of the chosen substrate must be determined in order to evaluate residence time.

**Permeability Constraints**

An initial permeability of between $5 \times 10^{-4}$ centimeters per second (cm/sec) and $1 \times 10^{-5}$ cm/sec is common for the compost substrates used for the first generation of SRB bioreactors. When initial permeability was high it typically dropped rapidly following system start-up due to settling and compaction. Therefore, a value of no greater than $1 \times 10^{-4}$ cm/sec should be assumed for compost substrates. The composted substrate in an SRB bioreactor should be at least 2 feet thick, but should not exceed about 3 to 4 feet in thickness because the substrate tends to compact with depth and can become too low for effective treatment. Additionally, short-circuiting due to preferential flow paths can develop when the permeability becomes too low relative to the pressure head of water in the system (Filipek et al. 1992; Cevaal and Whiting 1994).

The most recent generation of SRB bioreactors, such as that at the West Fork Unit of the Doe Run Mine use sawdust instead of compost for the majority of the organic substrate. This material has a significantly higher permeability, likely on the order of $10^{-2}$ cm/sec to $10^{-3}$ cm/sec. As with compost, it is also vulnerable to compaction, especially if heavy machinery is allowed to run over it during construction or maintenance. In general, it is believed that upflow reactors (i.e., systems which force water up through the contact media) tend to last longer than downflow systems because the upward flow tends to counter the tendency toward compaction and limit the resulting flow problems (e.g., development of preferential flow paths).
Flow Rate Variability

As evidenced in case studies, large flow variations can overwhelm anaerobic bioreactors if they are sized for lower flow rates. Conversely, if an SRB bioreactor is sized for maximum probable flow and the flow decreases significantly, some of the anaerobic substrate can become oxidized and release metals that had previously been precipitated. Release of metals is a potential problem especially in upflow bioreactors because the treated water will flush metals released during oxidation of the upper substrate as it leaves the bioreactor. If the flow and/or metal loading to the bioreactor increases significantly above the design flow, especially as in a spike increase event, the bacterial community in the bioreactor may suffer severe die-off and significantly reduce system effectiveness and require additional system maintenance (e.g., the microorganism consortium allowed to re-establish, the addition of new inoculum).

To accommodate flow variations into a system design, three options are typically used:

- Regulate flow to the bioreactor using a holding pond;
- Construct a series of bioreactor cells so that overflow from one cell can be treated in another; and
- Cover the top of the bioreactor cell(s) with a layer of geochemically inert gravel to accommodate head fluctuations due to flow variations.

Waste Generation

If it is required that spent substrate be removed from the site, the substrate should be discarded before concentrations of metals (e.g., cadmium, copper) become elevated and cause the substrate to fail leaching tests (i.e., toxicity characteristic leaching procedure [TCLP]) and be classified as a RCRA hazardous waste. To mitigate this potential cost the substrate should be tested periodically while the bioreactor is operating to monitor total metal concentrations and TCLP levels.

2.2.7 Liquid-Reactant (Semi-active) Bioreactors

SRB bioreactors suffer from three deficiencies that have recently been overcome by use of a liquid-reactant bioreactor. These deficiencies, and the corresponding effects of a liquid-reactant SRB, are as follows:

1) The solid reactant SRB bioreactor decreases in permeability with time because the organic reactant is also the physical matrix. As the matrix reacts, it breaks down into smaller pieces, the pore spaces decrease, and flow is restricted. In the liquid-reactant bioreactor, the matrix consists of nonreactive cobbles, which maintain a constant permeability over time as precipitated metals are removed through a filtration system.

2) The rate of the SRB reaction in the solid-reactant bioreactor cannot be controlled because the reactant substrate must be added in excess to allow for multi-year life. The sawdust and compost substrate in the bioreactor is a mix of several organic compounds. The SRB utilize only a subset of these compounds, each at a different rate. As the preferred SRB energy source becomes depleted, alternative compounds are used, but at a slower rate. Thus, the overall rate of sulfate reduction decreases with time. Also, as the temperature decreases from summer to winter, the reaction rate concurrently decreases.
By contrast, in the liquid-reactant bioreactor, an alcohol such as methanol, ethanol, or ethylene glycol (which are preferred energy sources for SRB) is added at a controlled rate based on the stoichiometric relation between the alcohol and the sulfate being reduced. Sodium hydroxide is also added to bring the pH to a level in which the SRB can thrive. The reaction rate can thus be better controlled.

3) In cold climates there is the possibility that the solid-reactant SRB bioreactor will freeze in the winter. The alcohol of the liquid-reactant bioreactor acts as an anti-freeze, keeping the bioreactor active and protecting the piping and valves.

These benefits of the liquid-reactant bioreactor can in some instances compensate for the added costs of operation and maintenance over a traditional SRB bioreactor.

Similar to the previous discussion on solid-reactant SRB bioreactors, sizing of the liquid reactant system is dependent on sulfate loading, metal loading, and residence time. However, in place of the permeability concerns inherent with the design of the solid-reactant system, liquid-reactant systems must also design around water acidity levels as well (Tsukamoto, undated).

Based on the available information, URS believes the liquid-reactant SRB bioreactor has significant potential for success. Experiences with bacterial water-treatment systems for sewage treatment applications, as well as experience with solid-reactant bioreactors could potentially supplement the success encountered in the field (see discussion below on Leviathan Mine research application) and lead to further applications of this technology.
Case Studies

The following are five short case histories excerpted from papers included in this report as Appendix D. The first case study is the Burleigh Tunnel SRB pilot cell designed by Camp Dresser McKee for the EPA and the Colorado Department of Public Health. It was one of the two longest continuously operated bioreactor systems identified during this review. The second, third, and fourth case studies presented are cells designed by Mr. Jim Gusek and coworkers at Knight Piesold and Co., who have designed some of the more long-lived and best-documented anaerobic bioreactors. The fifth case study presented is the Leviathan Mine site where Dr. Tsukamoto used a liquid-reactant SRB bioreactor to treat ARD.

The five case studies include the two longest operated SRB bioreactors for which information is available. Based on the literature review, no SRB bioreactor designed to treat ARD with heavy metals such as copper, zinc, or lead has operated more than 3½ or 4 years without either a decrease in substrate permeability or other problems resulting in significant reductions in treatment efficiency and either cessation of operation or system re-construction.

3.1 BURLEIGH TUNNEL PILOT CELLS

The Burleigh Tunnel site is at an elevation of 9,500 feet and has a long season of below-freezing temperatures. Two pilot SRB bioreactor cells, one upflow and one downflow, were designed to each treat 7 gpm of neutral-pH water with high zinc and cadmium concentrations draining from the Burleigh Tunnel in Colorado. The cells used first-generation bioreactor technology with a 4-foot deep substrate containing a mix of composted cow manure and wood chips with alfalfa hay, with a design permeability of $1.7 \times 10^{-4}$ cm/sec. The cells were installed below grade to limit effects due to freezing. The upflow system was operated for 3 years, beginning in fall 1993. Flow to the downflow cell was discontinued in the third year due to permeability losses that blocked flow.

Both cells were effective in removing zinc during their first year of operation, when treating their design flow. During the 1995 spring runoff, the upflow cell was inundated with 20 gpm of lower-pH, higher-zinc aerobic water. This water appeared to have changed the microbial ecology of the bioreactor. The cell never recovered and, thereafter, was only 50- to 60-percent efficient in treating zinc.

3.2 BREWER PAD 5 PILOT CELL

According to Gusek (2000), the Brewer Pad 5 pilot cell treated ARD with elevated iron, aluminum, and copper concentrations from a spent heap leach pad from 1993 through early 1995, after which time it was decommissioned. The cell was 2.4 feet deep and filled with a mixture of composted turkey manure, sawdust, phosphate rock reject (limestone), and cow manure SRB inoculum. Influent metal concentrations increased significantly over the design concentrations (e.g., iron by over an order of magnitude) throughout the life of the bioreactor. Therefore, the influent flow was gradually decreased from the design value of 5 gpm to 0.75 gpm by the end of cell’s life.

During its 18 months of operation, the cell suffered three episodes of invasion by volunteer vegetation at the cell surface. The vegetation caused oxidizing conditions within the cell which made the cell ineffective for copper removal. When the vegetation was eliminated and the cell refitted with a fresh source of native hay, the cell was able to remove iron, aluminum, and copper...
with nearly 100-percent efficiency. Following the vegetation removal, the bioreactor was able to treat the increasingly concentrated but lower-flow rate ARD for the approximate three months until the system was decommissioned in early 1995 to accommodate ongoing reclamation/closure activities at the site (Gusek, 2000).

3.3 FERRIS-HAGGERTY MINE/OSCEOLA TUNNEL

The Ferris-Haggerty Mine/Osceola Tunnel site is a high-elevation abandoned copper-mine site with ARD characterized initially as having neutral pH, low sulfate, elevated copper concentrations, and water temperature close to freezing. A pilot cell treated a flow of 3 to 5 gpm for two years beginning in the summer of 1997. The cell was enclosed in a shed to aid in water sampling during high-snowfall months. The 4-foot deep cell was filled with a mixture of softwood sawdust, hay, limestone, cow manure, and gypsum. The gypsum was added as a source of sulfate for the SRB.

During operation, copper removal efficiencies were acceptable even when the cell surface was 40-percent covered in ice and the influent changed to an acidic water with a pH between 3.5 and 4. The pilot cell operated successfully (i.e., 89- to 97-percent removal of copper) for two years, after which time it was decommissioned. In 1999, a second pilot-scale cell was built with a slightly different substrate mixture and buried below grade to more closely mimic the planned full-scale system (Gusek 2000). Data are not available for the second pilot scale cell, and no information is available regarding the implementation schedule for the full-scale system.

3.4 DOE RUN MINE

The two longest-running bioreactors are at the West Fork Unit of the Doe Run Mine site. They were designed to treat 1,200 gpm of pH 8 water containing elevated lead concentrations. The system, which included a 0.75-acre settling pond, two SRB cells (each of cell covering just under 1-acre), a 1.4-acre aerobic rock filter, and a 2.0-acre aeration pond. The system took about 4 months to construct. Each of the SRB cells has a substrate composed of approximately 60- to 70-percent sawdust, 20-percent limestone, and the remainder aged cow manure and alfalfa hay cubes. The cell permeability is significantly greater than the first-generation SRB bioreactors composed mainly of compost and is estimated to be significantly less than the design permeability of $5 \times 10^{-1}$ cm/sec.

The cells have been treating the mine drainage since 1996. During that time, the substrate in each cell has been removed and replaced twice. The first substrate removal was required to correct design changes that had not been previously tested. The distribution pipes had been connected to commonly shared layers of perforated high density polyethylene (HDPE) pipe and geonet materials sandwiched between layers of geofabric in an attempt to control sulfide production in hot weather by creating intentional short circuiting. In practice, the layered geotextiles appeared to trap hydrogen sulfide gas ($H_2S$), which caused gas build-up and permeability decreases in the cells. In June 1997, the substrate from the south anaerobic cell was excavated and the cell refilled without the geotextiles. The north cell was treated similarly in September 1997.

In the summer of 1998, some portions of the cells were intentionally by-passed to maintain lower sulfide concentrations. This mode of operation caused short-circuiting and plugging of the cells.
Also, changed operations within the mine resulted in the mine effluent water containing increasing concentrations of fine sediment, which tended to block flow within the cells. Therefore, the entire top 4 feet of substrate from the south cell was replaced in early 1999 with a new mixture containing more limestone than the original mix. In 2000, the same action was taken with the north cell. Since that time, the cells have received no special treatment during the summers and have operated without reported problems (Murphy 2002).

3.5 LEVIATHAN MINE

A liquid-reactant bioreactor was designed and constructed to treat ARD from an area of the Leviathan Mine site. This summary of the system is based on documentation included as Appendix E. The bioreactor system consists of 2 cells, which together were designed to treat up to 16 gpm of water with an average pH of 4, average iron concentration of 167 mg/l, and average concentrations of copper, nickel, and zinc on the order of 1 mg/l each. Each cell is 6 feet deep and contains approximately 1 foot of manure at the bottom. The remainder of cell one is filled with wood chips, while cell two is filled with 2- to 6-inch cobbles. Both cells were inoculated with an anaerobic horse manure culture. Influent ARD enters each cell at the surface, flows laterally across the cell, and eventually is collected in three loops of perforated pipe located at the cell bottom.

As a research application by the University of Nevada-Reno, methanol was added for the first 8 months of operation and then a mixture of methanol, ethanol, and ethylene glycol was added for the remainder of the experimental period. The application rates of the additives varied throughout the period, but were typically between 1 and 3 times the stoichiometric equivalent needed to reduce the sulfate in the influent solution. Initially, the inflow was kept below 0.8 gpm because the valves for inflow and outflow plugged frequently. At 8 months, a weir was added to control inflow, standpipes were added to control outflow, and the liquid organic mixture was changed. At 18 months, base was added and the flow increased from an average of about 0.4 gpm to an average of about 6.5 gpm, typically treating the entire flow of the source seep. For approximately the next 2 years, the bioreactor system operated effectively to remove nickel, copper, and zinc.
**SECTION FOUR**

Acid Rock Drainages at the Elizabeth Mine

This section describes the geologic setting of the Elizabeth Mine site and the geochemical conditions as currently defined.

### 4.1 GEOLOGY AND MINE FEATURES

The Elizabeth Mine is located in the towns of South Strafford and Thetford, Vermont and is the largest mining complex of the Vermont Copper Belt. Elizabeth Mine has a long history of copperas surface leaching operations and, in later years, milling of copper sulfide ore deposits. The ore deposit is characterized as a “besshi-type” massive sulfide with mineralization occurring as massive bands and disseminated sulfides. Pyrrhotite is a principal mineral in the deposit, with other minerals present including chalcopyrite, sphalerite, cubanite, and others.

Primary physical features at the site include two open cuts (north and south open cuts), two tailing impoundments referred to as Tailing Pile 1 and 2 (TP-1 and TP-2), and a leach pile referred to as Tailing Pile 3 (TP-3). TP-1 and TP-2 were generated as a waste product from on-site milling and floatation processes that occurred in the 1940s and 1950s and consist of approximately 35-acres of hydraulically deposited fine sand and silt mill tailing. The impoundments have a maximum thickness of about 100 feet and are underlain by either alluvium or fill.

TP-3 is irregularly shaped, side slope leach pile varying in thickness from several feet to tens of feet and covers approximately 12-acres. This historic side slope leach pile is comprised of a heterogeneous mixture of boulders, cobbles, gravels, sands, and silts consisting of leach ore, roasted ore, and spent ore.

### 4.2 GEOCHEMISTRY OF ACID ROCK DRAINAGE

The principal features at the site resulting in ARD include tailing piles TP-1 and TP-2 and a leach pile TP-3. Water discharging from TP-1, TP-2, and TP-3 is known to have a low pH and contains elevated metals due to the natural oxidization of the sulfides contained in the ore. This oxidizing process occurs when sulfide minerals are exposed to air and water and can be a natural phenomenon or the result of mining beneficiation activities. The discharges from TP-1, TP-2, and TP-3 exceed state and federal water quality standards.

At the Elizabeth Mine, the drainage from tailing piles TP-1, TP-2, and TP-3 demonstrate the wide range of water chemistries that can be classified as ARD. Discharges range from moderate pH, anoxic waters (with potential to acidify upon exposure to oxygen) with relatively low metal content associated with TP-1; to low-pH, highly oxic waters rich in a number of heavy metals associated with TP-3. Based upon USGS sampling and testing of seeps performed in April and August 1998, the drainage composition and condition encountered at three discrete sample locations at the Elizabeth Mine is as follows.

**Tailing Pile 1**

The seeps draining TP-1 tend to be anoxic, with measured DO concentrations averaging 0.87 mg/L for the two sampling periods. The measured DO concentrations were higher in the summer than in the spring and varied between the seeps tested, with the sample from the northwest corner of TP-1 containing higher DO than the other seep locations. Similarly, the ratio
of reduced iron (Fe$^{+2}$) to total iron was relatively high in these samples, ranging from 0.44 to 1.00. Iron was the dominant heavy metal in these waters, with a total iron concentration of 453 mg/L. Dissolved aluminum concentrations averaged 0.19 mg/L; copper 0.011 mg/L; zinc 0.055 mg/L; and manganese 4.0 mg/L. The pH of these seeps averaged 5.9. However, the calculated acidity was relatively high at approximately 850 mg/L CaCO$_3$ equivalent due to significant acidity contribution from dissolved iron species.

**Tailing Pile 2**

The seep draining from TP-2 had a DO concentration of 7 mg/L, (near saturation). The ratio of reduced Fe$^{+2}$ to total iron was 0.06 and the pH was 3.2; both parameters indicating oxidation of iron within this tailing pile. Due to the lower pH, heavy metal concentrations were higher in the TP-2 seep than those draining TP-1. The dissolved iron concentration was 1,000 mg/L; the dissolved aluminum concentration was 7.0 mg/L; copper 0.06 mg/L; zinc 1.4 mg/L; and manganese 7.3 mg/L. The acidity was calculated to be approximately 2,670 mg/L CaCO$_3$ equivalent.

**Tailing Pile 3**

The seeps from TP-3 had DO concentrations ranging from 7 to 9 mg/L (near saturation), ratios of reduced Fe$^{+2}$ to total iron of 0.24, and pH of 2.2; all components being similar to TP-2 and indicating oxidation of iron to be occurring within the waste rock pile TP-3. Due to the lower pH and the type of rock in TP-3, heavy metal concentrations were the highest monitored on site. The dissolved aluminum concentration was 225 mg/L; copper 165 mg/L; zinc 35 mg/L; and manganese 5.2 mg/L. Iron was less dominant in these seeps, averaging 505 mg/L. The acidity was calculated to be approximately 2,500 mg/L CaCO$_3$ equivalent, lower than acidity measured in the TP-2 seep due to lower concentrations of aluminum and iron.
Aerobic wetlands, ALDs, OLCs, settling ponds, SAPS, and solid-/liquid-reactant SRB bioreactors have all been used to treat ARD, either independently or in combination. Each has been successful under limited circumstances with the degree of success dependent upon the chemistry of the water to be treated and site characteristics. In most cases of ARD treatment, effective treatment systems use several treatment technologies in series to address the differing components of ARD (e.g., pH, heavy metal concentrations) and minimize the problems associated with any one component (e.g., managing unsteady flow rates).

The most common applications employ 1) addition of alkalinity using an ALD, OLC, or semi-active lime or caustic addition; 2) oxidation and settling ponds to remove iron and aluminum; 3) SRB anaerobic bioreactors to remove heavy metals as sulfides; and, 4) a series of aerobic wetlands to remove manganese, organics and BOD accumulated from the anaerobic treatment technologies. Typically, iron and aluminum are the metals of highest concentration in ARD. Removing the iron before treatment in an SRB bioreactor (Step 3) significantly decreases the required size of the SRB bioreactor. Removing aluminum before treatment in an SRB bioreactor significantly decreases maintenance requirements of the SRB bioreactor. Heavy metal-laden ARD without significant iron or aluminum can be treated directly with an SRB bioreactor, omitting Step 3.

5.1 CONCEPTUAL DESIGN

The discharge from TP-1 and TP-3 will require treatment to remove the dissolved metals and raise the pH. Historically, active lime precipitation treatment has been successfully used to raise the pH and remove metals from similar ARD discharges. The exact treatment chain is specific to the site conditions and dependant on the type and concentration of dissolved metals (water quality) and flow quantity.

To optimize the treatment process and minimize cost, it is beneficial to have a constant source inflow with consistent water chemistry. Variation in flow can require large storage areas to provide constant flow to the treatment cells. Variation in chemistry may require additional treatment steps and/or cells, increasing the overall cost. Preliminary indications suggest that the water discharging from the TP-1 toe is generally constant and appears to have a relatively consistent water quality. Conversely, the discharge from TP-3 is variable in both quantity and geochemistry. Flow is dependent on runoff potential and seasonal precipitation events, while metals concentrations may be dependent on frequency of runoff events (i.e., frequency of flushing events).

TP-1 Drainage Treatment Requirements

The likely treatment system to treat ARD emanating from TP-1, based on the current state-of-the-practice, includes:

1) Anaerobic addition of alkalinity using an ALD, or semi-active lime or caustic addition;
2) Oxidation and settling ponds to remove iron and aluminum; and
3) Aerobic wetlands with an algal mat to remove manganese and small amounts of copper and zinc.
The actual treatment chain and resultant size of treatment components is dependent on further site characterization, results of geochemical modeling and on-site pilot tests of systems. The actual size of the system and components will consider design constraints, site conditions, influent quantity and properties, and effluent discharge criteria.

The likely treatment system assumes, among other things, that treatment specific for heavy metals removal (e.g., copper, zinc) will not be required to meet discharge levels for TP-1 drainage.

**TP-3 Drainage Treatment Requirements**

The likely treatment system to treat ARD emanating from TP-3, based on the current state-of-the-practice, includes:

1. Addition of alkalinity using an OLC or semi-active lime or caustic addition;
2. Oxidation and settling ponds to remove iron and aluminum;
3. Solid or liquid-reactant SRB bioreactor(s) to remove heavy metals as sulfides; and,
4. Aerobic wetlands with an algal mat to remove manganese, organics and BOD accumulated from the anaerobic treatment technologies.

Based on the Leviathan Mine applications, although still considered in experimental stages, a liquid-reactant SRB bioreactors may potentially be useful in overcoming some of the inherent problems with the traditional SRB bioreactor for the application at TP-3. Continued evolution of the technology, and incorporation of engineering principles obtained from experiences with anaerobic digesters for sewage treatment and solid-reactant SRB bioreactors should improve their functioning and efficiency significantly.

Although system designers reference 30-year substrate life spans, no SRB bioreactor designed to treat heavy-metal laden ARD has yet operated continuously for more than about three to four years without requiring a change of substrate. Based on existing case histories of SRB bioreactor applications, URS believes the design life of the systems should be considered not to exceed four to five years to appropriately design and accurately evaluate system costs.

As with the TP-1 system, the actual treatment chain and resultant size of treatment components is dependent on further site characterization, geochemical modeling and pilot test results; and the final size of the system components will be based on design constraints, site conditions, influent quantity and properties, and effluent discharge criteria.

### 5.2 FEASIBILITY LEVEL COST ESTIMATE

Based on the TP-1 and TP-3 treatment systems outlined in the previous section, feasibility level cost estimates have been prepared to forecast anticipated expenditures for capital, and operation and maintenance. The estimates are based on available site information, URS experience with constructing and operating similar systems, and on information obtained from the literature review.
Literature-Based Cost Information

Limited cost information is available from other sites where systems are in place to treat hardrock ARD. For system components such as ALDs, OLCs, SAPS, and settling ponds, cost estimates vary widely between applications and no correlation can be drawn directly for the Elizabeth Mine site from application history at other sites. The actual costs are highly dependent on specific site conditions. Variables that control costs of these components include 1) the influent flow rate; 2) the influent acidity; and, for ALDs, OLCs, and SAPS, 3) material acquisition and transport costs (i.e., limestone).

