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# Recovery of Lithium From a Montmorillonite-Type Clay 

By R. H. Lien

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With an Appendix on Process Economics by D. A. Kramer


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## UNIT OF MEASURE ABBREVIATIONS USED IN THIS REPORT

| ${ }^{\circ} \mathrm{C}$ | degree Celsius | $\mathrm{L} /\left(\mathrm{m}^{2} \cdot \mathrm{~h}\right)$ | liter per square meter per hour |
| :--- | :--- | :--- | :--- |
| cm | centimeter | 1 b | pound |
| $\mathrm{d} / \mathrm{yr}$ | day peir year | M | thousand |
| ft | foot | $\mathrm{m}^{3}$ | cubic meter |
| $\mathrm{g} / \mathrm{L}$ | gram per liter | mi | mile |
| gal | gallon | min | minute |
| h | hour | mL | milliliter |
| $\mathrm{h} / \mathrm{d}$ | hour per day | MM | million |
| hp | horsepower | mm-diam | millimeter-diameter |
| in | inch | pct | percent |
| kg | kilogram | psi | pound per square inch |
| km | kilometer | ton/d | short ton per day |
| kW | kilowatt | W | watt |
| $\mathrm{kW} \cdot \mathrm{h}$ | kilowatt hour | wt pct | weight percent |
| L | liter | yr | year |
| $\mathrm{L} / \mathrm{kg}$ | liter per kilogram |  |  |

# RECOVERY OF LITHIUM FROM A MONTMORILLONITE-TYPE CLAY 

By R. H. Lien ${ }^{1}$<br>With an Appendix on Process Economics by D. A. Kramer


#### Abstract

The Bureau of Mines investigated a roast-leach process for recovering a marketable lithium product from a montmorillonite-type clay deposit located on the Nevada-Oregon border. The clay sample treated in the investigation contained 0.6 wt pct Li.

The lithium recovery process consisted of several unit operations. The lithium silicate compounds in the clay were converted to $\mathrm{Li}_{2} \mathrm{SO}_{4}$ by roasting a pelletized mixture of clay, limestone, and gypsum at $900^{\circ} \mathrm{C}$ in a direct-gas-fired rotary roaster. Water-leaching the calcine at 40 pct solids extracted the $\mathrm{Li}_{2} \mathrm{SO}_{4}$. Lithium recovery from the leach solution involved concentrating the solution by evaporation, adding $\mathrm{Na}_{2} \mathrm{CO}_{3}$ to the concentrated solution to precipitate $\mathrm{Li}_{2} \mathrm{CO}_{3}$, and filtering the slurry to obtain the product. The product filtrate was recycled to the evaporator following a crystallization step. About 80 pct of the lith... ium in the clay was recovered as 99 -pct-pure $\mathrm{Li}_{2} \mathrm{CO}_{3}$.

Process operating costs were estimated at $\$ 2.12 / 1 \mathrm{~b} \mathrm{Li}_{2} \mathrm{CO}_{3}$ produced; the current $\mathrm{Li}_{2} \mathrm{CO}_{3}$ selling price is $\$ 1.48 / 1 \mathrm{~b}$. Raw materials accounted for 30 pct of the total operating cost.


[^0]
## INTRODUCTION

The Bureau of Mines investigated several processes for recovering lithium from a clay deposit located on the Nevada-Oregon border. Development of a process to recover lithium from this nonconventional, domestic resource would help meet the Bureau's goal of developing technology to help the Nation maintain an adequate minerals base for future economic and strategic needs.

The United States, the world's largest producer and consumer of lithium minerals and chemicals, is self-sufficient in lithium. Nearly all the Nation's lithium is recovered from spodumene deposits in North Carolina and subsurface brines in Nevada.

The largest end use for lithium is in aluminum potlines. In the aluminum cells, $\mathrm{Li}_{2} \mathrm{CO}_{3}$ is added to reduce electricity consumption and fluorine emissions. The ceramics, air conditioning, grease, synthetic rubber, and pharmaceutical industries also use lithium chemicals. Recently, lithium has gained importance in areas such as (1) lowdensity aluminum-1ithium aircraft alloys, (2) lightweight batteries for use in electric automobiles and utility loadleveling purposes, and (3) nuclear fusion for use as a supply of tritium and as a coolant and heat transfer agent.
In the 1970's, several research and Government organizations discussed the possibility of domestic lithium shortages by the year $2000(\underline{1}-\underline{3}) .^{2}$ These predictions were based primarily on rapid expansion of the lithium batteries market, principally in the area of electric automobiles. The development of thermonuclear energy was expected to have an

[^1]impact on lithium reserves after the year $2000 .{ }^{3}$

Forecasts of lithium shortages prompted the U.S. Geological Survey (USGS), under a cooperative agreement with the Department of Energy, to search for alternative domestic lithium resources. The USGS identified a large deposit of lithiumbearing clay in the McDermitt caldera complex on the Nevada-Oregon border (fig. 1). The caldera complex, one of the largest in the world, comprises five overlapping and nested calderas (circular volcanic depressions). The principal clay deposit area measures 42 km long by 18 km wide.

The lithium-bearing clays are found primarily along the edge of the caldera in a crescent extending from the northeastern corner to the southwestern section. Lithium concentration in the clay is 0.1 to 0.36 pct in the northern deposits and 0.1 to 0.65 pct in the southern area.

These clay deposits can be considered a potential resource because of high lithium content in individual beds. Also, the beds have very little overburden. The amount of lithium in the caldera has been estimated as high as 10 MM tons.

The Bureau's Salt Lake City Research Center obtained about 6 M lb of clay from a discovery cut in the southwestern section of the caldera (fig. 1). A typical clay sample contained 0.6 wt pct lithium.

Bureau investigators studied several methods for extracting lithium from the McDermitt clay (5-9). The research included extensive $\bar{l}$ aboratory studies on a lime-gypsum roast process that converts
lithium demand picture. In 1980, the Bureau of Mines predicted that development of lithium resources such as McDermitt clays would not be needed until at least the year 2050 (4). However, the recent development of aluminum-lithium aircraft alloys, which cut aircraft weight, resulting in considerable fuel savings, may cause a substantial increase in the demand for lithium.


FIGURE 1. - Location of McDermitt caldera.
the lithium in the clay to a water-soluble product, without converting major contaminating elements such as aluminum and magnesium to water-soluble compounds (7). Water leaching extracts more than $8 \overline{0}$ pct of the lithium present in the ore, along with approximately equivalent percentages of the sodium and potassium. The leach solution is concentrated by evaporation, and a $\mathrm{Li}_{2} \mathrm{CO}_{3}$ product is
precipitated by adding soda ash ( $\mathrm{Na}_{2} \mathrm{CO}_{3}$ ). A process research unit (PRU) was built and operated to confirm the laboratory results and provide data for a cost evaluation.

This report presents the PRU operating procedures and conditions, test results, and material balances. A study or fac-tored-type capital and operating cost evaluation is also included.

## PROCESS DESCRIPTION

Figure 2 presents the generalized flowsheet for the PRU. The process involves several unit operations: feed preparation, roasting, leaching, evaporation, and crystallization.

A mixture of clay, limestone, and gypsum is pelletized, dried, and fed to the roasting furnace. Roasting the pellets at $900^{\circ} \mathrm{C}$ converts the 1ithium in the clay to water-soluble $\mathrm{Li}_{2} \mathrm{SO}_{4}$. Waterleaching the calcine produces a solution containing lithium, potassium, sodium,
and a small amount of calcium, all as sulfates. The leach solution, together with recycled product wash and mother liquor, is concentrated by evaporation. During this concentration operation, carbonate ion present in the recycle solutions causes calcium to precipitate as $\mathrm{CaCO}_{3}$, which is removed by filtration. The concentrated solution is then heated to bof.ling, and $\mathrm{Na}_{2} \mathrm{CO}_{3}$ is added to precipitate $\mathrm{Li}_{2} \mathrm{CO}_{3}$ product.


FIGURE 2. - Generalized process flowsheet. Material balances for numbered streams are presented in appendix A tables.

