7 Process Engineering
ABSTRACT: This paper presents some results of flotation modelling and simulation. The modernisation of technology is very slow in this field and the only way of contributing to the increment of extraction performances is to know the technological parameters and capacity reserves as precisely as possible. Some of the representative dependencies characterising the flotation process were simulated using MATLAB SIMULINK programs. Analysis and comparison of the results of simulations carried out enabled the determination and formulation of some of the flotation process characteristics.

1 THE STUDY OF A FLOTATION TWO-PHASE MODEL

The model considers the froth and pulp as two distinct phases dynamically equilibrated as given in Figure 1.

The notation in Figure 1 is as follows:

- \( M \) - floated material mass;
- \( V \) - pulp volume, excepting air;
- \( r, s \) - the ratio of the pulp volume (including air) to the airless volume, respectively and of the froth volume (including air) and the airless volume respectively;
- \( Q \) - volumetric flow rate, excluding air;
- \( C^* \) - concentration (ratio of the mass to volume unit, including the air in the pulp);
- \( a \) - speed constant, regarding the concentrate transfer from the pulp to the froth;
- \( b \) - speed constant, regarding the concentrate transfer from the froth to the pulp;
- \( p, s, c, t \) - indexes, referring to the pulp, froth, concentrate and waste.

The differential equation reflecting the pulp composition change for ideal mixing cells is:

\[
\frac{dM_p}{dt} = QC^* - a V_p C^* - b s V_s C_s + a s V_p C_p = \frac{dC^*}{dt} V_p, r \quad (0)
\]

The differential equation reflecting the change in the froth composition for the same cells is:

\[
\frac{dM_f}{dt} = -s V_s C_s - b s V_s C_s + a r V_p C_p = \frac{dC^*}{dt} V_f \quad (2)
\]

The transfer functions determined are given as follows:

\[
H_1(s) = \frac{M_p(s)}{Q(s)} = \frac{k_1}{T_p s + 1} \quad (3)
\]

\[
H_2(s) = \frac{M_f(s)}{C(s)} = \frac{k_2}{T_f s + 1} \quad (4)
\]

\[
H_3(s) = \frac{M_p(s)}{Q(s)} = \frac{k_3}{T_p s + 1} \quad (5)
\]

\[
H_4(s) = \frac{M_f(s)}{Q(s)} = \frac{k_4}{T_f s + 1} \quad (6)
\]

Where \( T_p \) and \( T_s \) are time constants; \( k_1, k_2, k_3, \) and \( k_4 \) proportionality constants depending on \( V_p, V_s, Q_p, Q_f \) following the same notation as in Figure 1.

Figure 1. Flotation cell block diagram.
Mineral flotation of the raw material exploited in the Baia-Mare area, processed in the mineral processing plant from this town was studied. A primary flotation cell was considered, for which we evaluated the characteristic values of die constants m the transfer functions were evaluated (Table 1).

Table 1. Values for time and proportionality constants in the transfer functions.

<table>
<thead>
<tr>
<th>Constant</th>
<th>Relation</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$T_p$</td>
<td>$V_p/Q_0$</td>
<td>0.004808</td>
</tr>
<tr>
<td>$T_1$</td>
<td>$V_1/Q_0$</td>
<td>0.025749</td>
</tr>
<tr>
<td>$k_1$</td>
<td>$T_p/C^*$</td>
<td>0.003468</td>
</tr>
<tr>
<td>$k_2$</td>
<td>$T_p/Q$</td>
<td>1.153920</td>
</tr>
<tr>
<td>$k_3$</td>
<td>$T_1/C^*$</td>
<td>0.013572</td>
</tr>
<tr>
<td>$k_4$</td>
<td>$T_1/Q$</td>
<td>6.179760</td>
</tr>
</tbody>
</table>

The response of the considered system was investigated, for the unit input signal of the feeding flow rate $Q$ and the concentration $C^*$, as well as for the "white noise" signal, in the case of $C^*$.

The results of the simulation are presented in Figure 2, Figure 3 and Figure 4.

Analysing the system response, it can be concluded that:

- the stabilising time for the pulp composition is 1.5-2 min., a sudden variation of the feeding flow rate or the feeding concentration being reflected in the system for about 2 min.; the concentration influence is more significant:
- the froth composition stabilisation time is about 6 min;
- the "white noise" stochastic variation of the flow rate influences the froth composition more.

Using the transfer functions which allow us to study the mechanism of the mineral particle transfer from the pulp to the froth and the reverse, the optimum time of the flotation in a cell can be determined, using the stabilisation time of the process.

Considering the estimating relations for the time and proportionality constants, it can be stated that these depend on the pulp flow rate, processed mineral composition, useful elements content in the raw material, weight extraction, aeration and dilution.

Our conclusion from the study performed is that; if a certain material and a given flotation technological hit the optimisation parameters are the aeration and the feedme dilution.
3 CONCLUSIONS

Flotation can be considered as physical and chemical separation process of solid products, by establishing a contact in three phases: the floating mineral, the liquid phase and air.

The great number of parameters involved in flotation, due to the high complexity of the process, require a new kind of approach to this process study.

The development of computer hardware and, most importantly, software, allows a new method of experimental research by simulation.

In order to simulate a primary flotation cell, a two-phase model can be adopted, considering the pulp and the froth as two distinct phases.

Use of MATLAB SIMULINK software enabled obtaining the process response when the feed flow rate and composition exhibited a sudden step or stochastic variation.

Using the transfer functions which allow study of the mechanism of the mineral particle transfer from the pulp to the froth and reverse, the optimum time of flotation in a cell can be determined, using the stabilisation time of the process.

Considering the estimating relations for the time and proportionality constants, it can be stated that these depend on: the pulp flow rate, processed mineral composition, useful elements content in the raw material, weight extraction, aeration and dilution.

The main conclusion of this study can be stated as; for a certain material and a given flotation technological line, the optimisation parameters are the aeration and the feeding dilution.

REFERENCES


ABSTRACT: Fine coal slimes (less than 0-0.2 mm) present great difficulty in cleaning. Their flotation ability depends on their size and physico-chemical characteristics. In particular, the degree of metamorphism of the coal, and the proportion and ash content of super-fine coal particles (less than 50 μm) produces quite a strong effect on the selectivity of flotation. The flotation properties of ultrafine slimes also depend on their nature: whether they are primary slimes in feed coal, or whether they form in circulating and slime water as a result of coarse and fine fraction washing. A series of investigations on the flotation of coals at different stages of metamorphism and with various characteristics of fine slimes was carried out. The effect of aerohydrodynamic and design parameters of multisectional column apparatus on the selectivity of flotation separation of coal slimes and fly ash of power plants was studied.

1 INTRODUCTION

Preparation plants have to deal with the problem of "fine slimes" in respect of their circulation in the process flow sheets and the difficulty of selective separation.

Of traditional cleaning methods of slimes less than 0.5 mm material, flotation is the most efficient. As the fines (<0.5 mm) content in the feed run-of-mine coal increases to 20-35%, the proportion of ultrafine material (<50 μm) in flotation feed reaches 40-60%, and in most cases the ash content of fine slimes less than 50 μm is 5-10% higher than that of fractions over 50 μm.

The problem of efficient separation of fine coal slimes arises when solving a range of non-traditional tasks, which include:

1) the production of super-pure concentrates to prepare water-coal fuel;
2) deep cleaning so as to desulphurize power-generating coals;

In the performance of these tasks, coal grinding to less than 0.2 mm is required to obtain adequate liberation of coal, and the 0-0.05 mm fraction content can be as high as 50-85%.

3) the recovery of carbon-bearing particles from the fly ash of power-generating plants for their subsequent utilization in construction materials production.

When coal dust is burnt, 50-75% of the fly ash of power-generating plants comprises particles under 0.05 mm in size, while the average particle size is less than 0.2 mm.

The aim of this work was to evaluate the effect of fine particles (less than 50 μm) of coals of different degrees of metamorphism on the selectivity of slimes separation during flotation, and to develop efficient processes and conditions for fine slimes cleaning.

A series of studies was carried out on the flotation of coals of different degrees of metamorphism with various characteristics of fine slimes. The effect of aerohydrodynamic and design parameters of multisectional column flotation machines on the efficiency of the desulphurization of anthracite slimes and separation of the pure mineral part (>95% ash) from the fly ash of power-generating plants was investigated.

2 SUMMARY OF THE STATE OF THE ART

Ultrafine particles (<0.05 mm) are characterized by specific physical and physico-chemical properties. A small particle size signifies their significant specific surface and higher adsorption ability. As failure during grinding most often occurs along micro-defects of grains and those in the crystal lattice, fine particles possess higher structural homogeneity, lower significance of external defects and higher significance of surface layers.

The presence of fine slimes in the slurry results in the following technological effects:
• contamination of froth product with fine fractions of waste rock;
• worsening of coarse grain flotation and deterioration of waste grade;
• higher reagent consumption;
• decrease of flotation rate.

In order to enhance the efficiency of the separation of coals containing fine slimes, the following methods are used:
• regulating reagents;
• fractional reagent feeding;
• special conditioning regimes (time and rate of mixing, density of slurry during its treatment with reagents);
• corresponding flotation machines allowing to variation in aerohydrodynamic conditions;
• special technologies (for example, flocculation-flotation method, etc.)

The applicability of a given method can be established by investigating the behaviour of fine slimes during flotation. Applicability also depends on requirements to separation indices, i.e., to concentrate and waste quality. The content of slimes (including so-called secondary slimes) in coal coming for cleaning varies from 10 to 35 % with an ash content of about 20-40%. These slimes contain proportions of 20 to 70% fine particles of less than 50 μm in size, with ash contents of 25-51%.

3 RESULTS OF INVESTIGATIONS

In order to have the opportunity to compare the flotation ability of different coals, the concept of "maximum flotation ability" is used. This is derived from the release-analysis curve based on the maximum recovery of combustible mass into froth product, and the separation of froth products into fractions with different ash contents by means of many cleaning stages. In this case, the flotation ability of coal slime is virtually determined by slime properties.

Screenings of differently metamorphosed coals with different proportions and ash contents of finely dispersed fraction (under 50 μm) were considered, and the indices of the studied coal separation were compared with the release-analysis. As a criterion of flotation ability, it was assumed that the coefficient of relative efficiency of maximum separation can be found from the formula:

$$E_o = \frac{E_{CM}^{CM}}{E_{CM}^{IM}} \times 100\%$$  \hspace{1cm} (1)

where $E_{CM}^{CM}$ - recovery of combustible mass to concentrate, %; and $E_{CM}^{IM}$ - recovery of incombustible mass to wastes, %.

$$E_{CM} = Y_c(100 - A_f) / (100 - A_f)$$  \hspace{1cm} (2)

$$E_{IC}^W = Y_w^* A_w / A_f$$  \hspace{1cm} (3)

where $Y_c$ - maximum yield of concentrate, %; $A_c$ - ash content of concentrate, %; $A_f$ - ash content of feed slime, %; $Y_w$ - yield of wastes, %; and $A_w$ - ash content of wastes, %.

The maximum concentrate yield was found using the curves of maximum flotation ability, including the middlings fractions yield.

The analysis of the data obtained demonstrated that it is not possible to draw a clear conclusion on the separate effect of ash content and fine slimes yield on the variation of efficiency index $E_o$. Assuming as a variable the weight content of ash in fine slimes equal to $Y_r* A_f / 100$, where $Y_r$ - content of 0-50 μm fraction in slimes, and $A_f$ - ash content in 0-50 μm fraction, we find the explicit effect of the degree of metamorphism on the results of maximum separation (Figure 1). Curve 1 characterizes coals of a moderate degree of metamorphism (coking, gas and gas-fat coals - C, G, GF), and curve 2 is characteristic of coals of a higher degree of metamorphism (weakly binding coal, anthracite - WB, A).

![Figure 1 Index of separation efficiency ($E_o$) of maximum flotation ability as a function of weight content of ash m - 50 μm fraction.](image)

The efficiency of separation decreases both at low and high values of $Y_r* A_f / 100$, and the spread in relative efficiency figures at maximum flotation ability is higher for coal of a moderate degree of metamorphism. The maximum values of $E_o$ correspond to 10-12%.

In order to optimize the flotation of coals with high proportion and high ash content of fine slimes, a number of technological parameters for various ash and fine slimes contents were studied. The effect of feed characteristics when changing the reagent regimes, flotation kinetics, hydrodynamic parameters of flotation machines and when using a
flocculation-flotation method was investigated, taking into account the combined effect of these factors on flotation figures.

A series of flocculation-flotation method investigations performed using latex flocculant BS-30F with coals of different degrees of metamorphism and particle size under 200 um revealed positive results with this method application for coking, fat and weakly binding coals.

These factors were investigated in more detail in the context of the solution of a series of practical tasks in order to enhance the efficiency of fine coal slimes flotation.

3.1 Flotation of coking coal slimes

First of all, consider a specific example of the effect of primary and secondary slimes on the flotation of difficult-to-clean coking coals at the preparation plant of Nerungrinsky open cast mine (Yakutsk coal basin).

As these coals are difficult to clean it is necessary to use a flow sheet including coarse fractions grinding to -30 mm, dense-media separation with recovery of middlings and fine slimes flotation also with recovery of middlings. The feed coal has a content of about 21% -0.5 mm slime, with an ash content of 17%. During cleaning using water recycling, the slimes proportion reaches 35%, and the finely dispersed slimes proportion is as high as 30-40% with a 30-39% ash content.

Figure 2 shows $E_0$ and concentrate ash content variation as a function of flotation time in kinetic tests with one-time (curves 1 and 3) and fractional (curves 2 and 4) reagent feeding.

The indices of the relative efficiency of flotation of slimes were found:
- from screenings (0-0.5 mm) of coking coal with 0-50 mm material content $Y_r = 18.1\%$ with ash content of this fraction $A_0 = 18.8\%$ and with initial ash content of screenings $A_F = 15.3\%$ - $Eo^{(1)}$;
- from flotation feed with ash content $A_F = 20.5\%$ with 0-50 um material share $Y_r = 40\%$ with ash content $A_F = 36.2\%$ - $Eo^{(2)}$.

Size grade distribution by products of flotation (yield and ash $Y^h A_{i=0}$ for screenings; $Y^o$, AIKF) for flotation feed was studied, including the recovery of concentrate and middlings at the first stage and recovery of middlings and wastes at the second stage. The analysis of the results obtained demonstrates that a high content and high ash content of fine slimes disturb the selectivity of coarser particle separation (Figure 3), though the overall efficiency of separation of slimes subjected to flotation remains practically the same.

3.2 Flotation of fat and weakly binding coals

It is undesirable to increase the content of fine particles when processing coal with low ash content. The high content of fine particles increases the ash content of concentrate, which reduces the quality of coke. Also, the ash content of middlings increases, which reduces the quality of fines.

Figure 3 shows the index of separation efficiency in narrow particle sizes and distribution of yield and ash of feed material by size $Eo^{(1)}$, $Y^{(1)}$, $A^{(1)}$, -screenings; $Eo^{(2)}$, $Y^{(2)}$, $A^{(2)}$, -flotation feed.

Figure 4 shows the variation in the relative efficiency index of separation into three products with recovery of concentrate and middlings at the first stage and recovery of middlings and wastes at the second stage for coking coal screenings (0-0.5 mm) with different contents of 0-50 um material ($Y^o$) and different ash contents of this material ($A^o$) as a function of the weight content of fine particles ($Y^o A^o/100$) in slimes.

The calculation of the indices of maximum flotation ability and flotation kinetics revealed the specificity of the flotation behaviour of slimes of the
studied coal. It was found that it is possible to enhance separation by fractional feed of the reagents. In this case, the efficiency of separation was 10% higher than when multiple cleaning stages were used.

Figure 4. Indices of relative efficiency of separation into two EQ(2) and three EQ(3) products

Figure 5. Indices of relative efficiency for different separation methods

3.2 Power-generating coals preparation

In Russia, thermal power stations are fueled with practically uncleaned coal, but in order to increase power plants' competitive ability and solve environmental problems, preliminary coal preparation is necessary. This problem is most urgent in the Rostov region, where thermal power plants are fueled with anthracites having a 30-35% ash content and 1.5-2.5% sulphur content.