The estimated cost for constructing an aerobic wetland is largely a function of the type of microbial mat used. URS estimates that construction costs for an unlined wetland using inoculated hay bales can range from $30,000 to $40,000 per acre. For comparison, a similar system designed using a proprietary microbial mat is estimated to cost more than $500,000 per acre, based on vendor costs of $12 per square foot.

Cost estimates for constructing and operating a solid-phase SRB bioreactor with polishing cell are presented by Gusek in his Design Example No. 1 (Gusek, 2002b). The design includes 1.7-acres of 3-feet deep SRB cells and 0.25-acres of polishing cell. He estimates a cost of between $30,000 to $50,000 to complete bench and pilot studies, and $315,000 for final system design and construction.

Gusek estimates annualized operation and maintenance, laboratory analytical, and substrate replacement costs to be approximately $24,000. The estimate is based on a 30-year operational life and includes a replacement cost of approximately $110,000, or approximately $65,000 per acre. If a four-year operational life (replacement period) is used, which is consistent with current field operability findings, the annualized operation and maintenance, laboratory, and substrate replacement costs would be approximately $52,500, or $40,000 per acre per year, bringing the total operation and maintenance costs to approximately $105,000 annually per acre.

The treatment system installed at the West Fork Unit of the Doe Run Mine (a case study discussed in Section 3.4) includes a 0.75-acre settling pond, two SRB cells, a 1.4-acre aerobic rock filter, and a 2.0-acre aeration pond. Gusek estimated the system cost at approximately $700,000, including engineering and permitting (Gusek et al. 2000). According to Gusek (2002), the rock filter and aeration pond required significant cutting and filling of the subgrade and required membrane liners, increasing construction costs to about twice what he suggests they should typically be. Based on information provided by Gusek (2002), URS estimates the actual construction cost of the SRB bioreactor portion of the system to be approximately $200,000 per acre (excluding permitting), slightly more than the design example provided above (Gusek, 2002b).

Cost estimates for designing, constructing, and operating a liquid-substrate bioreactor were presented by Gusek (2002). The system included a 4-acre, 6-foot deep liquid-substrate bioreactor combined with a 9-acre aerobic polishing cell (allowing for manganese and BOD removal). The system cost was about $1,360,000 for design and construction. Assuming that the aerobic cells cost about $270,000 to construct, and design costs are 10-percent, the liquid-substrate bioreactor cost about $950,000, or about $240,000 per acre.

Conversely, Tsukamoto (2002) estimated that the cost to construct two liquid-reactant SRB bioreactor cells treating up to 16 gallons per minute at Leviathan mine (discussed in Section 3.5)
was about $120,000. The alcohol and sodium hydroxide costs were approximately $0.75 and $0.22 per 1,000 gallons treated, respectively. For this system, reported operating material (i.e., chemical) costs were typically less than $5,000 per year. According to Tsukamoto, after the initial frequent monitoring phase during acclimation, long-term monitoring frequency was reduced to approximately one to two times per month.

**Elizabeth Mine Cost Estimate**

Feasibility level cost estimates for the likely treatment systems for TP-1 and TP-3 (as outlined above) are presented in Table 1 (Capital Costs) and Table 2 (Operation and Maintenance Costs). The costs are based on URS’ experience with the design of ARD treatment systems and on the information and case studies identified during the literature review. Implementation of the TP-1 and TP-3 systems will require additional data collection and further detailed design analyses. Capital costs and operation and maintenance costs for the TP-3 system are provided for both a solid-reactant and liquid-reactant SRB.

Capital costs, as described in Table 1, are estimated to be $268,500 for the TP-1 system, $1,452,400 for TP-3 with a solid-reactant SRB, and $1,688,400 for TP-3 with a liquid-reactant SRB.

Total annual operation and maintenance costs, as described in Table 2, are estimated to be $48,500 for TP-1, $120,000 for TP-3 with a solid-reactant SRB, and $132,000 for TP-3 with a liquid-reactant SRB.

These estimates assume, among other things, that algal mats can be created at the site. Included are costs for engineering and design, pilot testing, and construction oversight. Due to the preliminary nature of the estimates we have also included a contingency of 30 percent, which includes both design and reserve contingencies.

The operation and maintenance costs were developed by estimating the weekly labor required per year. We assumed that one technician and one laborer would be required for field activities. As noted, the operation and maintenance cost estimates include on-site sludge disposal, monitoring, analysis, and analytical testing, but not oversite or data interpretation. Component replacement costs (i.e., 5-year SRB replacement) are included in the estimate for the solid-phase reactant system. Liquid reagent costs are included in the estimates for the liquid-phase reactant system. Other assumptions incorporated into the estimate are provided on Table 2.


Filipek, L.H., Chao, T.T., and Carpenter, R.H., 1981, Factors affecting the partitioning of Cu, Zn, and Pb in boulder coatings and stream sediments in the vicinity of a polymetallic sulfide deposit: Chemical Geology, v. 33, p. 45-64.


Gusek, James, 2002, personal communication, Knight Piesold and Co.


Murphy, Dennis, 2002, personal communication, Doe Run Mine.


Tables
### TABLE 1
FEASIBILITY LEVEL CAPITAL COST ESTIMATE
PASSIVE/SEMI-ACTIVE TREATMENT SYSTEM, TP-1 AND TP-3
ELIZABETH MINE, VERMONT

<table>
<thead>
<tr>
<th>Description</th>
<th>Approximate Area (Acres)</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TP-1 Treatment System</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Anoxic Limstone Drain</td>
<td>0.01</td>
<td>$35,400</td>
</tr>
<tr>
<td>Oxidizing Ponds</td>
<td>0.5</td>
<td>$98,300</td>
</tr>
<tr>
<td>Aerobic Wetlands (1)</td>
<td>0.24</td>
<td>$8,500</td>
</tr>
<tr>
<td><strong>TP-1 Subtotal</strong></td>
<td></td>
<td>$142,200</td>
</tr>
<tr>
<td>Engineering and Design (10%)</td>
<td></td>
<td>$14,200</td>
</tr>
<tr>
<td>Pilot Testing (20%)</td>
<td></td>
<td>$31,300</td>
</tr>
<tr>
<td>Construction Oversight (10%)</td>
<td></td>
<td>$18,800</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td></td>
<td>$206,500</td>
</tr>
<tr>
<td>Contingency (30% of Subtotal)</td>
<td></td>
<td>$62,000</td>
</tr>
<tr>
<td><strong>Program Cost</strong></td>
<td></td>
<td>$268,500</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Description</th>
<th>Approximate Area (Acres)</th>
<th>Cost</th>
<th>Cost (4)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TP-3 Treatment System</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Open Limstone Channel</td>
<td>0.5</td>
<td>$28,000</td>
<td>$28,000</td>
</tr>
<tr>
<td>Oxidizing Ponds</td>
<td>0.25</td>
<td>$57,500</td>
<td>$57,500</td>
</tr>
<tr>
<td>Anaerobic Bioreactor (2)</td>
<td>2.5 (3)</td>
<td>$676,400</td>
<td>$726,900 (5)</td>
</tr>
<tr>
<td>Aerobic Wetlands (1)</td>
<td>0.2</td>
<td>$7,500</td>
<td>$7,500</td>
</tr>
<tr>
<td><strong>TP-1 Subtotal</strong></td>
<td></td>
<td>$769,400</td>
<td>$819,900</td>
</tr>
<tr>
<td>Engineering and Design (10% for solid, 20% for liquid)</td>
<td></td>
<td>$76,900</td>
<td>$164,000</td>
</tr>
<tr>
<td>Pilot Testing (20%)</td>
<td></td>
<td>$169,300</td>
<td>$196,800</td>
</tr>
<tr>
<td>Construction Oversight (10%)</td>
<td></td>
<td>$101,600</td>
<td>$118,100</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td></td>
<td>$1,117,200</td>
<td>$1,298,800</td>
</tr>
<tr>
<td>Contingency (30% of Subtotal)</td>
<td></td>
<td>$335,200</td>
<td>$389,600</td>
</tr>
<tr>
<td><strong>Program Cost</strong></td>
<td></td>
<td>$1,452,400</td>
<td>$1,688,400</td>
</tr>
</tbody>
</table>

(1) Includes locally developed algal mat - increase treatment area to 0.6 acres if no algal mat is used
(2) Cost can vary from less than $165,000/acre to over $240,000 per acre depending on the design constraints, local site conditions, influent properties and effluent discharge criteria. Actual cost should be verified by pilot testing.
(3) Liquid reactant will reduce the area, increase the depth and increase the unit cost
(4) Shows only cost changes due to liquid bioreactor
(5) Assumes two reactant storage tanks and simple shed with electricity
TABLE 2
FEASIBILITY LEVEL OPERATION AND MAINTENANCE COST ESTIMATE
PASSIVE/SEMI-ACTIVE TREATMENT SYSTEM, TP-1 AND TP-3
ELIZABETH MINE, VERMONT

<table>
<thead>
<tr>
<th>Work Items</th>
<th>Work Weeks /Year</th>
</tr>
</thead>
<tbody>
<tr>
<td>TP-1 Passive Treatment System</td>
<td></td>
</tr>
<tr>
<td>Clean Horizontal Drains</td>
<td>2</td>
</tr>
<tr>
<td>Maintain ALD</td>
<td>1</td>
</tr>
<tr>
<td>Clean Settling Ponds (1)</td>
<td>6</td>
</tr>
<tr>
<td>Maintain Aerobic Wetlands</td>
<td>2</td>
</tr>
<tr>
<td><strong>Subtotal TP-1</strong></td>
<td><strong>11</strong></td>
</tr>
</tbody>
</table>

**Annual Labor Cost**

<table>
<thead>
<tr>
<th>Item</th>
<th>Rate</th>
<th>Hours</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>One Technician</td>
<td>$40 per hr</td>
<td></td>
<td>$17,600</td>
</tr>
<tr>
<td>One Laborer</td>
<td>$25 per hr</td>
<td></td>
<td>$11,000</td>
</tr>
<tr>
<td>One Truck (Wet)</td>
<td>$15 per hr</td>
<td></td>
<td>$6,600</td>
</tr>
<tr>
<td>Small tools and equipment</td>
<td>$500 per mth for 2.5 mths</td>
<td></td>
<td>$1,300</td>
</tr>
<tr>
<td>Backhoe Rental (Wet)</td>
<td>$20 per hr for 11 wks</td>
<td></td>
<td>$8,800</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td></td>
<td></td>
<td><strong>$45,300</strong></td>
</tr>
</tbody>
</table>

**Material Cost**

(Assume approximately 1% of labor) $300

**Annual Monitoring**

(Assume approximately 10% of labor) $2,900

**Total (5)** $48,500
TABLE 2
FEASIBILITY LEVEL OPERATION AND MAINTENANCE COST ESTIMATE
PASSIVE/SEMI-ACTIVE TREATMENT SYSTEM, TP-1 AND TP-3
ELIZABETH MINE, VERMONT

<table>
<thead>
<tr>
<th>Work Items</th>
<th>Work Weeks /Year</th>
<th>Solid-react(3)</th>
<th>Liquid-react</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>TP-3 Passive Treatment System</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maintain OLC</td>
<td>3</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>Clean Settling Ponds (1)</td>
<td>4</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>Maintain Anaerobic Bioreactor (2)</td>
<td>4</td>
<td>12</td>
<td></td>
</tr>
<tr>
<td>Maintain Aerobic Wetlands</td>
<td>2</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td><strong>Subtotal TP-3</strong></td>
<td>13</td>
<td>21</td>
<td></td>
</tr>
</tbody>
</table>

**Annual Labor Cost**

<table>
<thead>
<tr>
<th>Item</th>
<th>Rate</th>
<th>Solid-react</th>
<th>Liquid-react</th>
</tr>
</thead>
<tbody>
<tr>
<td>One Technician</td>
<td>$40 per hr</td>
<td>$ 20,800</td>
<td>$ 33,600</td>
</tr>
<tr>
<td>One Laborer</td>
<td>$25 per hr</td>
<td>$ 13,000</td>
<td>$ 21,000</td>
</tr>
<tr>
<td>One Truck (Wet)</td>
<td>$15 per hr</td>
<td>$ 7,800</td>
<td>$ 12,600</td>
</tr>
<tr>
<td>Small tools and equipment</td>
<td>$500 per mth for 3 mths</td>
<td>$ 1,500</td>
<td></td>
</tr>
<tr>
<td>Small tools and equipment</td>
<td>$500 per mth for 5 mths</td>
<td>$ 2,500</td>
<td></td>
</tr>
<tr>
<td>Backhoe Rental (Wet)</td>
<td>$20 per hr for 21 wks</td>
<td>$ 16,800</td>
<td></td>
</tr>
<tr>
<td>Backhoe Rental (Wet)</td>
<td>$20 per hr for 13 wks</td>
<td></td>
<td>$ 10,400</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td><strong>59,900</strong></td>
<td><strong>$ 80,100</strong></td>
<td></td>
</tr>
</tbody>
</table>

**Material Cost**

<table>
<thead>
<tr>
<th>Item</th>
<th>Cost</th>
<th>Solid-react</th>
<th>Liquid-react</th>
</tr>
</thead>
<tbody>
<tr>
<td>Solid-reactant substrate replacement</td>
<td>$ 55,000</td>
<td>(3)</td>
<td></td>
</tr>
<tr>
<td>Liquid-reactant ethanol and NaOH</td>
<td>$ 43,700</td>
<td></td>
<td>(4)</td>
</tr>
</tbody>
</table>

**Annual Monitoring**

<table>
<thead>
<tr>
<th>Item</th>
<th>Cost</th>
<th>Solid-react</th>
<th>Liquid-react</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Assume approximately 15% of labor)</td>
<td>$ 5,100</td>
<td></td>
<td>$ 8,200</td>
</tr>
</tbody>
</table>

**Total (5)**

<table>
<thead>
<tr>
<th>Cost</th>
<th>Solid-react</th>
<th>Liquid-react</th>
</tr>
</thead>
<tbody>
<tr>
<td>$ 120,000</td>
<td>$</td>
<td>$ 132,000</td>
</tr>
</tbody>
</table>

(1) Cost includes on-site sludge disposal
(2) Assumes flushing, monitoring, and addition/replacement of substrate.
(3) Assumes replacement of solid substrate every 5 years. @ half original reactor cost of $676K.
(4) Assumes 2 times stoichiometric ratio for ethanol and Tsukamoto’s base reagent costs for ethanol and NaOH. (Est. 1000 gal ethanol needed per week for TP-3.)
(5) Cost includes monitoring and analytical testing, but not oversight or data interpretation.
Figures
Determine flow rate
Analyze water chemistry
Calculate loadings

Net acid water

Determine O2 content
Ferrous/ferric iron ratio

Net alkaline water

DO < 2 mg/L
Fe⁺³ < 10%
Al⁺³ < 25 mg/L

Anoxic limestone drain (ALD)
Net alkaline water
Net acid water

Aerobic or anaerobic wetland, or SAPS
Strip DO, precipitate Fe⁺³

LOW flow (<200 L/min)

pH > 4.5
Aerate

SETTLED POND

DO > 5 mg/L
Fe⁺³ > 25%

HIGH flow (>200 L/min)

pH < 4.5

SETTLING POND

OPEN LIMESTONE CHANNEL (OLC)

Does water meet effluent limits?

YES

Discharge

NO

Chemical treatment or recirculate through ALD, SAPS, OLC, wetlands, etc.

Aerobic wetland

Anaerobic wetland, or SAPS

NOTE:
Adapted from Hedan et al. 1994.

LEGEND
L/min: liters per minute
mg/L: milligrams per liter
DO: dissolved oxygen
SAPS: successive alkalinity-producing systems

DRAFT
NOTE:

TYPICAL CROSS-SECTION VIEW OF A SUCCESSIVE ALKALINITY PRODUCING SYSTEM (SAPS) TREATMENT COMPONENT

PONDED WATER

DIRECTION OF FLOW

ORGANIC COMPOST

LIMESTONE

IMPERMEABLE BARRIER AND OR EFFLUENT PIPES

DRAFT
Appendix A
Bibliography
Passive Treatment
Recent BIBLIOGRAPHY

(Some articles listed under multiple headings.)

Overview/Review


ALDs/OLCs


Aerobic Wetlands


SAPS


Anaerobic Bioreactors

Solid Substrate – Field Case Studies (Typically integrated with other passive methods such as ALDs and/or aerobic wetlands)


Gusek, J., Wildeman, T., Mann, C., Murphy, D. Operational results of a 1,200-gpm passive bioreactor for metal mine drainage, West Fork, Missouri. ??


Tyrrell, WR; Mulligan, DR; Sly, LI; Bell, LC. Trialing wetlands to treat coal mining wastewaters in a low rainfall, high evaporation environment. Int. Conf. on Wetland Systems for Water Pollution, Vienna (Austria), 15-19 Sep 1996. Haberl, R; Perfler, R; Laber, J; Cooper, P (eds). WETLAND SYSTEMS FOR WATER POLLUTION CONTROL 1996, 1997, pp. 293-299, Water Science and Technology, vol. 35, no. 5.

In-Situ – Case Studies


Laboratory Studies
Chen, B-Y. Metal toxicity and biosorption studies of pure culture isolates of sulfate reducing bacteria in acid mine-drainage treatment systems. Annual West Coast Conference on Contaminated Soils and Water, San Diego, CA (USA), 20-23 Mar 2000.


Steed, VS; Suidan, MT; Gupta, M; Miyahara, Takashi; Acheson, CM; Sayles, GD. Development of a sulfate-reducing biological process to remove heavy metals from acid mine drainage. Water Environment Research Vol. 72, no. 5, pp. 530-535. Sep-Oct 2000.

Tyrrell, WR; Mulligan, DR; Sly, LI; Bell, LC. Trialing wetlands to treat coal mining wastewaters in a low rainfall, high evaporation environment. Int. Conf. on Wetland Systems for Water Pollution, Vienna (Austria), 15-19 Sep 1996. Haberl, R; Perfler, R; Laber, J; Cooper, P (eds). WETLAND SYSTEMS FOR WATER POLLUTION CONTROL 1996, 1997, pp. 293-299, Water Science and Technology, vol. 35, no. 5.


Liquid Reagent

Steed, VS; Suidan, MT; Gupta, M; Miyahara, Takashi; Acheson, CM; Sayles, GD. Development of a sulfate-reducing biological process to remove heavy metals from acid mine drainage. Water Environment Research Vol. 72, no. 5, pp. 530-535. Sep-Oct 2000.


Unknown
Schrecengost, MA; George, B; Morrow, TO; Turner, AM. Longevity and long-term effectiveness of passive treatment systems in removing metals and acidity from acid mine discharges. 2002 Annual Meeting of the North American Benthological Society, Pittsburgh, PA (USA), 28 May-Jun 1 2002.

Appendix B
Overview Papers
Appendix B
Overview Papers


Skousen
1998
Overview of Passive Systems for Treating Acid Mine Drainage
Overview of Passive Systems for Treating Acid Mine Drainage

Jeff Skousen
West Virginia University

Introduction

Active chemical treatment of AMD to remove metals and acidity is often an expensive, long term liability. In recent years, a variety of passive treatment systems have been developed that do not require continuous chemical inputs and that take advantage of naturally occurring chemical and biological processes to cleanse contaminated mine waters. The primary passive technologies (Figure 1) include constructed wetlands, anoxic limestone drains (ALD), successive alkalinity producing systems (SAPS), limestone ponds, and open limestone channels (OLC).

Natural wetlands are characterized by water-saturated soils or sediments with supporting vegetation adapted to reducing conditions in their rhizosphere. Constructed wetlands are man-made ecosystems that mimic their natural counterparts. Often they consist of shallow excavations filled with a flooded gravel, soil, and organic matter to support wetland plants, such as Typha, Juncus, and Scirpus sp. Treatment depends on dynamic biogeochemical interactions as contaminated water travels through the constructed wetland. ALDs are abiotic systems consisting of buried limestone cells that passively
generate bicarbonate alkalinity as anoxic water flows through. SAPS combine treatment concepts from both wetlands and ALDs. Oxygenated water is pre-treated by organic matter removing \( \text{O}_2 \) and \( \text{Fe}^{+3} \), and then the anoxic water flows through an ALD at the base of the system. Limestone ponds are ponds built over the upwelling of a seep and the seep is covered with limestone for treatment. OLCs are surface channels or ditches filled with limestone. Armoring of the limestone with Fe hydroxides decreases limestone dissolution by 20 to 50%, so longer channels and more limestone is required for water treatment.

At their present stage of development, passive systems can be reliably implemented as a single permanent solution for many types of AMD and at a much lower cost than active treatment. Relative to chemical treatment, passive systems require longer retention times and greater space, provide less certain treatment efficiency, and are subject to failure in the long term. However, many passive systems have realized successful short-term implementation in the field and have substantially reduced water treatment costs at many mine sites (Faulkner and Skousen 1994). Current research seeks to understand the dynamically complex chemical and biological mechanisms that occur within passive systems and which are responsible for AMD treatment.

Selection of an appropriate passive system is based on water chemistry, flow rate and local topography and site characteristics (Hyman and Watzlaf 1995), and refinements in design are ongoing. Figure 2 (adapted from Hedin et al. 1994) summarizes current thinking on the appropriate type of passive system for various conditions. In general, aerobic wetlands can treat net alkaline water; ALDs can treat water of low Al, Fe\(^{3+}\), and DO; and SAPS, anaerobic wetlands and OLCs can treat net acidic water with higher Al, Fe\(^{3+}\), and DO. As scientists and practitioners improve treatment predictability and longevity of passive systems, they will be able to treat the more difficult waters of high acidity and high Al content.
Natural Wetlands

Huntsman et al. (1978) and Wieder and Lang (1982) first noted amelioration of AMD following passage through naturally occurring Sphagnum bogs in Ohio and West Virginia. Studies by Brooks et al. (1985), Samuel et al. (1988), and Sencindiver and Bhumbla (1988) documented similar phenomena in Typha wetlands. Although evidence suggests that some wetland plants show long term adaptation to low pH and high metal concentrations, AMD eventually degrades the quality of natural wetlands, which is contrary to federal laws designed for wetland protection and enhancement. Such regulations do not govern use of artificially constructed wetlands for water treatment, leading to the suggestion that these engineered systems might provide low cost, low maintenance treatment of AMD (Kleinmann 1991). Over a thousand wetlands have since been constructed to receive AMD from both active mines and abandoned mine lands.