The product filtrate contains about $3 \mathrm{~g} / \mathrm{L}$ Li. To recover this lithium, the solution is cooled to crystallize $3 \mathrm{~K}_{2} \mathrm{SO}_{4}$ $\cdot \mathrm{Na}_{2} \mathrm{SO}_{4}$ and $\mathrm{Na}_{2} \mathrm{SO}_{4} \cdot 10 \mathrm{H}_{2} \mathrm{O}$, and then recycled to the evaporation step.

Removing calcium from the leach solution as $\mathrm{CaCO}_{3}$ prevents calcium
contamination of the product. The solution recycled to the evaporation step contains sufficient carbonate ion, as either $\mathrm{Li}_{2} \mathrm{CO}_{3}$ or excess $\mathrm{Na}_{2} \mathrm{CO}_{3}$, to precipitate over 99 pct of the calcium contained in the leach solution.

## MATERIALS

Table 1 shows the composition of a typical clay sample. X-ray diffraction studies, together with the clay's chemical properties, indicated that the clay is chiefly montmorillonite. The lithium occurs in the clay mineral hectorite [ $\left.\mathrm{Na} 0 .{ }_{33}(\mathrm{Mg}, \mathrm{Li})_{3} \mathrm{Si}_{4} \mathrm{O}_{10}(\mathrm{~F}, \mathrm{OH})_{2}\right]$ and an

TABLE 1. - Composition of McDermitt clay, percent

| A1 | 3.1 | Li.......... | 0.6 |
| :---: | :---: | :---: | :---: |
| Ca | 1.8 | Mg. . . . . . . . | 9 |
| F. | 2 | Na. | . 58 |
| Fe | 2.5 | Total $\mathrm{SiO}_{2} \ldots$ | 53 |
| K. | 3.7 | Free $\mathrm{SiO}_{2} .$. | 15 |

unnamed clay mineral similar to hectorite but containing more aluminum (10).

Mineralogical studies showed that the lithium is evenly disseminated throughout the clay; thus, physical beneficiation techniques would not upgrade the lithium content.

The feed for the roast was a mixture of clay, limestone, and agricultural-grade gypsum. The $\mathrm{Li}_{2} \mathrm{CO}_{3}$ was precipitated from the concentrated solution by adding dense soda ash $\left(\mathrm{Na}_{2} \mathrm{CO}_{3}\right)$.

Simulated solutions used in laboratory evaporation and product purification tests were made up with reagent-grade chemicals.

The feed for the roast was ground and mixed in the ball mill (40-cm-diam by 60-cm-length) shown in figure 3 . The minus 100-mesh mixture was pelletized with water in the drum pelletizer (50-cm-diam) shown in figure 4.

The pellets were roasted in a directfired rotary roaster. The roaster, shown in figure 5, was fired with natural gas and had a working capacity of $0.014 \mathrm{~m}^{3}$. The roast product (calcine) was leached with water in the 208-L baffled, polyethylene vessel shown in figure 6. A propeller mixer, driven by a $0.75-\mathrm{hp}$ direct-drive motor, agitated the slurry.

Figure 7 shows the evaporation, filtration, and product precipitation equip-ment. To recover the leach solution, the slurry from the leach tank was transferred by gravity flow to a pan filter with a surface area of $0.8 \mathrm{~m}^{2}$. The


FIGURE 3. - Ball mill.
filter cloth was a medium-weight cotton twill. A tubing pump transferred the leach solution to a $208-\mathrm{L}$ stainless steel tank for concentration. The tank was equipped with a 9,000-W over-the-side immersion heater. The concentrated solution was filtered on a tabletop Buchner funnel to remove $\mathrm{CaCO}_{3}$. A tubing pump transferred the concentrated solution to the product precipitation unit, which consisted of a $24-\mathrm{L}$ stainless steel vessel equipped with a $1,000-W$ over-the $\cdots$ side immersion heater. An impeller mixer, driven by a $0.25-h p$ direct-drive motor, agitated the solution. The $\mathrm{Li}_{2} \mathrm{CO}_{3}$ product was recovered and washed on a Buchner funnel.

The product filtrate was cooled by refrigeration to crystallize $\mathrm{Na}_{2} \mathrm{SO}_{4} \cdot 10 \mathrm{H}_{2} \mathrm{O}$ and $3 \mathrm{~K}_{2} \mathrm{SO}_{4} \cdot \mathrm{Na}_{2} \mathrm{SO}_{4}$.


FIGURE 4. - Drum pelletizer.


FIGURE 5. - Direct-fired rotary roaster.


FIGURE 6. - Leach tank.


## OPERATING CONDITIONS AND TEST RESULTS

Process unit operations were studied in a PRU and in laboratory tests to determine operating conditions that would (1) maximize lithium recovery, (2) minimize process operating costs, and (3) produce a high-purity product.

Appendix A presents a material balance for a typical PRU test using a 5:3:3 ratio of clay, limestone, and gypsum; a 5:2:2 ratio might also be considered since reagent and utility costs would be lower

## FEED PREPARATION

The McDermitt clay contains lithium principally as hectorite. To convert the lithium to $\mathrm{Li}_{2} \mathrm{SO}_{4}$, the clay was mixed with limestone and gypsum and roasted.

Feed preparation entailed grinding and mixing the ingredients (clay, limestone, and gypsum) for 1 h in a ball mill. The resultant mixture ( 80 pct finer than 200 mesh) was pelletized with water to produce nominal 6.5 -mm-diam pellets. These pellets contained up to 20 pet moisture and were dried at $70^{\circ} \mathrm{C}$ before roasting

## ROASTING

Roasting studies consisted of tube fur. nace tests, batch tests conducted in an electrically heated muffle furnace, and continuous tests made in a gas-fired ro-* tary roaster. The tests were conducted to determine the reaction mechanism and to establish optimum roasting conditions (7).

## $\underline{\text { Roasting Mechanism }}$

The pelletized mixture of clay, limestone, and gypsum was roasted to convert lithium in the clay to water-soluble $\mathrm{Li}_{2} \mathrm{SO}_{4}$. Data obcained from exploratory tube furnace tests (7) indicated that $\mathrm{Li}_{2} \mathrm{SO}_{4}$ was formed by the following reactions (for simplicity, lithium-silicate minerals such as hectorite are represented by $\mathrm{Li}_{2} \mathrm{Si}_{2} \mathrm{O}_{5}$ )

$$
\begin{align*}
& \mathrm{CaSO}_{4} 2 \mathrm{H}_{2} \mathrm{O}+\mathrm{SiO}_{2} \\
& \longrightarrow \mathrm{CaSiO}_{3}+\mathrm{SO}_{2}+1 / 2 \mathrm{O}_{2}+2 \mathrm{H}_{2} \mathrm{O}  \tag{1}\\
& \mathrm{Li}_{2} \mathrm{Si}_{2} \mathrm{O}_{5}+\mathrm{SO}_{2}+1 / 2 \mathrm{O}_{2} \\
& \longrightarrow \mathrm{Li}_{2} \mathrm{SO}_{4}+2 \mathrm{SiO}_{2} \tag{2}
\end{align*}
$$

Limestone limited the back reaction of free $\mathrm{SiO}_{2}$ with $\mathrm{Li}_{2} \mathrm{SO}_{4}$. Free silica appeared to react with CaO to form a calcium silicate.

## Batch Tests

Extensive batch testing was conducted In an electrically heated muffle furnace to determine the effects of charge composition, roasting time, and temperature on lithium extraction (7-8). Specifically, the following conditions were investigated: (1) preroast mixtures (clay-limestone-gypsum) ranging from 5:0:6 to 5:6:0, (2) roasting times of 1 to 4 h , and (3) roasting temperatures of $750^{\circ}$ to $1,050^{\circ} \mathrm{C}$.

Reagent-grade $\mathrm{CaCO}_{3}$ and $\mathrm{CaSO}_{4} \cdot 2 \mathrm{H}_{2} \mathrm{O}$ were used in these tests. The ingredients were pelletized and roasted. The resultant calcines were water-leached to determine lithium extraction.

Data presented in table 2 show the effect of chaige composition on lithium extraction. These tests were conducted at $1,000^{\circ} \mathrm{C}$ for 1 h . Of the compositions tested, a 5:3:3 mix was optimum with a lithium extraction of 88.4 pct. Lithium extraction from clay roasted alone (5:0:0) was only 0.1 pet.