The possibility of reducing ash and sulphur contents and increasing the calorific value of coal fuel by means of preparation was considered in a specific example of coals burnt at the "Novocherkasskaya" State Regional Power Station and coals being mined from the new deposit at Sadkinskaya mine (Rostov region).

The anthracite culm of 0-0.5 mm coming to the "Novocherkasskaya" State Regional Power Station for burning has an ash content of 31.7% and sulphur content of 1.9%.

The flotation ability of 0-0.5 mm feed slime and that of slime ground to 0-0.2 mm and 0-0.1 mm was determined. The efficiency of separation (Eo) at maximum flotation ability of the studied fractions is practically the same, while in kinetic tests at a fine slime flotation rate 2 times lower the efficiency of separation Eo(tm) sharply decreases.

The relative separation efficiency figure for 0-0.1 mm slimes when using the flocculation-flotation method is Eo = 65.8% at 72.3% concentrate yield with an ash content of 11%. Although the sulphur content in the concentrate remains high (S(Conc) = 1.8%), the total annual sulphur emissions drop by 24% and ash and slag waste generation decreases by 3.6 times owing to reduced specific fuel consumption for the production of 1 MW-h of electric power.

The possibility of reducing ash and sulphur contents and increasing the calorific value of coal fuel by means of preparation was considered in a specific example of coals from the new deposit mined at Sadkinskaya mine (Rostov region). Studies of washability of slimes with a high content of high ash 0-50 μm material (Y = 42.5%, Ar = 40.6%) were performed for the purpose of desulphurization.

It was found that the most promising solution is the use of column multi-section machines in which separation efficiency is enhanced owing to the selection of optimum aerohydrodynamic conditions. The data obtained show that during flotation in column machines, sulphur recovery to concentrate is lower than in a mechanical flotation machine at equal levels of combustible mass recovery and concentrate ash content (Figure 6).
Figure 6. Sulphur and combustible mass recovery to concentrate under different separation conditions.

Table 1. Results of column desulphurization by flotation separation

| Coal slime (0-0.25 mm) of m. Sudkinskaya | N | Yield, % | Ash, % | Sulphur, % | Ash, % | Sulphur, % | Eo, % | Eo
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
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</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>68.7</td>
<td>13.2</td>
<td>0.9</td>
<td>64.87</td>
<td>3.64</td>
<td>84.42</td>
<td>42.50</td>
<td>57.42</td>
</tr>
<tr>
<td>2</td>
<td>78.4</td>
<td>17.3</td>
<td>1.0</td>
<td>72.63</td>
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<td>91.64</td>
<td>52.30</td>
<td>49.15</td>
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<td>3</td>
<td>68.2</td>
<td>14.6</td>
<td>0.8</td>
<td>62.96</td>
<td>3.53</td>
<td>83.18</td>
<td>40.41</td>
<td>55.54</td>
</tr>
<tr>
<td>4</td>
<td>69.2</td>
<td>14.2</td>
<td>0.9</td>
<td>63.57</td>
<td>2.57</td>
<td>85.08</td>
<td>44.03</td>
<td>53.35</td>
</tr>
</tbody>
</table>

- Optimal flotation results: short flow rate = 1.3 cm/sec, air flow rate during rougher and scavenger flotation = 1.6 l/min, air flow rate during cleaning = 2.8 l/min.
- Consumption of agents: Rougher flotation - keroseen 800 g/t, MIBK 200 g/t.
- Scavenging - keroseen 400 g/t, MIBK 150 g/t.
- Concentrate cleaning (1 and 2) - keroseen 150 g/t, MIBK 50 g/t.

3.3 Production of super-pure concentrates to prepare water-coal suspensions

The problem of enhancement of fine fraction flotation becomes more acute for the development of processes of super-pure concentrate production to prepare water-coal fuel. Consider some results of coals flotation at "Cherkasovskaya" preparation plant (weakly binding coal, Kuznetsk basin).

For pure coal liberation, the degree of grinding was chosen so as to obtain the maximum recovery of combustible mass with minimum ash content of the concentrate. The indices of flotation separation of coal ground to different sizes show that coal grain opening takes place at fine grinding to -50 Jm of 85% of the material and when the ash content of this material is close to that of the feed material - 17.1%.

The application of the flocculation-flotation method and concentrate recleaning at quite fine grinding makes it possible to optimize the separation process, increasing the relative efficiency index by 10% (Figure 7).

Figure 7. Variation of efficiency index Eo when using flocculation-flotation separation method.

3.4 Recovery of organic matter from fly ash of power-generating plants

The problem of coal particle recovery from the fly ash of power-generating stations is a separate issue as we deal here with a different coal material that passes
through the stage of burning. Nevertheless, it is a very interesting task related to the environmental and economical problems of recovering the pure mineral part of ash (with carbon content (LOI) of less than 5%) to use it for the production of construction materials and recycling recovered coal matter for burning.

Detailed investigations of fly ash separation were carried out at different power stations. Operating conditions of separation have been developed that include a conditioning regime at a high rate of dense slurry mixing (solid content 200g/l) with the addition of a collector, using a 3-section column flotation machine, making it possible to reproduce in one apparatus the flow sheets of rough, control and cleaner flotation, fractional reagent feeding, counter-current and straight-current slurry and air movement in different sections of the machine, with aeration conditions (consumption and dispersion of air bubbles).

The optimum results of fly ash flotation for a specific case at the "Novocherkasskaya" State Regional Power Station, which burns anthracites, (Rostov regions) are summarized in Table 2.

### Table 2. Results of column flotation separation fly ash.

<table>
<thead>
<tr>
<th>N</th>
<th>Concentrate</th>
<th>Yield %</th>
<th>Ash, %</th>
<th>Ash, %</th>
<th>Ash, %</th>
<th>LOI %</th>
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</thead>
<tbody>
<tr>
<td>1^*</td>
<td>20.4</td>
<td>40.3</td>
<td>79.6</td>
<td>96.6</td>
<td>3.4</td>
<td>81.76</td>
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<tr>
<td>2</td>
<td>17.2</td>
<td>37.3</td>
<td>82.8</td>
<td>95.3</td>
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<td>71.57</td>
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<td>3</td>
<td>21.59</td>
<td>48.6</td>
<td>78.61</td>
<td>95.6</td>
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<tr>
<td>4</td>
<td>20.50</td>
<td>41.50</td>
<td>79.3</td>
<td>96.3</td>
<td>3.7</td>
<td>80.70</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>90.00</td>
<td>72.18</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Optimal results. Slurry preparation prior to rougher flotation is carried out at a stirring speed of 2500 rpm and conditioning time of 3 min.

**Fly ash of Novocherkasskaya PS**

- Slurry density (solid matter content) - 200 g/l;
- Slurry flow rate in compartment - 1.3 clamp;
- Air consumption - 0.5 k/min (at each flotation stage, in any compartment);
- Consumption of agents: Rougher flotation - lecithine 160 g/l, MIBK 350 g/l;
- Scavenging - lecithine 1400 g/l, MIBK 250 g/l;
- Concentrate cleaning - lecithine 700 g/l, MIBK 250 g/l.

### CONCLUSIONS

New flotation flow sheets and conditions have been developed that facilitate solutions to the problems below.

1. Desulphurization of fine fractions (0-0.25 mm) of anthracites in the Rostov region (specific case of Sadinskaya mine coals);
2. Intensification of finely dispersed slimes flotation with production of super-pure concentrates to prepare water-coal fuel (specific case of weakly binding coals from Kuznetsk);
3. Recovery for use of coal particles from the fly ash of electric power stations.

### REFERENCES

Complex Coal Processing of Kazakhstan Deposits

DA. Kunoyev Institute of Mining, Ministry of Power, Industry and Trade, Almaty, Republic of Kazakhstan

ABSTRACT: This report reviews the uses of complex coal processing in Kazakhstan. Complex coal processing is necessitated by the deficiency of oil and gas, and methods of complex coal processing are reviewed - by the presence of various grades of coal with different organic parts, such as bitumen, resin, humus-wax, and chemical elements, including non-ferrous metals and rare earth elements.

1 INTRODUCTION

As it is known, about 30 groups and 44 subgroups of coal can be distinguished under the international classification of 1957 and the classification accepted in CIS countries. All these coals differ in their physical properties and elemental composition. Naturally, therefore, the methods of both use and processing vary: technological, energy, chemical, metallurgical, construction and other.

In conditions of deficiency of known oil and gas reserves (a projected 40-50 years of consumption) in comparison with coal (200-250 years), there is an especially urgent need to replace oil and gas with coal for power, in the chemical industry and in other areas. In particular, the world, including Kazakhstan, is passing to the use of substitutes of natural gas and synthetic liquid combustibles obtained by processing coal by gasification and hydrogenation (liquation). The production of petrol, fuel, jet and boiler combustibles from coal, by both direct liquation and synthesis, and gas obtained from coal by plasma-chemical methods, is most urgent.

Coal, like other mineral resources, in geomechanical terms consists of a set or complex of minerals of other complex connections of chemical elements. In total, coal contains about 50 chemical elements, which form its organic and inorganic parts, water and volatile substances. The organic part contains bitumens and humic acids. Coal has also more than 30 microelements, equal to 5600 g/t of coal, costing about 230 dollars according to the prices of 1988 (Mironov, 1991; Yudovitch, 1989; Drozhzhin, 1952).

Now coal is processed and is applied in the following ways.

1. Coal was first used only as a combustible for municipal-household needs, then coal was used for burning in boiler installations, in the fire chambers of ships and trains, in reflective furnaces, etc.
2. The next stage of coal use was the production of electric power, and to the present day the main consumer of coal in Kazakhstan is the electrical power industry.
3. Now coal is widely used in metallurgy in the form of coke, half-coke and ferrous coke.
4. For almost one century, coal has been bricked from chaff or coal dust in various countries of Europe, America and Asia. Dressing of coal by hardening, carbonization, pelleting (granulation, bricketing) and thermal processing is one of the methods of processing and using complex coal (Taits & Antonova, 1985).
5. Coal, after processing, is used as an agglomeration (caking) combustible for obtaining iron-ore concentrates, carbonate filler, electrode items and foundry coke, and for the production of carbide, calcium and electric corundum.
6. Coal ashes in waste-free production are used for the production of construction materials, including lime, cement and brick.
7. Hard coal and anthracite are mainly used for the production of coal adsorbents and active coal.
8. In various branches of the chemical industry, coal is processed by gasification, extraction, supercritical dissolution of coal, etc. Thus, it enables the production of more than 50 basic chemical products (synthesis - gas, montan wax, paraffin, motor and jet combustible, mineral fertilizers, plastic, etc.) in addition to numerous derivative substances.
9. Out of the total known reserves of power sources, 80-90 % - is coal and only 10-20 % is pe-
terolium and gas (Zhukov et al., 1990). Therefore, the problem of replacing oil and gas with coal and products obtained from processing coal is urgent. So, for example, the gas obtained by coal gasification in ground and underground gas generators is used for production of (Zhukov et al., 1990):

- Substitute of natural gas (SNG).
- Synthesis - gas (S-G) for production of motor fuel and other products in the chemical industry.
- Fuel gases for technology and power generation.
- Restoring gas for metallurgical purposes.
- Carbonization of brown coal.

10. Use of coal instead of oil and gas is possible in the following ways:

- Burning coal instead of oil (black oil), for example, in boiler installations.
- Combination of energy of coal with other types of energy, for example, nuclear energy.
- Development of gaseous and thermal networks.
- Production of motor fuel from coal.
- Use of natural gas substitute to conserve natural gas and supplement its use.
- Use of synthesis - gas from coal instead of products of oils for the chemical industry.
- Production of motor fuel and chemical products from coal after petroleum resources are exhausted.

11. The following microelements are now extracted from coal: uranium, germanium, gallium, and vanadium. It is also possible to extract molybdenum, rhenium, silver, scandium and other rare earth elements (Mironov, 1991).

12. Water alkaline solutions from brown and stone oxide coal extract also contain humic acids and humâtes. Humus coals are used in agriculture in two ways (Soprykin, 1984): first, in the production of physiologically active preparations (PAP) for stimulation of growth of plants, increasing the productivity and efficiency of animal industries; second, in the production and application of organic materials.

In Kazakhstan, coal is used for the production of briquettes, absorbers, restoring agents, humus preparations for stimulation of plant growth and increasing the productivity of animal industries, and rare earth elements, non-ferrous and precious metals are extracted from coal and its ash. Coal ash is used for the production of construction materials - cement, lime and bricks. Projects of coal gasification and hydrogénation have also been developed.

The benefits of complex coal processing are numerous. Coal gasification and hydrogénation produce cheaper and top-quality synthetic liquid combustible, which can replace oil in power, chemistry and in other areas. The production and use of coal bricks reduces losses of coal caused by the underburning of small coal pieces, and it also prevents self-burning.

2 CONCLUSIONS

The application of physiologically active preparations made of coal humus and microelements contained in coal now produces increased productivity in agricultural cultures of up to 30%. The use of other products from coal results in low-waste and waste-free production and increases the level of environment protection.

The rational area of application for coal gasification and hydrogénation is the processing of brown coal, and also long-flame gaseous hard coals to synthesis - gas and synthetic liquid combustible with application in areas of production from local coals. Hard coal and anthracite are economically more expedient for use in traditional areas such as energy production, metallurgy, construction and in the chemical industry.

REFERENCES


Application of Aeration to Complex Sulphide Ore and its Effects on Circuit Performance

B. Aksani & M. N. Mian
Çayeli Bakır isletmeleri, Çayeli, Rize, Turkey

ABSTRACT: Clastic ore is one of the complex sulphide ores found in the Çayeli deposit. It is the most difficult ore to treat and is processed in separate campaigns. The copper concentrates produced contain 19-20% copper and 9-10% zinc. In order to improve copper concentrate quality, a decision was made to try aeration. Laboratory-scale aeration and flotation tests were carried out in order to optimise aeration conditions. Based on the laboratory tests, aeration was put into practice with existing aeration cells in the plant. In this study, the laboratory and plant results are presented, and the effect of aeration on the plant metallurgy is discussed.

1 INTRODUCTION

The Çayeli ore deposit is volcanogenic massive sulphide. Clastic ore is one of the ore types found in the Çayeli deposit and is characterized by an abundance of fine-grained (less than 50 μm), generally anhedral, sulphide debris. The ore consists of rounded pyrite, chalcopyrite and sphalerite clasts in a matrix of fine-grained sphalerite, pyrite, chalcopyrite and barite. The clast size varies from 5 mm to greater than 50 mm, but the sphalerite clasts are generally larger than the pyrite and chalcopyrite clasts. Fine complex intergrowths are common. Most of the pyrite is frambooidal to colloform, and sometimes occurs in aggregates, but may also occur as fine (20-50 μm) subhedral/euhedral crystals. Typical grades of the clastic ore are 3% copper and 13% zinc. The ore is not only fine-grained, but also contains abundant fine mineral intergrowths and inclusions, and requires fine grinding.

Flotation collectors, modifiers and grinding medium can substantially lower the oxygen content of the pulp (Jones and Woodcock, 1984). A lack of oxygen in the pulp sometimes adversely affects the floatability of the minerals. Sometimes aeration of the pulp restores the oxygen concentration. Air acts as a flotation reagent when it is used before the conditioning step with reagents either improving or reducing mineral floatability. For example, flotation tests on chalcopyrite and galena from Black Mountain ore indicated that aeration of the pulp resulted in improvement of copper metallurgy (Graham and Heathcote, 1982; Ross and Van Deventer, 1985). Pyrite depression is a typical example of the depressive effect of aeration (Konigsman, 1973; Kristall et al., 1994; Houot & Duhamel, 1990). Some successful plant applications of aeration have been reported for pyrite depression (Konigsman, 1973; Carlson & Muir, 1976; Spira & Rosenblum, 1978).

The clastic ore contained a significant amount of pyrite (>50%). Therefore, it was decided to apply aeration to the clastic ore in order to improve concentrate quality by enhancing pyrite depression. In order to test this hypothesis, laboratory flotation tests were carried out. In this study, the laboratory tests and plant trial results are presented and the impact of the aeration on metallurgy is discussed.