Constructed Wetlands

Mechanisms of metal retention within wetlands listed in their order of importance include: 1) formation and precipitation of metal hydroxides, 2) formation of metal sulfides, 3) organic complexation reactions, 4) exchange with other cations on negatively-charged sites, and 5) direct uptake by living plants. Other mechanisms include neutralization by carbonates, attachment to substrate materials, adsorption and exchange of metals onto algal mats, and microbial dissimilatory reduction of Fe hydroxides and sulfate.

The way in which a wetland is constructed ultimately affects how water treatment occurs. Two construction styles currently predominate: 1) "aerobic" wetlands consisting of Typha and other wetland vegetation planted in shallow (<30 cm), relatively impermeable sediments comprised of soil, clay or mine spoil, and 2) "anaerobic" wetlands consisting of Typha and other wetland vegetation planted into deep (>30 cm), permeable sediments comprised of soil, peat moss, spent mushroom compost, sawdust, straw/manure, hay bales, or a variety of other organic mixtures, which are often underlain or admixed with limestone. In aerobic wetlands, treatment is dominated by processes in the shallow surface layer. In anaerobic wetlands, treatment involves major interactions within the substrate.
Aerobic wetlands are generally used to collect water and provide residence time and aeration so metals in the water can precipitate (Pictures 1 and 2). The water in this case usually has net alkalinity. Iron and Mn precipitate as they oxidize, and the precipitates are retained in the wetland or downstream. Wetland species are planted in these systems for aesthetics and to add some organic matter. Wetland plants encourage more uniform flow and thus more effective wetland area.

Because of their extensive water surface and slow flow, aerobic wetlands promote metal oxidation and hydrolysis, thereby causing precipitation and physical retention of Fe, Al, and Mn hydroxides. The extent of metal removal depends on dissolved metal concentrations, dissolved oxygen content, pH and net alkalinity of the mine water, the presence of active microbial biomass, and detention time of the water in the wetland. The pH and net acidity/alkalinity of the water are particularly important because pH influences both the solubility of metal hydroxide precipitates and the kinetics of metal oxidation and hydrolysis. Metal hydrolysis produces H\(^+\), but alkalinity in the water buffers the pH and allows metal precipitation to continue. Inorganic oxidation reaction rates decrease a hundred-fold with each unit drop in pH, but microbial oxidation may increase these. Following Fe oxidation, abiotic hydrolysis reactions precipitate Fe hydroxides. Therefore, aerobic wetlands are best used with water that contains net alkalinity to neutralize metal acidity. Abiotic Mn oxidation occurs at pH >8 while microorganisms are thought to catalyze this reaction at pH >6 (Wildeman et al. 1991). Manganese oxidation occurs more slowly than Fe and is sensitive to the presence of Fe\(^{2+}\), which will prevent or reverse Mn oxidation. Consequently in aerobic net alkaline water, Fe and Mn precipitate sequentially, not simultaneously, with the practical result that Mn precipitation occurs (if at all) mainly in the later stages of wetland flow systems, after all Fe is precipitated.
Brodie and co-workers at the Tennessee Valley Authority (TVA) have reported extensively on their use of aerobic wetlands to treat AMD (Brodie 1993). A typical staged design might include an anoxic limestone drain (ALD, see next section) to passively add alkalinity to the source AMD, a settling basin to hold precipitated Fe flocs, followed by two or three aerobic wetland cells to sequentially remove additional Fe and Mn. Nine TVA wetlands receive moderate quality AMD (pH range of 4.1 to 6.3; total Fe <70 mg/L; total Mn <17 mg/L; total Al <30 mg/L; net alkalinity 35 to 300 mg/L as CaCO$_3$), which require no further post-system treatment of water exiting the wetlands. Four TVA wetlands treat water with high Fe (>170 mg/L) and no net alkalinity. Two of these systems require NaOH treatment to comply with NPDES effluent limits, while two others use ALDs for further treatment of the effluent. A final TVA wetland system receives low Fe (<0.7 mg/L) and Mn (5.3 mg/L) and is ineffective in Mn removal. Based on their experience with these systems since 1985, Brodie (1993) suggests that staged aerobic wetland systems can accommodate Fe loads of up to 21 grams/m$^2$/day even in the absence of excess alkalinity. Manganese loads up to 2 grams/m$^2$/day can be accommodated, if alkalinity is present. Hedin et al. (1994) recommend for net alkaline water that wetlands be sized using 10 grams/m$^2$/day for Fe and 0.5 grams/m$^2$/day for Mn.

Analysis of 73 sites in Pennsylvania suggested that constructed wetlands are the best available technology for many postmining ground water seeps, particularly those of moderate pH (Hellier et al. 1994). However, those sites with net acidic discharges have a much lower successful treatment efficiency. For example, the Rougeux #1 site has a flow of 5.2 gpm and influent chemistry of 2.9 pH, 445 mg/L acidity as CaCO$_3$, Fe 45 mg/L, Mn 70 mg/L, and Al 24 mg/L. After flowing through a two-celled aerobic wetland, pH increased to 3.2, acidity was decreased by 43%, Fe by 50%, Mn by 17%, and Al by 83%. The wetland cost about $15/m$^2$ to build in 1992 and was severely undersized. Although there is improvement in the water, the wetland effluent did not conform to effluent limits. Two other wetlands constructed on the site show similar results.

**Anaerobic wetlands** encourage water passage through organic rich substrates, which contribute significantly to treatment (Pictures 3 and 4). The wetland substrate may contain a layer of limestone in the bottom of the wetland or may mix the limestone among the organic matter. Wetland plants are transplanted into the organic substrate. These systems are used when the water has net acidity, so alkalinity must be generated in the wetland and introduced to the net acid water before dissolved metals will precipitate. The alkalinity can be generated in an anaerobic wetland system in two ways (Hedin and Nairn 1990). Certain bacteria, Desulfovibrio and Desulfotomaculum, can utilize the organic substrate (CH$_2$O, a generic symbol for organic carbon) as a carbon source and sulfate as an electron acceptor for growth. In the bacterial conversion of sulfate to hydrogen sulfide, bicarbonate alkalinity is produced:

```markdown
[x] Image3.jpg (25370 bytes)
```
SO₄²⁻ + 2 CH₂O = H₂S + 2 HCO₃⁻

Alkalinity can also be generated as the limestone under the organic material reacts with acidity in the wetland:

CaCO₃ + H⁺ = Ca²⁺ + HCO₃⁻

The limestone continues to react when kept in an anaerobic environment because ferrous iron is relatively soluble at pH 7 in anoxic water and ferrous hydroxide does not form and coat the limestone. If ferrous iron is oxidized, forming ferric iron, then the ferric iron can hydrolyze and form ferric hydroxide, which then coats limestone when pH is above 3.0. Bacterial sulfate reduction and limestone dissolution produce water with higher pH and add bicarbonate alkalinity for metal removal.

Anaerobic wetlands promote metal oxidation and hydrolysis in aerobic surface layers, but also rely on subsurface chemical and microbial reduction reactions to precipitate metals and neutralize acid. The water infiltrates through a thick permeable organic subsurface sediment and becomes anaerobic due to high biological oxygen demand. Several treatment mechanisms are enhanced in anaerobic wetlands compared to aerobic wetlands, including formation and precipitation of metal sulfides, metal exchange and complexation reactions, microbially generated alkalinity due to reduction reactions, and continuous formation of carbonate alkalinity due to limestone dissolution under anoxic conditions. Since anaerobic wetlands produce alkalinity, their use can be extended to poor quality, net acidic, low pH, high Fe, and high dissolved oxygen (>2 mg/L) AMD. Microbial mechanisms of alkalinity production are likely to be of critical importance to long term AMD treatment. However, Wieder (1992) documents that the mechanism and efficiency of AMD treatment varies seasonally and with wetland age. Like their aerobic counterparts, anaerobic wetlands are most successful when used to treat small AMD flows of moderate water quality. At present, the sizing value for Fe removal in these wetlands is 10 grams/m²/day (Hedin and Nairn 1992).

Sorption onto organic materials (such as peats and soils) decreased Fe from 32 mg/L to 5 mg/L (84%), Mn from 15 to 14 mg/L (7%), and total suspended solids from 32 to 12 mg/L (63%), but eventually all sorption sites on substrate materials are exhausted by continual introduction of metals in acid water (Brodie et al. 1988). Kleinmann et al. (1991) suggested adsorption of metals by organic substrates may compensate for limited initial biological activity during the first few months of
operation in a new wetland system. A field study, which examined five wetland substrate types over a 25-month period, also demonstrated that organic substrates were saturated after only one to seven months of AMD input at 9 to 17 mg Fe per gram substrate (Wieder 1993). Although some natural inputs of organic matter occur annually at plant senescence, the adsorption capacity of a wetland is limited by saturation of all exchange sites. Substantial artificial inputs of organic matter have been used as a successful strategy to temporarily renew this adsorption capacity, following an observed decline in wetland performance (Eger and Melchert 1992, Haffner 1992, Stark et al. 1995).

Insoluble precipitates such as hydroxides, carbonates, and sulfides represent a major sink for metal retention in wetlands. About 50 to 70% of the total Fe removed from AMD by wetlands is found as ferric hydroxides (Henrot and Wieder 1990, Calabrese et al. 1991, Wieder 1992). Ferric hydroxide formation depends both on the availability of dissolved oxygen and on the initial oxidation state of Fe in the AMD. Wieder (1993) reported significant retention of ferric hydroxides in surface sediments of anaerobic wetlands.

Up to 30% of the Fe retained in wetlands may be found as ferrous iron and may be combined with sulfides (Calabrese et al. 1991, McIntyre and Edenborn 1990, Wieder 1992). Iron mono and disulfides form as a result of H₂S formation by microbial sulfate reduction in the presence of an oxidizable carbon source. In addition to its metal removal potential, sulfate reduction consumes acid and raises water pH (Hedin and Nairn 1992, Rabenhorst et al. 1992).

Long term retention of Fe sulfides and Fe hydroxides in a wetland is not well understood. Under continued anoxic conditions and in the absence of soluble Fe³⁺, pyrite should remain stable. Calabrese et al. (1994) changed the influent of their anaerobic wetland from AMD to freshwater with no concomitant export of Fe²⁺. The effluent pH was >6 due to continued limestone dissolution.

Some workers have indicated that wetland systems can be seeded with specially designed and selected microorganisms (Davison 1993, Phillips et al. 1994) to introduce or re-establish microbial activity. However, experiments utilizing appropriate controls have not established the efficacy of this approach (Calabrese et al. 1994). Experience with bioremediation of other wastes suggests that selection and enrichment of naturally occurring microbial populations is a superior, more cost-effective approach (Alexander 1993).

In constructed wetlands, higher plants serve several purposes including: substrate consolidation, metal accumulation, adsorption of metal precipitates, stimulation of microbial processes, wildlife habitat, and aesthetics. Wetland plant species vary in their ability to accumulate metals (Fernandes and Henriques 1990). Some reports document elevated tissue concentrations (Spratt and Wieder 1988), while others show little metal accumulation (Folsom et al. 1981). On an annual basis, uptake by Typha accounted for less than 1% of the Fe removed by volunteer wetlands treating AMD (Sencindiver and Bhumbla 1988).

Several studies report on the effects of different plant species in wetlands. Early in the development of treating AMD with constructed wetlands, Sphagnum was the predominant wetland species. Sphagnum has a well documented capacity to accumulate Fe (Gerber et al. 1985, Wenerick et al. 1989). However, Spratt and Wieder (1988) found that saturation of Sphagnum moss with Fe could occur within one growing season. Some have indicated that metal retention over the long term is limited in some wetlands because organic matter inputs by wetland plants are limited (Kleinmann 1990). Many of the original constructed wetlands were vegetated with Sphagnum but few remained
effective. Cattails (*Typha*) have been found to have a greater environmental tolerance than *Sphagnum* moss (Samuel et al. 1988). One of the reasons is that cattails do not accumulate metals into their tissues through uptake. Algae and a few other wetland species have also received attention due to the observation that enhanced metal removal was associated with algal blooms (Hedin 1989, Kepler 1988, Pesavento and Stark 1986, Phillips et al. 1994). In Colorado, algal mixtures were found to aerobically remove Mn from mine drainage (Duggan et al. 1992), presumably due to elevated pH resulting from algal growth and the extra oxygen generated photosynthetically by the algae. Probably the most important role that wetland plants serve in AMD treatment systems may be their ability to stimulate microbial processes. Kleinmann et al. (1991) explain that plants provide sites for microbial attachment, release oxygen from their roots, and supply organic matter for heterotrophs.

Long term successful treatment by a staged anaerobic wetland has also been reported for slightly net acidic water (Fe 89 mg/L; net acidity 40 mg/L as CaCO$_3$) at the Simco constructed wetland near Coshocton, OH (Stark et al. 1994). The wetland, built in 1985, has improved in treatment efficiency over time, not requiring any chemical treatment since 1990. The density of cattail shoots has increased to a current density of 17 shoots/m$^2$. Success at the Simco wetland is attributed to the presence of moderate mine water quality (near neutral pH and Fe <100 mg/L), sound wetland design, periodic site maintenance, and high vegetative cover.

Five anaerobic wetland systems in WV receiving 4 to 98 L/min of net acid water (110 to 2400 mg/L as CaCO$_3$ and Fe from 10 to 376 mg/L) reduced acidity by 3 to 76% and Fe concentrations by 62 to 80% (Faulkner and Skousen 1994). These wetlands were generally much smaller in area than that recommended by early formulas published by the U.S. Bureau of Mines (Hedin 1989) based on iron loads. For example, one of these wetlands, Keister, reduced the acidity of a 17-L/min flow from 252 to 59 mg/L as CaCO$_3$ (76% reduction) and increased pH from 3.1 to 5.4. Iron was reduced from 23 to 9 mg/L (62%), Mn from 23 to 20 mg/L (11%), and Al from 27 to 13 mg/L (52%). The Pierce wetland used an organic substrate over limestone and treated a 98-L/min flow. Influent pH was 3.3, acidity was 118 mg/L as CaCO$_3$, Fe of 10 mg/L, Mn of 8 mg/l, and Al of 9 mg/L. Outflow pH was 4.4, acidity was reduced to 57 mg/L as CaCO$_3$ (52%), Fe decreased to 2 mg/L (80%), Mn was reduced by 11%, and Al by 25%.

A wetland system consisting of six wetland cells (total area of 2500 m$^2$) and a sedimentation basin each received a small flow (5 L/min) of AMD with pH of 3.0, acidity of 217 mg/L as CaCO$_3$, Fe of 27 mg/L, Al of 12 mg/L, and Mn of 2 mg/L (Hellier 1996). At this site in PA, the effluent after passing through the wetland was raised to pH 5.1, and the water contained a net acidity of 16 mg/L as CaCO$_3$, with about 46% iron removal, and 56% Al removal.

A 1022 m$^2$ surface flow wetland was constructed in KY to treat 37 L/min of AMD with a pH of 3.3, acidity of 2280 mg/L as CaCO$_3$, Fe of 962 mg/L, Mn of 11 mg/L, and Al of 14 mg/L (Karathanasis and Barton 1997). After construction in 1989, metal concentrations in the effluent were reduced during the first six months of treatment, however, the system failed thereafter due to insufficient wetland area and metal overloading. In 1995, a two-phase renovation project began incorporating the use of an ALD, and a series of anaerobic drains that promote vertical flow through limestone beds overlain by organic compost (much like a SAPS). Results to date indicate a pH of 6.4, slightly net alkaline water (15 mg/L as CaCO$_3$), Fe reduction of 96%, Mn removal of 50%, and Al by 100%.
A large anaerobic wetland located at Douglas, WV treated a 1000-L/min flow effectively for one year (Cliff et al. 1996). The influent pH was 3.0, with acidity of about 500 mg/L as CaCO₃, Fe of 30 mg/L, and Al of 40 mg/L. An average net alkalinity of 127 mg/L as CaCO₃ was realized in the effluent water. Four years after installation, the original acidity of 500 mg/L as CaCO₃ is being reduced to between 250 to 300 mg/L as CaCO₃. It has remained at this level of treatment for the past two years.

**Anoxic Limestone Drains**

Anoxic limestone drains (ALD) are buried cells or trenches of limestone into which anoxic water is introduced (Picture 5). The limestone dissolves in the acid water, raises pH, and adds alkalinity. Under anoxic conditions, the limestone does not coat or armor with Fe hydroxides because Fe²⁺ does not precipitate as Fe(OH)₂ at pH <6.0.

ALDs were first described by the Tennessee Division of Water Pollution Control (TDWPC) (Turner and McCoy 1990). TVA subsequently observed that AMD seeping through a coal refuse dam was being treated passively by limestone contained in an old haul road buried under the dam. Once the water containing excess alkalinity reached aerobic conditions at the ground surface, the metals oxidized and precipitated while the water remained near pH 6 (Brodie et al. 1990). TVA and TDWPC began building ALDs in 1989. Originally, ALDs were used for pre-treatment of water flowing into constructed wetlands. Brodie (1993) reported that ALDs improved the capability of wetlands to meet effluent limitations without chemical treatment. Since 1990, ALDs have also been constructed as stand-alone systems, particularly where AMD discharges from deep mine portals.

Longevity of treatment is a concern for ALDs, especially in terms of water flow through the limestone. If appreciable dissolved Fe³⁺ and Al³⁺ are present, clogging of limestone pores with precipitated Al and Fe hydroxides has been observed (Faulkner and Skousen 1994, Watzlaf et al. 1994). For waters with high sulfate (>1,500 mg/L), gypsum (CaSO₄) may also precipitate (Nairn et al. 1991). For an accepted design, no Fe³⁺, dissolved oxygen (DO), or Al³⁺ should be present in the AMD. Selection of the appropriate water and environmental conditions is critical for long term alkalinity generation in an ALD. The maximum alkalinity that ALDs may generate is about 300 mg/L as CaCO₃, although the specific level varies with water chemistry and contact time (Watzlaf and Hedin 1993).
Faulkner and Skousen (1994) reported both successes and failures among 11 ALDs treating mine water in WV. In all cases, water pH was raised after ALD treatment, but three of the sites had pH values <5.0, indicating that the ALDs were not fully functioning or that the acid concentrations and flow velocities were too high for effective treatment. Water acidity in these drains, varying from 170 to 2200 mg/L as CaCO$_3$, decreased 50 to 80%, but Fe and Al concentrations in the outflow also decreased. Ferric iron and Al$^{3+}$ precipitated as hydroxides in the drains. With Fe and Al decreases in outflow water, some coating or clogging of limestone is occurring inside the ALD (Picture 6) and the water breaks out at the front of the ALD (Picture 7).

At the Brandy Camp site in PA, an ALD was employed to treat AMD with a pH of 4.3, acidity of 162 mg/L as CaCO$_3$, Fe of 60 mg/L, Mn of 10 mg/L, and Al of 5 mg/L (Hellier 1996). After passage through the ALD, the effluent had a pH of 6.0, net alkalinity of 10 mg/L as CaCO$_3$, Fe of 50 mg/L, Mn of 10 mg/L, and Al of <1 mg/L. Most of the Fe and Mn passed through this system and precipitated in subsequent wetlands, while Al was precipitated inside the drain.

Since many sources of AMD have mixed amounts of Fe$^{3+}$ and Fe$^{2+}$ and some DO, utilization of an ALD under these conditions is not recommended. Current research involves pre-treating AMD by passing the water through a submerged organic substrate to strip oxygen from the water and to convert Fe$^{3+}$ to Fe$^{2+}$. The pre-treated water is then introduced into an underlying bed of limestone.
(see section 5.4; Kepler and McCleary 1994, Skousen, 1995). Like wetlands, ALDs may be a solution for treating specific types of AMD or for a finite period after which the system must be replenished or replaced.

Limestone has also been placed in 60-cm corrugated pipe and installed underground, and water is introduced into the pipe. Septic tanks have also been filled with limestone and AMD introduced into the tank. These applications have been used on steep slopes in lieu of buried cells or trenches, and on sites that have poor access and small water quality problems (Faulkner and Skousen 1995).

**Successive Alkalinity Producing Systems**

Successive alkalinity producing systems (SAPS) combine the use of an ALD and an organic substrate into one system (Kepler and McCleary 1994). Oxygen concentrations in AMD are often a design limitation for ALDs. In situations where DO concentrations are >1 mg/L, the oxygen must be removed from the water before introduction into an anoxic limestone bed. In a SAPS, acid water is ponded from 1 to 3 m over 0.2 to 0.3 m of an organic compost, which is underlain by 0.5 to 1 m of limestone (Picture 8). Below the limestone is a series of drainage pipes that convey the water into an aerobic pond where metals are precipitated. The hydraulic head drives ponded water through the anaerobic organic compost (Picture 9), where oxygen is consumed and ferric iron is reduced to ferrous iron. Sulfate reduction and Fe sulfide precipitation can also occur in the compost. Water with high metal loads can be passed through additional SAPS to reduce high acidity. Iron and Al clogging of limestone and pipes can be removed by flushing the system (Kepler and McCleary 1997). Data are still being gathered on the ability of SAPS to treat high Al water. Kepler and McCleary (1997) describe success with periodic flushing of Al precipitates from drainage pipes. One SAPS cited by them treat AMD containing 41 mg/L Al. However, Brodie (personal communication, 1997) described a SAPS receiving >40 mg/L Al at the Augusta Lake site in Indiana being plugged with Al precipitates after 20 months despite flushing. Successful SAPS have used mushroom compost, while other types of organic material have problems with plugging. Many possible variations in composition and thickness of organic matter, including the addition of limestone, desirability of promoting sulfate reduction, flow rates through organic matter, time schedule for replacement or addition of new organic matter, and precipitation of siderite in the limestone remain to be investigated.
Kepler and McCleary (1994) reported on initial successes for three SAPS in PA. The Howe Bridge SAPS reduced acidity from 320 mg/L to 93 mg/L as CaCO₃, and removed 2 mg/L ferric iron. The REM SAPS decreased acidity from 173 to 88 mg/L as CaCO₃, and exported more ferrous iron than entered. The Schnepp Road SAPS decreased acidity from 84 to 5 mg/L as CaCO₃, but removed all 19 mg/L ferric iron, with only 1 mg/L ferrous iron exiting the wetland.

Kepler and McCleary (1997) reported the use of SAPs in OH, PA, and WV. In all cases, Al in AMD precipitated in their systems. Their drainage design incorporates a flushing system called the 'Aluminator' (Picture 10). This allows for the precipitated Al to be flushed from the pipes thereby maintaining hydraulic conductivity through the limestone and pipes. One SAPS, Buckeye, received 3 L/min of very acid water (pH of 4.0, acidity of 1989 mg/L as CaCO₃), Fe of 1005 mg/L, and Al of 41 mg/L. Over a two-year period, the effluent had a pH of 5.9, net acidity concentration of about 1000 mg/L, Fe of 866 mg/L, and <1 mg/L Al. A second site, Greendale, treated a 25-L/min flow, and increased the pH from 2.8 to 6.5, changed the water from a net acid water (925 mg/L as CaCO₃) to a net alkaline water (150 mg/L as CaCO₃), Fe from 40 to 35 mg/L, and Al from 140 to <1 mg/L.