Table 3 shows the effect of roasting temperature on lithium extraction. In these tests, a $5: 3: 3$ mixture was calcined for 1 h at $750^{\circ}$ to $1,100^{\circ} \mathrm{C}$. The optimum extraction was attained at $1,000^{\circ} \mathrm{C}$. Roasting for periods longer than 1 h did not improve lithium extraction. Prolonged roasting for 4 h at elevated temperatures (above $850^{\circ} \mathrm{C}$ ) slightly reduced lithium recovery.

TABLE 2. ... Effect of charge composition on lithium extraction under static conditions ${ }^{1}$

| Weight ratio |  |  | Li extraction, |
| :--- | :---: | :---: | :---: |
| pct |  |  |  |

TABLE 3. - Effect of roasting temperature on lithium extraction under static conditions ${ }^{1}$

Li extraction, pct

| $750^{\circ}$ | C. | 59.4 |
| :---: | :---: | :---: |
| $800^{\circ}$ | C............. | 73.3 |
| $850^{\circ}$ | C. | 79.5 |
| $900^{\circ}$ | C. | 83.8 |
| $950^{\circ}$ | C. | 85.6 |
| 1,000 ${ }^{\circ}$ | C. | 88.4 |
| 1,050 ${ }^{\circ}$ | C. | 83 |
| 1, $100^{\circ}$ | C.............. | 65.7 |

Tests were also conducted to study the effect of naturally occurring limestone and gypsum on lithium recovery (7). In these tests, the quantities of limestone and gypsum mixed with the clay were calculated to produce an equivalent 5:3:3 mix. The equivalence was based on the CaO content of reagent-grade $\mathrm{CaCO}_{3}$ and $\mathrm{CaSO}_{4} \cdot 2 \mathrm{H}_{2} \mathrm{O}$. The mixtures were ground to minus 200 mesh, pelletized, and roasted at $1,000^{\circ} \mathrm{C}$ for 1 h . Lithium extractions ranged from 83 to 85.5 pct, compared with 88.4 pct for roasts using reagent-grade chemicals. The decrease in extraction
was attributed to the coarser particle size of the natural materials.

## Continuous Roast Tests

Larger scale testing was conducted in a gas-fired rotary roaster to study the dynamic variables affecting the calcine and to confirm laboratory test results.

Initially, a series of batch tests was conducted in the rotary roaster to determine optimum roast temperature and retention time (7). In these tests, small charges ( 500 g ) of pelletized $5: 3: 3$ mix were roasted, The test resilts showed $900^{\circ} \mathrm{C}$ and a $2-\mathrm{h}$ retention time to be optimum. The initial continuous roasting tests were conducted at these conditions.

The primary objective of the continuous testing was to generate calcine for use in PRU leach tests. An equivalent 5:3:3 mixture of clay, limestone, and gypsum was used because batch testing established this mix as optimum. The pelletized feed was charged to the roaster in 600-g increments every 5 min. Generally, each test produced 36 kg of calcine in about 6-1/2 h operating time.

The final phase of the continuous roast work involved investigating the effects of charge composition and roast temperature on lithium extraction. A series of tests was conducted in which various mixes were roasted. Lithium extraction was determined by water-leaching composite samples of the calcines.

Test results, presented in table 4 , show that lithium extractions of at least 80 pct were attained with a wide range of clay-limestone-gypsum ratios. Also, the data indicate that good lithium extraction was achieved over a temperature range of $850^{\circ}$ to $975^{\circ} \mathrm{C}$. The 5:3:3 mix was chosen as the basis for a cost evaluation because other mixtures, such as 5:2:2, had not been adequately studied at the time the evaluation was begun. Details are presented in appendix B.

TABLE 4. - Effect of charge composition and roast temperature on lithium extraction under dynamic conditions

| Weight ratio |  |  | Roast <br> temp, ${ }^{\circ} \mathrm{C}^{1}$ |  | Roasts <br> at given <br> conditions |
| :--- | :---: | :---: | :---: | :---: | :---: |
| Clay | $\mathrm{CaCO}_{3}$ | $\mathrm{CaSO}_{4} \cdot 2 \mathrm{H}_{2} \mathrm{O}$ | Li <br> ertraction, <br> pct ${ }^{2}$ |  |  |
| 5 | 1 | 1 | 900 | 1 | 59 |
| 5 | 1.5 | 1.5 | 900 | 1 | 80.5 |
| 5 | 1.5 | 1.5 | 975 | 1 | 81 |
| 5 | 2 | 1 | 900 | 1 | 81.6 |
| 5 | 2 | 2 | 850 | 1 | 84.6 |
| 5 | 2 | 2 | 900 | 3 | $84.3-86.9$ |
| 5 | 2 | 2 | 950 | 1 | 89.3 |
| 5 | 2 | 3 | 900 | 1 | 84.4 |
| 5 | 3 | 2 | 900 | 2 | $86.8-88.5$ |
| 5 | 3 | 3 | 900 | 17 | $83.3-86.6$ |
| 5 | 3 | 4 | 900 | 1 | 85.1 |

${ }^{1}$ Retention time was 2 h .
${ }^{2}$ For analysis, calcine samples were leached with water at 25 pct solids; the residues were then washed. Lithium extraction was based on calcine and residue analyses.

LEACHING

The objective of the PRU leach tests was to determine the relationship between leach-system variables and optimal lithium extraction. The following variables were studied: (1) leach pulp percent solids, (2) wash water recycle, (3) calcine particle size, and (4) leach time.

The calcines leached in these tests were generated by roasting 5:3:3 mixtures of clay, limestone, and gypsum. General$1 \mathrm{y}, 32 \mathrm{~kg}$ of calcine was water-leached in each test. The slurry was filtered to recover the leach solution. The filter cake was then washed and discarded.

## Solids Content and Wash Water Recycle

Studies were conducted to minimize the volume of leach solution produced and thereby reduce subsequent evaporation requirements. The volume of solution can be reduced by increasing the percent solids in the leach slurry and recycling the calcine wash water.

A series of $30-\mathrm{min}$ leach tests was conducted to study the effect of percent solids and wash water recycle on lithium extraction. The test results, presented in table 5, show that the calcine was leached effectively at 40 pct solids with

TABLE 5. - Effect of solids content and wash water recycle on lithium extraction, percent ${ }^{1}$

| Solids content, pct | Leached with-- |  |
| :---: | :---: | :---: |
|  | Fresh water | Recycled wash water |
| $25^{2}$ | 83.4 | 82.7 |
| $40^{3}$ | 81.4 | 83.6 |
| $50^{3}$ | 80.9 | 69.4 |
| 'Extractions based on lithium concen- |  |  |
| trations in calcine and leached residue. |  |  |
| 296-L leach and wash water volume. |  |  |
| 348 - L leach an | wash water | lume. |

recycled wash water. At 50 pct solids, the lithium extraction decreased. Since the wash water was recycled to the leach step, the volume of wash water used was equal to the volume of water required for the next leach.

## Calcine Particle Size and Leach Time

The calcine pellets do not break apart during the leach. If a coarse particle could be leached effectively, grinding requirements would be minimized. Therefore, a series of tests was conducted to study the effect of calcine particle size on lithium extraction.

The calcine was leached for 30 min at 40 pct solids using recycled wash water. The test results, given in table 6 , show that the 30 -min leach extracted the lithium equally well from all particle sizes tested。

TABLE 6. - Effect of calcine particle size on lithium extraction

| Li extraction, |  |
| :---: | :---: |
| Particle size | $p \mathrm{ct}{ }^{2}$ |
| Minus 100 mesh | 82.6 |
| Coarse-crushed pellets ${ }^{1}$. | 83.6 |
| Whole pellets.......... | 83.9 |
| ${ }^{1}$ Screen analysis of calcine | particles |
| was 70 pct plus 10 mesh. |  |
| ${ }^{2}$ Extractions based on lithiu | concen- |
| trations in calcine and leached | residue |

To determine the effect of leach time on lithium extraction, a series of tests was conducted with coarse-crushed and whole pellets. The pellets were leached at 40 pct solids in recycled wash water. The test results, presented in table 7 , show that no more than a 5 -min leach was required to extract the lithium from coarse-crushed pellets. However, whole pellets were not effectively leached in 5 min .