2 METHOD

2.1. Laboratory tests

The clastic ore samples used in the experimental study were taken from primary ball mill feed in September 1999. The grade of the ore was 4.8% copper and 10.4% zinc. According to previous plant experience, processing such ores caused selectivity problems in the copper circuit.

The ore samples were ground in a laboratory rod mill and a Humbold Wedag-type laboratory flotation machine was used in the aeration and flotation stages.

Sodium metabisulphite (Na2S2O5) was used in the laboratory tests as a sphalerite and pyrite depressant instead of the SO2 applied in the plant due to
handling difficulties. Aerophine (Cytec 3418A) was used as a collector.

2.2. Plant application

Aeration was first applied in the December 1999 clastic campaign and continued in the year 2000 campaigns with the existing aeration unit in the plant. This unit consists of three 45-m$^3$ cells equipped with an original Dorr Oliver rotor stator mechanism. The maximum air rate to the aeration unit is 6000 m$^3$/h.

The plant data were evaluated so as to determine statistically significant differences in the metallurgy with aeration. Results for the copper rougher, zinc rougher, and overall circuit metallurgy were used in these evaluations.

3 RESULTS

3.1 Laboratory test results

3.1.1. Effects of aeration

The test results showed that aeration improves copper and zinc grades and recoveries in the rougher stage (Table 1). Although zinc recovery increases with aeration, selectivity is better due to the increased copper recovery, especially at pH 11.5. As shown in Table 2, the main effect of aeration observed was the depression of pyrite particles, so the relative amount of sphalerite and chalcopyrite particles in the copper concentrate increased.

<table>
<thead>
<tr>
<th>pH</th>
<th>Aeration (minutes)</th>
<th>Cu%</th>
<th>Zn%</th>
<th>Cu recovery</th>
<th>Zn recovery</th>
</tr>
</thead>
<tbody>
<tr>
<td>12.0</td>
<td>20</td>
<td>1075</td>
<td>1271</td>
<td>82.09</td>
<td>48.18</td>
</tr>
<tr>
<td>12.0</td>
<td>20</td>
<td>1075</td>
<td>1271</td>
<td>82.09</td>
<td>48.18</td>
</tr>
<tr>
<td>11.5</td>
<td>20</td>
<td>1045</td>
<td>982</td>
<td>78.63</td>
<td>36.75</td>
</tr>
<tr>
<td>11.5</td>
<td>20</td>
<td>1045</td>
<td>982</td>
<td>78.63</td>
<td>36.75</td>
</tr>
<tr>
<td>12.0</td>
<td>5</td>
<td>538</td>
<td>78</td>
<td>53.60</td>
<td>27.19</td>
</tr>
</tbody>
</table>

Table 1. Effects of Aeration on Metallurgy

<table>
<thead>
<tr>
<th>pH</th>
<th>Aeration (minutes)</th>
<th>Cu recovery</th>
<th>Zn recovery</th>
<th>Fe recovery</th>
</tr>
</thead>
<tbody>
<tr>
<td>12.0</td>
<td>30</td>
<td>85.11</td>
<td>62.41</td>
<td>31.25</td>
</tr>
<tr>
<td>12.0</td>
<td>30</td>
<td>85.11</td>
<td>62.41</td>
<td>31.25</td>
</tr>
<tr>
<td>12.0</td>
<td>30</td>
<td>85.11</td>
<td>62.41</td>
<td>31.25</td>
</tr>
<tr>
<td>12.0</td>
<td>30</td>
<td>85.11</td>
<td>62.41</td>
<td>31.25</td>
</tr>
</tbody>
</table>

Table 2. Effects of Aeration on Pyrite Depression

3.1.2. Effects of pH during aeration

The effects of aeration pH on metallurgy are given in Figure 1. In these tests, the aeration pH was adjusted at the beginning of aeration with lime addition. An increase in aeration pH resulted in a decrease in copper and zinc recoveries. The copper grade tended to increase with pH. At pH 11.5, the lowest level of zinc recovery was obtained.

Figure 1. Effect of Aeration pH on Metallurgy.

3.1.3. Effects of aeration time

The effects of the duration of aeration on copper and zinc recovery are given in Figure 2. Copper recovery was improved with aeration. Zinc recovery was also improved with aeration but to a lesser extent. Aeration times of over 20 min increased zinc recovery. However, zinc recovery did not increase at pH 11.5.

Figure 2. Effect of Aeration time on Metallurgy.

Based on the test results, aeration was put into practice using the procedure applied in the laboratory (Figure 3).

3.2 Plant results

In the plant, grinding was performed without any chemical additions such as time or depressants. The secondary cyclone overflow was fed to the first aeration cell. Lime was added to the first cell and the pH was adjusted to 11.5. At the start of the plant
trial, the total air rate was adjusted to 4500 m³/h and then decreased to 2700 m³/h. It was observed that copper recovery in copper rougher deteriorated at high air rates probably due to oxidation of chalcopyrite. At the end of the aeration stage, the pH dropped to 10. After the aeration stage, the pulp was conditioned with SO₂ (-1 kg/t), followed by collector addition. The flotation pH was 7-7.5.

Figure 3. Experimental procedure.

3.2.1. Rougher copper and zinc concentrate

The effects of aeration on copper rougher and zinc rougher are given in Table 3. In this table, iron grade was used as an indication of pyrite content.

The results were analyzed using F and t-test statistics at a confidence level of 95% (Underhill & Bradfield, 1994). Statistical analysis showed that the zinc and iron grades were not significantly different. However, the increase in the copper grade in the application of aeration was statistically significant. The copper, zinc and iron recoveries were lower than those with the previous processing method and the decreases in recoveries were also statistically significant. Sphalerite and pyrite rejection improved with aeration, but chalcopyrite recovery dropped 4%.

The rougher zinc concentrate was affected positively by aeration. Both zinc grade and recovery improved, and these improvements were statistically significant. This was due to better zinc rejection in the copper rougher circuit and efficient pyrite depression.

Another indication of pyrite rejection was the reduced tonnage recovered in the rougher copper and zinc concentrate. The reduction in the copper rougher and zinc rougher concentrate tonnages were 5.7% and 4.7%, respectively.

Table 3. Copper Rougher and Zinc Rougher Performances with Aeration.

<table>
<thead>
<tr>
<th>Rougher</th>
<th>Concentrate Grade %</th>
<th>Recovery %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cu</td>
<td>Zn</td>
</tr>
<tr>
<td>Aeration</td>
<td>15.0</td>
<td>9.8</td>
</tr>
<tr>
<td>Previous</td>
<td>135</td>
<td>0.4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Rougher</th>
<th>Concentrate Grade %</th>
<th>Recovery %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cu</td>
<td>Zn</td>
</tr>
<tr>
<td>Aeration</td>
<td>2.84</td>
<td>30.7</td>
</tr>
<tr>
<td>Previous</td>
<td>2.44</td>
<td>27.1</td>
</tr>
</tbody>
</table>

3.2.2. Overall circuit metallurgy

The effects of aeration on overall copper and zinc circuit metallurgy are given in Table 4. The decreases in the copper grade, and copper, zinc and iron recoveries in the final copper concentrate were significant. Although a higher copper grade was not obtained, (he zinc and iron rejections increased with aeration.

Table 4. Copper and Zinc Circuit Performances with Aeration.

<table>
<thead>
<tr>
<th>Final Cu Concentrate</th>
<th>Concentrate Grade %</th>
<th>Recovery %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cu</td>
<td>Zn</td>
</tr>
<tr>
<td>Aeration</td>
<td>19.3</td>
<td>9.1</td>
</tr>
<tr>
<td>Previous</td>
<td>20.4</td>
<td>9.4</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Final Zn Concentrate</th>
<th>Concentrate Grade %</th>
<th>Recovery %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cu</td>
<td>Zn</td>
</tr>
<tr>
<td>Aeration</td>
<td>3.0</td>
<td>51.0</td>
</tr>
<tr>
<td>Previous</td>
<td>3.1</td>
<td>49.1</td>
</tr>
</tbody>
</table>

The decreases in copper grade and recovery in the final copper concentrate could be due to:

j. lower copper feed grades (3.77±0.17 in aeration and 4.16±0.23 in previous campaigns without aeration, at a confidence level of 95%).
Difficulties of circuit adjustments during short clastic campaigns; Lack of operator experience with the aeration scheme.

Another important feature observed in the application was the sensitivity of the copper rougher tail to air rate in the aeration cells. The increase in rougher tail resulted in a decrease in rougher recovery and copper circuit recovery. As mentioned above, clastic ore contains abundant fine mineral intergrowths and inclusions. As a result, chalcopyrite-pyrite locked particles might be depressed by oxidation due to its pyrite content and lost in the rougher tail.

The most pronounced effect of aeration was obtained in the zinc circuit. Both zinc grade and recovery increased significantly. The zinc concentrate quality was improved, zinc grade increased and the iron content of the concentrate decreased due to pyrite rejection. Zinc recovery in the final zinc concentrate also increased. Lower zinc recovery in the copper rougher improved zinc recovery in the zinc circuit. In addition, depressed pyrite particles in the copper rougher did not float in the zinc rougher and were rejected in the zinc rougher tail. Thus, the amount of pyrite particles recovered in the rougher zinc concentrate decreased and resulted in improved final zinc concentrate quality.

4 CONCLUSIONS

The zinc circuit metallurgy improved with aeration. Improvement in copper metallurgy was not obtained as expected with better zinc and pyrite rejection in the circuit. Due to the limited durations of the clastic campaigns and the ore variability, CBI intends to use aeration to investigate whether improvements in copper metallurgy can also be achieved with operating experience and circuit tuning.

ACKNOWLEDGEMENT

The authors wish to thank the managements of CBI A.S. for permission to publish this paper.

REFERENCES

ABSTRACT: The Çayeli concentrator designed nameplate capacity was 600,000 tonnes per year. Originally, three Rittershaus & Blecher pressure filters were installed: one for copper concentrate, one for zinc concentrate, and one standby for both copper and zinc concentrates. Mill throughput gradually increased after commissioning and was 896,700 tonnes in 1999. At these production levels, the filtration capacity created a bottleneck. In 1999, CBI decided to increase the mill capacity to one million tonnes per year and a fourth filter was installed in the year 2000 as part of the expansion plan. A LASTA M.C. automatic filter press was selected. This paper describes the selection, installation, commissioning, and operation of the new filter; and compares its performance with that of the old filters.

1 INTRODUCTION
Çayeli Bakır isletmeleri is a copper zinc mine at Çayeli on the eastern Black Sea coast in the province of Rize. The mine is located 7km from the coast. It is an underground operation of one million tonnes per year, producing copper and zinc concentrates. The concentrates are trucked 25 km to a port facility at Rize for shipment to customers.

2 SELECTION PROCEDURE
Originally, the CBI mill was equipped with 3 Rittershaus & Blecher (R&B) pressure filters. These filters are high-maintenance items and required dedicated mechanics to look after them as production increased. Initially, one operator was budgeted for the dewatering section, but due to the operational difficulties of these filters, this was increased to two operators per shift. In choosing a filter, the main selection criteria were low maintenance and operation costs, and the filter had to be simple to operate and robust in design. The capacity requirement was 400 dmt/day.

Quotations were requested from the Larox, Outokumpu, Bethlehem, and MC LASTA pressure filter manufacturers.

After preliminary evaluation of the quotations, a trip to North America was arranged so that these filters could be seen in operation and first hand information about their performance could be obtained.

Larox filters were observed in the Louvicourt and New Brunswick mines in Canada. The filters in Louvicourt were being used for filtering copper and zinc concentrates and the one in New Brunswick was being used for copper concentrate.

A Bethlehem filter was examined at Mason Metals, Chicago, USA. This filter is similar to the Larox. The material filtered was tin hydroxide slurry.

The MC LASTA filter was inspected at Kennecott Utah Copper, Salt Lake City, USA.

The Outokumpu Ceramic filter was rejected in the preliminary evaluation due to the high capital cost.

After the trip, a spreadsheet was prepared to compare the three selected bids for final evaluation based on our selection criteria.

The MC LASTA AUTOMATIC FILTER PRESS was selected. Industrial Process Machinery (IPM) offered a model MCFHC 1500 X 40/34 for 400 dmt/day operation. The number of plates offered was 34, while the frame capacity was 40 plates.

During discussions with the supplier, 6 more plates were added in order to utilise the maximum capacity of the machine. The nominal capacity with 40 plates is 470 dmt/day.

3 ENGINEERING AND INSTALLATION
The engineering contract was awarded to Foster Wheeler Bimas (FWB). The filter was a retrofit in place of two pilot-scale disc filters, which presented
an engineering challenge FWB completed the
detailed engineering for the installation, including
filter auxiliaries. In two months. The auxiliaries
were:
• one high pressure water pump for filter cloth
washing, feed core clean-up, channel cleans, etc.,
• one slurry feed pump, which was also retrofit
to the existing copper concentrate holding tank,
• one compressor, dedicated for use with the
Lasta filter,
• one transfer pump, to pump filtrate and filter
cloth washings.
FWB also prepared bid packages and helped in
the selection of the contractor. EPRO Construction
was awarded the project based on the CBI
competitive bidding process.
The scope of the contract included all civil, steel,
electrical and instrumentation works. A site project
superintendent was appointed from FWB to
supervise the contractor and to ensure that the
construction was done according to their design.
The construction contract was awarded in mid-April
and commissioning of the filter started in mid-
August 2000. The filter was fully commissioned by
the end of September 2000. The cooperation from
1PM during all phases of engineering and
construction was excellent. One person from 1PM
was on site during erection and two persons were on
site for commissioning. They programmed the PLC
and checked that the filter operation was
satisfactory. They trained the CBI operation and
maintenance staff. The training sessions were well
conducted and covered in detail all the finer points
of operation and maintenance of MC LASTA filters.
These points were found to be very beneficial,
ensuring smooth commissioning and trouble-free
operation of the filter.

4 COMMISSIONING
After the usual pre-commissioning checks and dry
runs, hot commissioning of the filter started on 19
August 2000. The filter exceeded its rated capacity
after one month of operation. The current
operational parameters are as follows:
Average Cycle Time: 11 minutes.
Average Number of Cycles: 120 per day.
Average Capacity: 518 dmt/day.
The filter is consistently exceeding its rated capacity
of 470 dmt and we are very satisfied with its
performance.

5 OPERATION
The filter press has a fully automated cycle and does
not have to be restarted after each cycle.
The press has three operational modes.
2. Semi Auto Mode.
3. Auto Mode (Operation Mode).

5.1 Manual Mode
All functions are manually controlled. This mode is
also called the maintenance mode and is required
during maintenance.

5.2 Semi Auto Mode
The press operates and stops at the end of each
phase of operation and requires restarting after each
phase. This mode is used to optimise the operating
conditions, especially during commissioning.

5.3 Auto Mode
The filter is started by simply pressing the start
button and it completes a full cycle automatically. It
does not need restarting at the end of each cycle. It
stops only when the "feed tank empty" alarm is
activated.

6 COMPARISON OF THE MC LASTA WITH
R&B PRESSES ACCORDING TO OUR
EXPERIENCE
6.1 Performance Comparison
6.1.1. Capacity (Number of Cycles)
R&B filters: The rated capacities are 390 dmt/day
for copper and 330 dmt/day for the zinc filters. The
average cycle time is 20-25 minutes. This is highly
dependent on cake discharge. The filling, membrane
squeezing and cake blow cycles take 12-15 minutes.
Cake discharging takes about half of the time
required for the rest of the steps. The plate-shifting
mechanism is a problem and the operator is required
to complete the cake discharge and clean the plates.
The plate-shifting mechanism chain and
connection pieces, shifting hook, and locking hook
(pawl) stick and plates often do not move.
Sometimes, due to these difficulties, it is only
possible to average 40-50 cycles per day.
In actual operation, only about 280 dmt/day is
obtained, based on availability due to operational
problems which are explained below.
MC LASTA filter: The Lasta filter average cycle
time including the cake discharge, is about 11
minutes. This goes down to as low as 10.5 minutes
when the filter cloths are new and goes up to 12.5
minutes as the cloths get older. There is no operator
assistance required for discharging the cake. The
average number of cycles per shift is 38-40. Forty
cycles per shift can be achieved without difficulty. This figure has been as high as 45 per shift. The average capacity is over 500 dmt/day.