At the Brandy Camp site in PA, a SAPS was employed to treat AMD with a pH of 4.3, acidity of 162 mg/L as CaCO₃, Fe of 60 mg/L, Mn of 10 mg/L, and Al of 5 mg/L (Hellier 1996). After passage through the SAPS, the effluent had a pH of 7.1, net alkalinity of 115 mg/L as CaCO₃, Fe of 3 mg/L, Mn of 10 mg/L, and Al of <1 mg/L. The system effectively increased alkalinity, but retained most of
the Fe and Al inside the system. Longevity of treatment is a major concern for ALDs, especially in terms of water flow through the limestone. Eventual clogging of the limestone pore spaces with precipitated Al and Fe hydroxides, and gypsum (CaSO$_4$) is predicted (Nairn et al. 1991). For optimum performance, no Fe$^{+3}$, dissolved oxygen (DO), or Al should be present in the AMD. Selection of the appropriate water and environmental conditions is critical for long-term alkalinity generation in an ALD.

**Limestone Ponds**

Limestone ponds are a new passive treatment idea in which a pond is constructed on the upwelling of an AMD seep or underground water discharge point (Picture 11). Limestone is placed in the bottom of the pond and the water flows upward through the limestone (Faulkner and Skousen 1995). Based on the topography of the area and the geometry of the discharge zone, the water can be from 1 to 3 m deep, containing 0.3 to 1 m of limestone immediately overlying the seep. The pond is sized and designed to retain the water for 1 or 2 days for limestone dissolution, and to keep the seep and limestone under water. Like ALDs, this system is recommended for low DO water containing no Fe$^{3+}$ and Al$^{3+}$. However, the advantage of this system is that the operator can observe if limestone coating is occurring because the system is not buried. If coating occurs, the limestone in the pond can be periodically disturbed with a backhoe to either uncover the limestone from precipitates or to knock or scrape off the precipitates. If the limestone is exhausted by dissolution and acid neutralization, then more limestone can be added to the pond over the seep. Three limestone ponds have been installed but no information is available on their treatment.

**Open Limestone Channels**

Open limestone channels (OLCs) introduce alkalinity to acid water in open channels or ditches lined with limestone (Ziemkiewicz et al. 1994). Acid water is introduced to the channel and the AMD is treated by limestone dissolution (Picture 12). Past assumptions have held that armored limestone (limestone covered or coated with Fe or Al hydroxides) ceased to dissolve, but experiments show that coated limestone continues to dissolve at 20% the rates of unarmored limestone (Pearson and McDonnell 1975). Recent work has demonstrated that the rate for armored limestone may be even higher (Ziemkiewicz et al. 1997). The length of the channel and the channel gradient, which affects turbulence and the buildup of coatings, are design factors that can be varied for optimum performance (Picture 13). Optimum performance is attained on slopes exceeding 20%, where flow velocities keep precipitates in suspension, and clean precipitates from limestone surfaces. In appropriate situations,
OLCs are being implemented for long term treatment. Utilizing OLCs with other passive systems can maximize treatment and metal removal (Picture 14).

Among the questions still to be investigated are the behavior of OLCs in waters of different pH and high heavy metal loads (like metal mine drainage), possible interactions of slope with water chemistry, and the possible importance of limestone purity.

Ziemkiewicz et al. (1997) found armored limestone in a series of laboratory experiments was 50 to 90% as effective as unarmored limestone in neutralizing acid. Seven OLCs in the field reduced acidity in AMD by 4 to 205 mg/L as CaCO$_3$, at rates of 0.03 to 19 mg/L per meter of channel length. The highest removal rates were with channels on slopes of 45 to 60% and for AMD with acidity of 500 to 2600 mg/L as CaCO$_3$. For example, the Eichleberger OLC was 49 m long on a slope of 20%, and received about 378 L/min of 510 mg/L acidity as CaCO$_3$. After flowing down the channel, the acidity was decreased to 325 mg/L as CaCO$_3$ (36% decrease). The PA Game Commission OLC was only 11 m in length on a 45% slope, and received 484 L/min of 330 mg/L acidity as CaCO$_3$. The water acidity at the end of the channel was 125 mg/L as CaCO$_3$ (62% decrease).
Three OLCs were installed in the Casselman River Restoration project (Ziemkiewicz and Brant 1996). One OLC, 400 m long on a 8% slope, received 60 L/min of pH 2.7 water, 1290 mg/L as \( \text{CaCO}_3 \) acidity, 622 mg/L Fe, 49 mg/L Mn, and 158 mg/L Al. The effluent pH over a two year period was 2.9, acidity was 884 mg/L as \( \text{CaCO}_3 \) (31% decrease), Fe was 210 mg/L (66% removal), Mn was 42 mg/L (14% decrease), and Al was 103 mg/L (35% decrease).

At the Brandy Camp site in PA, a 15-m-long OLC on a 10% slope was employed to treat AMD with a pH of 4.3, acidity of 162 mg/L as \( \text{CaCO}_3 \), Fe of 60 mg/L, Mn of 10 mg/L, and Al of 5 mg/L (Hellier, case study 1997). After passage through the OLC, the effluent had a pH of 4.8, net acidity of 50 mg/L as \( \text{CaCO}_3 \), Fe of 17 mg/L, Mn of 8 mg/L, and Al of 3 mg/L. The OLC removed 72% of the Fe and about 20% of the Mn and Al from the water.

**Bioremediation**

Bioremediation of soil and water involves the use of microorganisms to convert contaminants to less harmful species in order to remediate contaminated sites (Alexander 1993). Microorganisms can aid or accelerate metal oxidation reactions and cause metal hydroxide precipitation. Other organisms can promote metal reduction and aid in the formation and precipitation of metal sulfides. Reduction processes can raise pH, generate alkalinity, and remove metals from AMD solutions. In most cases, bioremediation of AMD has occurred in designed systems like anaerobic wetlands where oxidation and reduction reactions are augmented by special organic substrates and limestone. In a few cases, substrates have been incorporated into spoils to aid in in-situ treatment of water by the use of indigenous microorganisms.

A mixture of organic materials (sawdust and sewage sludge) was emplaced into a mine spoil backfill to stimulate microbial growth and generate an anoxic environment through sulfate reduction. The results of the organic matter injection process caused no change in water pH, about a 20% decrease in acidity (1500 to 1160 mg/L as \( \text{CaCO}_3 \), and a similar decrease in Fe, Mn, and Al. The results indicate that the process works, but improvements in organic material injection and the establishment of a reliable saturated zone in the backfill are needed for maximum development (Rose et al. 1996).

The Lambda Bio-Carb process is a bioremediation system utilizing site-indigenous, mixed microorganism cultures selected for maximum effectiveness (Davison 1993). On a field site in PA using this bioremediation process, Fe in AMD was decreased from 18 mg/L to < 1 mg/L, Mn declined from 7 mg/L to 2 mg/L, and pH increased from around 6.0 to 8.0.

The Pyrolusite process uses selected groups of microorganisms growing on limestone to oxidize Fe and Mn into their insoluble metal oxides (Vail and Riley 1997). On a field site in PA using a limestone bed inoculated with microorganisms, Fe was decreased from 25 mg/L to < 1 mg/L, Mn went from about 25 mg/L to < 1 mg/L, while pH and alkalinity in the effluent were increased.

**Diversion Wells**

The diversion well is a simple device initially developed for treatment of stream acidity caused by acid rain in Norway and Sweden (Arnold 1991). It has been adopted for AMD treatment in the eastern USA. A typical diversion well consists of a cylinder or vertical tank of metal or concrete, 1.5-
1.8 m in diameter and 2-2.5 m in depth, filled with sand-sized limestone (Picture 15). This well may be erected in or beside a stream or may be sunk into the ground by a stream. A large pipe, 20-30 cm in diameter, enters vertically down the center of the well and ends shortly above the bottom (Picture 16). Water is fed to the pipe from an upstream dam or deep mine portal with a hydraulic head of at least 2.5 m (height of well). The water flows down the pipe, exits the pipe near the bottom of the well, then flows up through the limestone in the well, thereby fluidizing the bed of limestone in the well. The flow rate must be rapid enough to agitate the bed of limestone particles. The acid water dissolves the limestone for alkalinity generation, and metal flocs produced by hydrolysis and neutralization reactions are flushed through the system by water flow out the top of the well. The churning action of the fluidized limestone also aids in limestone dissolution and helps remove Fe oxide coatings so that fresh limestone surfaces are always exposed. Metal flocs suspended in the water are precipitated in a downstream pond.

Arnold (1991) used diversion wells for AMD treatment in PA and reports that three wells increased pH from 4.5 to 6.5, with corresponding decreases in acidity. For example, one diversion well located at Lick Creek treats about 1000 L/min of slightly acid water. After passing through the diversion well, the pH changes from 4.5 to 5.9 and the net acid water (8 mg/L as CaCO₃) changes to net alkaline water (6 mg/L as CaCO₃). Similar results are found for several other sites in PA.

Diversion wells have also been constructed in the Casselman River Restoration Project (Ziemkiewicz and Brant 1996). This large diversion well has a retention time of about 15 min for a 360-L/min flow of moderately acid water. The diversion well reduces the acidity from 314 to 264 mg/L as CaCO₃, Fe from 83 to 80 mg/L, and Al from 24 to 20 mg/L.
At the Galt site in WV, a diversion well changes a 20-L/min flow from a pH of 3.1 to 5.5, acidity from 278 to 86 mg/L as CaCO$_3$, Fe from 15 to 2 mg/L, and Al from 25 to 11 mg/L (Faulkner and Skousen 1995).

**Limestone Sand Treatment**

Sand-sized limestone may also be directly dumped into AMD streams at various locations in watersheds (Picture 17). The sand is picked up by the streamflow and redistributed downstream, furnishing neutralization of acid as the stream moves the limestone through the streambed. The limestone in the streambed reacts with the acid in the stream, causing neutralization. Coating of limestone particles with Fe oxides can occur, but the agitation and scouring of limestone by the streamflow keep fresh surfaces available for reaction.

The WV Division of Environmental Protection treats 41 sites in the Middle Fork River, including the headwaters of 27 tributaries (Zurbuch 1996). The first year's full treatment was based on four times the annual acid load for non-AMD impacted streams and two times the load for AMD tributaries. During subsequent years, limestone sand application will be an amount equal to the annual acid load, or about 2,000 tons/yr. About 8,000 tons of limestone were deposited among the 41 sites in 1995. Water pH has been maintained above 6.0 for several miles downstream of the treatment sites. The anticipated precipitate coating of the limestone was not observed. It is predicted that treating the river with limestone sand will be necessary three times a year to maintain water quality for fish populations.

**Acknowledgments**

This paper is the second section in "Acid Mine Drainage Control and Treatment," a chapter in a new book entitled "Reclamation of Drastically Disturbed Lands," being prepared by the American Society for Agronomy and the American Society for Surface Mining and Reclamation. The anticipated release date for this book is 1998. The author thanks Tiff Hilton, Ben Faulkner, Alan Sexstone, Paul Ziemkiewicz, Robert Darmody, John Sencindiver, Tim Phipps, Jerry Fletcher, Keith Garbutt, Bill Hellier, Art Rose and two anonymous reviewers for helpful comments during the review process. Acid mine drainage research at West Virginia University is supported by grants from the College of Agriculture and Forestry, the National Mine Land Reclamation Center, the USDI Bureau of Mines, the West Virginia Division of Environmental Protection, and from funds appropriated by the Hatch Act.
References
MEND Report 3.14.1
1996, Revised 1999
Review of Passive Systems for Treatment of Acid Mine Drainage

Click here to view MEND Report

Warning: There is no link back to this report and Bookmarks from the MEND Report.
Please use the Go To Previous View arrow to come back or bookmark this page in your browser.
C. Zipper and C. Jagie
2001
Passive Treatment of Acid-Mine Drainage with Vertical-Flow Systems
Introduction

The purpose of this publication is to present guidance for the design and construction of vertical-flow systems for the passive treatment of acid mine drainage (AMD). The term "passive treatment" refers to methods of treating AMD that rely on biological, geochemical, and gravitational processes. Passive systems do not require the constant input of chemical reagents that are characteristic of "active" AMD.
treatment.

This publication is intended to help potential users determine whether or not a vertical-flow passive treatment system should be considered for a specific AMD discharge. Should the reader decide to proceed, the reader is encouraged to engage the services of a professional with passive-system design and construction experience, especially if the system is intended to meet specific effluent criteria.

These guidelines reflect results of recent research and current practices. AMD treatment technology is developing rapidly as more is learned about how these systems function. Prior to engaging in a vertical-flow AMD-treatment project, readers are advised to access the on-line version of this publication, available through both Virginia Cooperative Extension http://www.ext.vt.edu/resources/ and the Powell River Project http://als.cses.vt.edu/prp/, for reference to updated design guidance as it becomes available.

Return to Table of Contents

Overview of Vertical Flow Systems

Vertical-flow systems have also been given a variety of names over the years such as: SAPS (for "successive alkalinity producing systems," Kepler and McCleary, 1994), RAPS (for "reducing and alkalinity producing systems," Watzlaf and others, 2000), and APS (for "alkalinity producing systems," Skousen and Ziemkiewicz, 1995). In 1987, A.C. Hendricks developed a vertical-flow system at Galax, Virginia, to treat the effluent from a long-abandoned pyrite mine (Hendricks, 1991). In 1990, Westmoreland Coal Company and A.C. Hendricks developed a vertical-flow system in Wise County, Virginia, working through Powell River Project (Duddleston and others, 1992). Kepler and McCleary (1994) developed similar systems in Pennsylvania. They are also largely responsible for the widespread use of vertical-flow systems in northern Appalachia, and the development of several design advances.

When properly designed, constructed, and maintained in appropriate situations, vertical-flow treatment offers advantages relative to other means of treating AMD. Unlike active treatment, vertical-flow systems do not require the purchase of chemical reagents or storage of chemical reagents on site. Although vertical-flow systems do require more area and volume than active systems sized with equivalent treatment capacity, they require far less area than other "wetland" systems. Vertical-flow systems are generally ineffective in removing Mn, but passive treatment methods for removing Mn from mine-discharge waters are currently being developed (Kerrick and Horner, 1998; Brent and Ziemkiewicz, 1997; Sikora and others, 1996). These systems can, however, be very effective in pre-treating AMD prior to an active treatment finishing process, which may reduce the total costs of meeting regulatory standards.

Even where Mn is not a problem, vertical-flow treatment systems should not be considered as either a stand-alone or a walk-away AMD-treatment solution. This publication describes how vertical-flow treatment can be integrated with other passive-treatment elements to provide AMD treatment, and it presents guidelines for vertical-flow system design.

Although vertical-flow systems do require periodic attention and maintenance, they can be maintained on a week-to-week basis with less time and expense than conventional active systems. Operators should expect, however, that a vertical flow system in a long-term application will require renewal via replacement of major system elements. Current design practice assumes 20- to 25-year
lifespans for these systems. As of this writing, at least two vertical-flow systems have operated successfully over periods approaching 10 years (Pine Branch in Virginia, and Howe Bridge in western Pennsylvania) without requiring renewal of major system elements. Many other systems have operated successfully over shorter periods, while still others have failed to meet treatment goals due to inadequate design.

**Acid Mine Drainage**

Acidic mine drainage (AMD) is an environmental pollutant of major concern in mining regions throughout the world. AMD occurs as a result of the oxidation of sulfide minerals when they are exposed to oxygen and water during the mining process. In coal-mining areas, the most common of these minerals is pyrite (FeS$_2$). The process for AMD formation is commonly represented by the following reactions:

$$\text{FeS}_2(s) + 3.5 \text{O}_2 + \text{H}_2\text{O} \rightarrow \text{Fe}^{2+} + 2 \text{SO}_4^{2-} + \text{H}^+ \ (1)$$

$$\text{Fe}^{2+} + 0.25 \text{O}_2 + \text{H}^+ \rightarrow \text{Fe}^{3+} + 0.5 \text{H}_2\text{O} \ (2)$$

$$\text{Fe}^{3+} + 2 \text{H}_2\text{O} \rightarrow \text{FeOOH} (s) + 3 \text{H}^+ \ (3)$$

The process is initiated with the oxidation of pyrite and the release of ferrous iron (Fe$^{2+}$), sulfate, and acidity (Eq. 1). The sulfide-oxidation process is accelerated by the presence of Thiobacillus bacteria. Ferrous iron then undergoes oxidation forming ferric iron (Fe$^{3+}$) (Eq. 2). Finally, Fe$^{3+}$ reacts with H$_2$O (is hydrolyzed), forming insoluble ferric hydroxide (FeOOH), an orange-colored precipitate, and releasing additional acidity (Eq. 3). The FeOOH formation process is pH-dependent, and occurs rapidly when pH is greater than 4.

**Passive Treatment of AMD**

Passive treatment systems are typically modeled after wetlands and other natural processes, with modifications directed toward meeting specific treatment goals. Early research included investigations of natural Sphagnum sp. peat wetlands that were receiving AMD (Weider, 1982). These systems were able to raise pH and lower iron concentrations without visible deterioration.

**Aerobic Wetlands**

One of the first designs put into use was a shallow (± 1 foot), surface flow wetland planted with cattails (Typha sp.) (Hedin and others, 1994a; Skousen and others, 1998; Skousen and others, 2000). Substrates for these wetlands varied from natural soils to composted organic matter. These "aerobic" wetlands aerated the mine waters flowing among the vegetation. This allowed for the oxidation of
Fe$^{2+}$ and its subsequent deposition as FeOOH. Aerobic wetlands are typically used to treat mildly acidic or net-alkaline waters containing elevated Fe concentrations. Published design criteria for Fe removal are up to 310 mg/day per square-foot on sites where the discharge is intended for regulatory compliance, and up to 620 mg/day per square-foot where regulatory compliance is not an issue (Hedin and others, 1994a). Where waters are net-alkaline and Fe is not a problem, aerobic wetlands have also proved capable of removing Mn, but very large areas are needed. Use of aerobic wetlands for Fe removal generally causes pH to decline due to the generation of proton acidity by Fe hydrolysis (Eq. 3) (Skousen and others, 1997).

**Anaerobic Wetlands**

Modifications of the aerobic wetland design were made to raise water pH and increase metal precipitation. These included the addition of a bed of limestone beneath an organic substrate (Hedin and others, 1994a). This design encouraged the generation of bicarbonate alkalinity (HCO$_3^-$) by both anaerobic microbial sulfate reduction (Eq. 4, with CH$_2$O representing biodegradable organic compounds) and limestone dissolution (Eq. 5).

$$2 \text{CH}_2\text{O} + \text{SO}_4^{2-} \rightarrow \text{H}_2\text{S} + 2 \text{HCO}_3^- \quad (4)$$

$$\text{CaCO}_3 + \text{H}^+ \rightarrow \text{Ca}^{2+} + \text{HCO}_3^- \quad (5)$$

The bicarbonate neutralizes the acidity of the AMD, thereby raising pH (Eq. 6) and increasing the precipitation of acid-soluble metals such as Fe.

$$\text{HCO}_3^- + \text{H}^+ \rightarrow \text{H}_2\text{O} + \text{CO}_2^\text{(aq)} \quad (6)$$

Anaerobic wetlands have proved capable of removing Fe and producing alkalinity. Hedin and others (1994a) reported average Fe removal rates of up to 1300 mg/day per square-foot, but these systems are limited in capability to raise pH, especially where Fe is present. The primary factor limiting their effectiveness is the slow mixing of the alkaline substrate water with acidic waters near the surface. This slow mixing can be overcome by constructing very large wetlands to provide long retention times (Skousen and others, 1997). This demand on land area is a major impediment to the increased use of these systems by mine operators with limited space for wetland construction.

Current guidelines for the construction of anaerobic wetlands advocate use of a 1- to 2- foot layer of organic matter over a 0.5- to 1- foot bed of limestone with a surface water depth of 1 to 3 inches. At water levels deeper than 2 to 3 inches, growth of wetland vegetation is hindered. The organic matter must be permeable to water and biodegradable; spent mushroom compost has been used successfully at a number of sites in northern Appalachia. For greater effectiveness, limestone may be mixed in with the organic matter. Cattails (Typha sp.) may be planted throughout the wetland to supply additional organic matter for heterotrophic bacteria and to promote metal oxidation with the release of oxygen from their root system (Skousen and others, 1997). Available guidelines for system sizing recommend planning for acidity removal rates 100 mg/day per square-foot for systems designed to achieve regulatory compliance, and up to 200 mg/day per square-foot where regulatory compliance is not a concern. For a more thorough review of anaerobic wetlands, see either Hedin and others (1994a) or Skousen and others (2000).
Anoxic Limestone Drains

One method used to reduce wetland size is pre-treatment of the AMD using anoxic limestone drains (ALDs). ALDs are limestone-filled trenches that can rapidly produce bicarbonate alkalinity via limestone dissolution. They are installed at the point of discharge to capture the AMD subterraneously. ALDs are capped with clay or compacted soil to prevent AMD contact with oxygen (Hedin and Watzlaf, 1994). The acidic water flowing through trench dissolves the limestone and releases bicarbonate alkalinity (Eq. 5). These systems have demonstrated capabilities to raise the alkalinity and/or neutralize acidity by as much as 300 mg/L (CaCO$_3$ equivalent) with retention times of only 14 - 23 hours (Hedin and Watzlaf, 1994, Hedin and others, 1994a), although net-alkalinity generation rates of 150 to 250 mg/L are more typical. The effluent is discharged into a settling pond to allow for acid neutralization, pH adjustment, and metal precipitation. ALD pretreatment of AMD allows for the construction of smaller, more effective treatment systems due to the decreased metal loadings and increased alkalinity of the ALD effluent discharged into them.

ALDs, however, are not capable of treating all discharges. Significant concentrations of Al or Fe$^{+3}$ in the discharge can cause an ALD to clog with metal-hydroxides once a pH of 4.5 or above is reached (Hedin and others, 1994a). When excess Fe$^{+3}$ is present in the AMD, or is allowed to form from Fe$^{+2}$ due to the presence of O$_2$, formation of solid FeOOH can occur within the ALD (Eq. 3). Ideally, Fe$^{+3}$, Al, and dissolved O$_2$ concentrations of waters being treated by an ALD would all be below 1 mg/L. However, AMD is not always ideal. Skousen and others (2000) state that ALDs have been used successfully for AMD with dissolved oxygen concentrations of up to 2 mg/L and Al concentrations of up to 25 mg/L, when less than 10 percent of total Fe in the Fe$^{+3}$ form. If Al is present at a concentration greater than 1 mg/L and waters in the ALD reach a pH of 4.5 or above, Al will precipitate as Al(OH)$_3$. Both FeOOH (eq. 3) and Al(OH)$_3$ precipitation generate acidity.

$$\text{Al}^{3+} + 3 \text{H}_2\text{O} \rightarrow \text{Al(OH)}_3 (s) + 3 \text{H}^+ \quad (7)$$

ALD systems will also fail if Fe$^{3+}$ precipitates on the limestone surface, thus limiting its dissolution, a process known as "arming." In low dissolved oxygen ("anoxic") environments, the Fe$^{2+}$ form of iron predominates and does not form a coating on the limestone or interfere with limestone dissolution (Hedin and others, 1994a; Watzlaf, 1997). A thorough reference for the design and sizing for ALDs can be found in Hedin and Watzlaf (1994).