TABLE 7. - Effect of leach time on lithium extraction

| Particle size | Leach <br> time, <br> min | Li extraction, <br> pct |
| :---: | :---: | :---: |
| Coarse-crushed | 5 | 84.2 |
| pellets.......... | 15 | 84.2 |
|  | 30 | 83.6 |
|  | 60 | 83.1 |
| Whole pellets...... | 5 | 70.7 |
|  | 30 | 83.9 |
|  | 60 | 84.2 |

Although the pellets did not break apart during the leach, prolonged agitation generated fines which affected filtration rates. The data presented in table 8 show that filtrate rates decreased with increased leach time. Also, the
rates decreased as calcine particle size increased. For 30-min leaches, the whole pellet slurry filtered slowly because the filter cloth was biinded with fines. As calcine particle size decreased, the fines tended to remain on top of the filter cake, allowing faster filtration.

TABLE .. - Effect of calcine particle size and leach time on filtrate rates

| Calcine particle <br> size | Leach <br> time, <br> min | Filtrate rate, <br> $\mathrm{L} /\left(\mathrm{m}^{2} \cdot \mathrm{~h}\right)$ |
| :---: | :---: | :---: |
| Minus 100 mesh... | 30 | 86 |
| Coarse-crushed |  |  |
| pellets......... | 5 | 155 |
|  | 15 | 111 |
|  | 30 | 68 |
| Whole pellets.... | 50 | 67 |
|  | 30 | 150 |

TRates calculated from data obtained by filtering leach slurry on process pan filter.

The test results indicate that the calcine particle size used in the leach would be determined by evaluating the operating costs, including grinding, filtration, and agitation costs.

## Summary of Leach Test Results

The PRU results show a maximum lithium extraction from the calcine of 82 to 84 pct. Based on test data, the preferred extraction procedure involved leaching coarse-crushed pellets with recycled wash water for 5 min at 40 pct solids. Leaching at ambient temperature produced a solution containing 2.5 to $3.0 \mathrm{~g} / \mathrm{L} \mathrm{Li}$.

## EVAPORATION

As figure 2 shows, the evaporator was fed with leach solution and solution recycled from the previous test. The recycled solution (mother liquor plus product wash) accounted for about 20 pct of the total volume in the evaporator.

In addition to concentrating the solution, calcium was removed from the leach
solution in this step of the process. The leach solution was saturated with $\mathrm{CaSO}_{4}$ (about $0,6 \mathrm{~g} / \mathrm{L} \mathrm{Ca}^{2+}$ ). Extensive laboratory testing showed that reducing the calcium ion concentration to about $0.015 \mathrm{~g} / \mathrm{L}$ prevented calcium contamination of the product.

The evaporation procedure involved the following steps:

1. The solution (leach plus recycle) was evaporated to about 50 pct of its original volume and then filtered to remove $\mathrm{CaCO}_{3}$. Carbonate ion (approximately $15 \mathrm{~g} / \mathrm{L}$ ) present in the recycled solution precipitated over 99 pct of the calcium contained in the leach solution.
2. The filtrate was returned to the evaporator. Evaporation continued until the solution was reduced to 20 pct of its original volume.
3. The hot concentrated solution, containing 12 to $13 \mathrm{~g} / \mathrm{L} \mathrm{Li}$, was transferred to the product precipitation step. Generally this concentrated solution was cloudy because a small amount of $\mathrm{Li}_{2} \mathrm{CO}_{3}$ precipitated during evaporation.

## PRODUCT PRECIPITATION

Lithium recovery involved heating the concentrated solution to boiling and adding a stoichiometric amount of $\mathrm{Na}_{2} \mathrm{CO}_{3}$ to precipitate a $\mathrm{Li}_{2} \mathrm{CO}_{3}$ product. The objective of this step was to recover a product of at least $99-$ pct purity.

Initially, the product was recovered from the hot solution by vacuum filtration and then dried. This procedure yielded a product of about $80-\mathrm{pct}$ purity with the principal contaminants being $\mathrm{Na}_{2} \mathrm{SO}_{4}$ and $\mathrm{K}_{2} \mathrm{SO}_{4}$. Numerous tests were conducted in the PRU and laboratory (using PRU leach solution) to investigate product purification techniques. Test results were erratic because precise control of solution concentration was difficult; therefore, simulated solutions were used to study operating variables.

A series of laboratory tests was conducted using $1-\mathrm{L}$ batches of simulated concentrated solution (made up with reagent chemicals) containing $97 \mathrm{~g} / \mathrm{L} \mathrm{Li}_{2} \mathrm{SO}_{4}$, $158 \mathrm{~g} / \mathrm{L} \mathrm{K}_{2} \mathrm{SO}_{4}$, and $87 \mathrm{~g} / \mathrm{L} \mathrm{Na} 2 \mathrm{SO}_{4}$. Adding a stoichiometric amount of $\mathrm{Na}_{2} \mathrm{CO}_{3}$ to the
hot solution precipitated $\mathrm{Li}_{2} \mathrm{CO}_{3}$. Product filtration and washing procedures were then studied. From the test results presented in table 9, the following observations were made:

1. Pressure filtration yielded a product of higher purity than vacuum filtration by reducing the moisture content of the filter cake.
2. On the pressure filter, 4 to 6 L of wash water per kilogram of dry product was required to produce a 99-pct-pure product. A much higher volume of water was needed to produce a comparable product by vacuum filtration.
3. For pressure filtration, wash water volumes above 6 L per kilogram of dry product did not further improve product purity. Also, single-stage washing was as effective as either multistage washing or product reslurry.
4. Adding $\mathrm{Na}_{2} \mathrm{CO}_{3}$ as a saturated solution rather than as a dry powder had little effect on product purity. However, this procedure generated a coarse grainy product in contrast to the fine powdery product obtained by adding dry $\mathrm{Na}_{2} \mathrm{CO}_{3}$.

The wash water and product filtrate recovered in these tests contained 14 to $16 \mathrm{~g} / \mathrm{L} \mathrm{Li}{ }_{2} \mathrm{CO}_{3}$. The wash was recycled to the evaporator. After a crystallization step, the product filtrate was also recycled.

## CRYSTALLIZATION

In addition to residual $\mathrm{Li}_{2} \mathrm{CO}_{3}$, the product filtrate contained high concentrations of $\mathrm{K}_{2} \mathrm{SO}_{4}$ and $\mathrm{Na}_{2} \mathrm{SO}_{4}$ (over 150 $\mathrm{g} / \mathrm{L}$ of each). If the solution is to be recycled, the buildup of these salts must be prevented.

Laboratory and PRU tests showed that the most effective method for reducing the sulfate concentration involved crystallizing the salts by chilling the solution to between $0^{\circ}$ and $-4^{\circ} \mathrm{C}$ (below $-4^{\circ} \mathrm{C}$, the solution freezes). The mother liquor, which contained $70 \mathrm{~g} / \mathrm{L} \quad \mathrm{Na}_{2} \mathrm{SO}_{4}$ and $100 \mathrm{~g} / \mathrm{L} \mathrm{K}_{2} \mathrm{SO}_{4}$, was recovered by either vacuum or pressure filtration. Pressure filtration tended to reduce lithium loss by decreasing the amount of mother liquor present in the filter cake.