6.1.2 Labour Requirements

R&B filters: CBI employs 2 operators per shift to operate these filters due to frequent cloth changes (average 2200 cycles for one set of cloth) and plate-moving problems. The seal surfaces on the sides of the plates and around the feed opening require continuous cleaning after each cycle due to the lack of an auto cloth wash provision. An auto cloth wash after each cycle is not possible with the existing design. This is explained in more detail in the maintenance comparison section.

MC LASTA filter: The press requires no operator assistance during normal operation. Operator assistance is required only during alarm status or to change filter cloths. Cloth life is about 5000 cycles. Two sets of cloth have been changed to date. The first set was changed at 4500 cycles and the second at 5000 cycles.

6.1.3 Difficulties in Plate Shifting

R&B filters: The plates are hung on a rail with carriages. The carriages are equipped with hooks. There is a continuous running chain mechanism with shifter hooks attached. These shifter hooks catch the locking hooks (paws) on the plate carriages and move the plates. This complex system causes frequent plate-shifting problems. This reduces the performance of the press. When plate-shifting problems start, the operator needs to climb on top of the filter to release the plates and maintenance assistance is required to change the damaged connection pieces.

MC LASTA filter: The Lasta filter plate-moving mechanism is very simple. The plates are located on two rails. There is a replaceable polyethylene wear pad attached to the plates and this acts as a cushion, making movement easier. The top parts of the rails are also replaceable. The top surface of the rails needs regular greasing and the shifter operators on each shift do this. The rail surfaces are easily accessible.

The plates are connected to each other at 4 corners by a piece of chain. The head plate on the side of the cylinder acts as a drive head. When the cylinder opens, the plates pull each other, leaving space between plates which is the same as the length of the chain connecting the plates. This helps the discharge of the concentrates and ensures equal space between plates for effective washing of the cloth after cake discharge.

When the filter closes, the cylinder pushes the head plate in the opposite direction and the head plate pushes the rest of the plates, which sit freely on rails. The gap between the plates starts to close, first from the cylinder side, and it is progressively closed as the cylinder moves to the feed side.

6.1.4 Vibration Bars for Cake Removal

R&B filters: No vibration bars in design.

MC LASTA filter: When cake sticks to the cloth, the cloth support bar is depressed due to the weight of the sticky material and blocks the photo beam. This activates a vibration cycle and shakes all the plates from both sides. The vibration cycle is also an indication of possible cloth problems, i.e., holes, blinding, etc.

6.1.5 Cake Discharge System

R&B filters: The cake falls with its own weight by gravity, and there are no additional features to help discharge the cake.

MC LASTA filter: The weight of the cake on the cloth rapidly compresses the springs which support the cloth support bars. The bars come to a sudden stop, which creates a shear effect between the cake and the filter cloth. This helps removal of the cake from the filter cloth.

6.2 Maintenance Comparison

6.2.1 Filter Plate Repairs

R&B filters: These filters require extensive maintenance on the plates for several reasons. The plates undergo continuous wear on the seal surfaces due to the lack of a good seal. This is due to the absence of an auto cloth wash after each cycle. The wear surfaces are repaired by plastic welding and filling. After a while, this is no longer possible and the whole plate is taken out and sent away for resurfacing.

In general, plate repair is a continuous process. The plates are collected on site when they cannot be repaired with plastic welding, and are sent to Germany for resurfacing. CBI is looking for a local workshop to repair these plates.

Damage occurs on the membrane plates. The rubber membranes are attached to the membrane plates by insertion of the sealing ring into the channel provided on the peripheries of the plates. These channels are damaged by compression in time and membranes cannot be inserted anymore.
Figure 1. Vibration mechanism of MC Lasta filter

Figure 2. Cake Discharge of MC Lasta filter
MC LASTA filter: There are no membrane plates. Only chamber plates are provided. The air entry and filtrate discharge ports are replaceable units called filtrate rejecters. When worn, they are replaced. Their wear is also an indication of holes in the filter cloth.

To date, no plates have been damaged due to any other factors.

6.2.2 Cloth Change Comparison

R&B filters: The cloth is a single piece. Changing it is a time-consuming and fairly laborious job. One part of the cloth has to be rolled, squeezed and passed through the feed opening of the plate so that it can be put on the other side of the plate. Similarly, removing the old cloth is not easy as the cloth becomes harder and very difficult to roll back through the feed opening. It is common practice to cut the cloth when removing it from the plates.

MC LASTA filter: The cloth change is very easy. There are two pieces, which are hung on each side of a plate by hanging bars. The filter cloth assembly can be prepared in advance of cloth change. The odd-numbered plate cloth is equipped with a feeding device which is attached to the cloth before it is hung on the bars.

6.2.3 Plate-Moving and Shifting Mechanism

R&B filters: The chains located inside the traverse are equipped with shifter hooks, which glide over the carriages of the plate pack when the press is closed. When the press is opened, one of the shifter hooks engages, due to its own weight, on the last plate and unlocks this plate. The shifter hook now moves the plate suspended from the carriage fitted with offset rollers. This action is repeated until the last plate is opened. The offset roller movement is hindered by dirt accumulating inside the traverse rails, which are difficult to reach and clean. If the locking hooks of the carriages are not caught by the shifter hook of the chain in the proper order, men a group of plates shift together. This overloads the locking hook and the drive motor of the chain mechanism. The locking hook bends or the chain drive motor trips due to overloading. In short, the plate-shifting mechanism is complex.

MC LASTA filter: The plate-moving mechanism is rather simple. The plates are connected to each other by chains at four corners. When the cylinder retracts, it moves the head plate backwards and the head plate pulls the rest of the plates one by one. The plates move freely on top of two side rails. The greasing of the rails makes it easy.

6.2.4 Washing System Maintenance

R&B filters: The washing mechanism is a separate unit. It has a trolley with a moving chain and washing device. A geared motor is provided for the longitudinal and the wash travel drives. Both the washing device and the plates have to move to wash the cloth. The problems with plate moving continue during cloth washing. Each washing takes about 45 minutes if there are no problems with plate movements or washing device movements. The cloth wash is a manual process. The filter has to be stopped and the control unit is changed to cloth wash mode. It is not practical to wash cloth after each cycle since one wash takes 45 minutes.

![Filter cloth of MC Lasta](image)
In addition to these problems, the washing device catches the filter cloth and sometime damages it while the sprays are in upward motion.

MC LASTA filter: The cloth wash is part of the auto cycle; it takes 40 seconds to wash all 41 plates and 40 chambers. Each plate is equipped with three cloth wash sprays as an integrated part of the cloth support bar. Water from the spray water header is connected to the cloth support bars on the even-numbered plates by hoses. Wash nozzles direct wash water over the cake side of the filter cloth.

6.3 Safety and Environment

6.3.1 Leaks and Noise from Filter Plates

R&B filters: Sealing surfaces deteriorate due to poor cloth washing, especially at the bottom of the plates. Cake remnants deteriorate the sealing surfaces. When the filter comes to the cake blow cycle, the compressed air leaks through these surfaces, creating dust and a shrill whistling noise. The cloth is also damaged by these surfaces.

MC LASTA filter: There is no sealing problem. The effective cloth-washing system after each cycle cleans all the surfaces of the plate. There are no leaks and there is no whistling.

6.3.2 Position and Access to Plates

R&B filters: The working platform is level with the bottom of the plates. This is necessary due to the operator assistance required in cake discharge. In order that the top of the plates can be reached, a ladder is provided. The operator or maintenance personnel have to put on a safety belt while working on top of the filter.

MC LASTA filter: The working platform is located on top of the plates. The support rails, spray nozzles and cloth change bars are all accessible without the need for any safety equipment. One operator can change a filter cloth.

7 CONCLUSIONS

The selection of the MC LASTA filter was the right decision for Çayeli. The filter has exceeded our expectations. We recommend this filter for base metal concentrates where simplicity of operation and maintenance are key factors.

REFERENCES

Rittershaus & Blecher Filter Operation Manual
MC Lasta Filter Operation and Maintenance Manuals.
ABSTRACT: In this study, the adsorption of quaternary amines onto original and heat-treated sepiolite was investigated as a function of several variables. Adsorption of dodecyltrimethylammonium bromide (C_{12}H_{25}N(CH_3)_3Br) and hexadecyltrimethylammonium bromide (C_{16}H_{33}N(CH_3)_3Br) onto sepiolite at a constant temperature of 25°C exhibits typical isotherms with plateaus. The adsorption densities, as represented by mole per m^2 of the adsorbate, reveal that although heat activation results in a several-fold increase in the specific surface area, practically, it produces no appreciable difference in the adsorption values.

1 INTRODUCTION

Abatement of mining and mineral processing wastes without harming the environment is an important technical problem in the development of the mining industry. The use of flotation reagents such as cationic-amine-type reagents in the flotation of silicates, oxides and salt-type minerals results in the contamination of process water. These aliphatic and aromatic compounds are also used in the production of pigments, rubber and pesticides, leading to substantial organic wastes. The adsorption process is usually applied to those wastes that cannot be removed by biological techniques. Activated carbon is the most common adsorbent used in such processes. Sepiolite, which was used in this investigation, is cheaper and is considered an alternative to activated carbon. It is not only used in the abatement of organic compounds, but is used for inorganic metal cations as well.

Sepiolite exhibits a variety of attractive properties, such as high specific surface area, high porosity and surface activity, and the ability to form high and stable viscosities at low solids concentrations. Sepiolite is therefore used in a range of areas where sorptive, catalytic and rheological properties are exploited. The sorption ability of sepiolite is mainly ascribed to its high surface area. Thus, it is commonly used in oil refining, wastewater treatment, in the removal of odor, drug and pesticide carriers, paper, detergent, etc.

Similar to other chain-type silicate minerals, it contains a continuous two-dimensional tetrahedral sheet but without continuous octahedral sheets. This unique fibrous structure with interior channels of 3.6x10.6 Å in size permits adsorption of organic and inorganic ions (Gonzalez-Pradas et al. 1990, Singer and Huang 1989). Sepiolite has a total specific surface area of 800-900 m^2/g, of which meroretically 400 m^2/g is external and 500 m^2/g is internal (Alvarez 1984, Sema and Van Scoyoc 1978). The penetration of organic molecules into the channels depends upon the polarity, ability to make hydrogen bonds, and the match of the size and shape of the molecule (Çetişli 1989, Hoyo et al. 1993).

It was therefore the objective of this study to investigate the ability of the uptake of typical quaternary amine surfactants, dodecyltrimethylammonium bromide (DTAB) and hexadecyltrimethylammonium bromide (HTAB), onto original and heat-treated sepiolites and elucidate the adsorption mechanism.

2 EXPERIMENTAL

2.1 Materials

The sepiolite sample used in this study was obtained from the Mayas Mining Company in Sivrihisar, Turkey. The cation ratios of sepiolite calculated using oxygen as the basis of 33 (Jones and Galan 1984) yielded the data presented in Table I.

This brown sepiolite, ground to minus 65 μm, produced an average particle size (d50) of 199 μm as determined by Zetasizer. The XRD together with the chemical analysis results given elsewhere indicate that calcite and dolomite are the major impurities...
accompanying sepiolite (Sabah et al. 1997). The specific surface areas of the original and heat-treated sepiolite at 300°C as determined by BET technique are 68 m²/g and 253.3 m²/g respectively.

<table>
<thead>
<tr>
<th>Cation Ratios</th>
<th>Tetrahedral Layer</th>
<th>Octahedral Layer</th>
<th>Cations Between Sheets</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si</td>
<td>11.84</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ti</td>
<td>0.01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Al</td>
<td>0.41</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mg</td>
<td>8.23</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fe³⁺</td>
<td>0.12</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ca</td>
<td>0.19</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Na</td>
<td>0.05</td>
<td></td>
<td></td>
</tr>
<tr>
<td>K</td>
<td>0.10</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Cation ratios of sepiolite.

Dodecyltrimethylammonium bromide (DTAB), C₁₂H₂₅N(CH₃)₃Br and hexadecyltrimethylammonium bromide (HTAB), C₁₆H₃₃N(CH₃)₃Br were used as the cationic surfactants. Both were specified to be of high purity. Distilled and deionized water with a conductivity value of 2x10⁻⁶ mhos/cm was used in all experiments. The experiments were conducted at 25°C. pH was adjusted using either HCl or NaOH solutions.

2.2 Methods

Adsorption tests were conducted in 20- or 40-ml glass vials. 500 mg of sepiolite sample was mixed in 10 cc or multiples of this with a solid-to-liquid ratio of 0.05. The vials were shaken for 2 h on a shaker and centrifuged for 15 min. The supernatant was analyzed for the cationic surfactant by a two-phase titration technique originally applied to anionic surfactants using dimidium bromide and disulfme blue as indicators. This technique is based on the formation of a complex between an anionic surfactant (sodium dodecylsulfate) and cationic reagent (DAH). This complex is soluble in chloroform and changes from blue to pink in the presence of indicators (Powers 1970, Reid et al. 1967).

The adsorption density was calculated by the following formula (1).

\[ r = \frac{(C_i - C_r) \cdot V}{m \cdot S \cdot 1000} \]  

where \( C_i \) and \( C_r \) = the initial and residual concentrations (moles/l); \( m \) = the amount of solid (grams); \( V \) = the volume of the solution (ml); \( S \) = the specific surface area of sepiolite (m²/g); and \( F \) = the adsorption density (M/m²).

In order to increase the surface area and induce physico-chemical changes in the structure of the sepiolite, heat activation was performed. In the heat activation, 20 g of sepiolite was placed in a muffle furnace for a period of 10 hours.

3 RESULTS AND DISCUSSION

3.1 Adsorption of Surfactants on Original Sepiolite

Figure 1 shows the adsorption isotherms of DTAB (C₁₂) and HTAB (C₁₆) at 25±2°C onto sepiolite. Plateau adsorption values for the quaternary amines in the initial concentration range of 10⁻³ - 10⁻⁶ M/l are \( r_{DTAB} = 6.18 \times 10^{-6} \) M/m² for DTAB and \( r_{HTAB} = 6.68 \times 10^{-6} \) M/m² for HTAB. The residual concentration values for these surfactants at the onset of the plateau in the adsorption isotherms for DTAB and HTAB respectively were found to be \( C_D = 3.9 \times 10^{-2} \) M/l, and \( C_D = 1.22 \times 10^{-3} \) M/l. By plugging these values in Equation 2 the cross-sectional areas for each reagent can be calculated, as 27 Å² and 25 Å² for DTAB and HTAB, respectively. If the cross-sectional area of the polar part of DTAB and HTAB is 25 Å², as reported by Harkins and Florence (1938), the degree of coverage for DTAB and HTAB on the original sepiolite is 0.1. This reveals that DTAB and HTAB adsorbs onto original sepiolite in the form of a monolayer. This has been verified by the following equation:

\[ Y \cdot A = \frac{10^{20}}{r_{max}} \cdot A \]  

where \( T_{max} \) denotes adsorption in the plateau region (M/m²); and \( A \) represents the Avagadro number (6.02x10²³).

The adsorption isotherms of the original sepiolite/quaternary amines system given in Figure 1 are characterized by three distinct regions with different slopes. While the adsorption of amines shows a gradual increase in the first region, the increase in the second region is rather sharp. Despite significant differences in the rising part of the adsorption isotherms for two surfactants of different chain length onto sepiolite adsorbent with 68 m²/g of specific surface area, the adsorption densities overlap at the onset of the plateau (Region III). The solution concentration in Region III reaches saturation, which indicates that both quaternary amines concomitantly attain a region of micellar interactions and in turn point out the differences in the adsorption mechanisms of DTAB and HTAB molecules in each region.
The data in Figure 1 also show that the cationic surfactant with 16 CH_2 groups exhibits, much higher adsorption densities than that with 12, CH_2 groups. This is particularly pronounced in the region of hemimicelle formation, where the slope of the isotherms is steeper. Although this finding is in agreement with the surface activity of the respective surfactants, it does disagree with the assumption that the molecules penetrate into the channels of sepiolite.