Vertical Flow Systems

Vertical flow systems combine the treatment mechanisms of anaerobic wetlands and ALDs in an attempt to compensate for the limitations of both (Kepler and McCleary, 1994). The basic elements of these systems are similar to the anaerobic wetland, but a drainage system is added within the limestone layer to force the AMD into direct contact with both the organic matter and the limestone.

The three major vertical-flow system elements are the drainage system, a limestone layer, and an organic layer. The system is constructed within a water-tight basin, and the drainage system is constructed with a standpipe to regulate water depths and ensure that the organic and limestone layers remain submerged. As the AMD waters flow downward through the organic layer, two essential functions are performed: dissolved oxygen in the AMD is removed by aerobic bacteria utilizing biodegradable organic compounds as energy sources, and sulfate-reducing bacteria in the anaerobic
zone of the organic layer generate alkalinity (Eq. 4). Low DO concentrations, biodegradable carbon, and the presence of dissolved sulfate are necessary for sulfate-reduction to take place. An organic layer capable of removing DO to concentrations below 1 mg/L is essential to prevent limestone armoring. In the limestone layer, \( \text{CaCO}_3 \) is dissolved by the acidic, anoxic waters moving down to the drainage system, producing additional alkalinity. The final effluent is discharged from the drainage system standpipe into a settling pond to allow acid neutralization and metal precipitation prior to ultimate discharge.

In order to avoid clogging of the limestone layer with Fe\(^{+3} \) and Al precipitants (Eqs. 3 and 7), a valved flushing pipe is typically included as a part of the drainage system (Kepler and McCleary, 1997). When opened, this valved drain discharges at a lower elevation than the standpipe. Head pressure (usually, 6 to 10 feet) caused by the standing water in the system moves waters through the system rapidly, flushing the gel-like forms of Al and Fe ("floc") that accumulates in the drainage pipes and limestone pores. Opening this valve periodically removes the loose metal hydroxide floc and discharges it into the settling pond.

Current practices include a limestone layer of 2 to 3 feet in depth, an organic layer of 6 to 12 inches in depth, and a standpipe and basin capable of maintaining a 3- to 5-feet deep body of water above the organic layer (Skovran and Clouser, 1998; Kepler and McCleary, 1994). Building systems with 3 feet or more of standing water over the mulch layer provides sufficient head-pressure and aids flushing.

For severe AMD discharges, several vertical-flow systems can be linked in series to generate alkalinity successively until the treatment goals are reached.

**Open Limestone Channels**
Where AMD must be conveyed over some distance prior to or during treatment, use of open channels lined with limestone has been shown to be an effective mechanism for removing Fe and generating small amounts of alkalinity (Ziemkiewicz and others, 1997). Even though the limestone in such channels typically becomes armored with Fe, research indicates that the armored limestone retains some treatment effectiveness. Open limestone channels are most effective when placed on slopes of greater than 20%, as the abrasive action of fast-moving water tends to dislodge the armoring Fe. Open limestone channels can be effective as one element of a passive treatment system, but typically are not relied upon for stand-alone AMD treatment.

Return to [Table of Contents](#)

**Developing a Passive Treatment Strategy**

**AMD Characterization**
The design of all passive treatment systems starts with characterization of influent AMD chemistry and flow. Prior to designing a passive treatment system, a complete characterization of influent AMD is needed to determine which type of system is appropriate and how to design that system to meet treatment goals.

Regular sampling over at least a 12-month period is recommended to account for the variations that may occur or in response to seasonal changes or storms. At a minimum, all water samples should be analyzed for pH, total Fe, Mn, Al, \( \text{SO}_4^{2-} \), total alkalinity and total acidity. Additional analyses,
including Fe$^{+2}$, Fe$^{+3}$, and dissolved oxygen, are necessary if an anoxic limestone drain is being considered as a treatment option. Dissolved oxygen and pH should be measured on-site. Dissolved oxygen measurements are sensitive, and experience by the sampling personnel is necessary to obtain an accurate reading. Samples designated for metals analysis should be filtered at the time of sampling to remove particulate matter and acidified to pH<2 (APHA, 1985). Acidity and alkalinity samples should be placed on ice immediately following sampling and analyzed within 24 hours. Flow measurements should also be taken on all sampling dates.

Based on water-sampling data, a "design" water quality condition should be established. This will generally be the worst-case condition, as defined relative to regulatory standards, if the AMD discharge is intended to achieve regulatory compliance. The water sampling procedures should assure that a variety of weather and climate conditions are represented, to ensure that the resultant data provide a realistic assessment of the design conditions. Average and maximum influent flow should also be estimated for use in the design process.

**Passive Treatment System Selection**

The selection of a passive treatment system is governed by influent water quality and site characteristics. The diagram in Figure 2 illustrates a decision process for selecting an appropriate system for a given discharge (Hedin and others, 1994a; Skousen and others, 1998). For net alkaline discharges containing elevated concentrations of Fe, no additional alkalinity additions are needed. The only conditions necessary to complete treatment are an oxidizing environment and sufficient residence time to allow for metal oxidation and precipitation. These conditions can be provided by a settling pond; if sufficient area is available, the settling pond may be followed by an aerobic wetland.

The treatment of net acidic drainage can be handled in a number of ways depending on influent chemistry. If the influent quality is suitable for an ALD, an ALD can be employed as a pretreatment method. A post-ALD settling pond or aerobic wetland is required to allow for the oxidation and precipitation of metals.

Acidic waters that are not suitable for ALDs can be treated with either an anaerobic wetland or a vertical flow system. Due to the potentially large demands on land area of anaerobic wetlands, they are usually only practical for low-flow situations. For systems that receive water that has a pH greater than 4, settling ponds may precede an anaerobic wetland cell to remove significant quantities of Fe. The remaining discharges can be treated using a vertical-flow system.

Return to [Table of Contents](#)

**Vertical-Flow System Design**

Vertical-flow passive treatment systems are able to neutralize acidity and promote metal precipitation in difficult treatment situations. Due to the active mixing of the AMD with the limestone, acid neutralization is more rapid in vertical-flow systems than in anaerobic wetlands, and thus vertical-flow systems require shorter residence times and smaller surface areas. These systems are not stand-alone, but require the addition of an oxidation/settling pond at the effluent point to allow for the precipitation and storage of the metals in solution. For discharges containing significant quantities of Fe$^{+3}$, vertical-flow systems should be preceded by either a settling pond or an aerobic wetland if sufficient land area is available. The removal of such metals prior to vertical-flow treatment will lengthen the system's useful life and reduce necessary maintenance by limiting accumulation of
metal-hydroxide precipitants on the organic matter surface. A settling pond should also precede the 
system if incoming waters contain sediment.

Guidelines reported by Skovran and Clouser (1998), Skousen and others (1998) and Kepler and 
McCleary (1994) provide a general form for design of these systems.

**Sizing the Limestone Layer**

Although per-unit-area Fe removal-rates are typically applied in designing "wetland" treatment 
systems, per-unit-area rates of alkalinity generation by installed vertical-flow systems vary. Research 
has demonstrated the presence of relationships between influent water quality, AMD residence time 
in the limestone layer, and alkalinity generation (Jage and others, 2000). The primary factor 
governing alkalinity generation by most vertical-flow systems is the rate at which limestone dissolves 
relative to the rate at which AMD moves through the system. Residence-time of the AMD in the 
limestone layer is one factor governing alkalinity generation. Limestone dissolves most rapidly during 
the first few hours of AMD contact. As the waters in contact with the limestone become saturated 
with dissolved Ca$^{2+}$ and HCO$_3^-$, the rate of limestone dissolution slows considerably. Another factor 
governing the rate at which limestone dissolves is pH; at lower pH, CACO$_3$ dissolves more rapidly.

Research has developed a model that can be used to estimate vertical flow systems' alkalinity 
generation rates as a function of AMD residence time within the limestone layer and AMD 
concentrations of Fe and non-Mn acidity (Jage and others, 2001; see Eq. 9 below). Given the volume 
and quality of the AMD to be treated, this model can be used to estimate the size of the limestone 
layer required to generate a given quantity of alkalinity. We recommend that the model be used to 
provide a design minimum, but that systems be constructed larger than indicated by the model 
whenever possible. Increasing the size of a vertical-flow system will increase the probability of 
successful treatment.

**Calculating a Preliminary Limestone Volume**

The first step in the system sizing process is to determine the size of the limestone layer and number 
of vertical-flow cells needed for adequate treatment. We recommend that the system be sized to 
generate alkalinity sufficient to offset incoming non-Mn acidity, plus additional alkalinity so as to 
achieve a factor of safety. We recommend sizing the system to generate at least 100 mg/L of 
alkalinity, over and above the amount needed to offset influent non-Mn acidity when sufficient land 
area is available, so as to provide a reasonable probability of successful treatment. Non-Mn acidity 
can be calculated as:

$$\text{Non-Mn Acidity} = \text{Acidity} - 1.818 \times \text{Mn} \quad (8)$$

where,

Acidity = Total Acidity (mg/L as CaCO$_3$) of the design influent water quality

Mn = Manganese concentration (mg/L) of the design influent water quality

Non-Mn Acidity = acidity derived from Al, Fe, H+ and other ions (mg/L as CaCO$_3$)

Once the design rate of alkalinity generation has been determined, the limestone residence time of a 
system can be estimated using the equation below:
Alknet = 99.3 * log 10(t_r) + 0.76 * Fe + 0.23 * Non-Mn Acidity - 58.02 (9)

where,

Alknet = net alkalinity to be generated (mg/L as CaCO_3)

Fe = total iron concentration (mg/L) of the design influent water quality

Non-Mn Acidity = non-manganese acidity (mg/L as CaCO_3 - see equation 8)

\( t_r \) = average residence time in the limestone layer (hours).

Equation 9 ("the model") can be solved mathematically for \( t_r \), if the reader is so inclined, or the reader may choose to estimate a residence time that may be achievable based on site conditions and use the model to determine whether or not a system built with such a residence time is likely to be adequate for treatment of the design AMD discharge. Figure 3 represents Equation 9's predictions for a sample influent water quality.

This model is intended for application to systems built with high-calcium limestone in the 4-to-6 inch size range. High-calcium limestone contains more than 90% CaCO_3 and is more soluble than limestone that contains appreciable quantities of Mn. The CaCO_3 residence time that results from solving Equation 9 should be adjusted to account for limestone dissolution as the system ages (see below). The model was developed by analyzing data from vertical flow systems receiving influent waters with Fe concentrations less than 300 mg/L and non-Mn acidity concentrations of less than 500 mg/L (Jage and others, 2001). The model has not been tested for drainages where Al concentrations exceed 60 mg/L.

Equation 9 is not expected or intended to give precise results. Figure 4 shows the relationship of predicted alkalinity generation to observed generation for 18 vertical flow cells. All values plotted are system averages over periods exceeding one year. The plots for Howe Bridge and the Oven Run systems were calculated from system averages published by Watzlaf and others (2000), while all other system averages were calculated from monthly observations.

For the data set of 179 observations used to derive the above predictive model (Jage and others, 2001), the standard deviation of the difference between observed and predicted values is about 50 mg/L. This deviation between predicted and observed values is the justification for suggesting sizing the system to generate alkalinity in excess of the anticipated need to offset incoming non-Mn acidity, especially if the system is intended to achieve regulatory compliance. Larger systems will provide an increased factor of safety, and are likely to operate successfully for a longer time prior to requiring major renovation.

Figure 5 demonstrates that alkalinity generation rates vary considerably between systems. Two of the systems exhibited average alkalinity generation rates in excess of 300 mg/L and less than 2500 mg/day per square foot. In both of these systems, average residence times exceeded 300 hours. One southern West Virginia system exhibited an average alkalinity generation rate of approximately 7000 mg/day per square foot; this system had a relatively short residence time, a rich organic layer more
than one meter in thickness, and received AMD with pH's that are favorable to sulfate-reducing bacteria, in the 4-to-5 range. This system generated alkalinity most rapidly during the early years of its operation; during its third year, its performance declined considerably.

These data demonstrate Equation 9 may be used to provide design guidance, but does not provide precise predictions. Vertical-flow systems' performance exhibits considerable variation in the field, on a month-to-month basis as well as between locations (Figure 6). Building a system with a larger residence time will increase the probability of successful treatment.

For highly acidic AMD discharges, the above sizing method may generate an estimated residence time of several hundred hours. If area limitations prevent construction of such a large system, treatment may be provided as two or more successive vertical-flow cells separated by a settling pond. For example, considering the influent water chemistry represented by Figure 3: a limestone residence time on the order of 1000 hours would be required in a single cell to generate 300 mg/L acidity, whereas two cells in series, each with a 30-hour residence time and separated by an settling pond, may be capable of generating a comparable amount of alkalinity. As a conservative design principle, we recommend that residence times of less than 15 hours should be avoided and longer residence times should be preferred. At low residence times (that is, at rapid rates of AMD movement through the vertical-flow system), the organic matter layer within the vertical flow system may begin to limit system performance. At very rapid rates of water movement, the permeability of the organic layer may become limiting especially if Fe is being precipitated on its surface. Also, as the organic layer ages, its capacity to remove oxygen from the AMD will decline. Therefore, with all other things being equal, larger systems with slower residence times can be expected to maintain performance for longer periods than smaller systems with short residence times.

The residence time calculated with Equation 9 is an estimate of the time that the AMD should reside in the limestone layer to achieve desired results. In order to size the system, the residence time must be converted into a limestone layer volume (\(V_{ls}\), expressed in cubic feet):

\[
V_{ls} = \frac{8.02 \times Q \times t_{r}}{V_{v}} \quad (10)
\]

where,

\(Q\) = influent flow (gallons per minute)

\(t_{r}\) = residence time in limestone (hours)

\(V_{v}\) = bulk void volume of limestone expressed as a decimal (e.g., 50% = 0.5)

A reasonable estimate of the bulk-void volume of 4-to-6 inch limestone is about 50%. For common unit-conversion factors, see Table 1.

**Adjusting Limestone Volume to Account for Loss over Time:**
An additional volume of limestone should be added to compensate for limestone dissolution over the design life of the SAPS based on a method defined by Hedin and Watzlaf (1994) for ALDs. The additional volume of limestone needed (\(V_{ls+}\), expressed as cubic feet) over the design size can be
calculated as:

\[ V_{ls} = \frac{0.044 \times Q \times C \times T}{x} \]  

where,

- \( Q \) = influent flow (GPM)
- \( C \) = predicted net alkalinity generation (mg/L)
- \( T \) = design life (years)
- \( x \) = \( \text{CaCO}_3 \) content of limestone, expressed as a decimal (e.g., 90% = 0.9)

This volume of limestone should be added to the amount needed to attain the design residence time (Table 2). By placing additional limestone in the vertical flow system, the design residence time will be maintained even as some limestone is dissolved by AMD moving through the system. Common practice is to design the limestone layer for 20- to 25-year lifetimes. High-calcium limestone should be used to construct the limestone layer; use of dolomitic limestones should be avoided. High-calcium limestone is more soluble than dolomitic limestone.

The Organic Layer

The organic layer is the most vulnerable of the major system elements and is critical to long-term performance. In addition to harboring sulfate-reducing bacteria, the organic layer removes dissolved oxygen and promotes reducing conditions necessary to prevent limestone armoring. The removal of dissolved oxygen, however, is directly related to water temperature and the AMD residence time in the organic matter. In order to ensure that the vertical-flow system performs well year-round and to prevent performance degradation due to limestone armoring, the organic layer must be sized adequately to ensure cold weather performance. Permeability is also a key property of the organic layer.

Well-weathered organic bark material has been used successfully in one high-residence-time Virginia system, but this system receives a relatively high-quality influent. Bark materials tend to be permeable, but they have a relatively low biochemical oxygen demand due to high proportions of large, woody debris that are not readily broken down by the bacteria. Other materials that are more easily processed by aerobic bacteria, such as composted manure or spent mushroom compost, should allow for shorter organic matter residence times given the greater bioavailability of the organic compounds in these materials. In practice, a variety of materials have been used successfully including well-decomposed wood mulch, spent mushroom compost, composted manure, and mixtures of composted materials with less-expensive organic sources such as rotting hay. Mixing organic-layer materials with limestone is not recommended, due to the potential for metal-hydroxide floc precipitation within the small pores of the mulch layer where flushing will not be effective.

For most systems, organic layer depths of 12 to 18 inches should be adequate. Deeper substrates can be problematic due to the low permeability of organic matter, especially as it begins to decompose. Shallower substrates should be avoided due to the risk of creating zones of preferential flow that would allow oxygenated water to reach the limestone layer. In designing systems with relatively rapid
movement of water through the organic layer due to exceptionally short residence times and/or large limestone-layer depths, the permeability of the organic-layer material should be tested.

Care should be used in installing the organic layer to assure that the material is well mixed and to assure uniform distribution and depth of material across the limestone-layer surface. Once the organic layer is in place, any activity causing compaction, such as walking or driving equipment on the surface of the system, should be avoided. Such activity may also result in the creation of zones of preferential flow. These areas will cause surface waters that are moving toward the drain to "short circuit" the system and decrease treatment effectiveness.

Drainage

The subsurface drainage system should be constructed from schedule 40 perforated PVC piping; piping diameter should be determined based on the design flow. Generally speaking, drainage diameters less than 6 inches should be avoided, due to the potential for metals precipitation and sediment accumulation within the drainage structure. Hole diameters should not be less than 1/2 inch, and 1 inch is preferred. The holes of tile drains, if used in vertical flow cells, should be enlarged. Adequately sized holes will help to ensure that plugging by floc precipitants does not occur. Typical layouts for the drain are 'T' or 'Y' shaped, and located in the lower 12 in of the limestone layer (Figure 7). Drains should be designed and oriented so as to promote full utilization of the limestone volume. The drains are joined to an effluent standpipe that is elevated to maintain a constant head of water above the organic substrate. The effluent should cascade into the adjacent settling pond in order to oxygenate the waters and promote the precipitation of the metals in solution.

A flushing system should also be included to maintain maximum treatment efficiency. This consists of a valved discharge port connected to the drainage network located at a level below the height of the effluent standpipe (Figure 8). When the valve is opened, the head of water in the vertical-flow system causes a rapid drawdown of the system, which removes the metal-hydroxide floc that can accumulate in the limestone layer. The flushing system outlet should discharge the floc into the settling pond to allow for the collection and removal of the precipitants. This drawdown process can require 10 to 15 minutes but should be continued until the discharge waters run clear.

A settling pond located after the vertical flow cell is crucial to effective treatment performance. For effective treatment prior to discharge, the settling pond is a necessity. The effluent of the vertical flow cell must undergo oxidation, pH adjustment, and subsequent precipitation of insoluble metal complexes before being discharged. The settling pond allows for these processes to take place in a controlled setting where the precipitates can be dredged and disposed of in a proper manner. Settling ponds should be large enough to allow for the accumulation of the precipitated metals with recommended residence times of at least 2.8 days (Skovran and Clouser, 1998).

Construction

Because Skovran and Clouser (1998) have produced a detailed guide to construction practice, only a small amount of construction guidance will be repeated here. Vertical-flow system construction requires the excavation of a basin, and a compacted clay or plastic liner to prevent seepage of untreated AMD into the groundwater. Side embankments may be constructed with 2:1 interior slopes and 3:1 exterior slopes with 8 - 10 ft top widths (Figure 8). Skovran and Clouser (1998) also recommend a minimum of 12 inches of freeboard above design high water to an emergency spillway in order to maintain system integrity.

Skovran and Clouser (1998) recommend considering public safety when designing the basin. Some
developers have chosen to encircle vertical-flow systems with chain link fences and post warning signs, in an effort to discourage uninvited visitors that might be attracted by the open waters. Configuring the basin to include a shallow-water bench area adjacent to the bank can enhance safety. Such a shallow bench separates the deep-water pool from bankside walking areas; in the event of accidental entry into the pool (e.g., someone falls in), the shallow bench will aid a quick exit. Depending on the AMD’s acidity, the shallow bench may also become populated with cattails and other wetland vegetation, making entry to the deeper pool appear more difficult.

In long-term applications, the system basin should be situated so as to allow access by mechanical equipment, such as a back hoe or a small loader, to aid eventual system renewal.

**Operation, Maintenance, and Renewal**

Once the installation is complete, influent and effluent water-chemistry and flow monitoring should continue to allow assessment of system performance. Availability of adequate background data will enable informed decisions regarding maintenance if water treatment performance begins to deteriorate. Drainage systems should be flushed periodically; common practice is about one flush per month, but the frequency should be determined based on the rate of Al and Fe accumulation.

Both organic matter and limestone are consumed by vertical-flow system operation, so degradation of performance over time should be expected. When the operator determines that the time for renewal is at hand, the first step would be to drain the system and excavate the organic layer. Depending upon the degree of limestone armoring, the system operator may wish to either remove and replace the limestone, or add some additional limestone prior to re-installation of the organic layer. If substantial quantities of Fe-precipitate are deposited in the limestone layer due to organic layer deterioration, then the drainage system may also require replacement.