TABLE 9. - Results of product purification tests using simulated solutions

| Wash water volume, mL | Water per dry product, L/kg | Product analysis, wt pct ${ }^{1}$ |  | Comments ${ }^{2}$ |
| :---: | :---: | :---: | :---: | :---: |
|  |  | K | Na |  |
| Vacuum filtration: ${ }^{3}$ |  |  |  |  |
| 0................. | 0 | 4.1 | 3.9 | No wash. |
| 300................. | 5.7 | . 86 | . 6 | Single wash. |
| 600................. | 12.8 | - 4 | . 3 | Do : |
| 700. . . . . . . . . . . . . | 15.3 | . 12 | . 14 | Do. |
| 800................ | 17.5 | . 16 | . 16 | Do. |
| Pressure filtration: ${ }^{4}$ |  |  |  |  |
| 0................. | 0 | 2.3 | 2 | No wash. |
|  | 0 | 1.9 | 1.8 | $\mathrm{Na}_{2} \mathrm{CO}_{3}$ added as saturated solution; no wash. |
|  | 0 | 1.6 | 1.5 | Do. |
| 100................ | 2 | . 3 | . 3 | Single wash. |
| 200................. | 4.1 | .17 | . 17 | Do. |
| 300................. | 6.1 | .16 | . 15 | Do. |
|  | 6.3 | .16 | . 1 | Product reslurried with 200 $\mathrm{mL}, \mathrm{filtered}$, then washed with 100 mL . |
|  | 6.6 | .26 | . 18 | $\mathrm{Na}_{2} \mathrm{CO}_{3}$ added as saturated solution; single wash. |
|  | 6.2 | . 16 | .16 | 3 separate $100-\mathrm{mL}$ washes. |
| 600................ . | 12.9 | . 18 | .16 | Single wash. |
|  | 13.2 | . 14 | . 14 | 6 separate $100-\mathrm{mL}$ washes. |
| 800................. | 17.7 | . 18 | . 2 | Single wash. |

${ }^{T} \mathrm{~K}$ and Na present as sulfates; product also contained traces of Ca ( $0.02 \mathrm{wt} \mathrm{pct)}$ and Mg ( 0.01 wt pct ), probably present as carbonates.
${ }^{2}$ Except as indicated, $\mathrm{Na}_{2} \mathrm{CO}_{3}$ was added to the concentrated solution as dry powder.
${ }^{3}$ Carried out on a laboratory-size Buchner funnel.
${ }^{4}$ Conducted with a tabletop filter connected to a 45 -psi air supply.

The filter cake was a mixture of glauber salts $\left(\mathrm{Na}_{2} \mathrm{SO}_{4} \cdot 10 \mathrm{H}_{2} \mathrm{O}\right)$ and glaserite $\left(3 \mathrm{~K}_{2} \mathrm{SO}_{4} \cdot \mathrm{Na}_{2} \mathrm{SO}_{4}\right)$.

Laboratory tests showed that, if desired, glaserite and glauber salts could be recovered separately by a two-step crystallization procedure. At solution temperatures down to about $17^{\circ} \mathrm{C}$, glaserite crystallized. The salt was recovered by vacuum filtration and analyzed as 33 wt pct $\mathrm{K}, 8$ wt pct Na , and $<0.1$ wt pct Li. Further cooling of the solution (to as low as $-4^{\circ} \mathrm{C}$ ) crystallized glauber salts. These salts were recovered by pressure filtration and dried. The dried salts contained 28 wt pct $\mathrm{Na}, 6$ wt pct $K$ (a small amount of glaserite crystallized with the $\mathrm{Na}_{2} \mathrm{SO}_{4} \cdot 1 \mathrm{IOH}_{2} \mathrm{O}$ ), and 0.15 wt pct Li.

OVERALL LITHIUM RECOVERY

PRU roast-leach test results (tables 57) indicate 82- to 84-pct Li extraction as optimum. Treating the leach solution by the methods specified resulted in 95to $98-$ pct recovery of the contained $1 i t h-$ ium. Losses occurred in $\mathrm{CaCO}_{3}$ filtration (0.5-pet loss) and in the crystallization step (2- to 5-pct loss depending on the filtration method used to separate the mother liquor from the salts). Overall, 78 to 82 pet of the lithium contained in the clay was recovered as 99-pct-pure $\mathrm{Li}_{2} \mathrm{CO}_{3}$.

## MATERIAL BALANCE AND ECONOMIC EVALUATION

Appendix A shows a material balance for a typical PRU test using a 5:3:3 mixture of clay, limestone, and gypsum. The Bureau's Process Evaluation Office at Avondale, $M D$, prepared a cost evaluation based on the PRU material balance, filtration rate data, flowsheet, and process description supplied by the Salt Lake City Research Center.

The process evaluation, presented in appendix $B$, estimates the operating cost of the process at $\$ 2.12 / 1 \mathrm{~b} \mathrm{Li}_{2} \mathrm{CO}_{3}$ produced. (The current selling price of $\mathrm{Li}_{2} \mathrm{CO}_{3}$ is $\$ 1.48 / 1 \mathrm{~b}$ ). The evaluation identifies the high-cost areas of the process as reagents, utilities, and de-preciation. A summary of capital and operating costs for a plant processing 1,000 ton/d of McDermitt clay is presented in tables $B-2$ and $B-3$.

Table B-3 shows raw materials and utilities to be the high-cost areas of the process. Reagent costs are high because of the plant location, which necessitates shipping large quantities of limestone and gypsum long distances. Reagent costs could be reduced if (1) sources of limestone and gypsum in close proximity to the plant were identified and developed and (2) the quantity of reagents used in the roast feed was reduced. (Table 4 indicates that a 5:2:2 ratio of clay, limestone, and gypsum is as effective as a 5:3:3 mix.) In addition, high fuel costs associated with roasting would be reduced by decreasing the quantity of limestone and gypsum in the roast feed.

## DISCUSSION

PRU studies demonstrated a process for recovering lithium from the lithiumenriched clays of the McDermitt caldera. The process involves the following operations: (1) roasting a mixture of clay, limestone, and gypsum, (2) leaching the calcine with water, (3) concentrating the leach solution by evaporation, and (4) adding $\mathrm{Na}_{2} \mathrm{CO}_{3}$ to the concentrated solution to precipitate $\mathrm{Li}_{2} \mathrm{CO}_{3}$. About 80 pct of the lithium in the clay was recovered as 99-pct-pure $\mathrm{Li}_{2} \mathrm{CO}_{3}$ 。

Based on the $\$ 1.48 / 1 \mathrm{~b} \mathrm{Li} 2 \mathrm{CO}_{3}$ selling price, the process is not economical because of high reagent, utility, and depreciation costs. Reagent and utility costs associated with feed preparation and roasting can be reduced by using a 5:2:2 roast mix rather than 5:3:3; also, a reduction in reagent shipping charges, through reduced reagent requirements and/ or development of closer reagent sources, would significantly decrease process operating costs. However, these savings would not make the process economic.

## REFERENCES

1. Chilenskas, A. A., G. J. Bernstein, and R. O. Ivins. Lithium Requirements for High-Energy Lithium-Aluminum/IronSulfide Batteries for Load-Leveling and Electric-Vehicle Applications. Ch. in Lithium Resources and Requirements by the Year 2000, ed. by J. D. Vine. U.S. Geol. Surv. Prof. Paper 1005, 1976, pp. 5-9.
2. Cooper, J. F., I. Y. Borg, L. G. O'Conne11, E. Behrin, B. Rubin, and H. J. Wiesner. Lithium Requirements for Electric Vehicles Using Lithium-Water-Air

Batteries. Ch. in Lithium Resources and Requirements by the Year 2000, ed. by J. D. Vine. U.S. Geol. Surv. Prof. Paper 1005, 1976, pp. 9-12.
3. Vine, J. D. The Lithium-Resource Enigma. Ch. in Lithium Resources and Requirements by the Year 2000, ed. by J. D. Vine. U.S. Geol. Surv. Prof. Paper 1005, 1976, pp. 35-37.
4. Searls, J. P. Lithium. Ch. in Minerals Facts and Problems. BuMines Bul1. 671, 1980, pp. 521-534.
5. Davidson, C. F. Recovery of Lithium From Clay by Selective Chlorination. BuMines RI 8523, 1981, 19 pp. 6. . Recovery of Lithium From Low-Grade Ores. U.S. Pat. 4,285,914, Aug. 25, 1981.
7. Edlund, V. E. Lime-Gypsum Processing of McDermitt Clay For Lithium Recovery. BuMines RI 8832, 1983, 15 pp.
8. May, J. T., C. F. Davidson, and V. E. Edlund. Extracting Lithium

From McDermitt Clay. Ch. in Light Metals 1982. TMS-AIME, Warrendale, PA, pp. 1159-1168.
9. May, J. T., D. S. Witkowsky, and D. C. Seide1. Extracting Lithium From Clays by Roast--Leach Treatment. BuMines RI 8432, 1980, 16 pp.
10. Glanzman, R. K., J. J. Rytuba, and J. H. McCarthy, Jr. Lithium in the McDermitt Caldera, Nevada and Oregon. Energy, v. 3, No. 3, 1978, pp. 347-353.