The contribution of CH_2 groups to the free energy of adsorption in Region H was assessed by modifying the assumptions used by Fuerstenau and his co-workers for an alumina/alkylsulfonate system as follows (Somasundaran and Fuerstenau 1966, Wakamatsu and Fuerstenau 1968):

\[
C_s = C_D \cdot \exp \left( \frac{-Z \psi_{ST} + n \phi}{kT} \right)
\]

where \( C_s \) and \( C_D \) = surface and equilibrium concentrations; \( Z \) = valency of the adsorbate species; \( e \) = fundamental unit of charge; \( \psi_{ST} \) = Stern layer potential; \( n \) = number of straight chain alkyl groups in the surfactant; \( \phi \) = free energy per mole of CH_2 groups for hemimicellization; and \( k \) = Boltzman constant.

Under the conditions of constant adsorption density, the above equation can be rewritten for dodecyl and hexadecyl amines as

\[
\frac{C_{D-16}}{C_{D-12}} = \exp \left( \frac{Z \psi_{ST-16} - Z \psi_{ST-12}}{kT} \right) \cdot \left( \frac{16 \cdot 12}{12} \right)^{n/2}
\]

Since at pH 8.5 the ionic strength is maintained roughly constant, under these conditions one can write

\[
\psi_{ST-16} = \psi_{ST-12}
\]

Equation 4 is then reduced to

\[
\ln \frac{C_{D-16}}{C_{D-12}} = \frac{4 \phi}{kT}
\]

Substituting the values for bulk concentrations corresponding to an adsorption density of \( 10^{-6} \) M/m^2 in Equation 5:

\[
10^{-6} \cdot \frac{6 \times 10^{-5}}{2.28 \times 10^{-3}} = \frac{4 \phi}{kT}
\]

\( \phi = -540 \text{ Cal/mol CH}_2 \text{- group} \),

This value for free energy contribution per CH_2 group (-1 kT) is in good agreement with the values reported by Fuerstenau and his associates for alumina/alkylsulfonate (Somasundaran and Fuerstenau 1966, Wakamatsu and Fuerstenau 1968). These results clearly demonstrate that, regardless of the mechanism in Region 1, the chains of the molecules associate in Region II and enhance the adsorption process depending on the chain length.
3.2 Effect of Heat Treatment on Surfactant Adsorption

The adsorption isotherms of heat-treated sepiolite at 300°C with an increased specific surface area of 235.5 m²/g/a for DTAB and HTAB surfactants at 25°C are illustrated in Figures 2a and 2b. As is evident from the figure, the adsorption of heat-treated sepiolite exhibits a definite reduction at all concentrations due to the loss of zeolitic and adsorbed water molecules upon heat treatment and the inability of amine molecules to form mainly hydrogen bonding. It is to be noted that while the adsorption densities between the original and heat-treated sepiolites in M/g exhibit marginal differences, those in M/mo show considerable differences because there is more than a four-fold increase in surface area upon heat treatment, but there is no extra adsorption. Although the increase in surface area upon increasing the temperature is in line with the data in the literature (Jimenez-Lopez et al. 1978, Hibino et al. 1995), the decrease seen in the adsorption needs to be elaborated further.

It is also evident that the residual concentrations in the plateau region are almost the same, indicating that the saturation adsorption is possibly governed by the formation of CMC. The CMC values in water for DTAB and HTAB are 1.56x10⁻⁵ M (Tottewill 1967) and 1.9x10⁻⁵ M (Okuda et al. 1987). However, it is well known that in the presence of mono- and multivalent ions, CMC values undergo significant reduction (Shinoda 1963). In a typical sepiolite supernatant, about 20 ppm of Mg and 40 ppm of calcium is released into the solution. It appears that the Cr values of about 3.8x10⁻⁵ M obtained at the plateau values in Figure 2 are close to the CMC of the system and in line with the above explanation.

The parking areas of DTAB and HTAB molecules on heat-activated sepiolite respectively yield 96 Å² and 75 Å². The surface coverage of DTAB on heat-activated sepiolite gives a surface coverage of 0 = 0.26, whereas that of HTAB yields 0 = 0.30. These values indicate that both amines adsorb onto die surface and channels of the heat-activated sepiolite in the form of patches and islands, but not of layers.

4 CONCLUSIONS

Examination of the slopes of adsorption isotherms for both DTAB and HTAB/original sepiolite systems at 25°C reveals that adsorption densities are high even at low concentrations. This shows that adsorption of amine molecules not only occurs in the external surfaces but also in the internal channels depending upon the position of water molecules, i.e., in die edges and corners of the sepiolite.

The adsorption isotherms of DTAB and HTAB/original sepiolite systems at 25°C show that HTAB with 16 CH₂ groups exhibits higher adsorption densities than DTAB with 12 CH₂ groups. This finding is particularly pronounced in the region of hemimicelle formation where the curves become steeper. Moreover, despite considerable differences in the rising part of the adsorption isotherms for two surfactants of different chain length onto a sepiolite adsorbent, the adsorption densities almost coincide at the onset of plateau (Region III), where the solution concentration reaches saturation. This emphasizes the crucial finding that it is adsorption and not absorption that is the driving force in the uptake of surfactant molecules.

In spite of a several-fold increase in the specific surface area of sepiolite upon heat activation, the adsorption in M/g was found to remain almost constant, whereas in M/m was found to decrease significantly. This was attributed to the increase in the number of micropores which exclude the cationic surfactant and lead to the partial collapse of the magnesium layer in the octahedral sheet. The removal of zeolitic water upon heat treatment is also instrumental in the loss of hydrogen bonds for primary amines. In conclusion, heat activation of sepiolite appears to be detrimental as far as cationic surfactants are concerned. The question of whether this can be extended to all organic compounds requires further studies on other organic compounds.

REFERENCES


Powers, G.W. 1970. The volumetric determination of organic sulfates or sulfonates by the double indicator method. Tulsa, USA; Communication C-225, Amoco Production Co.


Figure 2. Adsorption isotherms of DTAB (a) and HTAB (b) in original and heat-activated sepiolite (Natural pH of 8.5, solution temperature 25°C, S/L=0.05, Cond. Time: 2h).
The Application of Colemanite Addition to Floor Tile Glazes

A. Yamık, İ. Bentli, C. Karagüzel, M. Çınar & B. Cengiz
University of Dumlupinar, Department of Mining Engineering, Kütahya, Turkey

ABSTRACT: Colemanite ore, a borate mineral, has been used in ceramic glaze recipes due to its low melting point, low solubility and its smoothness on unleaded glazes. In this study, colemanite was added at 2%-10% to a standard glaze recipe and six different recipe samples, including the standard recipe, were prepaired. These recipes were applied to industrial-grade ceramic floor tiles. Various physical tests were done on ceramic floor tiles fired at 1180 °C in intensity ovens. The results indicate that the physical properties of the floor tiles did not change considerably and that the surface quality and smoothness of the floor tiles improved with 6%-8% colemanite addition.

1 INTRODUCTION

Ceramic products can be defined as substances produced from inorganic industrial minerals mixed in appropriate proportions, formed into desired shapes and fired in an oven with or without glazes. Ceramic floor tiles, which have a wide variety of applications, consist of a base body overlaid with a hygienic and decorative thin layer of glaze (Arcasoy, 1983). The durability of the surface for its application can be obtained by means of a suitable glaze recipe (Kartal, 1998).

One of the main reasons for the use of glaze recipes in floor tiles is to make the porous and uneven surface much smoother. In this way, it is much easier to clean die floor tile surfaces. Another reason is to give an aesthetic and decorative appearance to the ceramic floor tiles. In addition, resistance to chemical reactions is provided by increasing the strength and surface hardness of the ceramic floor tiles (Yamik et al. 1999, Özalşan 1993).

The objective of applying the engobe process to ceramic floor tiles is to fill cavities, and obtain a smooth surface and conformity between the ceramic body and its glaze. Engobe is the undercoat material, which covers unwanted stains on the ceramic body, provides conformity between the ceramic body and its glaze by forming a thin layer between them, and improves the appearance of motif applications.

The objective of this study was to investigate the physical properties of floor tiles produced by the addition of colemanite to glaze recipes. The test results are compared with the TS-EN 176 standard for ceramic industrial-grade floor tiles.

Borate mineral affects the melting point of other oxide minerals such as lead oxide, which has a very low melting point. Both borate (B2O3) and lead oxide (PbO) are contained in glazes which have low melting points. The main advantages of having B2O3 in ceramic glazes are the following (Kartal, 1998):

- It has a very low melting point;
- Gives better flux properties;
- Decreases thermal expansion;
- Provides resistance to acids;
- Provides better glazing properties by decreasing viscosity;
- Gives more surface brightness.

2 EXPERIMENTAL STUDY

2.1 Samples and procedures

The glaze recipe samples prepared for the tests contained sodium and potassium feldspars, zircon, clay, zinc oxide, talc, corundum, wollastonite and colemanite minerals. The composition of the engobe, which is spread underneath the glaze, included clay minerals, frit, quartz and zircon (Ünaydin, 1993).

For the test, six different glaze recipes were prepared: one standard sample and five samples with colemanite contents of 2%-10%. The chemical compositions of the recipe samples are given in table 1. The seger formula parameter of the standard glaze recipe was determined from chemical analysis:

- 0.082Na2O, 0.037K2O, 0.27CaO, 0.417ZnO
- 0.614Al2O3, 0.007Fe2O3, 2.493SiO2, 0.004TiO2
- 0.125ZrO2 mole. The glaze recipes were prepared from this seger formula by the addition of various amounts of colemanite. The effects of colemanite addition on glaze quality were investigated.
Table 1. Glaze recipe samples with various amounts of colemanite addition.

<table>
<thead>
<tr>
<th>Industrial mineral</th>
<th>Composition (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Standard Recipe</td>
</tr>
<tr>
<td>Na-Feldspar</td>
<td>26</td>
</tr>
<tr>
<td>K-Feldspar</td>
<td>11</td>
</tr>
<tr>
<td>Quartz</td>
<td>12</td>
</tr>
<tr>
<td>Zircon</td>
<td>16</td>
</tr>
<tr>
<td>Clay</td>
<td>6</td>
</tr>
<tr>
<td>Zinc oxide</td>
<td>2</td>
</tr>
<tr>
<td>Talc</td>
<td>4</td>
</tr>
<tr>
<td>Corundum</td>
<td>9</td>
</tr>
<tr>
<td>Wollastonite</td>
<td>13</td>
</tr>
<tr>
<td>Colemanite</td>
<td>—</td>
</tr>
</tbody>
</table>

All of the glaze recipes from wet grinding were sieved under a size of 45 μm. The glaze recipe samples were applied to the ceramic floor tiles by a laboratory pistol (Mercan, 1993). The unbaked floor tiles used in the tests had dimensions of 20x20 cm and were prepared under industrial conditions. The glazing was performed on the ceramic body after the engobe, with a constant density and quantity applied. The glazed floor tiles were fired in a roller-type high-intensity oven. The sample preparation method, firing time, amount of engobe and glaze contents were kept constant. For each test procedure, measurements were performed on eight different samples, and the average of these measurements was taken for all tests. The flow sheet of the experimental study is shown in Figure 1.

2.2 Tests applied to glaze samples

Several tests were conducted on the six glaze recipe samples in order to determine the effect of colemanite addition.

2.2.1 Density test on glaze recipe samples

The density test of the glaze recipe samples was carried out using a pycnometer. The results of the density tests are shown in Figure 2 and Table 2. It can be seen from Figure 2 that the density of the samples increased with the addition of colemanite.

Figure 2 The effect of colemanite amount on the density of glaze recipe samples.

Table 2. Density values of glaze recipe samples with various amounts of colemanite addition

<table>
<thead>
<tr>
<th>Glaze Samples</th>
<th>Stand</th>
<th>N1</th>
<th>N2</th>
<th>N3</th>
<th>N4</th>
<th>N5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density (gr/l)</td>
<td>1710</td>
<td>1690</td>
<td>1702</td>
<td>1704</td>
<td>1705</td>
<td>1705</td>
</tr>
</tbody>
</table>

2.2.2 Sieve analysis of glaze recipe samples

The glaze recipe samples were sized with a 45-μm sieve. It was observed from the sieve analysis that the oversize proportion did not vary significantly, being in the range of 1.5-1.9%.

2.2.3 Fluxing time of glaze test

The fluxing time of the glaze recipe samples was determined with a ford-cup viscosimeter with a hole diameter of 3mm and volume of 100 ml. The results for the fluxing time of the glaze tests are given in Table 3 and Figure 3. It can be seen that the fluxing...
time of the glaze increases with increases in colemanite addition.

Table 3. Fluxing time values of glaze recipe samples with various amounts of colemanite addition.

<table>
<thead>
<tr>
<th>Glaze Samples</th>
<th>Stand</th>
<th>N1</th>
<th>N2</th>
<th>N3</th>
<th>N4</th>
<th>N5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluxing Time (second)</td>
<td>12.4</td>
<td>13.9</td>
<td>17.2</td>
<td>24.9</td>
<td>32.3</td>
<td>34.5</td>
</tr>
</tbody>
</table>

Figure 3. The effect of colemanite addition on fluxing time of glaze recipe samples.

2.3 Physical tests done on glazed ceramic floor tiles

The ceramic floor tiles, in industrial condition, were first engobed, then glazed with various glaze recipes containing various amounts of colemanite, and finally fired in ovens. Various physical tests were performed on these samples.

2.3.1 Water absorption test

The water absorption results for die glazed floor tiles with various amounts of colemanite are shown in Table 4 and Figure 4. It can be seen from Figure 4 that die water absorption values of the glazed floor tiles are very close to each other.

2.3.2 Strength test

The strength values of the glazed and fired floor tile samples with various amounts of colemanite are shown in Table 5 and Figure 5. It was found that the strength of fired floor tile samples is not affected significantly since it is directly related to the composition of the ceramic bodies and the firing temperature.

Table 4. Water absorption values of the glazed floor tile samples with various amounts of colemanite.

<table>
<thead>
<tr>
<th>Glazed Floor Tile Samples</th>
<th>St.</th>
<th>N1</th>
<th>N2</th>
<th>N3</th>
<th>N4</th>
<th>N5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water absorp. (%)</td>
<td>2.9</td>
<td>2.9</td>
<td>2.8</td>
<td>2.9</td>
<td>2.8</td>
<td>2.9</td>
</tr>
</tbody>
</table>

Figure 4. The effect of colemanite content in glazed floor tile samples on water absorption.

Table 5. Strength of fired and glazed floor tile samples with various amounts of colemanite.

<table>
<thead>
<tr>
<th>Glazed Floor Tile Samples</th>
<th>Strength of fired Tiles (kg/cm²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>St.</td>
<td>340</td>
</tr>
<tr>
<td>N1</td>
<td>337</td>
</tr>
<tr>
<td>N2</td>
<td>340</td>
</tr>
<tr>
<td>N3</td>
<td>338</td>
</tr>
<tr>
<td>N4</td>
<td>338</td>
</tr>
<tr>
<td>N5</td>
<td>337</td>
</tr>
</tbody>
</table>

Figure 5. The effect of colemanite content on strengths of glazed and fired floor tile samples.

2.3.3 Harkot and Autoclave tests

The Harkot test was performed on the floor tile samples to determine their strength properties under instantaneous temperature variation. An autoclave test was conducted to evaluate the reaction of the floor tile samples under water vapor pressure conditions. The results of the Harkot and autoclave tests for all samples are shown in Table 6. They were found to be satisfactory.

Table 6. Results of Harkot and autoclave tests on glazed floor tile samples with various amounts of colemanite.

<table>
<thead>
<tr>
<th>Glazed Floor Tile Samples</th>
<th>Harkot</th>
<th>Autoclave</th>
</tr>
</thead>
<tbody>
<tr>
<td>St.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>N1</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>N2</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>N3</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>N4</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>N5</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

2.3.4 Stain test

Glazed ceramic floor tiles are considered to be of first-grade quality when all stains are removed from
them. All the ceramic floor tile samples were found to be first grade after stain tests were performed.