**Table 1. Common conversion factors for use in vertical-flow system design.**

<table>
<thead>
<tr>
<th>Units</th>
<th>Equivalent units</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 gallon</td>
<td>0.134 cubic feet</td>
</tr>
<tr>
<td>1 gallon per minute</td>
<td>8.02 cubic feet per hour</td>
</tr>
<tr>
<td>1 cubic foot of 4-to-6 inch limestone</td>
<td>100 pounds (approximate)</td>
</tr>
<tr>
<td>1 pound CaCO₃</td>
<td>454,000 mg of alkalinity (as CaCO₃)</td>
</tr>
</tbody>
</table>

**Table 2. Example Calculation**

1. Determine design water quality and flow from field measurements: Fe concentration = 40 mg/l
   Acidity = 200 mg/l as CaCO₃
   Mn concentration = 25 mg /l
   Flow = 10 gallons per minute
2. Calculate non-Mn acidity using Equation 8. 154.6 mg/l as CaCO₃
3. Determine design rate of alkalinity generation: Designer wishes to 250 mg/l alkalinity, as achieve a high probability of success, so the system is designed to CaCO₃ generate excess alkalinity.
4. Determine residence time necessary to achieve 250 mg/l of 270 hours alkalinity production, given the design water quality. Insert all parameters into equation 9; solve for residence time (Note that figure 3 was produced using influent data identical to this example).
5. Determine preliminary volume of 4-to-6 inch high-calcium 43,308 cubic feet limestone necessary to achieve desired residence time, using equation 10. (This estimate assumes bulk void volume = 50%).
6. Determine design lifetime of system. 20 years
7. Determine additional limestone needed to offset amount dissolved 2,444 cubic feet over design life, using Equation 11. (This estimate assumes CaCO3 content of limestone = 90%).
8. Determine total volume of limestone needed (add 7 + 8). 45,752 cubic feet
9. Determine if project is worth further investigation:
   - Assuming a limestone layer of 4 feet and sloping sides, this project will require an area on the order of 12,000 square feet for the vertical flow cell alone.
   - If this amount of flat area is not available but some smaller areas separated by elevation differences, are available: recalculate the required area as two cells, each designed to produce 125 mg/l alkalinity or more per day, separated by a settling pond.
   - If either option looks feasible, involve a party experienced in passive treatment.

Return to Table of Contents

**Acknowledgments**

Thanks to Jeff Skousen (West Virginia University), Art Rose (Pennsylvania State University), and George Watzlaf (U.S. Department of Energy) for their assistance to the research that supports these recommendations, and to Jeff Skousen for his thorough review. Thanks to Albert Hendricks for his guidance and assistance throughout. Thanks to the coal-mining firms that assisted that research. The research was funded by Powell River Project. (http://als.cses.vt.edu/PRP/)

Return to Table of Contents

**Literature Cited**


Return to Table of Contents

Powell River Project / Virginia Cooperative Extension Publications

Reclamation Guidelines


Restoring Forests on Surface-Mined Land J.A. Burger and JL. Torbert. VCE 460-123.  
http://www.ext.vt.edu/pubs/mines/460-123/460-123.html

Establishment and Maintenance of Quality Turfgrass on Surface Mined Land. John R Hall III. VCE 460-126.


http://www.ext.vt.edu/pubs/mines/460-130/460-130.html

Reclamation of Coal Refuse Disposal Areas. Daniels, W.L., Barry Stewart, and Dennis Dove. VCE 460-131.  


Information for the Virginia Coalfields


Recovery of Native Plant Communities after Mining. K. Holl, C. Zipper, and J. Burger. VCE 460-140.  

Return to Table of Contents

View this document in PDF format

Visit Virginia Cooperative Extension
J.J. Gusek

20002

Sulfate-reducing Bioreactor Design and Operating Issues
SULFATE-REDUCING BIOREACTOR DESIGN AND OPERATING ISSUES: IS THIS THE PASSIVE TREATMENT TECHNOLOGY FOR YOUR MINE DRAINAGE?

James J. Gusek, P.E.
Knight Piésold and Co.
1050 Seventeenth Street, Suite 650
Denver, CO 80265
e-mail jgusek@knightpiesold.com

ABSTRACT

There are basically two kinds of biological passive treatment cells for treating mine drainage. Aerobic Cells containing cattails and other plants are typically applicable to coal mine drainage where iron and manganese and mild acidity are problematic. Anaerobic Cells or Sulfate-Reducing Bioreactors are typically applicable to metal mine drainage with high acidity and a wide range of metals. Most passive treatment systems employ one or both of these cell types. The track record of aerobic cells in treating coal mine drainage is impressive, especially back in the eastern coalfields. Sulfate-reducing bioreactors have tremendous potential at metal mines and coal mines but have not seen as wide an application.

This paper presents the advantages of sulfate-reducing bioreactors in treating mine drainage, including the ability to work in cold, high altitude environments; handle high flow rates of mildly affected ARD in moderate acreage footprints; treat low pH acid drainage with a wide range of metals and anions including uranium, selenium, and sulfate; accept acid drainage-containing dissolved aluminum without clogging with hydroxide sludge; have life-cycle costs on the order of $0.50 per thousand gallons; and be integrated into “semi-passive” systems that might be powered by liquid organic wastes.

Sulfate reducing bioreactors might not be applicable in every abandoned mine situation. A phased design program of laboratory, bench, and pilot scale testing has been shown to increase the likelihood of a successful design.

Additional Key Words: Constructed wetlands, acid mine drainage, heavy metals, sulfate reduction
INTRODUCTION

It has been over 20 years since the pioneering work of a group of researchers at Wright State University documented water quality improvements in a natural *Sphagnum* bog in Ohio that was receiving low pH, metal laden water (Huntsman, et al., 1978). Independently, a group at West Virginia University found similar results at the Tub Run Bog (Lang, et al., 1982). Subsequently, researchers, practitioners, and engineers focused on developing the promising technology of using “constructed wetlands” to treat acid mine drainage (AMD) or acid rock drainage (ARD). But the term “wetland,” besides carrying legal and regulatory baggage, does not quite describe structures like “anoxic limestone drains” or “successive alkalinity producing systems;” hence, the term “passive treatment” was coined.

The design of passive treatment systems basically entails the selection of treatment “modules” appropriate to the geochemistry of the mine drainage. As shown in Figure 1, there are two geochemical “zones” in a natural wetland ecosystem. The lower, oxygen-depleted zone is where sulfate-reducing bacteria thrive. The focus of this paper will be the design of passive treatment modules that capitalize on the geochemical reactions typically found in the anaerobic zone of natural systems.

![Figure 1. Typical Natural Wetland Geochemical Zones](image)

Definition of Passive Treatment

There are many technologies for treating AMD/ARD. To properly focus the discussion, the following definition of passive treatment is proposed:

*Passive treatment* is a process of sequentially removing metals and/or acidity in a natural-looking, man-made bio-system that capitalizes on ecological and geochemical reactions. The process requires no power and no chemicals after construction and lasts for decades with minimal human help.
It is a *sequential* process because no single treatment cell type works in every situation or with every AMD/ARD geochemistry. It is an *ecological/geochemical* process because most of the reactions (with the exception of limestone dissolution) that occur in passive treatment systems are biologically assisted. Lastly, it is a *removal* process because the system must involve the filtration or immobilization of the metal precipitates that are formed. Otherwise, they would be flushed out of the system, and the degree of water quality improvement would be compromised.

A truly passive system should also function for many years, without a major retrofit to replenish construction materials, and be able to function without using electrical power. Benning and Ott (1997) described a volunteer passive system outside of an abandoned lead-zinc mine in Ireland that has apparently been functioning unattended for over 120 years. Ideally, a passive treatment system should be designed to last for at least several decades.

**METAL REMOVAL AND OTHER BIO-GEOCHEMICAL MECHANISMS IN PASSIVE TREATMENT SYSTEMS**

Many physical, chemical, and biological mechanisms are known to occur within passive treatment systems to reduce the metal concentrations and neutralize the acidity of the incoming flow streams. Notable mechanisms include:

- Sulfide and carbonate precipitation catalyzed by sulfate-reducing bacteria (SRB) in anaerobic zones
- Hydroxide and oxide precipitation catalyzed by bacteria in aerobic zones
- Filtering of suspended material
- Metal uptake into live roots and leaves
- Adsorption and exchange with plant, soil, and other biological materials.

Remarkably, some studies have shown that plant uptake does not contribute significantly to water quality improvements in passive treatment systems (Wildeman, et al., 1993). However, plants can replenish systems with organic material and add aesthetic appeal. In aerobic systems, plant-assisted reactions appear to aid overall metal removal performance, perhaps by increasing oxygen and hydroxide concentrations in the surrounding water through photosynthesis-related reactions and respiration in the plant root zone. Plants also appear to provide attachment sites for oxidizing bacteria/algae. Research has shown that microbial processes are a dominant removal mechanism in passive treatment systems (Wildeman, et al., 1993). One anonymous researcher considered a passive treatment system as a "bioreactor with a green toupee," referring to the substrate where most of the bioreactions occur and the collection of plants that grow on top of the treatment cells.

**Sulfate Reducing Bioreactors**

Sulfate reduction has been shown to effectively treat AMD/ARD containing dissolved heavy metals, including aluminum, in a variety of situations. The chemical reactions are facilitated by the bacteria *desulfovibrio* in sulfate-reducing bioreactors as shown in Figure 2 in schematic form and the photo in Figure 3.
The sulfate-reducing bacterial reactions (equation 1) involve the generation of:

- Sulfide ion (S$^{2-}$), which combines with dissolved metals to precipitate sulfides (equation 2)
- Bicarbonate (HCO$_3^-$), which has been shown to raise the pH of the effluent
The sulfate reducing bacteria produce sulfide ion and bicarbonate in accordance with the following reaction (Wildeman, et al., 1993):

\[ \text{SO}_4^{2-} + 2 \text{CH}_2\text{O} \rightarrow \text{S}^2 + 2 \text{HCO}_3^- + 2 \text{H}^+ \]

1)

The dissolved sulfide ion precipitates metals as sulfides, essentially reversing the reactions that occurred to produce AMD/ARD. For example, the following reaction occurs for dissolved zinc, forming amorphous zinc sulfide (ZnS):

\[ \text{Zn}^{2+} + \text{S}^2 \rightarrow \text{ZnS} \]

2)

Suspected geochemical behavior of aluminum in sulfate reducing bioreactors has just recently been documented (Thomas and Romanek, 2002). It is suspected that insoluble aluminum sulfate forms in the reducing environments found in sulfate-reducing bioreactors, perhaps in accordance with the following reaction which is one of many possible:

\[ 3\text{Al}^{3+} + \text{K}^+ + 6\text{H}_2\text{O} + 2\text{SO}_4^{2-} \rightarrow \text{KAl}_3(\text{OH})_6(\text{SO}_4)_2 \text{ (Alumite)} + 6\text{H}^+ \]

3)

The key conditions for SRB health are a pH of 5.0 (maintained by the SRB itself through the bicarbonate reaction and/or the presence of limestone sand), the presence of a source of sulfate (typically from the AMD/ARD), and organic matter ([CH₂O] from the substrate). Sulfate-reducing bioreactors have been successful at substantially reducing metal concentrations and favorably adjusting pH of metal mine drainages.

**FLOW CHART FOR PASSIVE TREATMENT SYSTEM DESIGN**

In the late 1980s, the design methods for aerobic passive treatment cells for iron removal were still under development. Brodie (1991) sorted out the empirical relationships in a milestone design flow chart that provided the foundation for a more comprehensive design flow chart subsequently developed by Hedin and Nairn at the former U.S. Bureau of Mines as shown in Figure 4.

This figure, in one form or another, continues to guide engineers and practitioners in the passive treatment cell design process. It has been modified by the author to include the passive treatment of heavy metal-bearing AMD/ARD based on observations since 1988. The sulfate-reducing bioreactor as shown reflects where this particular technology fits in the design philosophy. Although the technology is well suited for AMD/ARD with net acidity and/or heavy metals, it can also be effectively applied to net alkaline water sources as indicated by the arrow drawn from the settling pond on the left hand side of the flow chart.
Figure 4. Flow Chart for Selecting a Passive AMD Treatment System Based on Water Chemistry and Flow (Adapted from Hedin, et al., 1994).

**PHASED DESIGN PROTOCOL**

There is no "cookbook" design manual for passive treatment systems although the design flow chart above is a safe starting point. A phased approach design project is recommended; it typically begins in the laboratory with static tests, graduating to final testing phases (bench and pilot) performed at the site on the actual AMD/ARD. Bench scale testing will determine if the treatment technology is a viable solution for the AMD/ARD and will narrow initial design variables for the field pilot. A proper bench scale test will certainly reduce the duration of the more costly field pilot test. Field pilot test duration can range from days, to months, to years, depending on the nature of the technology. Depending on the nature of the equipment and personnel needed, significant costs may be incurred during the field pilot tests – about $500 to $1,000 per week, mostly for sampling and analysis. Compare this to $5,000-$10,000 per week
for active treatment pilot tests. More detailed descriptions of laboratory, bench, and pilot tests are provided in Gusek (2001).

ADVANTAGES OF SULFATE-REDUCING BIOREACTION

As shown in Figure 4, sulfate-reducing bioreactors can be applied in a number of different AMD/ARD situations. While most passive treatment systems (both aerobic zone and anaerobic zone types) offer simplicity of design and operation and economic advantages over active/chemical treatment, sulfate-reducing bioreactors have advantages worth considering.

- No aluminum plugging
- Can easily handle low flow net acidic water or high flow net alkaline water
- Uses waste organic materials
- Resilient to loading and climate variations
- Consumes sulfate; capable of treating selenium and uranium
- Generates more net alkalinity in effluent
- Burial to minimize vandalism
- Opportunities for community involvement in organic procurement
- Might be able to construct them in abandoned underground mines

Brief discussions of these issues follow.

No Aluminum Plugging

When AMD/ARD attacks clay-bearing formations at mining sites, significant amounts of dissolved aluminum can be created. The geochemistry of aluminum is complex, and this can cause problems in passive treatment systems. The formation of the mineral gibbsite [Al (OH)₃] is especially problematic as it is a gelatinous solid. Gibbsite tends to form in limestone-dominated passive treatment cells, and the sludge tends to plug the void spaces between the limestone rock, becoming a major maintenance problem. While the precise mechanisms are just beginning to be understood (Thomas and Romanek, 2002), the precipitation of gibbsite is avoided in SRB cells. It is suspected that unidentified alternative aluminum compounds form in the SRB cells instead of gibbsite, and these compounds are less prone to plugging. Several case histories of SRB passive treatment projects that involved treating ARD with high aluminum concentrations are provided in Gusek and Wildeman (2002)

Use of Waste Materials in Construction

Organic materials are a key component in the formulation of the substrate of sulfate-reducing bioreactors. Often these materials are considered waste materials and can be obtained for little or no purchase cost. The only expense incurred might be in their transport to the treatment site. If the site is in a remote forest environment, some of the materials such as wood chips and sawdust might be generated onsite or from local sources. A short list of organic waste materials, both solid and liquid, that might be candidates for use in a sulfate-reducing bioreactor is provided below. The list is not necessarily all inclusive as specialty wastes unique to different locales might be available.
• Wood chips
• Sawdust
• Yard waste
• Mushroom compost
• Animal manure
• Partially treated sewage?

• Hay and straw (spoiled)
• Cardboard?
• Waste alcohols including antifreeze
• Waste dairy products
• Sugar cane processing residue
  (Bagasse)

Using liquid organic wastes poses a specific opportunity and challenge. These materials are typically very biodegradable and as such are considered “candy” to sulfate-reducing bacteria. Thus, they are consumed quickly and need to be replenished on a nearly continual basis. This is not consistent with the strict definition of passive treatment cited earlier. However, since these materials might be stored in tanks or fed continuously from offsite sources through pipelines, systems using these waste organic sources would be considered “semi-passive” in nature. Such cells are often called “enhanced sulfate-reducing bioreactors” due to the boost provided by the liquid organic material. When alcohol is the chosen enhancer, the technique has sometimes been called “bugs on booze.”

Resilient to Loading and Climate Variations

If properly designed, sulfate-reducing bioreactors can be resilient to metal-loading variations. Pilot scale tests are the best venue for establishing the expected operating ranges of flow and metal concentrations and the reactions of the SRB cells to those varying conditions. For example, a pilot SRB cell at a lead mine in Missouri was sized for 25 gpm. Once steady state operation was observed for many months, the flow was increased to nearly double the design rate. The SRB cell began to show evidence of stress (i.e., decreased metal removal efficiency) after several months of exposure to the higher flow (Gusek, et al., 1998). Not all SRB cells might be this resilient, but this observation allowed engineers to include a significant factor of safety in the design of the full-scale system (1,200 gpm capacity) at this site.

Low temperature operation is a major concern at some sites, especially in the mountainous states in the west and Appalachia. Pilot cell data at the Ferris Haggerty Copper Mine/Osceola Tunnel Site in Wyoming at elevation 9,500 feet suggests that sulfate reduction rates decline in cold weather, but the decrease is not significant enough to render the design concept untenable. At this site, the typical water temperature is about 4°C. Winter operational data revealed that the cell continued to function at temperatures less than 1°C, and the sulfate reduction rate was estimated to be about 0.24 moles per day per cubic meter (m/d/m³) (Gusek, 2000). Compared to the benchmark design value of 0.3 m/d/m³, this constitutes a 20 percent decrease.

Sulfate-Reducing Bioreactors Consume Sulfate; Selenium and Uranium Reduced

Sulfate is a component of AMD/ARD that may be receiving more regulatory attention. It contributes to the total dissolved solids (TDS) concentration. But unlike other TDS constituents such as sodium, chlorine, and calcium, it is not conservative and can be mitigated in sulfate-reducing bioreactors. No other passive treatment technique has this capability as its primary function. Some sulfate reduction is typically observed in Successive Alkalinity Producing
Systems (SAPS) (see Kepler and McCleary, 1994), but their primary function is to add alkalinity through limestone dissolution.

While sulfate-reducing bioreactors are naturally efficient at consuming sulfate, the geochemical conditions generated in a typical cell are also conducive to reducing selenium from the dissolved state to elemental selenium; this is facilitated by selenium-reducing bacteria. They are also effective in reducing uranium from the oxidized state to form insoluble uranium oxide similar to the way that some natural uranium deposits formed.

Burial to Minimize Vandalism

Any passive treatment system might be a target for vandalism. Because neither plants nor air are required for the sulfate-reducing bioreactors to function, they can be buried beneath a veneer of rock and soil provided that the feed water plumbing to the cell is not compromised. Settlement of the organic substrate needs to be considered in the design if burial is being considered. However, most organic substrate designs typically include a large component of wood chips or sawdust, which do not readily compress under minor surcharge loads developed by soil/rock covers. This aspect of the design should ideally be evaluated at the pilot stage of the design effort.

Underground In-Mine Treatment Systems

As stated above, one of the beauties of SRB systems is that they do not require plants to operate. All that is needed is a carbon source and an SRB arranged in a manner that encourages bacterial growth in concert with managed loading of AMD/ARD. In areas where land surface favorable to passive treatment system construction is at a premium due to steep terrain or the encroachment of civilization, building passive treatment systems in abandoned underground mine voids (using the mine void itself as the containment “vessel”) is an attractive possibility that has been realized in only one study at a metal mine in Montana (Canty, 1999).

Two challenges to overcome to implement this technology include the placement of large volumes of solid organic matter into mine voids through boreholes and the procurement of inexpensive organic material like forestry or paper waste and animal manure (SRB inoculum). The introduction of animal manure (even in small amounts) into ground water (i.e., a mine pool) will be a regulatory hurdle that may prove to be difficult to surmount. Carefully controlled field tests in small mines will probably be required.

Low Flow Net Acidic Water or High Flow Net Alkaline Water

At a given flow rate, the footprint of a sulfate-reducing bioreactor is governed by the mineral acidity of the AMD/ARD. The higher the acidity, the larger the surface area is required per unit of flow. The land area available for the system may be limited, especially for high flows of net alkaline AMD/ARD. In this situation, the surface area of the SRB cell might be as small as 10 square feet per gpm of flow. Thus, a net alkaline flow of 2,000 gpm might require as little as 20,000 square feet or about half an acre of cell. Cell depth will be a function of metal load.
Conversely, a very acidic AMD/ARD source might require a similar area to treat a significantly less flow rate. For example, a flow of only 30 gpm of AMD with over 2,000 mg/L of acidity would require nearly 3 acres of surface area. However, there are no other technologies capable of passively treating AMD/ARD with this aggressive a chemistry.

**Added Net Alkalinity in Effluent**

Sulfate-reducing bioreactors are typically sized to deliver treated water with low concentrations of metals and a near neutral pH. However, experience has shown that SRB cell effluents typically contain excess alkalinity at concentrations above those expected from SAPS units or anoxic limestone drains. This excess alkalinity is therefore available to ameliorate acidity contributions that might be impacting the receiving stream far removed from the original passive treatment site.

**New Opportunities for Community Involvement**

The construction of passive treatment systems is an ideal way to make the most of community volunteerism. The transplanting of wetland vegetation is the most common activity in which volunteers can become involved with passive treatment projects. However, the collecting of organic materials for sulfate-reducing bioreactor substrate opens an entirely new opportunity for local community organizations to release pent-up volunteerism. Some pet owners are often hard pressed to find useful and environmentally sound ways to dispose of significant amounts of manure (e.g., horse). Homeowners could divert tree trimmings or yard waste away from the local landfill and into a community stockpile of wood waste to be mulched (but not composted) and used in a nearby sulfate-reducing bioreactor. Farmers would have a place to dispose of moldy hay. Community events similar to paper drives could be used to collect materials in advance of a project. This not only lowers the cost of the project but also provides additional community buy-in.

**SULFATE-REDUCING BIOREACTOR DESIGN EXAMPLES**

**Design Example No. 1**

This is a hypothetical abandoned underground coal mine in Appalachia with a relatively small mine pool. The site is adjacent to a fresh water lake. The flow varies through the year, but the AMD chemistry is fairly constant. SAPS had been considered at this site but rejected due to the elevated aluminum concentration. Pertinent design parameters are listed below.

- 67 gpm peak flow
- pH = 2.5
- Fe = 152 mg/L (ferric iron)
- Aluminum = 30 mg/L
- Acidity = 500 mg/L
- 990 moles of Fe per day
The hypothetical passive treatment system will include two sulfate-reducing bioreactors (each treating 50 percent of the flow) to raise the pH, produce net alkalinity, to remove nearly 100 percent of the aluminum and 95 percent of the iron. The system would comprised the following components:

- 1.7 acres of SRB cell 3 feet deep
- 0.25 acres of aerobic polishing cell

The costs of developing this design from initial concept to complete construction include:

- $30,000 to $50,000 for bench and pilot studies
- $315,000 design and construction (assuming no donated materials or labor)

The 8,250 cubic yards of organic substrate originally installed would require replacement every 20 to 30 years. The substrate typically comprises about 33 percent of the construction cost. This would be about $110,000 or less depending on the availability of local materials and in-kind donations. This and other maintenance costs are summarized on an annual basis in the table below. Some of these costs might be minimized through volunteer labor and other contributions.

<table>
<thead>
<tr>
<th>Maintenance Item</th>
<th>Annual Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Replace Substrate</td>
<td>$3,569</td>
</tr>
<tr>
<td>Dispose Substrate (20% of replacement cost.)</td>
<td>$714</td>
</tr>
<tr>
<td>Weekly inspection &amp; pipe clean?</td>
<td>$5,000</td>
</tr>
<tr>
<td>Flushing for aluminum buildup</td>
<td>$0</td>
</tr>
<tr>
<td>Sampling/lab costs lump sum</td>
<td>$15,000</td>
</tr>
</tbody>
</table>

The life cycle cost of this treatment (includes capital and operating cost) is about $0.70 per thousand gallons treated.