APPENDIX A.--MATERIAL BALANCE FOR PRU TEST ${ }^{1}$
TABLE A-1. - Feed preparation

| Stream | Li | K | Na | Ca | Mg | $\mathrm{SO}_{4}$ | $\mathrm{CO}_{3}$ | $\mathrm{H}_{2} \mathrm{O}^{2}$ | Other | Total |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Wt, g: |  |  |  |  |  |  |  |  |  |  |
| 1. | 108.1 | 647.3 | 115.9 | 327.8 | 1,645.15 | 12.75 | 254.9 | 842.5 | 14,255.6 | 18,210 |
| 2. | 0 | 8.63 | 4.32 | 2,568.02 | 10.79 | 5,729.49 | 0 | 2,148.29 | 320.46 | 10,790 |
| 3. | 0 | 11.46 | 6.88 | 4,389.2 | 150.13 | 3.44 | 6,898.9 | 0 | 0 | 11,460 |
| $4^{3}$. | 108.1 | 667.39 | 127.1 | 7,285.02 | 1,806.07 | 5,745.68 | 7,153.8 | 2,990.79 | 14,576.06 | 40,460 |
| wt pct: | 0.59 | 3.55 | 0.64 | 1.8 | 9 | 0.07 | 1.4 | 4.6 | 78.28 | 100 |
| 2. | 0 | . 08 | . 04 | 23.8 | . 1 | 53.1 | 0 | 19.91 | 2.97 | 100 |
| 3. | 0 | . 1 | . 06 | 38.3 | 1.31 | . 03 | 60.2 | 0 | 0 | 100 |
| $4^{4} \ldots \ldots$. | . 27 | 1.65 | . 31 | 18.01 | 4.46 | 14.2 | 17.68 | 7.39 | 36.03 | 100 |

${ }^{1}$ Streams are identified in figure 2.
${ }^{2}$ The clay-limestone-gypsum mixture was pelletized with 6.587 L water. The pellets were dried at $70^{\circ} \mathrm{C}$ to drive off this water.
${ }^{3}$ The weight of stream 4 represents the combined weights of streams $1-3$.
${ }^{4}$ The composition of stream 4 represents the overall composition of streams 1-3.
TABLE A-2. - Roast

| Stream | Li | K | Na | Ca | Mg | $\mathrm{SO}_{4}$ | $\mathrm{CO}_{3}$ | F | $\mathrm{H}_{2} \mathrm{O}$ | Other | Tota= ${ }^{1}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Wt, g: |  |  |  |  |  |  |  |  |  |  |  |
| 4. | 108.1 | 667.39 | 127.1 | 7,285.02 | 1,806.07 | 5,745.68 | 7,153.8 | 509.8 | 2,990.79 | 14,066.26 | 40,460 |
| 5. | 108.1 | 667.39 | 127.1 | 7,285.02 | 1,806.07 | 5,493.93 | 0 | 276.5 | 0 | 16,015.89 | 31,780 |
| wt pet: |  |  |  |  |  |  |  |  |  |  |  |
| 4. | . 27 | 1.65 | .31 | 18.01 | 4.46 | 14.2 | 17.68 | 1.26 | 7.39 | 34.77 | 100 |
| 5........ | . 34 | 2.1 | . 4 | 22.92 | 5.68 | 17.29 | 0 | . 87 | 0 | 50.4 | 100 |

${ }^{1}$ The difference in total weight between the streams was accounted for in the roaster offgas. Excluding combustion gases, the offgas contained $167.83 \mathrm{~g} \mathrm{SO}_{2}, 41.96 \mathrm{~g} \mathrm{O}_{2}, 5,246.12 \mathrm{~g} \mathrm{CO} 2,2,990.79 \mathrm{~g} \mathrm{H}_{2} \mathrm{O}$, and 233.3 g F .

TABLE A-3. - Leach


NAp Not applicable.
${ }^{1}$ The filter cake was washed with 48 L water.
${ }^{2}$ The total volume of stream 6 was 48.04 L solution with a specific gravity of 1.025 .
${ }^{3}$ The total volume of stream 8 was 33 L solution with a specific gravity of 1.05 .
TABLE A-4. - Evaporation

| Stream | Li | K | Na | Ca | Mg | $\mathrm{SO}_{4}$ | $\mathrm{CO}_{3}$ | $\mathrm{H}_{2} \mathrm{O}^{1}$ | Total |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Wt, g: |  |  |  |  |  |  |  |  |  |
| 8................. | 90.4 | 428.99 | 99 | 19.82 | 0.99 | 1,405.97 | 0.0 | 32,605 | 34,650.17 |
| 9. | . 45 | 1 | . 5 | 19.79 | . 99 | 2.27 | 34.06 | 25.6 | 84.66 |
| 10. | 119.3 | 681.97 | 237.86 | . 098 | ND | 2,007.13 | 91.38 | 7,813.98 | 10,952.17 |
| 14. | 7.09 | 10.57 | 10.57 | . 014 | ND | 35.07 | 30.41 | 2,798.28 | 2,892 |
| 16. | 22.26 | 243.41 | 128.79 | . 054 | ND | 568.36 | 95.48 | 5,217.7 | 6,276 |
| $\begin{array}{r} \text { wt pct: } \\ \text { 9... } \end{array}$ | . 53 | 1.18 | . 59 | 23.38 | 1.17 | 2.68 | 40.23 | 30.24 | 100 |
| $\mathrm{g} / \mathrm{L}$ : |  |  |  |  |  |  |  |  |  |
| 8.................. | 2.74 | 13 | 3 | . 6 | . 03 | 42.61 | 0 | NAp | NAP ${ }^{2}$ |
| 10. | 13.42 | 76.71 | 26.26 | . 01 | <. 01 | 225.77 | 10.33 | NAp | Nap ${ }^{3}$ |
| 14. | 2.5 | 3.73 | 3.73 | . 005 | <. 01 | 12.37 | 10.7 | NAP | NAP ${ }^{4}$ |
| 16................. | 4.11 | 44.99 | 23.81 | . 01 | <0.1 | 105.06 | 17.65 | NAp | NAP ${ }^{5}$ |

NAp Not applicable. ND Not determined.
${ }^{1}$ Solution was concentrated by evaporating approximately 32.8 L water.
${ }^{2}$ The total volume of stream 8 was 33 L solution with a specific gravity of 1.05 .
${ }^{3}$ The total volume of stream 10 was 8.89 L solution with a specific gravity of 1.232 .
${ }^{4}$ The total volume of stream 14 was 2.835 L solution with a specific gravity of 1.02 .
${ }^{5}$ The total volume of stream 16 was 5.41 L soluticn with a specific gravity of 1,16 .