2.3.5 Test for acid-base resistance
The glazed ceramic floor tile samples were subjected to acidic and basic mediums so as to determine their susceptibility to acids and bases. The glazed floor tile samples were subjected to 3% HCl and 30 gr/lt KOH. The results indicate that the floor tile samples are resistant to acidic and basic mediums.

2.3.6 Surface corrosion test
The ceramic floor tiles samples were found to be suitable for use in workplaces and markets after standard surface corrosion tests.

2.3.7 Surface hardness test
A surface hardness test was performed to measure the resistance of the ceramic floor tile samples against scratching. The surface hardness of the floor tile samples was found to be level 6 on the Mohs scale.

2.3.8 Fired color test
Fired color test results for the glazed ceramic tile samples with various amounts of colemanite are given in Table 7. It may be seen that they are very close to each other. In this table, L, a and b correspond to the degree of white, green and yellow, respectively.

<table>
<thead>
<tr>
<th>Glazed Floor Tile Samples</th>
<th>Color L</th>
<th>N1</th>
<th>N2</th>
<th>N3</th>
<th>N4</th>
<th>N5</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>86.6</td>
<td>85.62</td>
<td>85.5</td>
<td>86.3</td>
<td>86.74</td>
<td>85.56</td>
</tr>
<tr>
<td>A</td>
<td>1.1</td>
<td>1.38</td>
<td>1.44</td>
<td>1.45</td>
<td>1.46</td>
<td>1.54</td>
</tr>
<tr>
<td>B</td>
<td>6.8</td>
<td>6.65</td>
<td>6.65</td>
<td>6.00</td>
<td>6.12</td>
<td>6.40</td>
</tr>
</tbody>
</table>

3 RESULTS AND CONCLUSIONS
In this study, the physical properties of ceramic floor tile samples prepared by adding various amounts of colemanite to standard glaze recipes were investigated in relation to its appropriateness for ceramic bodies. The following results were obtained:

1. The density of the glaze recipe samples increased as the amount of colemanite increased.
2. The oversize percentage in the glaze recipe samples was found to be between 1.5% and 1.9%.
3. The fluxing time of the glaze recipe samples increased with increasing in the colemanite concentration. The standard glaze recipe samples without colemanite had a fluxing time of 12.3 seconds, while the new glaze recipe samples with colemanite had a fluxing time of 34.5 seconds.
4. The water absorption values of the glazed floor tile samples were found to be in the range of 2.8-2.9%. This is consistent with standard TS-EN 176 for ceramic floor tiles.
5. It was determined that the strength of the fired and glazed floor tile samples was between 337 and 340 kg/cm². This is also consistent with standard TS-EN 176.
6. The results of the Harkot and autoclave tests on the glazed floor tile samples are positive.
7. The stain test and resistance to acid-base test results are consistent with standard TS-EN 176.
8. The corrosion and surface hardness tests on the floor tile samples indicate that they can be classified as rank 4 and level 6 according to standard TS-EN 176 for surface corrosion and the Mohs scale, respectively.
9. The fired color tests showed that the floor tile samples with various amounts of colemanite were similar to the standard floor tiles.
10. The floor tile samples with contents of 6%, 8%, and 10% colemanite had less cavities and much smoother surfaces than the floor tile samples without colemanite, which had dull surfaces. Therefore, the glaze recipes with contents of 6%-8% colemanite should be utilized. This will enable the large potential borate reserves of Turkey to contribute significantly to its economy.
11. Optical and electron microscope measurements of the floor tile samples should be done for future work. This will ensure that fine details and deficiencies caused by engobe, glaze or body are determined more effectively.

REFERENCES
TS-EN 176, Standards for Ceramic Floor Tiles, Turkey Standards Institute.
Productivity of Brown Coal for Briquette Production

D.A. Kunoyev Institute of Mining, Ministry of Power, Industry and Trade, Almaty, Republic of Kazakhstan

ABSTRACT: This paper discusses such problems as the necessity of bricketting coal, the efficiency of complex coal processing, and the basic consumer properties and evaluation criteria of briquettes. Also outlined are methods which enable the production of briquettes of high quality, results for tested briquettes made in laboratory conditions, and recommendations for manufacturers.

1 INTRODUCTION

During production, transportation and storage, brown coal is crushed by 20 to 80% of its original volume and turns into coal dust or chaff. Thus, during layer burning of raw coal in furnaces, and in small and average-size boilers, up to 40% of this dust drops unburned through grid-irons together with ashes, and up to 20% of small pieces are lost through pipes together with smoke gases. Up to 30% of the total coal is lost in these ways. Thus, for every 3 tons of coal, 1 ton is not used, but instead pollutes the environment. In addition, pulverised coal frequently causes spontaneous combustion in storage. Therefore, bricketing of dust with particle sizes of 6-8 mm is an important issue.

It is more effective to make coal bricks together and in conjunction with other coal production: synthesis - gas, synthetic combustible, absorbents, re-storing agents, etc. Complex production of coal products reduces the costs of the crushing of coal up to the required standard, and the heavy hydrocarbons obtained during coal gasification and hydrogenation are used in the production of briquettes as binding substances. It is also possible to respond actively to fluctuations of coal product sales in the market, by changing the volume of briquette production and, for example, liquid combustible from the same volume of coal with the purpose of stabilizing the volumes of production, transportation and crushing.

The production of briquettes is carried out with known techniques, but for each type of coal under optimum technological modes and parameters, which need to be determined by research and development.

The basic consumer properties and seams are made in laboratory conditions for the solution of these problems. Briquettes are tested for durability, heat-generation ability and water-resistance. For these purposes, the theoretical bases of coal bricketting with and without binding substances are developed. The technique of production and tests of experimental briquettes from coal dust are developed, and the laboratory equipment and devices are assembled.

The research conducted has shown that the best binding materials are bitumen and wastes of cellulose-paper production, but bricketing brown coal is possible without binding substances. The optimum humidity of briquettes is within the limits of 14-18%. At a specific pressure of 0 to 130 MPa, the strongest briquettes are obtained. For production of briquettes at indicated pressures, the design and creation of a special rolling type pressing is necessary. On the basis of the results of the research, the technological principles for designing briquette factories are developed, in which technological, economic and ecological demands, and also the need for safe technology in briquette factory processing of coal dust to briquettes (small coal pieces) of Kýyk-tinsk B3 grade brown coal are fulfilled.

The evaluation criteria for briquettes are its heat-generating ability (calorie content), durability during transportation, storage and burning, and also atmosphere-resistance (in particular water-resistance). These properties of briquettes depend on the material and granulometric structure of the bricketted coal, the properties of the binding substance and other components added for the improvement of separate qualities of briquettes. The properties and quality of briquettes also depend on the production...
conditions and, in particular, the pressing of briquettes, i.e., on the "know-how" of briquettes.

Therefore the basic purpose of the research carried out in the D. A. Kunayev Institute of Mining is the development of the optimum "know-how" for briquettes for various grades of coal extracted in the Republic of Kazakhstan. For this purpose, the following problems are being tackled:

1) Determination of optimum granulometrics of coal structure for bricketing.
2) Establishment of optimum binding substance and other necessary components.
3) Establishment of optimum amount of water added for bricketing.
4) Determination of optimum pressure and temperature of pressing.
5) Determination of optimum size of the briquettes for their various applications.
6) Determination of optimum methods of coal drying and cooling of briquettes.
7) Development of technology for packing briquettes for loading, transportation and storage.

The experimental briquettes are made from various grades of coal from different deposits.

2 CONCLUSIONS
The suggested principles and the "know-how" of coal brick production in conjunction with production of other coal products results in increased quality of briquettes, reduction in cost, increased profitability of production and combustible consumption, increased competitiveness of briquettes in consumers markets, and reduced anthropogenous load on the environment by prevention of self-burning and contamination of the bio-sphere with small coal pieces.

The technique of bricketing in conjunction with the production of other coal products is more effective in brown coal processing, as it is richer in organic materials, in humuses and pitch, and is crushed more than other types of coal. The coal bricks are intended mainly for municipal-household use and for consumption in trains, schools, hospitals and military houses.
A New Microscale Flotation Cell: Combination of Canadian Column and Partridge-Smith Cell

A. Özkan & M. Yekeler
Department of Mining Engineering, Cumhuriyet University, Sivas, Turkey

ABSTRACT: Flotation is one of the most important physicochemical processes for mineral separations and other recovery operations. Flotation machines have been developed since the beginning of the 19th century and are still under intensive research and development. The devised cell is a combination of the Canadian Column Flotation Cell and Partridge-Smith Cell. The materials used for the construction of the newly-designed cell are cheap and available laboratory accessories and aquarium materials. The cell functions well in terms of its scale, the small samples required and control. It can be used anywhere in the laboratory for research and in the classroom for demonstrations of experiments. Some of the data obtained using this cell by the flotation method are in good agreement with the contact angles method measured independently on the same minerals. The measurements obtained in our experiments are comparable to those previously published for the same minerals used.

1 INTRODUCTION

Flotation is one of the most important physicochemical separation processes, used largely in mineral separation operations. This method is not only used in mineral concentration and recovery operations, but also in water treatment and purification, the recycling of secondary materials, and the recovery of ionic and colloidal materials from aqueous solutions (Yarar, 1988a).

Flotation is a process for separating finely-ground minerals or solids dispersed in water (their slurry densities are usually 10-30% by weight) from associated worthless minerals or gangues. It is based on the adhesion of the minerals to air bubbles that are generated in the pulp. These air bubbles carry the floatable minerals or solids to the flotation cell surface, where they are taken as valuable solids or minerals referred to as 'concentrates'. The solids or minerals which do not float with the air bubbles stay in the pulp or at the cell bottom. They are known as 'tailings' (Weiss, 1985).

Flotation machines are made of a cell equipped with air, solution and mineral entry parts and exit parts. These machines perform the following functions:
- mixing and agitation of the slurry,
- aeration and promotion of particle-bubble collisions,
- formation of a froth layer and its removal for product recovery,
- continuous discharge of tailings.

A new version of an old flotation cell has led to the development of column flotation cells. These cells perform a good job of separating the highly hydrophobic minerals from their associated hydrophilic minerals (Yarar, 1988a).

2 THEORETICAL CONSIDERATIONS

The wettability characteristics of mineral surfaces can be defined in terms of their critical surface tension of wetting values (\(\gamma_c\)) to achieve selectivity in froth flotation, adhesion and welding. Therefore, the \(\gamma_c\) values of minerals can be used as a measure of their flotation behavior and response. The floatability or wettability characteristics of solids or minerals are estimated quantitatively by a number of experimental and empirical techniques. This quantifying parameter mentioned above is die critical surface tension of the wetting (\(\gamma_w\)) values of minerals or solids (Yekeler & Yarar, 1994a). The Zisman contact angle measurement technique and the flotation method are die two major techniques used to determine the \(\gamma_c\) values of minerals among these experimental and empirical techniques (Kelebek, 1987, Yarar, 1988b, Parekh & Apian, 1974, Williams & Fuerstenau, 1986, Yekeleri Yarar, 1994b). The Zisman method, which uses a plot of \(\cos \theta\) (where \(\theta\) is the contact angle) against \(\gamma_w\) (where \(\gamma_w\) is the solution surface tension)
tension), gives a line that intercepts the x-axis at \( Y_c = Y_{LV} \). As can be seen from Figure 1, at \( Y_c > Y_{LV} \) the liquid spreads on the solid (mineral), while at \( Y_c < Y_{LV} \) a contact angle, \( \theta > 0 \) is established in the solid/liquid/vapor interfacial region (Yarar & Aksu, 1997).

The flotation method, on the other hand, estimates the \( Y_c \) value of any mineral by plotting % recovery (\%R) versus solution surface tension (\( Y_{LV} \)) with the extrapolation of the linear part of the \( R-Y_{LV} \) curve to the surface tension axis in order to obtain an intercept at \( \%R = 0 \) as given in Figure 2.

3 THE NEWLY-DESIGNED CELL

The devised cell is a combination of the Modified Partridge-Smith Cell (Partridge & Smith, 1971) and the Canadian Column Cell (Yoon, 1994) as shown in Figure 3. The cell is made from a graduate cylinder and various plastic materials. The total volume of the cell is 255 cm³, the internal diameter is 36 mm and the height is 250 mm. There is an air-sparging system, which is used in aquariums as air stone, at the bottom of the cell. There are two channels (canals) to the microflotation column cell above the air-sparging system: one is the solution/liquid entry channel, the other one is the froth (floatable material) collecting channel, which is made of "plastic" material.

The main function of the air-sparging system is to produce enough micro-bubbles to adhere to the mineral particles.

In order to produce the micro-bubbles required, there is an air-producing system (called the 1st air compressor, shown in Figure 3) which is turned on by button S3. The air flow rate to the cell is 200 cm³/min, and this rate is adjusted by an air valve in the control panel. The liquid or solution is fed to the cell by a liquid reservoir which is made from a glassware separating funnel available in any laboratory. The feed part is a removable unit and it is taken apart from the cell after the feeding process is completed. In order to push the liquid to the cell, the 2nd air compressor, controlled by the S2 button, is employed to send the air at a feed rate of 75 cm³/min liquid flow. Both the air and liquid flow rates are adjusted by air and liquid valves located in the control panel shown in Figure 3. The 1st and 2nd air compressors, valves, plastic hoses and small motors are die same as those used in aquariums.

The main advantages of this newly-designed microscale flotation cell are as follows:

- The cell represents both the functions of the Canadian Column flotation and Modified Partridge-Smith Cell in small scale.
- It needs small amount of sample, solution, chemicals and air to perform tests.
- The cell works well in terms of sparger design, aeration rate, froth-collecting zone and operating pressure.
- It is easy to use, control and carry anywhere in the laboratories or classroom for demonstration.

The materials used for the constructing of the cell and its operating system are cheap and easily obtainable materials.
Figure 3 The new microscale flotation cell, which is a combination of the Canadian column and Partndge-Smithcell

3.1 Operating the system

The cell is operated at very low pulp density dilutions, which requires small amounts of sample to be separated from each other. After putting the solution in the reservoir, the S2 button is turned on to push the solution from the reservoir into the cell by means of pressured air. The SI button is used when the micro-bubbles are needed to float the particles from the particles that are unfloatable for the separation processes. Therefore, the floatable solids, known as the froth, are collected from the collecting channel as the 'floatable product' shown in Figure 3.

3.2 Experimental test results

3.2.1 Flotation recoveries of some minerals

Four minerals were tested in the new cell. These minerals are from different sources, being naturally floatable and non-floatable minerals. Two grams of sample was tested with 3 minutes of conditions and 1 minute of flotation. The floatable material (froth) was collected, filtered, dried and weighed in order to determine the % recovery values [(% Recovery = Floatable weight / Feed) x 100].

These minerals were floated alone, in binary mixtures, without chemical reagents and with chemical reagents. The results are outlined in Table 1.

These results are comparable with the values in the literature (Kelebek & Smith, 1985).

3.2.2 Determination of the $\gamma_c$ values of minerals

Methanol solutions of different concentrations (%w/w) were prepared and used to obtain wettability data for construction of $\cos \theta$ versus $\gamma_LV$ and % Recovery versus $\gamma_LV$ plots for talc and sulfur minerals. The solutions used in all the experiments were prepared with mono distilled water. The surface tensions of the solutions were measured by the drop-weight method (Padday, 1968). The surface tensions of methanol solutions determined against methanol concentration are shown in Figure 4, with a comparison of the experimental data with data in the literature (Weast, 1987). The measured values of the surface tensions of the methanol solutions (by %v/v) were very close to the data in the literature.

The contact angle measurements were made with a Contact Angle Goniometer (Rame-Hart Inc., Model 100 NRL) in order to construct Zisman's wettability data. From these data, as shown in Figure 5, the $\gamma_c$ values of talc and sulfur were obtained; they were found to be 31 and 26, respectively. The $\gamma_c$ values of these minerals were also found to be 30 mN/m for talc and 28 mN/m for sulfur, using the same solutions as for the flotation medium in our newly-designed cell. The flotation results from the new cell are shown in Figure 6.
The designed cell is a combination of the Canadian column flotation and Partridge-Smith Cell.