**Design Example No. 2**

This is another hypothetical abandoned underground coal mine in Appalachia but with a relatively large mine pool covering over 100,000 acres. The site contributes nearly 50 percent of the metal load to a nearby river. The flow is relatively steady through the year, and the AMD chemistry is constant as well. The site has only 6 acres available for construction of a main treatment system, but there are no restrictions on effluent polishing. This is a major project due to the flow rate. Pertinent system design parameters are listed below.

- 3,000 gpm from a deep mine pool
- Sulfate = 1000 mg/L (50 effluent goal)
- pH = 5.5
- Fe$^{2+}$ = 150 mg/L
- Al = 2 mg/l
• Mn = 2.7 mg/L (0.05 effluent goal)
• Acidity = 50 mg/L ("Hot Acidity")

The 6-acre restriction eliminates a standard sulfate-reducing bioreactor. However, an enhanced sulfate-reducing bioreactor (ESRB) is feasible due to the steady availability of a waste alcohol product and other factors. The enhancement allows the footprint of the ESRB cell to shrink to easily fit in the space available. The ESRB effluent will have a neutral pH and some excess alkalinity. However, it will also have elevated biological oxygen demand (BOD) and manganese, which require further polishing. Key features of this hypothetical system include:

• 4 acres of enhanced sulfate-reducing bioreactor cell 6 feet deep
• 9 acres of aerobic polishing cell (for Mn and BOD treatment)

The costs of developing this design from initial concept to complete construction include:

• $200,000 for bench and pilot studies
• $1.36MM design and construction

The operating cost of the enhanced sulfate-reducing bioreactor (including paying $2.00 per gallon for the alcohol) is $674,000 per year or $0.43 per 1000 gallon treated. The system effluent would meet drinking water standards. To be conservative, the above cost assumes that the substrate in the ESRB be replaced every 20 to 30 years due to metal sulfide precipitate buildup.

**Design Example No. 3 - Do SRBs Need More Room?**

This design example compares the area requirements for using a standard aerobic wetland and a standard sulfate-reducing bioreactor to treat a relatively large net alkaline flow. The design assumptions are listed below.

• 3,000 gpm from a deep mine pool
• pH = 6.5
• Fe+2 = 50 mg/L (817,560 grams/day or 14,638 moles per day)
• Net alkaline
• No manganese
• 10 acres available for main treatment cells

If an aerobic wetland dominated by cattails and other vegetation was designed on the standard assumption of 11 grams/day per square meter of iron loading criteria (which was established by U.S. Bureau of Mines researchers), approximately 18 acres of wetland habitat would be needed.

A sulfate-reducing bioreactor with an identical treatment capacity would cover 8 acres (probably split into four 2-acre cells plumbed in parallel). The cells would be 7.5 feet deep, and the AMD/ARD would enter them at the bottom and exit at the top. This upflow configuration allows the top of the cell to function as a primary dissolved oxygen polishing cell. The remaining
2 acres available would be fitted with a final aerobic polishing cell to complete the facility. In this situation, both cell types would work geochemically, but only one – the sulfate-reducing bioreactor – would be feasible.

SUMMARY

Sulfate-reducing bioreactors are not the only type of passive treatment technique available to the design engineer, and they are not applicable in every situation. However, they can handle a wide variety of flows and AMD/ARD chemistries in hostile cold climates, and they can treat aluminum-bearing AMD/ARD without plugging. Furthermore, they can generate excess alkalinity in their effluent that further enhances the quality of the receiving stream.

Sulfate-reducing bioreactors typically require large amounts of organic materials that are usually considered waste. Enhanced SRB cells can consume liquid organic wastes like antifreeze or cheese whey.

While not readily practiced, it may be feasible to build them in mine voids to provide in situ treatment at sites with limited land area.

LITERATURE CITED


Appendix B
Overview Papers

Pamela Milavec and D. Seibert
2002
Pennsylvania’s Efforts to Address Operation, Maintenance and Replacement of AMD Passive Treatment Systems
ABSTRACT

An increasing number of watershed groups, as well as many Federal, State and local agencies, have become active in watershed restoration over the past several years. As a result, a large number of restoration projects are being funded and constructed. In Pennsylvania, a total of nearly $93 million of public money has been spent on all types of watershed restoration projects since 1988. A portion of this funding has gone to construct 153 AMD passive treatment systems statewide. Funds have come from a variety of sources, including the Office of Surface Mining's Title IV program, the Natural Resources Conservation Service's (NRCS) PL-566 program, Pennsylvania's Growing Greener program and the Environmental Protection Agency's 319 Non-Point Source program. Water quality and aquatic habitat improvements are occurring as these projects are implemented. The need for long-term operation, maintenance and replacement (O, M & R) has been increasingly recognized as a requirement to ensure the success of watershed restoration projects. The failure to maintain the systems being constructed could have detrimental impacts to watersheds that are beginning to support an increasing number of stream uses. As a result of growing concern over this issue, the PA Department of Environmental Protection (DEP) established a workgroup to provide recommendations to address this need. The workgroup consisted of individuals from Federal, State and local governments, as well as private consultants and watershed group officials. All had extensive experience in the operation and maintenance of watershed restoration projects, both mining and non-mining related. Recommendations were finalized by late 2001.

The NRCS and DEP have taken a lead role in implementation of these recommendations, including the development of maintenance plans and agreements, working with local watershed groups to provide routine maintenance and trouble-shooting to solve problems that arise. Their experiences are providing a greater understanding of the efforts needed to keep systems functioning properly.

INTRODUCTION

In Pennsylvania, watershed groups and various government agencies started constructing passive treatment systems to treat abandoned mine drainage (AMD) in the early 1990's. The design of these systems has evolved from simple, aerobic wetlands to complex vertical flow alkalinity generating systems with mechanisms to flush accumulated metals. While the design improvements have resulted in the ability to treat highly acidic discharges, the resulting systems have required significantly more effort to keep them operating effectively. Early vertical flow systems that did not have adequate flushing mechanisms have started to fail due to plugging by metals. Additional problems have surfaced at older systems, including leaks, short-circuiting
and metals accumulation. All these experiences have pointed to the need to address long-term operation, maintenance and replacement (O, M & R) of AMD passive treatment systems.

The need to address O, M & R became even more pressing as additional funding was made available to build these types of systems. Early on, funds were provided by the EPA 319 Non-Point Source program, the Natural Resources Conservation Service (NRCS) Rural Abandoned Mine Program and P.L. 566 program and the Federal Office of Surface Mining's Ten Percent Set Aside and Appalachian Clean Streams Initiative (ACSI) programs. These funds continue to increase, while significant additional funding has surfaced with the establishment of Pennsylvania's Growing Greener program in 1999.

In June 2000, State Representative Sam Smith provided remarks at a statewide AMD conference that emphasized the need to address O, M & R in watershed restoration work. Many others working in the AMD treatment field were recognizing the same need. As a result, the PA Department of Environmental Protection (DEP) established a workgroup in early 2001 to develop recommendations to address this problem. The workgroup met throughout 2001 and finalized recommendations to DEP Secretary David Hess in November, 2001. These recommendations are provided in Attachment A. Since that time, much effort has gone into implementation of the recommendations. The DEP and NRCS have been leaders in that effort, along with several strong watershed groups and knowledgeable consultants.

IMPLEMENTATION

A major focus of implementation of the workgroup's recommendations has been the educational aspect. Members of the workgroup spoke at various conferences and Growing Greener training workshops in order to develop an awareness of the need to address O, M & R. Also, several changes were implemented in the Growing Greener program (the single most significant funding source) as a result of the recommendations. They included: developing an O, M & R fact sheet, providing detailed information in the application packets about the need to address O, M & R in the application, changing project score sheets to emphasize the need for O, M & R on implementation projects, and requiring the development of an O, M & R plan for all construction projects. In addition, a separate funding category has been developed to address funding of O, M & R projects with significant costs, starting with the 2003 round. The State Legislature passed Legislation in late June that continues Growing Greener through 2012 and provides a permanent funding source for the program.

The development of O, M & R plans has become a major focal point for projects constructed by the NRCS, under their P. L. 566 program, and the DEP's Bureau of Abandoned Mine Reclamation (BAMR), which uses OSM and Growing Greener funds to construct treatment facilities. These plans emphasize the partnerships with local watershed groups that are needed to provide for necessary O, M & R. O, M & R plans are developed for all NRCS projects and all new BAMR projects. The NRCS requires local sponsors in counties where the projects are located to accept the responsibility for all O, M & R needs. Operation and maintenance agreements and plans are prepared for the sponsors. NRCS staff provides consultation and technical input when significant maintenance is needed. BAMR looks upon the local groups to provide for routine monitoring, operation and minor maintenance requirements. Training is provided to the local groups where needed. BAMR is responsible for more significant maintenance needs and for eventual system replacement.
A major issue with regard to O, M & R is monitoring, particularly water sampling and laboratory analysis. Monitoring of the treatment system efficiency is very important – it provides feedback for future design improvements and signals when systems are not operating properly and may need maintenance. Typically, local groups and BAMR staff collect the samples and DEP's laboratory provides the analysis. This has proven to be a significant portion of the expenses involved with O, M & R. Determining how best to deal with this cost is an ongoing issue within DEP that has yet to be resolved. The DEP has recently convened a small workgroup to address this.

TECHNICAL ISSUES

A number of difficult technical issues have arisen with regard to the long-term operational efficiency of AMD passive treatment systems. The most complex issue deals with metals precipitation within the systems, particularly vertical flow systems that treat net acid water. Vertical flow systems typically consist of a layer of standing water over organic material (usually mushroom compost), which is underlain by limestone with a pipe collection system below the limestone. In theory, vertical flow systems are expected to keep ferrous iron in the ferrous state and to reduce existing ferric iron to a ferrous state as a result of a lack of oxygen in the system (the compost layer strips oxygen present in the water as a result of the biological oxygen demand of the compost). Ferrous iron is expected to pass through the limestone and precipitate in a subsequent settling basin. Aluminum, which comes out of solution as the pH increases, whether or not oxygen is present, is expected to precipitate within the compost and/or limestone layer. Flushing is expected to remove precipitated aluminum. However, field evidence indicates that most systems retain some iron within the limestone. In a few cases, iron has precipitated on top of the compost layer, causing the system to plug. While flushing these systems provides visual evidence of both iron and aluminum removal from the systems, a recent study has indicated that only a small percentage of retained metals is being flushed from the system in two systems studied (Watzlaf, 2001). The long-term implication of this is unknown, but efforts are underway to determine the best way to design the flushing systems to maximize removal from the limestone.

A less frequent problem occurs when ferric iron precipitates on top of the compost in vertical flow systems, as the pH starts to increase. When this occurs, it eventually reduces the permeability of the compost until water cannot flow vertically through the system. Since the solubility of iron is pH dependent and precipitation occurs more rapidly as the pH increases (Hem, 1992), the pH of the raw water that is being treated must be carefully considered when designing systems with high iron levels. Creating larger precipitation ponds and wetlands before the vertical flow systems can reduce the amount of ferric iron reaching the vertical flow system.

Accumulation of aluminum in the top 6 inches of the rock layer in vertical flow systems can create hydraulic conductivity problems within the limestone. As mentioned earlier, oxygen is not needed to precipitate aluminum - there only needs to be an increase in pH to above 4.5 (Hem, 1992) for precipitation to occur. In many systems, this pH increase occurs in the upper areas of the limestone rock column, or even within the compost. The initial precipitation of aluminum creates a very loose, jelly-like precipitate \((2\text{Al(OH)}_3)\) that is easily dislodged and flushed from the rock. This precipitate is easily flushed from the system when aggressive flushing systems are initially designed in the system. Frequent flushing is recommended, at least quarterly, to keep the aluminum purged from the system. Field observations indicate that if the
aluminum is allowed to accumulate in the rock to point where hydraulic conductivity is reduced, the flushing of aluminum becomes more difficult. Also, aluminum precipitate seems to harden and take on a paste consistency with time and becomes harder to flush.

Other problems encountered with vertical flow systems include short-circuiting caused by the development of preferential flow paths through the compost layer. This has especially been evident on several systems where the compost was washed out immediately below the influent point to the system. Dispersing the inflow through a manifold rather than at a single, point-source location seems to help this. The configuration of the piping system underlying the limestone rock can also impact short-circuiting through the system. Long narrow systems that have continuous pipes running along the longitudinal axis of the system seem to short circuit through the piping system. This apparently results from the water entering the piping near the influent end of the system and traveling through the piping rather than through the limestone. Segmenting can alleviate this, or zoning the piping system by designing incremental breaks in the pipes to force the flow of water into the limestone (Danehy et al., 2002). Consideration of a system with a lower length to width ratio during the design phase also may be important in reducing short-circuiting.

The most frequently used source of organic material in vertical flow wetlands is spent mushroom compost. The function of the compost is to reduce ferric iron to ferrous iron and provide a medium for biological activity. When new systems are first put into service the mushroom compost can create very high biological oxygen demands (BOD) on the stream receiving the treated water. If aquatic life is present in the stream at the time the system goes on line, the BOD may kill much of the aquatic life present in the stream. To manage this potential problem, it is best to install a valve on the inflow pipe to the system and limit the amount of water entering the system. This allows for a slow flushing of the BOD out of the system and prevents an aquatic kill on the receiving stream.

OVEN RUN CASE STUDY

Stony Creek is a historically AMD impacted watershed located primarily in Somerset County, PA. The first major source of AMD to Stony Creek was a tributary known as Oven Run. The Oven Run AMD Abatement Project was developed under an NRCS PL-566 plan, the first in the country approved to address AMD. Six project areas were identified for abatement in the plan. Five of the six project sites required passive treatment using vertical flow systems. With completion of the first five projects by the fall of 1999, the Stony Creek turned from net acidic to net alkaline for a distance of 22 miles. The project has been very successful and has restored a recreational fishery in a stream long considered to be dead. The amazing success of this project has occurred even though three of these systems have had O & M related problems, which will be discussed below. Although the Somerset County Commissioners signed an agreement to be the responsible O & M entity under the PL-566 Plan, both BAMR and the NRCS have taken the lead role in addressing the technical problems that have arisen.

Oven Run Site A, the sixth and final site to be constructed, was just completed in the spring of 2002 using NRCS and Growing Greener funds. It consists of a passive treatment system that incorporates many design features developed as a result of lessons learned from the sites constructed earlier. Specifically, aerobic wetlands and limestone filter dams were used to remover as much ferric iron as possible prior to the vertical flow wetland. Also, an aggressive flushing system was constructed. Initial indications are that this site is functioning as designed.
Routing water sampling of the system is planned starting in the fall of 2002, after the system has stabilized.

Oven Run Site B was completed in 1999 by BAMR, using OSM ACSI and Ten Percent Set Aside funds. BAMR has continued to provide all monitoring, operation and maintenance at this site, although turning routine operation and maintenance to the Somerset Conservation District is a future possibility. Site B has a design flow of 350 gpm, although the system has successfully treated flows as high as 450 gpm. The influent has acidity between 500 and 600 mg/l, total iron between 55 and 85 mg/l and aluminum between 35 and 45 mg/l. The system consists of a deep mine drainage collection trench, a vertical flow wetland, settling pond, second vertical flow wetland and second settling pond. Until spring of 2000, the system effluent was net alkaline, with very low metals concentrations. After flows increased in the spring of 2000, effluent quality degraded to the point that net acid water was being discharged with a pH in the low 4’s and elevated iron and aluminum. This situation continued, even after flows subsided. Site inspectors had observed an opening in the compost directly below the point of influent discharge into the first vertical flow system. BAMR’s construction crew was brought in to construct an inflow manifold system to spread influent over a larger area and prevent the development of preferential flow paths through the compost. The expected water quality improvements did not occur, although this vertical flow system is generating about 200 – 250 mg/l of alkalinity, which is about all that can be expected. Now, concerns are being directed to the second vertical flow pond, which has steadily lost the ability to generate the additional alkalinity needed to fully neutralize the raw water. Both vertical flow systems are scheduled to be drawn down in the summer of 2002 to evaluate the development of preferential flow paths and take remedial action. One possibility is that the very aggressive flush system has actually pulled compost down into the limestone, causing the compost layer to be too thin above the pipes and allowing the development of preferential flow paths. The aggressiveness of the flush system is due to the large head differential between the system and the flush discharge point and the separation of the flushing system into 3 cells, allowing greater velocities when each is flushed individually. There is no evidence of metals plugging in either vertical flow wetland and the system continues to remove over 90% of the iron, 70 to 80% of acidity and about 60% of the aluminum (see Figure 1).
Oven Run Site C was completed in 1997 and consisted of backfilling a 5,000 foot long highwall using OSM Title IV funds. A small, but highly acidic, discharge from the open pit was largely eliminated. A minor seep that remains is directed to the Oven Run Site B system. Since backfilling this highwall, which was hydrologically connected to the deep mine discharge at Site B, maximum flows at Site B have been no more than half the flow measured prior to completion of this project. This flow reduction is a very important secondary benefit to completion of the backfilling project.

Oven Run Site D was completed in September of 1995 by NRCS, using PL-566 and section 319 funds. Site D treats approximately 100 gpm of moderately acidic deep mine drainage. Since beginning operation as the first completed system on Oven Run, the system worked well. In sequential order the system is composed of an initial settling pond, aerobic wetland, vertical flow wetland, settling pond, aerobic wetland, vertical flow wetland and a final settling pond. A rise in the water level in the first vertical flow wetland was noticed after the second winter of operation.

A limited flushing system in this system allowed for some flushing of accumulated metals from the system. Initially, the limited flush lowered the water level back to original levels. In subsequent years, the water level raised every winter when flow rates increased. Each time the system was flushed, the effectiveness of the flush decreased. A layer of iron was accumulating on top of the compost in the vertical flow ponds that limited the flow through the system. The system continued to treat the water flowing through it, but the quantity of water flowing through the system was controlled by the permeability of the iron on top of the compost. In the winter of 1999, some of the water began to flow through the emergency spillway and was not treated in the vertical flow wetland. This condition continued through the spring of 2002. At this point in time, approximately 50 GPM would pass through the system; any flow above this amount would bypass the system through the emergency spillways.
To remedy this situation, the Somerset County Conservation District applied for and received a DEP Growing Greener grant in the spring of 2002. To limit the iron accumulation on the compost, 3 rock filter dams were placed in the wetlands preceding the vertical flow ponds. The rock filter dams will eliminate short-circuiting of flow through the wetland and will increase the detention time in the wetland. An aggressive bottom flushing system was also installed to eliminate inert materials left behind by limestone dissolution and to remove metal accumulation in the rock layer. As part of the bottom flushing system, piping was extended to the surface of the compost and capped with threaded pipe caps. These vertical pipes will act as ports to flush accumulated iron from the surface of the compost. The caps will remain on the vertical pipes during normal operations of the system and be removed prior to flushing for surface iron removal.

Oven Run Site E was completed in 1997 with NRCS funds, and treats two moderately acidic deep mine discharges with a configuration very similar to Site B (two vertical flow wetlands and two settling ponds). Plugging of the vertical flow wetlands started to occur relatively soon after commencement of operation, which was very puzzling to the system designers. The problem was eventually determined to be the result of the limestone used in the vertical flow wetlands. While technical specifications called for a required calcium carbonate percentage, they did not specify rock meeting soundness standards as specified by the PA Department of Transportation (PA DOT, 2000). Initially, the rock produced good water quality, but the rock quickly began to deteriorate. After 4 years, hydraulic conductivity through the system was lost to the point that the system is not providing treatment. Designs have been prepared to replace the degraded rock and install an aggressive flushing system. Late summer of 2002 construction is planned for the upgrading of this system.

Oven Run Site F was completed in September of 2000 with NRCS funds. The design of this system was completed with much more knowledge about the operation and maintenance concerns of vertical flow wetlands. Experience at site D taught us that routine flushing from the bottom of the vertical flow wetland was necessary to keep the systems functioning over time. This system treats 300 GPM of acid mine drainage and consists of a vertical flow wetland, settling pond, vertical flow wetland and a final settling pond. A very intensive flushing system was designed for the vertical flow wetlands at this site. The vertical flow wetlands have a grid of 6 inch, perforated pipes on 11 foot centers, under the limestone. These pipes outlet to two, 12 inch pipes that are at opposite ends of the vertical flow wetland. The flow out of the system when these two pipes flow is approximately 5000 GPM per pipe, at the beginning of the flush. Experimenting with the sequencing of opening valves and the duration of flushes has provided some empirical observations of the flushing events at this site.

Observations were made while opening only one valve at a time, with alternate opening and closing, after 30 minutes of flow, and opening both valves at the same time until the vertical flow wetland was drained. With both flushing scenarios, the flush water was initially very turbid. With the alternate opening and closing of valves, the effluent water began to clear and become less turbid after 30 minutes of flushing. The amount of turbidity decreased with each successive alternate opening and closing of valves. When both valves were opened simultaneously and the water was allowed to flow until the system was drained, the effluent water remained very turbid during the entire time of the flush, approximately 3 hours. Since this system was constructed with an aggressive flushing system in 2000, quarterly flushing has been completed. There has been no indication of rising water levels in this system as were noted at Site D, where only a limited flushing system was designed. These empirical observations need
to be followed by more controlled flushing experiments that involve water quality testing during
the flush to calculate quantities of metals flushed from the system.

CONCLUSIONS

The future of passive treatment of AMD is dependent upon resolution of the long term O,
M & R issues related to these systems. Implementation of the DEP Workgroup
recommendations is underway and starts to resolve these issues, particularly from an
administrative standpoint. However, resolution of many of the technical concerns may be more
difficult. One of PA’s leading AMD treatment consultants, Hedin Environmental, received a
Growing Greener grant to evaluate design and operation of flushing facilities. This work was
initiated with a workshop attended by a number of government and consulting personnel to
gather and organize existing data on vertical flow systems in order to design the most efficient
systems (Workshop Proceedings, 2002). The technical issues fall under several general
categories that are discussed below.