TABLE A-5. - Product precipitation

| Stream | Li | K | Na | Ca | $\mathrm{SO}_{4}$ | $\mathrm{CO}_{3}$ | $\mathrm{H}_{2} \mathrm{O}^{1}$ | Total |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Wt, g: |  |  |  |  |  |  |  |  |
| 10. | 119.39 | 681.97 | 237.86 | 0.098 | 2,007.13 | 91.83 | 7,813.98 | 10,952.17 |
| 11. | 0 | 0 | 321.7 | . 146 | 0 | 419.83 | 0 | 741.68 |
| 12. | 88.59 | . 76 | . 76 | . 09 | 2.52 | 379.81 | 104.43 | 576.96 |
| 13. | 23.62 | 670.64 | 548.23 | . 14 | 1,969.54 | 101.44 | 7,709.55 | 11,023.2 |
| 14. | 7.09 | 10.57 | 10.57 | . 014 | 35.07 | 30.41 | 2,798.28 | 2,892 |
| wt pct: |  |  |  |  |  |  |  |  |
| 11. | . 0 | . 0 | 43.4 | . 02 | . 0 | 56.6 | . 0 | 100 |
| 12. | 15.35 | .13 | . 13 | . 016 | . 44 | 65.83 | 18.1 | 100 |
| $\mathrm{g} / \mathrm{L}$ : |  |  |  |  |  |  |  |  |
| 10........... | 13.42 | 76.71 | 26.76 | . 01 | 225.77 | 10.33 | NAp | Nap ${ }^{2}$ |
| 13.......... | 2.74 | 77.8 | 63.6 | . 016 | 228.48 | 11.76 | NAP | Nap ${ }^{3}$ |
| 14........... | 2.5 | 3.73 | 3.73 | . 005 | 12.37 | 10.7 | NAP | NAp ${ }^{4}$ |

NAp Not applicable.
${ }^{1}$ Product was washed with approximately 2.8 L hot water.
${ }^{2}$ The total volume of stream 10 was 8.89 L solution with a specific gravity of 1.232 .
${ }^{3}$ The total volume of stream 13 was 8.62 L solution with a specific gravity of 1.279 .
${ }^{4}$ The total volume of stream 14 was 2.835 L solution with a specific gravity of 1.02 .
TABLE A-6. - Crystallization

| Stream | Li | K | Na | Ca | $\mathrm{SO}_{4}$ | $\mathrm{CO}_{3}$ | $\mathrm{H}_{2} \mathrm{O}$ | Total |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Wt, g: |  |  |  |  |  |  |  |  |
| 13. | 23.62 | 670.64 | 548.23 | 0.14 | 1,969.54 | 101.44 | 7,709.55 | 11,023.1 |
| 15. | 1.36 | 427.23 | 419.44 | . 086 | 1,401.18 | 5.96 | 2,491.85 | 4,747 |
| 16........... | 22.26 | 243.41 | 128.79 | . 054 | 568.36 | 95.48 | 5,217.7 | 6,276 |
| wt pct: 15............. | . 03 | 9 | 8.84 | . 002 | 29.52 | . 12 | 52.49 | 100 |
| $\mathrm{g} / \mathrm{L}:$ |  |  |  |  |  |  |  |  |
| 13. | 2.74 | 77.8 | 63.6 | . 016 | 228.48 | 11.76 | NAp | NAp ${ }^{1}$ |
| 16............ | 4.11 | 45 | 23.81 | . 01 | 105.06 | 17.65 | NAp | $N A p^{2}$ |

NAp Not applicable.
${ }^{1}$ The total volume of stream 13 was 8.62 L solution with a specific gravity of 1.279 .
${ }^{2}$ The total volume of stream 16 was 5.41 L solution with a specific gravity of 1.16 .

## APPENDIX B.--PROCESS ECONOMICS

By D. A. Kramer ${ }^{1}$

## PROCESS DESCRIPTION

The proposed process is designed to recover lithium, in the form of $\mathrm{Li}_{2} \mathrm{CO}_{3}$, from McDermitt clay. As-mined clay is crushed to minus $3 / 16$-in and mixed with minus 3/16-in limestone and gypsum in a 5:3:3 ratio (clay-limestone-gypsum). These three materials are ground to minus 100 mesh, pelletized with water, and dried. Dried pellets are roasted at $900^{\circ}$ C to react the gypsum with the lithium, producing lithium sulfate. Leaching the roasted pellets with water at ambient temperature extracts the soluble lithium sulfate and any other soluble sulfates formed during roasting. The leach slurry is filtered to separate the insoluble residue from the leachate.

Leachate is concentrated from $3 \mathrm{~g} / \mathrm{L} \mathrm{Li}$ to $13 \mathrm{~g} / \mathrm{L} \mathrm{Li}$ by evaporation, precipitating calcium impurities as a carbonate at the same time; these are separated from the solution by filtration. Mixing the concentrated solution with soda ash at $95^{\circ} \mathrm{C}$ precipitates lithium as $\mathrm{Li}_{2} \mathrm{CO}_{3}$. This slurry is filtered, and the solids are washed and dried to yield a $\mathrm{Li}_{2} \mathrm{CO}_{3}$ product. Total recovery is about 83 pet of the lithium in the clay.

After lithium precipitation, the remaining solution is cooled to $0^{\circ} \mathrm{C}$ to crystallize $\mathrm{Na}_{2} \mathrm{SO}_{4}$ and $\mathrm{K}_{2} \mathrm{SO}_{4}$ impurities that are leached from the clay. The crystal slurry is filtered, the crystals are discarded, and the mother liquor is recycled to the evaporator.

The proposed plant is designed to process 1,000 ton/d of McDermitt clay with the composition shown in table B-1. The material balance for the proposed process is detailed in figures B-1 through B-6. (Units are tons per day.)

[^2]TABLE B-1. .. Composition of McDermitt clay (dry basis), weight percent

| Li | 0.6 |
| :---: | :---: |
| K。 | 3.7 |
| Na | . 6 |
| Mg . | 9 |
| Fe . | 2.5 |
| Ca | 1.8 |
| A1 | 3.1 |
| F. | 2 |
| $\mathrm{SiO}_{2}$ | 53 |
| Other | 23.7 |
| Total | 100.0 |

## ECONOMICS

Estimated capital and operating costs are based on the preceding process description.

## Capital Costs

The capital cost estimate is of the general type called a study estimate by Weaver and Bauman. ${ }^{2}$ This type of estimate, prepared from a flowsheet and a minimum of equipment data, can be expected to be within 30 pct of the actual plant cost.

The estimated capital cost on a second quarter 1984 basis (Marshall and Swift [M and S] index of 781.7) for a plant processing 1,000 ton/d of McDermitt clay is about $\$ 106$ million, as shown in table B-2. Most of the plant is designed to operate 3 shifts per day, 7 days per week, $330 \mathrm{~d} / \mathrm{yr}$. Some of the clay preparation facilities only operate 1 shift per day, 7 days per week. The remainder of the time is for scheduled and unscheduled downtime.

[^3]
## TABLE B-2. - Estimated capital cost ${ }^{1}$

Fixed capital:
Feed preparation section. ..... \$15,570,700
Roasting section. ..... 9,316,500
Leaching section. ..... 20,250,400
Evaporation section. ..... 9,347,300
Lithium recovery section. ..... 2,081,900
Crystallization section ..... 3,420,800
Tailings pond ..... 1,701,400
Steamplant ..... 406,800
Subtotal
6,209,600
Plant facilities: 10 pct of above subtotal
7,451,500
Plant utilities, 12 pct of above subtotal.
75,756,900
Escalation costs during construction ..... $6,112,100$
Total plant cost 81,869,000
Land cost.81,869,000
Interest during construction period. ..... 14,095,800
Fixed capital cost ..... 95,964,800
Working capital:
Raw material and supplies ..... 1,081,500
Product and in-process inventory ..... 2,993,800
Accounts receivable ..... 2,993,800
Available cash ..... $\frac{2,302,300}{9,371,400}$
Capitalized startup costs. Subtotal

| 959,600 |
| ---: |
| $10,331,000$ |
| $106,295,800$ |

${ }^{1}$ Basis: $M$ and $S$ equipment cost index of 781.7.

Equipment costs used in this estimate are based on informal cost quotations from equipment manufacturers and on capacity-cost data. Cost data are brought up to date by the use of inflation indexes. In developing the plant capital costs, corrosion-resistant materials of construction were used where appropriate. For example, the tanks are rubber-lined to withstand the corrosive properties of the sulfate solution. The tailings pond is designed with a $2-y r$ capacity. It is assumed that after 2 yr the mine site will be developed to dis pose of the remainder of the processing residue as backfill.

Factors for piping, etc., except for the foundation and electrical factors, are assigned to each section, using as a basis the effect fluids, solids, or a
combination of fluids and solids may have on the process equipment. The foundation factor is estimated for each piece of equipment individually, and a factor for the entire section is calculated from the totals. The electrical factor is based on the motor horsepower requirements for each section. A factor of 10 pct, referred to as miscellaneous, is added to each section to cover minor equipment and construction costs that are not shown with the equipment listed.