The materials used for the construction of the newly-designed cell are cheap and available laboratory accessories and aquarium materials.

The cell needs small amount of sample, solutions, chemicals and air to perform the experiments. It also works well in terms of the sparger design, aeration in our newly-designed flotation cell, indicating that the new cell performs well.

### Table 1. The recoveries of some minerals obtained in the designed cell.

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Feed Sample, g</th>
<th>Recovery, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfur</td>
<td>-38</td>
<td>97.3</td>
</tr>
<tr>
<td>Calcite</td>
<td>-38</td>
<td>40.0</td>
</tr>
<tr>
<td>Talc</td>
<td>-38</td>
<td>46.0</td>
</tr>
<tr>
<td>Galena Concentrate</td>
<td>-20</td>
<td>6.6</td>
</tr>
</tbody>
</table>

#### III. Binary Mixtures (1:1) with No Chemical Reagents

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Feed Sample, g</th>
<th>Recovery, %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sulfur + Calcite</td>
<td>-38</td>
<td>43.3</td>
</tr>
</tbody>
</table>

*Sodium chlorite (100 mg/L)

**Potassium acryl xanthate (250 mg/L)**

**CONCLUSIONS**

The combined data obtained from the contact angle measurements and flotation tests are outlined in Table 2. As it can be seen, the \( \gamma_c \) values obtained by goniometer are very close to fee values obtained in our newly-designed flotation cell, indicating that the new cell performs well.
rate, froth-collecting zone and operating pressure. Therefore, it is easy to use, control and carry this cell anywhere in the laboratory or classroom for demonstrations.

The quantitative test results obtained from this cell are in good agreement with those obtained with other techniques and equipment.

REFERENCES


Concentration Studies on Chromite Tailings by Multi Gravity Separator

Ş.G.Özkan & B.Ipekoğlu
Istanbul University, Engineering Faculty, Mining Eng. Dept., 34850, Avcilar, Istanbul, Türkiye

ABSTRACT: In this study, a representative sample was obtained from Uckopru mine in Türkiye in order to beneficiate fine-sized chromite tailings. Physical and chemical analyses showed that while the whole tailings contained 12.80% Cr2O3 (100% by weight), the particle size fraction of minus 38 microns contained 25.20% Cr2O3 (33.20% by weight) and the particle size fraction of plus 38 microns contained 6.64% Cr2O3 (66.80% by weight). The mineralogical investigations revealed that the sample contained chromite and serpentine group minerals as its main components. For concentration purposes, a Multi Gravity Separator was used for chromite fines in the experiments. The best results using the MGS device for the sample of minus 38 microns were obtained under the following conditions: angle, 2°; frequency, 5.7 cpm; amplitude, 15 mm; and drum speed, 230 rpm. These conditions gave a chromite grade of 47% Cr2O3 with fairly high recovery of 72%.

1 INTRODUCTION

It is a well-known fact that Türkiye is one of the major chromite producers in the world. Approximately a million tons of chromite ore are mined and processed for further metallurgical treatment and use in the country. Large amounts of fine-sized chromite tailings naturally occur during production and processing, and this causes severe environmental hazards as well as source losses due to the high chromite content.

There have been numerous research studies, Guney (1990), Guney et al. (1991), (1992), (1993), Ozdag et al. (1993), (1994), Ucbas and Ozdag (1994), Guney et al. (1994), Gui et al. (1995), Cicek et al. (1998), Sönmez and Turgut (1998), Gence (1999), on the beneficiation of these tailings in recent years. The Uckopru Mine of Eti Holding in particular was thoroughly investigated with regard to roughly a million tons of fine-sized chromite tailings that have been stored there for about a 100 years, and several treatment techniques have been developed for fine tailings, such as column flotation and gravity concentration devices.

For the reasons given above, beneficiation experiments were carried out on samples of fine-sized chromite tailings from Eti Holding’s Uckopru Mine by the Mining Engineering Department of Istanbul University.

2 MATERIAL AND METHODS

In this study, about 120 kg of representative sample was obtained from Eti Holding in order to evaluate fine-sized chromite tailings. It was determined by physical and chemical analyses that while the whole tailings sample contained 12.80% Cr2O3 (100% by weight), the particle size fraction of minus 38 microns contained 25.20% Cr2O3 (33.20% by weight) and the particle size fraction of plus 38 microns contained 6.64% Cr2O3 (66.80% by weight). The sample preparation flowsheet is given in Figure 1 in detail. The complete mineralogical investigations showed that the sample contained chromite and serpentine group minerals (crizotile, antigorite, olivine, pyroxene and tremolite) as its main components. The results of the complete chemical analyses according to particle size classification in a 38-micron sieve are also given in Table 1. Moisture analysis showed that the sample contained roughly 8% moisture in the form it was obtained and that it could be dried and lose its moisture in a 24-hour period. In order to determine the particle size distribution of the sample prior to concentration, a screening analysis was performed and the results are shown in Table 2.
3 EXPERIMENTAL STUDIES

As a method, a Multi Gravity Separator (MGS) was used for concentration purposes. The MGS is an efficient device that can easily separate two minerals from each other provided that there is a reasonable difference between their specific gravities. The MGS is generally used for treatment of fines with a maximum particle size of about 100 microns.

A laboratory type of MGS with a length of 0.6 m and radius of 0.5 m was used in the experiments. The MGS is made up of a slightly tapered open-ended drum that rotates in a clockwise direction, producing 6-24 g of gravitation, and is shaken in a sinusoidal form in an axial direction. The parameters affecting the efficiency of separation of the MGS are the drum speed (130-280 rpm), tilt angle (0-9°), shaking amplitude (10-25 mm), shaking frequency (4-6 cps), amount of wash water (0-10 litres per minute) and feed pulp density (20-50% solid by weight).

![Sample preparation flowsheet.](image)

Figure 1. Sample preparation flowsheet.

<table>
<thead>
<tr>
<th>Component</th>
<th>CaO</th>
<th>Fe₂O₃</th>
<th>SiO₂</th>
<th>Al₂O₃</th>
<th>MgO</th>
<th>CuO</th>
<th>Weight %</th>
</tr>
</thead>
<tbody>
<tr>
<td>-38 microns</td>
<td>6.64</td>
<td>15.60</td>
<td>28.03</td>
<td>1.51</td>
<td>45.07</td>
<td>0.35</td>
<td>66.80</td>
</tr>
<tr>
<td>&gt;38 microns</td>
<td>25.20</td>
<td>14.10</td>
<td>19.80</td>
<td>5.40</td>
<td>31.60</td>
<td>1.10</td>
<td>33.20</td>
</tr>
<tr>
<td>Total</td>
<td>12.80</td>
<td>15.10</td>
<td>25.30</td>
<td>2.80</td>
<td>40.60</td>
<td>0.60</td>
<td>100.00</td>
</tr>
</tbody>
</table>

Table 1. Complete chemical analysis results according to particle size of the sample
weight) (Chan et al. 1990, Gence 1999, Yüce 1994). A Mozley C-705 model hydrocyclone was used to determine and optimize the vortex diameter, apex diameter and feed pressure for classification purposes, while the solid-liquid ratio was fixed at 30% during all experiments. Prior to the concentration experiments, the sample was classified according to particle size by means of a hydrocyclone in regard to several parameters in order to categorize the sample into two different particle size fractions, i.e., plus 38 and minus 38 microns.

The concentration experiments failed for the plus 38-micron particle size, whose weight was found to be about 67% of the whole sample, due to an inadequate degree of liberation, and this fraction was stored for further size reduction and concentration studies.

The MGS device gave fairly promising results for the chromite tailing samples at particle size fractions of minus 38 microns due to the fact that there was an appropriate degree of liberation and rich chromite content.

Several experiments were carried out to optimize the parameters of the MGS device for the sample at the particle size fraction of minus 38 microns, and the stable amplitude and frequency values were 15 mm and 5.7 cpm, respectively. The best results using MGS for the particle size fraction of minus 38 microns were obtained under the following conditions; an angle of 2° and drum speed of 230 rpm. These conditions gave a chromite grade of 47% C12O3 with fairly high recovery of 72%.

The graphs below (Figures 2 and 3) show the results of the optimization studies for comparison of the conditions. All experiments showed that before the application of any concentration method dependent on gravity, such as MGS, the particles should be thoroughly classified by hydrocyclone and the parameters for different cut sizes for hydrocyclone should be optimized. It is recommended that different fine particle size fractions, which could be obtained by changing the hydrocyclone parameters, should be tested with MGS for further beneficiation studies.

![Figure 2 Results of MGS experiments according to drum speed at an angle of 2°.](image)

![Figure 3. Results of MGS experiments according to angle at a drum speed of 230 ipm.](image)
DISCUSSION AND CONCLUSION

Beneficiation experiments were carried out on samples of chromite tailings from Eti Holding’s Uçkopru Mine. It was determined by chemical analyses that while the whole tailings sample contained 12.80 % Cr₂O₃, the particle size fraction of minus 38 microns contained 25.20 % C₉O₆ and contained 12.80 % Cr₂O₃, the particle size fraction of plus 38 microns contained 6.64 % Cr₂O₃. The complete mineralogical investigations showed that the sample contained chromite and serpentine group minerals (crizotile, antigorite, olivine, pyroxene and tremolite) as its main components.

For classification purposes, a Mozley C-705 model hydrocyclone was used in the present study. The cycloning tests were used to optimize the vortex diameter, apex diameter and feed pressure, while the solid-liquid ratio was fixed at 30 % during all experiments.

For concentration purposes a MGS device was utilised, and this separator gave fairly promising results for the chromite tailing samples at particle size fractions of minus 38 microns due to the presence of an appropriate degree of liberation and rich chromite grade.

The best results using MGS for the particle size fraction of minus 38 microns were obtained under the conditions: angle, 2°; frequency, 5.7 cpm; amplitude, 15 mm; and drum speed, 230 rpm. These conditions gave a chromite grade of 47 % C₉O₆ with fairly high recovery of 72 %.

When the tests were performed for plus 38-micron fractions, the results were considerably lower than the test results for the minus 38 microns; therefore, the coarse fractions were stored for future evaluations after size reduction.

Finally, all experiments showed that before the application of any concentration method dependent on gravity, such as MGS, the particles should be thoroughly classified by hydrocyclone and the parameters for different cut sizes for hydrocyclone should be optimized. It is recommended that different fine particle size fractions, which could be obtained by changing the hydrocyclone parameters, should be tested using MGS for further beneficiation studies.

ACKNOWLEDGMENTS

"This work was supported by the Research Fund of Istanbul University. Project number: 1207/070998"
Influence of Coal Face Mechanization on the Properties of Çayirhan Lignite

A. Özder
General Directorate of Turkish Coal Enterprises, Ankara, Turkey

G. Özbayoğlu
Department of Mining Engineering, Middle East Technical University, Ankara, Turkey

ABSTRACT: The influence of coal face mechanization on the properties of r.o.m. coal - size, ash, sulfur, and moisture contents, volatile matter and calorific values - were investigated at Çayirhan (Ankara). Representative samples were obtained from r.o.m. coal and channel samples from two seams of panel A03 of Çayirhan Underground Coal Mine and chemically and physically analyzed. In addition, samples from roof and floor rock layers, which were unavoidably mixed with r.o.m. coal during mining operations, were taken and analyzed. The results show that the amount of fines and the ash, sulfur, and moisture contents of r.o.m coal in mechanized panels of Çayirhan Underground Coal Mine have increased, while the calorific value has decreased.

1 INTRODUCTION

In order to improve working conditions and safety, increase production levels and decrease unit cost of production, it is essential to make use of up-to-date coal mining techniques, machinery and equipment in underground coal mines.

In current conditions, Turkey is not able to meet the demand for energy using only domestic sources. In the current situation, only around 33% of Turkey’s energy needs are fulfilled by domestic sources. Turkey, with an annual population growth rate of 1.5% and an average annual growth rate in electricity required of 8.5%, is likely to be more and more dependent upon imported energy sources in the next few years (Özder et al, 1999). In addition, Turkish Coal Enterprises foresee that most of the open pit reserves in the country will have been mined within the next 10 years. In order to attain planned levels of coal production, underground coal reserves gain special attention. According to TKI, in the next 10 years and beyond, more coal is to be mined from underground mines. In addition to all this, due to the growing pressure resulting from environmental concerns, all products must be cleaner and more environmentally friendly than the ones produced in the past.

Therefore, to meet the demand of 2x150 MW power plants, the first fully mechanized panel went into operation in Çayirhan Underground Coal Mine in 1986.

Despite the fact that the study presented here was carried out in 1990, due to the rapid depletion of open-pit reserves and the necessity of designing more underground mines in the next coming years, the subject is still important and valid. All of the results of the analyses show that coal samples from mechanized faces are inferior to channel coal samples. Therefore, the results are significant for mines faced with transition from open-pit coal mining to underground coal mining.

2 GENERAL INFORMATION ABOUT THE MINE

2.1 Location of the mine

The area under consideration lies within the boundaries of the Ankara and Bolu provinces. Çayirhan is 125 km away from Ankara. The altitude of the region is approximately 1000 m.

2.2 Coal seam

Çayirhan Underground Coal Mine consists of two overlying lignite seams separated by an innerburden (interbedded siltstone layer) of 80 cm in average thickness. The upper seam is around 1.70 m thick and the lower seam is 1.80 m thick. A simplified section, showing the seams and the innerburden strata is shown in Figure 1.
2.3 Mining method

At the panel where this study was conducted (A03), the whole face section was mined as two longwall panels (upper face and lower face) working simultaneously. The interbedded siltstone layer was not mined but left at the goaf. The designed average distance between the upper and lower faces is around 30 m. The seams are located at a depth of 150-200 m from the surface with an inclination of 3°-45°.

A gradual transition from a conventional mining system to a semi-mechanized and then fully-mechanized underground coal mining system was first realized at Çayırhan Underground Coal Mine, formerly a subsidiary of the state-owned Turkish Coal Enterprises but recently privatized. Most of the modernization and transition of the mining system took place between 1982 and 1988. The first fully mechanized panel started to mine lignite in 1986. With the conventional mining system, coal production was carried out manually, using picks, shovels, and blasting. The production rates were relatively low (225 tons/shift).

In Çayırhan Underground Coal Mine, a semi-mechanized panel was in operation between 1983 and 1985. At the face, extraction and the loading of coal onto the conveyor was carried out by hobel. Despite its unsuitable design and die fact that it was taken from Turkish Hard Coal Enterprises (another state-owned company) only on a trial basis, coal production increased to 460 tons/shift.

Finally, with fully mechanized panels, cutting and loading of the coal from the face was carried out by double-drum ranging arm shearer loaders. Face support was achieved with shield-type hydraulic supports. This system has raised the coal production to 1240 tons/shift.

2.4 Change in r.o.m. production levels by year

At the time when this study was conducted (1990), mined coal was not subject to any washing processes. It was only screened to +46 mm, -46+32 mm, and -32 mm. If we define fine coal as -32 mm and lump coal as +32 mm, Figure 2 below shows the fine and lump percentages according to year.

Table 1 lists the results of the analysis of saleable coal by year. Figure 3, Figure 4 and Figure 5 show the changes of the properties of saleable coal by year.

As it may be seen from the tables and figures presented in this section, there is a gradual decrease in the calorific value and lump percentage of the produced coal and an increase in the ash, total sulfur and fines contents caused by the shift in mining method from a conventional to mechanized system. To determine why all these changes occurred, the experimental procedure described below was carried out.
3 EXPERIMENTAL PROCEDURE

Representative samples of in-situ and r.o.m. coals were collected from the interlaying rocks that were mixed with mined coal during coal-cutting operations. Fig.6 shows the general view of the coal seams, and innerburden layers mixing with mined coal when excavated.

Region i. Roof of the siltstone layer. The shearer unavoidably cuts this region during coal cutting. The thickness of this layer depends upon the geologic conditions and operator's ability. Since there is no geologic disturbance, nothing is mixed from the main roof. This is because it is very strong and the shearer operator can follow this region visually.