Use of Mushroom Compost:

The use of mushroom compost in vertical flow wetlands to promote biological activity
and facilitate the conversion of ferric iron to ferrous iron is a widely accepted practice (although,
more recent thinking is that this conversion is not a complete one; ferric iron is observed being
flushed from these systems). Mushroom compost is also used in anaerobic wetlands to
encourage sulfate reduction reactions and in aerobic wetlands to promote plant growth. When
AMD systems are constructed using mushroom compost, sample analyses have shown a
significant biological oxygen demand (BOD) present during the first week to 3 months the
system is in operation. This BOD can present devastating impacts on receiving streams if any
aquatic life is present. Frequently, the acid water being treated has already devastated aquatic
life to the level that there is little need for concern. However, in situations where aquatic life is
present or where there are downstream municipal water supplies, the management of the outflow
of BOD from the system is imperative. The simplest method to control the initial flow of BOD
is to strategically place piping and valves in the system so that the majority of the AMD can be
bypassed around the system while a small flow is allowed to move through the system. The
small flow will, with time, reduce the BOD to levels that are not harmful to aquatic life. Once
this condition is reached, the flow of AMD through the system can gradually be increased until
all of the AMD is flowing through the system. Recent trends have been toward a reduction in
compost from early designs, where compost thickness was as high as two feet. Experience on
the NRCS Oven Run sites indicates that 6 inches of compost is adequate to promote biological
activity and facilitate conversion of ferric iron to ferrous iron. Larger quantities of compost take
longer to dissipate the BOD and long-term odor problems can be a concern in residential areas.
Another consideration, however, is that too little compost may contribute to short-circuiting, as
well as possible rapid depletion of fine limestone within the compost and a possible loss of
biological activity once the pH drops. BAMR has recently dropped compost specifications from
2 feet to 1 foot to reduce BOD generation while still providing adequate compost to address the
other concerns.
Short Circuiting:

In Oven Run, we have learned that the flow through systems with under draining flush capabilities may short circuit in a number of ways. Preferential flow patterns may develop as a result of shifting compost below inflow points and possibly even above flush pipes on systems with aggressive flushing. Other short circuiting may occur directly through the flushing pipes. In long, narrow systems, with flush piping extending the full length of the system, there is a high potential for water to enter the flush pipes at the inflow end of the system and travel through the pipe instead of through the rock. This short-circuiting does not allow adequate detention time in the limestone rock for treatment of AMD. Short-circuiting can be reduced using a couple different methods. Distribution manifolds that evenly distribute the water throughout the vertical flow wetland prevent water from entering at one point and entering the flush system. This also keeps compost from being shifted immediately below the inflow point. Another method of preventing short-circuiting is to incrementally cap the longitudinal flush pipes so water is forced to leave the flush pipes and flow through the limestone rock (compartmentalizing the flush systems). On larger flow discharges, the use of both of these techniques is recommended to maximize water to rock contact. Minimizing the length to width ratio also should be considered during design.

Flushing:

Experience in Oven Run and other watersheds over the last seven years has clearly shown that vertical flow wetlands treating highly mineralized AMD that are not designed and operated with an aggressive flushing system will experience plugging with time. In vertical flow wetlands that treat acid mine drainage with dissolved iron and aluminum, the accumulation of precipitated metals in the rock will fill the rock voids, eventually plugging the system, and cease to treat the mine drainage. Bottom flushing of vertical flow wetlands has proven to be an effective way of removing accumulated precipitates from these systems. Long duration, high volume flushing performed on a regular schedule has maintained hydraulic conductivity through the Oven Run Site F system. The system has continued to produce high quality water since construction. Figure 2 illustrates how the water quality improves as it moves through the system.

Some system designers have recommended limiting the number of holes in the perforated pipe to increase the velocity of flow through each perforation in the pipe. The number of perforations in the pipe is the limiting factor determining the quantity of water leaving the system. There are some concerns with this type of design. One concern is that the distance between holes in the pipe becomes too great and the area of influence of the perforation is less than the distance between perforations. In this case, there potentially could be “dead areas” between perforations where little flow occurs, which would eventually cause the rock to plug with metal precipitate. Another concern is that limestone rock placed directly on the perforated pipe could greatly reduce the capacity of the perforations if a rock lodges in a perforation. If this happens, the distance between perforations becomes even greater, increasing the potential for "dead" areas. Other designs currently being evaluated include those using multi-tiered flush piping and those that do not use compost at all. Determining the best design for these systems is an ongoing process. Hedin Environmental has retained an engineer to evaluate current system designs and make recommendations for future designs. A draft paper, currently being circulated for comment, recommends increasing flush velocities to assist in the removal of retained solids by dividing the under drain systems into multiple cells and designing the header pipes for gravity flow to provide for even flow distribution (Langese, 2002).
Rock Quality:

The quality of the limestone rock used in vertical flow wetlands is somewhat dictated by the local commercially available sources of limestone. In western Pennsylvania, there are several sources of limestone that perform well over long periods of time. It is important to specify stone that is durable and will not deteriorate when exposed to acid. Specifying rock that meets the soundness standards specified by the Pennsylvania Department of Transportation (PADOT) should ensure the integrity of the system with regard to ability to transmit water. Early in the implementation of the Oven Run Site E project, the soundness of the limestone was not specified by NRCS. Rock was provided that met the specified chemical standards, but was not from an approved PADOT quarry. This is believed to have caused the system to fail.

It is also important to specify a minimum of 80% calcium carbonate as determined by ASTM C-114. It is important to note that calcium carbonate equivalent (CCE) is not specified but elemental calcium carbonate is. Also, the maximum content of elemental magnesium should not exceed 2% and inert material (e.g. silica) should be less than 15%.

The above discussions should help to emphasize the need to have a knowledgeable, reliable entity to provide for long-term O, M & R. The failure to do so will result in the eventual failure to adequately treat AMD and the loss of millions of dollars worth of public investment. Gains in the restoration of aquatic habitat will also be lost. The design of these systems is expected to continue to evolve as more is learned about long-term operation. Passive treatment
of AMD is expected to be an important aspect of watershed restoration into the foreseeable future. O, M & R improvements will ensure the continued success of this restoration work.

REFERENCES


An increasing number of watershed groups, as well as many Federal, State and local agencies, have become active in watershed restoration over the past several years. As a result, a large number of restoration projects are being funded and constructed. Water quality and aquatic habitat improvements are occurring as these projects are implemented. The need for long-term operation, maintenance and replacement (O, M & R) has been increasingly recognized as a requirement to ensure the success of watershed restoration projects. In Pennsylvania, a total of nearly $93 million of public money has been spent on these projects since 1988 (see attachment A). The establishment of the Growing Greener grant program has greatly accelerated this effort. The failure to maintain the systems being constructed under these projects could have detrimental impacts to watersheds that are beginning to support an increasing number of stream uses. As a result of growing concern over this issue, the Department's Greener Team established a workgroup to provide recommendations to address this need. This workgroup consists of individuals from Federal, State and local governments, as well as private consultants and watershed group officials. All have had extensive experience in the operation and maintenance of watershed restoration projects.

It is important to note that, for the purpose of defining needs and determining costs, the workgroup defined long-term O, M & R as system operation and maintenance, plus one system replacement at the end of the design life of the project. Also of note is a decision by the group to include all publicly funded watershed restoration projects constructed through 2001, when determining costs associated with existing systems.

**O, M & R Plan**

An O, M & R Plan, developed by the project sponsor, is an integral part of providing for operation and maintenance of watershed restoration projects. The basic elements of an O, M & R plan include: a written agreement with the entities responsible for O, M & R, identification of tasks to be completed, development of a schedule and determination of responsible parties and costs. Plans must become a "deliverable" of all new implementation grants. For existing projects that have no O M & R plan, site-specific plans will need to be developed prior to receiving funds to address O, M & R.

Operation, maintenance and replacement concerns should begin at the initial site inventory of a project and continue through all phases of project development. Water quality information, along with flow measurements, should be looked at critically with respect to future operation and maintenance. If a site requires intense operation and maintenance to function, the sponsors of
the project need to understand the intensity and potential cost. Once the decision is made to move ahead with project design, the focus should be to make the operation and maintenance of the system as easy as possible. Prior to project implementation, the sponsors need to understand what it is they need to do and at what frequency. Additional training may need to be provided to facilitate a more detailed understanding of operation and maintenance.

In developing an O, M & R plan, the following should be considered:

**Operations** - Sponsors need to demonstrate an understanding of, and the ability to perform, routine duties, such as:

- Inspections (including water sampling and flow measurements);
- Litter control;
- Vegetation control;
- Mechanical maintenance (including flushing);
- Insect and vector control;
- Physical stability and erosion control.

**Maintenance** - Sponsors need to demonstrate an understanding of, and the ability to perform, more intensive items that may take considerable dollars and time to accomplish, such as:

- Removal and disposal of accumulated precipitate or sediment;
- Maintenance of channels;
- Industrial cleaning of pipes;
- Repairing damage after major storm events;
- Repairing cracks or leaks;
- Adding limestone, compost, sand or gravel;
- Repairing vandalism damage;
- Adjusting grade or outlet structures.

**Replacement** - Systems have a designed life expectancy; once that design life is exceeded, much of the system will need to be recharged or replaced. Replacement will involve much of the same effort originally needed to construct the system. Changes in technology and water quality and quantity will need to be considered to determine if the size and/or design of the system must be changed. Replacement considerations include:

- Estimating BMP (Best Management Practice) design life;
- Determining replacement responsibility, including a successor, in the event of the original project sponsor's inability to carry out these responsibilities;
- Determining approximate costs for the following possible needs: removing accumulated sediments, replacing defective valves, water control structures, re-sizing the system to accommodate changed water quality or quantity, recharging organic matter layer on wetlands, recharging limestone rock.

An O, M & R Plan should include:
- Narrative describing O, M & R needs and identifying responsible parties
- Signed maintenance agreement with all parties, including property owners
- O, M & R Site Map that includes BMP's, flushing points, monitoring points (water sample locations, benchmark cross sections, etc.)
- Site specific instructions
- "As-built" plans

Long-term Cost Analysis

Long-term costs are analyzed in many business and government applications. The starting point for most analyses is a spreadsheet that projects costs over the lifetime of the BMP. Costs are often divided into tasks such as site inspections, sample collection, sample analysis, sludge management, flushing, and reconstruction. A long-term cost spreadsheet should be developed for all projects early in the planning process. The construction of this spreadsheet will help sponsors to recognize long-term responsibilities and also encourage them to identify mechanisms that will legitimately lessen the long-term costs of their projects.

The workgroup collected information on long-term costs of BMPs by reviewing existing policies, interviewing technology experts, and by analyzing current cost data. The workgroup did not find established O, M & R cost estimates for mine drainage treatment systems, so estimates were developed from Department, Natural Resources Conservation Service (NRCS), non-profit and consultant experiences. These data were used to develop spreadsheets that projected long-term O, M & R costs for specific mine drainage BMPs. Development of the spreadsheets required assumptions about the time period over which to project costs and whether to include a BMP replacement in the extrapolation. The workgroup decided to analyze 25 years of costs and to include one replacement in the calculation.

The spreadsheets were analyzed in two ways. The first method was to calculate the present value of the long-term costs. The method requires financial assumptions about rates of inflation and investment return. The result of this calculation is a sum of money that, if the financial assumptions are realized, will yield proceeds adequate to cover all anticipated long-term costs. Our analyses assumed a 3% inflation rate and a 6% rate of return, or a net rate of return of 3%. While this may seem conservative, it is consistent with long-term economic trends in the U.S. It is also consistent with similar analyses of long-term AMD treatment costs being conducted by the Department for permitted mines. The present value analysis yields a sum of money that can be placed into perspective by comparing it to the BMP’s original construction cost. On average, the present value of the long-term O, M, & R costs were approximately 60% of the construction costs. Thus, if the Department wanted to fully fund a $100,000 passive treatment project, it should plan on placing $60,000 in an interest-bearing account at the time of construction. If the cost projections and financial assumptions are correct, no more funds should be required for 25 years.

A second analytical method calculated the annual costs of an on-going O, M & R program. Instead of paying all the anticipated long-term costs in the first year, only those expenses anticipated for the current year would be paid. Each year, for 25 years (the workgroup’s
analytical timeframe), the annual O, M & R costs would be paid. We calculated the average annual cost by summing all anticipated O, M & R costs and dividing by the analytical period (25 years). This average O, M & R cost was related to the construction cost to calculate the O, M & R factor. On average, most AMD passive treatment technologies had an average annual O, M & R factor of 4%. Thus, a $100,000 project would require an average of $4000 per year in annual costs. The actual costs would vary widely because major maintenance costs and replacement costs – both high expense items – occur infrequently and generally toward the end of the BMP lifetime. The factor does not account for inflation. Since most watershed restoration projects have been constructed relatively recently, not accounting for inflation shouldn't be a problem at this time. If the Department decides to fund O, M & R using the O, M & R factor, it should regularly adjust the base value of construction to account for inflation in the future.

The estimated O, M & R factors varied with the type of watershed restoration project. Table 1 shows the range in factors. Most of the BMPs being implemented with Growing Greener funds are in the 3-5% range. While the AMD factors were based on actual experiences of several workgroup members, the non-AMD estimates were derived from informal surveys of the following sources: NRCS, the Center for Watershed Protection, the Keystone Stream Team, PA DEP, MD DOE, Universities and consultants. These sources were able to provide good input concerning O & M (particularly the NRCS, which just completed an evaluation of agricultural BMP's by June C. Grabemeyer, Agriculture Economist, East Lansing, MI), but were less certain about replacement costs. The group decided that 4% was a good average O, M & R factor to use in estimating long-term costs for all types of restoration projects, for the purpose of estimating funding needs.

Table 1: Average O, M & R Factor for Watershed Restoration BMP's

<table>
<thead>
<tr>
<th>Agricultural BMPs</th>
<th>4%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stream Restoration BMPs</td>
<td>4%</td>
</tr>
<tr>
<td>Stormwater Management BMPs</td>
<td>3%</td>
</tr>
<tr>
<td>AMD Vertical Flow Systems</td>
<td>5%</td>
</tr>
<tr>
<td>AMD Anoxic Limestone Drain Systems</td>
<td>4%</td>
</tr>
<tr>
<td>AMD Compost Anaerobic wetlands</td>
<td>4%</td>
</tr>
<tr>
<td>AMD Pyrolusite© Systems</td>
<td>3%</td>
</tr>
<tr>
<td>AMD Open Limestone Channels</td>
<td>1%</td>
</tr>
</tbody>
</table>

The workgroup broke down the long-term O, M & R factor into cost categories. For a passive treatment system that has a 5% annual factor, system reconstruction accounted for 40% of the costs, routine operations (inspections, sampling, flushing) accounted for 20%, water sample analyses accounted for 10%, and general and unscheduled maintenance and repairs accounted for 30%. This breakdown was valuable because it showed that well-organized project sponsors should be able to cover up to 60% of the estimated O, M & R costs by assuming all or part of the non-replacement responsibilities.

The workgroup decided to use the O, M & R factor method to analyze costs and make recommendations concerning the amount of funds needed to address O, M & R on a long-term
basis. This determination was made based on feedback received from Executive Staff and others that up-front, lump sum funding of O, M & R was not likely to be pursued by the Department.

A difficult issue within the analysis of long-term costs was the cost of lab analyses of water monitoring samples collected. The water sampling cost analysis was based on the Department’s cost of $65 per sample (approximate cost of the Bureau of Abandoned Mine Reclamation's 711 Standard Analysis Code, used for routine AMD samples). Private laboratories experienced with AMD analysis can provide reliable analyses for $15-35 per sample, although inexperienced private labs sometimes provide inconsistent results. Two possible options were discussed with regard to sample analyses. One is for the Department to consider certifying private laboratories for AMD analysis and encourage watershed groups to use private labs, thereby decreasing long-term costs. Another option is for the Department to develop a regular funding source for analyses of watershed samples currently being collected under Mineral Resource Management's collector numbers. Costs can be reduced by determining a Standard Analysis Code that provides the minimum number of parameters needed to evaluate system performance. The advantages of this option are that lab results would be made available more easily to the Department and the quality assurance issues are addressed. However, it may be possible to address the quality assurance issue with private labs through a certification process. The workgroup has decided to recommend both options so that watershed groups can utilize what works best on an individual basis. The workgroup believes that the Bureau of Mining and Reclamation's existing SOAP/ROAP certification process is the best vehicle to use to certify private labs.

**Implementation (Funding Engine)**

Various funding options were reviewed by the workgroup to provide for the sustainability of existing and future facilities that benefit the general public and improve the water resources of the Commonwealth.

True sustainability needs local community ownership and involvement. Public-private partnering develops healthy interdependence (working relationship) between state agencies and the watershed residents, including volunteers, students, service groups, private industry, environmental professionals, and other interested parties.

Some project sponsors have developed and are implementing long-term plans; however, many groups currently do not have the means or ability to do this.

The workgroup developed recommendations for a support strategy to enable groups to provide for long term O, M & R. It includes the following:

- **Commonwealth**: develop a source of funding and create a grant funding category for the O, M & R of existing and future construction projects;
- **Sponsor**: provide available resources for total or partial O, M & R;
- **Other**: provide additional O, M & R support by use of the Bureau of Abandoned Mine Reclamation (BAMR) construction/maintenance crews, the 12th Congressional District Equipment Center, and local/private industry.
Funding Options:

The workgroup calculated the approximate initial annual funding needed to address long-term O, M & R at $1.86 million, using the following method. This amount, discussed in both options below, has been calculated by determining the cost of providing for 50% of the average 4% O, M & R factor of $93 million for existing projects. This amount is expected to cover major maintenance (10%, or approximately 1/3 of the expected total maintenance costs) and replacement (40%) needs. It is expected that watershed groups and their local partners, Department assistance with lab costs and BAMR and 12th Congressional District Equipment Center assistance with maintenance will make up the remaining 50% of the O, M & R factor.

The following are two alternatives developed by the workgroup as possible solutions to the funding challenges associated with long-term O, M & R. One of these alternatives, or a combination thereof, may ultimately be seen as the appropriate funding solution.

Option 1: Funding O, M & R on an annual-basis (“pay as you go”)

- Up to 10% of Growing Greener funds are earmarked for funding of O, M & R projects; the amount not spent for O, M & R is released to provide additional new project funding.
- The Secretary's approval is needed if demand is such that more than 10% of Growing Greener funds are necessary.
- Some of the 10% is held back for emergency O, M & R projects, with this money released for new project funding at the end of the fiscal year.
- The delivery system would be the existing Growing Greener Grant Center, using an additional funding category on the grant application form.

Advantages:

With this option, if the O, M & R amount is not fully requested, then the balance would be available for funding new projects. At the current Growing Greener funding level of $50 million per year, it is expected that less than 10% of this amount will cover all major maintenance and replacement needs for the foreseeable future (expected to be about $1.86 million for existing projects).

Disadvantages:

This option requires the continuation of Growing Greener beyond year five. At this time, continuation is considered likely, but is not a certainty. Also, if Growing Greener is continued, the funding level may be reduced, thereby reducing the amount available for O, M & R. Another disadvantage is that it will take away from money to be spent on new projects, unless the Legislature authorizes increased Growing Greener funding to make up the difference.

Option 2: Funding O, M & R for the long-term (“set aside”)

- The PA legislature provides an annual budget appropriation for long-term needs (or, an existing funding source is found within the Department) at an initial rate of $1.86 million per year.
- The amount appropriated will need to increase annually based upon the amount spent on
construction projects annually. For example, if $25 million worth of projects is constructed in 2002, there will be $93 million plus $25 million, or $118 million worth of constructed projects; therefore, $2.36 million will need to be appropriated for O, M & R the following year (50% of 4% factor multiplied by $118 million).

- The annual appropriation would be placed in a "set-aside" fund administered and managed by the Commonwealth. Applicants would apply for funds using the established Growing Greener framework. Any money left over at the end of the year would stay in the fund. The fund would be allowed to build up so that, when needs become greater (as systems need replaced or major floods or other catastrophes occur), the funds would be available to cover that need.

Advantages:

This option would leave the current project funding amounts for Growing Greener intact and would not be dependent upon the continuation of Growing Greener beyond year five. It would allow an accumulation of funds to deal with long-term needs that are expected to increase as systems age and need to be replaced.

Disadvantages:

This option would require legislative action to appropriate funds. It would require the establishment and administration of an interest-bearing fund. It would require tracking of implementation projects from all public funding sources in order to know how much new construction takes place on an annual basis, to determine funding amounts.

**Actions Needed by the Department for Implementation**

- Select a funding option and appropriate funds for O, M & R support of existing and future projects.
- Develop a fund/program management system, including a Growing Greener O, M & R project category and related activities (including changes to scoring and application guidance).
- Require the development of O, M & R plans prior to the provision of O, M & R funds for existing projects and as a deliverable under construction contracts for new projects.
- Provide O, M & R training for watershed groups via Growing Greener workshops and watershed conferences, with assistance from others.
- Improve DEP capacity to assist groups with O, M & R:
  - Provide improved capacity of BAMR's construction crews to assist with major maintenance.
  - Dedicate funds to support Mineral Resource Management sponsored lab analysis for watershed groups and determine an appropriate Standard Analysis Code.
  - Adopt SOAP/ROAP lab criteria and cost guidelines for watershed sample analysis.
# Appendix A

## Publicly Funded Restoration Projects

<table>
<thead>
<tr>
<th></th>
<th>NRCS</th>
<th>319</th>
<th>BAMR</th>
<th>WRPA</th>
<th>OSM</th>
<th>G2</th>
<th>Totals</th>
</tr>
</thead>
<tbody>
<tr>
<td>1988</td>
<td>$125,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$125,000</td>
</tr>
<tr>
<td>1989</td>
<td>$150,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$150,000</td>
</tr>
<tr>
<td>1990</td>
<td>$0</td>
<td>$75,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$75,000</td>
</tr>
<tr>
<td>1991</td>
<td>$75,000</td>
<td>$200,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$275,000</td>
</tr>
<tr>
<td>1992</td>
<td>$12,000</td>
<td>$225,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$237,000</td>
</tr>
<tr>
<td>1993</td>
<td>$0</td>
<td>$400,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$400,000</td>
</tr>
<tr>
<td>1994</td>
<td>$0</td>
<td>$675,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$675,000</td>
</tr>
<tr>
<td>1995</td>
<td>$152,066</td>
<td>$850,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$1,002,066</td>
</tr>
<tr>
<td>1996</td>
<td>$0</td>
<td>$1,000,000</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$1,000,000</td>
</tr>
<tr>
<td>1997</td>
<td>$183,959</td>
<td>$275,000</td>
<td>$1,502,626</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$1,961,585</td>
</tr>
<tr>
<td>1998</td>
<td>$34,314</td>
<td>$1,700,000</td>
<td>$1,664,737</td>
<td>$0</td>
<td>$0</td>
<td>$0</td>
<td>$3,399,051</td>
</tr>
<tr>
<td>1999</td>
<td>$274,454</td>
<td>$3,400,000</td>
<td>$2,470,041</td>
<td>$688,458</td>
<td>$262,240</td>
<td>$25,350,000</td>
<td>$32,445,193</td>
</tr>
<tr>
<td>2000</td>
<td>$109,284</td>
<td>$2,700,000</td>
<td>$643,873</td>
<td>$296,558</td>
<td>$567,800</td>
<td>$21,050,000</td>
<td>$25,367,515</td>
</tr>
<tr>
<td>2001</td>
<td>$200,000</td>
<td>$3,700,000</td>
<td>$1,390,401</td>
<td>$0</td>
<td>$321,400</td>
<td>$20,140,000</td>
<td>$25,751,801</td>
</tr>
</tbody>
</table>

$92,864,211