For each section, the field indirect cost, which covers field supervision, inspection, temporary construction, equipment rental, and payroll overhead, is estimated at 10 pet of the direct cost. Engineering cost is estimated at 10 pct, and administration and overhead cost is estimated at 5 pct of the construction
cost. A contingency allowance of 15 pct and a contractor's fee of 5 pct are included in the section costs.

The costs of plant facilities and plant utilities are estimated as 10 and 12 pct, respectively, of the total process section costs and include the same field indirect costs, engineering, administration and overhead, contingency allowance, and contractor's fee as are included in the section costs. Included under plant facilities are the cost of material and labor for auxiliary buildings such as offices, shops, laboratories, and cafeterias, and the cost of nonprocess equipment such as office furniture and safety, shop, and laboratory equipment. Also included are labor and material costs for site preparation such as clearing, grading, drainage, roads, and fences. The costs of water, power, and steam distribution systems are included under plant utilities. Cost for the plant owner's supervision is not included in the capital cost of the proposed plant.

Working capital is defined as the funds in addition to fixed capital, land investment, and startup costs that must be provided to operate the plant. Working capital, shown in table $\mathrm{B}-2$, is estimated from the following items: (1) Raw material and supplies inventory (cost of raw material and operating supplies for 30 days), (2) product and in-process inventory (total operating cost for 30 days), (3) accounts receivable (total operating cost for 30 days), and (4) available cash (direct expenses for 30 days).

Capitalized startup costs are estimated as 1 pet of the fixed capital costs. Land investment is not included in this estimate.

## Operating Costs

The estimated operating costs (table $B-3$ ) are based on an average of 330 days of operation per year over the life of the plant. The operating costs are divided into direct, indirect, and fixed costs.

Direct costs include raw materials, utilities, direct labor, plant maintenance, payroll overhead, and operating
supplies. Raw materials and utility requirements per pound of $\mathrm{Li}_{2} \mathrm{CO}_{3}$ are shown in table B-4. The direct labor cost is estimated on the basis of assigning 4.2 employees for each position that operates $24 \mathrm{~h} / \mathrm{d}, 7$ days per week and 1.4 employees for each position that operates $8 \mathrm{~h} / \mathrm{d}, 7$ days per week. The cost of labor supervision is estimated as 15 pct of the labor cost.

Plant maintenance is separately estimated for each piece of equipment and for the buildings, electrical system, piping, plant utility distribution systems, and plant facilities.

Payroll overhead, estimated as 35 pct of direct labor and maintenance labor, includes vacation, sick leave, social security, and fringe benefits. The cost of operating supplies is estimated as 20 pct of the cost of plant maintenance.

Indirect costs are estimated as 40 pct of the direct labor and maintenance costs. The indirect costs include the expenses of control laboratories, accounting, plant protection and safety, plant administration, marketing, and company overhead. Research and overall administrative costs outside the plant are not included.

Fixed costs include the cost of taxes (excluding income taxes), insurance, and depreciation. The annual costs of both taxes and insurance are each estimated as 1 pct of the plant construction cost. Depreciation is based on a straight-line, 20-yr period.

The estimated annual operating cost of a plant processing 1,000 ton/d of McDermitt clay is about $\$ 36$ million, as shown in table $\mathrm{B}-3$. This corresponds to a cost of about $\$ 2.12 / 1 \mathrm{~b} \mathrm{Li}_{2} \mathrm{CO}_{3}$. No charge has been included for the clay.

The estimated operating cost of \$2.12/ $1 b \mathrm{Li}_{2} \mathrm{CO}_{3}$ is about $1-1 / 2$ times the current selling price of $\$ 1.48 / 1 \mathrm{~b}$.

## ECONOMIC EVALUATION

After examining the high-cost areas in the proposed process, it appears that the largest cost factors are the limestone and gypsum costs. These costs contribute over 30 pct to the total operating cost. Raw material costs are high because of

TABLE B-3. - Estimated annual operating cost

|  | Annual cost | Cost per pound of $\mathrm{Li}_{2} \mathrm{CO}_{3}$ |
| :---: | :---: | :---: |
| Direct cost: |  |  |
| McDermitt clay at \$0.00/ton.............................. | \$0 | \$0.0 |
| Limestone at \$26.50/ton. | 5,771,700 | . 366 |
| Gypsum at \$25.50/ton | 5,225,700 | . 305 |
| Soda ash at \$90/ton..................................... | 1,277,100 | . 074 |
| Replacement balls for grinding at \$0.27/1b............ | 143,400 | . 008 |
| Chemicals for steamplant water treatment............... | 5,900 | . 001 |
| Total | 12,423,800 | . 724 |
| Utilities: |  |  |
|  | 793,000 | . 046 |
| Process water at $\$ 0.25 / \mathrm{Mgal} . . .$. ......................... | 75,400 | . 004 |
| Heavy oil at \$0.80/gal. | 7,892,400 | . 460 |
| Tota1................................................ | 8,761,700 | . 510 |
| Direct labor: |  |  |
| Labor at \$9/h.............................................. | 1,104,500 | . 064 |
| Supervision, 15 pet of labo | 165,700 | . 010 |
| Total.......................... . . . . . . . . . . . . . . . . . . . | 1,270,200 | . 074 |
| Plant maintenance: |  |  |
| Labor | 1,670,200 | . 097 |
| Supervision, 20 pct of maintenance labor............... | 334,000 | . 019 |
| Materials.................................................. | 1,670,000 | . 097 |
| Total | 3,674,400 | . 213 |
| Payroll overhead, 35 pct of above payroll................ | 1,146,000 | . 067 |
| Operating supplies, 20 pct of plant maintenance......... | 734,900 | . 043 |
| Total direct cost | 28,011,000 | 1.631 |
| Indirect cost, 40 pct of direct labor and maintenance... | 1,977,800 | . 115 |
| Fixed cost: |  |  |
| Taxes, 1 pct of total plant cost. | 818,700 | . 048 |
| Insurance, 1 pct of total plant cost................... | 818,700 | . 048 |
| Depreciation, 20-yr life.................................... | 4,798,200 | . 280 |
| Total operating cost.............................. | 36,424,400 | 2.122 |

TABLE B-4. - Raw material and utility requirements per pound of $\mathrm{Li}_{2} \mathrm{CO}_{3}$

Raw materials:
McDermitt clay..............ton.. 0.020
Limestone...................ton.. . 013
Gypsum......................ton.. . 012
Soda ash.....................ton.. . 001
Replacement balls
for grinding..................lb.. . 031
Utilities:
Electric power................kW•h.. 1.542
Process water..............Mgal.. . 018
Heavy oil...................gal.. . 575
the plant location. It has been assumed that the quantity of limestone needed would be available from a deposit in Lyon County, NV, and that the required quantity of gypsum would be available from deposits in either Pershing or Washoe County in Nevada. Shipping charges from these distances have been included in the costs of these raw materials. These shipping charges significantly increase the costs of what would otherwise be relatively inexpensive raw materials.

To minimize the impact of these costs, the ratios of limestone and gypsum to the
clay must be minimized without affecting in reducing the fuel requirements for lithium extraction. This would also aid roasting.


FIGURE B-1. - Feed preparation section (tons per day).


FIGURE B-2. - Roasting section (tons per day).


FIGURE B.3. - Leaching section (tons per day).


FIGURE B-4. - Evaporation section (tons per day).


FIGURE B-5. - Lithium recovery section (tons per day).


FIGURE B-6. - Crystallization section (tons per day).


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[^1]:    ${ }^{2}$ Underlined numbers in parentheses refer to items in the list of references preceding the appendixes.
    ${ }^{3}$ Estimates of lithium demands have been revised because the lithium batteries market has not expanded as anticipated. Also, fusion power programs have encountered technological as well as funding problems, which has further clouded the

[^2]:    ${ }^{1}$ Chemist, Avondale Research Center, Bureau of Mines, Avondale, MD (now commodity specialist, Bureau of Mines, Washington, DC).

[^3]:    ${ }^{2}$ Weaver, J. B., and H. C. Bauman. Cost and Profitability Estimation. Sec. 25 in Perry's Chemical Engineers' Handbook, ed. by R. H. Perry and C. H. Chilton. McGraw-Hill, 5th ed., 1973, p. 47.