Region ii. Floor of the siltstone layer or roof of the lower seam. The siltstone layer between the two seams is so weak that there is continuous siltstone mixing with the mined coal during coal cutting and moving of the shield supports.

Region hi. Floor clay. Due to operating conditions, the operator unavoidably cuts this layer of rock.

At boundary layers or contact zones (regions i, ii, iii), there are bands or zones approximately 15 cm thick which are rich in fossils and pyritic sulfur content. The importance of these bands stems from their mixing with the produced coal. In this way, the sulfur and ash content of the r.o.m. coal is increased.

To find the size distribution and sulfur content of these zones, the procedure below was used; it is not possible to take pure samples of rock layer cuttings from the faces during coal production due to the loading principles of the shearers. For approximation, samples from these regions were collected from a preparatory gallery next to the production panel using a road header. All of the three regions were cut by road header and the samples were subsequently collected.

3.1 Results of analyses of coal samples

For comprehensive understanding of mined coal quality using the mechanized system, channel and r.o.m. coal samples were analysed and compared separately for both seams. The results are as follows:

<table>
<thead>
<tr>
<th></th>
<th>Moisture (%)</th>
<th>Ash (%)</th>
<th>Volatile matter (%)</th>
<th>Fixed carbon (%)</th>
<th>Total S (%)</th>
<th>Lower cal. val. (K.Cal/kg)</th>
<th>Upper cal. val. (K.Cal/kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A03 upper seam (as obtained)</td>
<td>22.86</td>
<td>27.14</td>
<td>29.38</td>
<td>20.62</td>
<td>3.72</td>
<td>2700</td>
<td>3033</td>
</tr>
</tbody>
</table>

Figure 4. Total sulfur content of saleable coal by year

Figure 5. Humidity and ash contents of saleable coal by year

Figure 6. General view of coal seams, and innerburden layers mixing with mined coal when excavated
3.2 Results of analysis of rock cuttings

The percent sulfur content of the rock cuttings was obtained 5.33 and the percent moisture content of the samples was 11. From the screen analysis results it was observed that the size of the rock cuttings generally varied between -50 mm and +0.5 mm.

3.3 Comparison of channel and r.o.m. coal samples of pane! A03 using screen analysis results.

3.3.1 Lower seam

Tables 6 and 7 list the results of the analysis of the lower seam channel and r.o.m. samples on an air-dried basis in different size fractions.

Figures 7, 8 and 9 compare the results for the channel and r.o.m. samples of the lower seam.

3.3.2 Upper seam

Tables 8 and 9 list the results of the analysis of the upper seam channel and r.o.m. samples on an air-dried basis in different size fractions.
Table 8. Results of analysis of upper seam channel samples

<table>
<thead>
<tr>
<th>Size(mm)</th>
<th>Wt(%)</th>
<th>Ash(%)</th>
<th>Total S(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>+100</td>
<td>17.13</td>
<td>28.92</td>
<td>4.12</td>
</tr>
<tr>
<td>-100+50</td>
<td>10.68</td>
<td>30.43</td>
<td>4.73</td>
</tr>
<tr>
<td>-50+18</td>
<td>28.42</td>
<td>31.28</td>
<td>4.26</td>
</tr>
<tr>
<td>-18+10</td>
<td>16.34</td>
<td>28.96</td>
<td>4.02</td>
</tr>
<tr>
<td>-10+0.5</td>
<td>26.32</td>
<td>25.72</td>
<td>4.52</td>
</tr>
<tr>
<td>-0.5</td>
<td>1.6</td>
<td>29.47</td>
<td>4.46</td>
</tr>
<tr>
<td>TOTAL</td>
<td>100</td>
<td>28.92</td>
<td>4.31</td>
</tr>
</tbody>
</table>

Figure 10. Comparison of size distributions of channel and r.o.m. samples of upper seam.

Figure 11. Comparison of ash percentages of channel and r.o.m. samples.

Figure 12. Comparison of sulfur contents of channel and r.o.m. samples.

4 DISCUSSION AND CONCLUSIONS

The analysis of the upper and lower seams in regard to channel and r.o.m. coal samples shows that due to mechanized coal cutting systems, the moisture, ash, and sulfur contents of the r.o.m. coal samples from Çayırhan Underground Coal Mine were greater than those of the channel samples.

The increase in the moisture content of the mined coal is mainly due to the use of water as a coolant for the cutting tools of the shearsers and for the suppression of dust.

The increase in the sulfur content of the mined coal is due to the mixing of the interbedded roof and floor rock cuttings, which have a pyritic sulfur content of 5.33%.

The increase in the ash content of the mined coal is due to the mixing and cutting of neighboring rock layers during coal mining. The increase in the ash content of the r.o.m. samples from the lower seam is greater than that of the samples from the upper seam. This is due to the weak character of the siltstone layer lying between the two seams. Since this weak layer constitutes the roof of the lower longwall face, a considerable quantity of siltstone particles are mixed during mining operations.

Referring to the statistics produced by the mine management (Table 1, Figures 2, 3, 4 and 5), it can be observed that in the transition from conventional to semi-mechanized and then to mechanized systems, the fines, ash and total sulfur contents of mined coal gradually increase and the calorific value decreases.

The mechanization of Çayırhan Underground Coal Mine has led to remarkable improvements in productivity, from 225 tons/shift to 1240 tons/shift, but the detrimental effects of this transition must be carefully considered in all mines faced with such a transition.

REFERENCES


ABSTRACT: The subject of this study is a novel process: the Dry Process (DP) for the concentration of boron ores. The conditions of the concentration of boron ores by the dry process were determined in laboratory and pilot-scale tests. In addition to this, the necessary requirements to apply this process in the plant were investigated. In this process, three types of boron ore were used and they were compared at each step. Before the start of the tests, for determination properties of representative samples of feeds taken from the concentrators (tincal, colemanite and ulexite), particle sizes and the B2O3 % distributions of feeds were investigated, and the B2O3 grade of the samples was 23-32 % as the grades in the plants. In tests conducted in optimum conditions, concentrates with 43-60 % B2O3, tailings with 3-22 % 6263 and 70-96 % recoveries were obtained. In the currently operated plant, the concentrate grade is 32-38 % B2O3, recovery is 80 %, and the tailing grade is 14-16 % B2O3. The investigations showed that boron ores might be calcined. It was understood that higher quality and dry, cleaner products might be obtained.

1 INTRODUCTION

The following ores are exploited by Eti Holding: colemanite and ulexite in Bigadiç, Balıkesir; colemanite in Emet, Kütahya and Kestelek, Bursa; and tincal ore in Kirka, Eskişehir.

Although the evaluation of boron production technology in Turkey is limited in quantity, it is a very valuable mine for countries' economic resources, especially for export. Currently, in boron concentration plants in Turkey, concentrates are produced that fulfill the minimum requirements of the world market in terms of concentration processes. In these plants, only a concentrate ore with content from colemanite-ulexite ore of 32 % B2O3 grade and tincal concentrate with a maximum 32 % B2O3 content from tincal ore with a B2O3 grade of 26 % have so far been produced.

2 PRELIMINARY RESEARCH

At the beginning of this research, all the boron ores had different ratios of crystal water. The occurrence of calcination events and the results of the heating process were investigated in a perlite mine operated in Cumaovasi, İzmir (Eti Holding). For this purpose, interesting results of calcination were first produced by simple tests conducted on tincal ore (Aytekin et al., 1992). Calcination was previously performed by preparing samples for the calcination and dry grinding processes rather than just obtaining concentrates for boron ores. However, it was found that there was no need for these processes when adopting wet grinding.

To date, research on the heating process done by different researchers (Aytekin et al., 1992a, b, c; Aytekin et al., 1993; Eric et al., 1982; Gundiler et al., 1972; Kaytaz et al., 1986; Sener et al., 1992) has been conducted on a pilot scale rather than as a laboratory test for the purpose of concentration. There have been several studies of the application of the calcination process instead of the wet process, which is used in ore production at present. These studies have mainly dealt with the beneficiation of rich tailings and concentrate beneficiation of low-grade ores produced at Kirka, Bigadiç and Emet. There is not a low market demand for these ores. Ores containing inherent crystal water lose it by heating, and as a result, calcination occurs. Due to this characteristic behaviour, the calcination process is applied to tincal, colemanite and ulexite ores (Akcil, 1994).

3 MATERIAL

Boron samples (tincal, colemanite and ulexite) were taken from Eti Holding’s Kirka and Bigadiç concentrator plant fine ore stocks with a weight of
250 kg for each ore. The coarsest sample size was homogenized; this size was 40-50 mm. The tinocal ore had an average content of 26 % B₂O₃, with different parts of the ore body having values of 24-29 % B₂O₃. Colemanite and ulexite had an average content of 32 % B₂O₃.

4 METHOD

Colemanite and ulexite ores calcine by losing crystal water during the heating process under specific conditions. However, tinocal ore behaves in a different manner. When tinocal ore is treated with a heating process, popping occurs and a porous surface is observed. As a result of ore processing, clay minerals agglomerate in different sizes and colors emerge, while gangue minerals are calcined and become hard. The more compacted part, which is made of clayey product and gangue minerals, gives a coarse-size product when squeezed under rollers, while softer parts, which are rich in B₂O₃, give a finer-size product. These two different products are classified by separation and concentrate is obtained.

5 EQUIPMENT USED

All die tests were conducted with the characteristics of a revolving preheating calcination furnace (originally designed by Aytekin, 1992).

The furnace consists of a cylindrical ceramic tube, which is used for moving the materials in pieces inside the furnace and for heating with rotation. This tube has a special position, either balancing its position angle or balancing its rotating velocity. Feeding is done with a vibration feeder. The tube conveying materials into the calcination furnace contains two temperature areas. The first part is 80 cm away from the feeding point of the material. This part, at the same time, is the first drying and calcination temperature point area. The second part is only 40 cm away. In this part, the material loses crystal water and calcination also takes place. A portable thermometer on the control panel is used for measuring the inner furnace temperature.

6 MAIN RESEARCH

The main parameters listed below, in addition to other secondary parameters, were investigated during the calcination tests:

- preheating and calcination temperatures,
- heating and calcination times,
- particle size of the feed.

As a result of the loss of crystal water in the structure of the boron ore under temperature, it calcined. Its expansion properties were determined and systematic calcination tests were conducted. The concentration treatment was completed by the separation of the large particle impurities which remained, such as clay, dolomite and calcite. The white mass should be subjected to further treatment such as screening or air separating. The parameters of calcination temperature, calcination time, particle size and the quantity of feeding, which affected the recovery obtained with this method, were investigated by doing different series of tests for the optimum calcination conditions of Kirka tinocal ore and Bigadiç colemanite-ulexite ores.

Various parameters were tested for separation of the expanded ore from solidified clay together with other gangue minerals after the calcination process, and as a result, the comminution of the end products on top of a uniform rubber plate by roller was preferred. Sieving was used to separate the boron concentrate and tailings. Boron powder is easily separated from solidified clay and gangue minerals after the comminution process. After the tests, the optimum conditions were determined from the results of chemical analysis.

7 APPLICATION OF DRY PROCESS ON BORON ORES

It was found that it was possible to concentrate the tinocal ore by the dry process (DP). The optimum
conditions were determined for the concentration of tincal ore by this process. Thus, systematic tests were carried out to determine the calcination temperature, time and particle size of the used ore. According to the results obtained in the tests, the sample was divided into different seven particle sizes: 50-25 mm, 25-12.5 mm, 12.5-9.51 mm, 9.51-4.76 mm, 4.76-3 mm, 3-1 mm, and -1 mm.

The concentrate grade and recovery obtained as a result of the calcination tests done on ore which was divided into different particle sizes at different times (5-25 mm.) and temperatures (300°C-500°C) were compared with regard to each parameter. First, tests carried out at the optimum temperature, which had been found to be the most suitable calcination temperature beforehand, were evaluated. When the ore was smaller than 9.51 mm with a calcination time of 15 minutes, it was observed that the concentrate recovery was the highest. After this value, a small decrease was observed. For 15 minutes of calcination, the concentrate grade was 46-56 % B_2O_3 with 87-95 % recovery and the tailing grade was 3-13 % B_2O_3.

Calcination tests were conducted on Bigadiç colemanite ore in order to find the optimum calcination conditions in the same manner as for the concentration of tincal ore by the dry process. The optimum conditions in terms of concentrate grade, recovery and tailing grade were found to be a calcination temperature of 500°C, a calcination time of 20 minutes and a particle size of less than 12.5 mm.

The results of tests in optimum conditions showed that calcination recovery was 92-94 %, the concentrate grade was 48-51 % B_2O_3 and the tailing grade was 9-11 % B_2O_3. The concentrate recovery of the Bigadiç concentrator was 80 %, the concentrate grade was 42 % B_2O_3 and the tailing grade was 16 % B_2O_3. Concentration by the calcination tests increased the concentrate recovery by 16 % and the concentrate grade by 19 %, and decreased the tailing grade by 37 %.

Calcination tests were conducted on Bigadiç ulexite ore in order to find the optimum calcination conditions. As a result of the tests, it was observed that the 25-1-mm particle size fraction was the optimum fraction, and the grade and recovery of ulexite concentrate from this fraction constituted the highest values. At the end of the tests, it was determined that the optimum calcination temperature and calcination time were 450°C and 25 minutes, respectively. In the tests carried out at the optimum calcination temperature, it was observed that the water in the chemical composition of the ulexite ore was not completely evaporated, and the surface of the ulexite ore was covered with a block glassy layer over this temperature.

According to these results, the grades and recoveries of the calcined product obtained in the tests carried out in optimum conditions were within the ranges of 43-44 % B_2O_3 and 94-95 %, respectively. In the currently operated plant, the concentrate grade is 36-38 % B_2O_3, concentrate recovery is 80 % and the tailing grade is 14-16 % B_2O_3.
During the concentration of boron ores by the dry process, a specially designed preheated calcination furnace was used. At the end of the tests carried out with this furnace, die optimum calcination conditions were found in terms of calcination temperature, calcination time and feed fraction. For classification, a screening method was chosen. For calcine, a 0.2-mm sieve was found to be suitable. In this classification process, undersize parts were taken as concentrate and oversize parts as tailing.

In the plant, which is located at Kırka, a concentrate with a grade of 32-34 % B2O3, at a recovery of 62-94 % and the tailing grade was 11-32 % B2O3. In optimum calcination conditions, the coarse particle fractions had concentrated B2O3. The tailing grade was the sample grade of the Bigadiç part over 12.5 mm had a grade of 43 % B2O3. This grade was the sample grade of the Bigadiç concentrate. In optimum calcination conditions, 80 % of the original sample mass was over 4.76 mm and had a grade of 32.33 % B2O3. Coarse particle fractions had concentrated B2O3.

In the tests conducted on colemanite ore in optimum conditions, 80 % of the original sample was over 4.76 mm and had a grade of 32.33 % B2O3. Coarse particle fractions had concentrated B2O3. The part over 12.5 mm had a grade of 43 % B2O3. This grade was the sample grade of the Bigadiç concentrator and this sample was used as the concentrate. In optimum calcination conditions, the grade of colemanite concentrate was 31-51 % B2O3 at a recovery of 62-94 % and the tailing grade was 11-32 % B2O3. In addition to the concentration process used in the Bigadiç concentrator to determine the concentration of colemanite by the dry process, laboratory studies were carried out. Finally, instead of a concentrator plant, a dry process plant was used.

In the tests done in optimum conditions, ulexite concentrates with grades of 38-44 %, tailings with grades of 10-13 % B2O3, and 91-95 % recovery were obtained. In the currently operated plant, the concentrate grade is 36-38 % B2O3. Concentrate recovery is 80 % and the tailing grade is 14-16 % B2O3.

The investigations showed that boron ores might be calcined. Final products of concentration with the calcination of boron ores are better than the product of currently operated plants. For this reason, it is concluded that it is more suitable to set up a plant operated with concentration by the dry process. The production of boron derivatives from the concentrate products obtained by the dry process is easier and cheaper technologically. In addition, with this process, the problems of tailing, storing and dewatering can be eliminated.

REFERENCES


