THE IMPACT OF HIGH-POWER ELECTROMAGNETIC PULSES ON THE SURFACE STATE AND TECHNOLOGICAL PROPERTIES OF THE CALCIFEROUS MINERALS

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Abstract

The modification of structural chemical properties of natural fluorite, scheelite and calcite under the impact of high-power electromagnetic pulses (HPEMP–treatment) were studied with help of adsorption of acid-base indicators. It was determined that the HPEMP–treatment during of 30 seconds (3 x 10³ pulses) resulted in the intensification of fluorite surface ‘electron-donating ability and acceptor properties of calcite and scheelite surfaces.

Single mineral flotation tests has allowed to determine that treatment by high-power electromagnetic pulses resulted in the improvement of the calciferous minerals floatability. The rising of the scheelite recovery was 10 – 12%, for fluorite it was 5 – 6%, for calcite 7 – 8%.

Key words: Calcite; Fluorite; Scheelite; Floatability; High-power electromagnetic pulses.

1. Introduction

A detailed study of the relationship between the parameters of the treatment by high-power electromagnetic pulses (HPEMP–treatment) and the surface properties of calciferous minerals is of high relevance for the development and refinement of the existing understanding of the processes involved in the interaction between strong electromagnetic fields and geomaterials of various origin, for the development of a scientific framework for the application of HPEMP–treatment for directed modification of the structure and properties of mineral, and for solving practical problems aimed primarily at improving of the efficiency of calciferous minerals processing [1 - 4].

The complexity of the flotation separation of calciferous minerals is determined by the similarity of the flotation properties of calcite, fluorite and scheelite. Therefore, increasing the flotation efficiency is an urgent technological task.

Traditionally, the collector for scheelite is technical oleic acid. Flotation is carried out in an alkaline medium with the addition of a depressant-liquid glass, which has a slight selective effect. Traditionally, to refine rough scheelite concentrates, a Petrov' method is used, in world practice. The final concentration of scheelite concentrates according to the Petrov' method is an effective, but expensive way. The method consists in processing a rough concentrate at a temperature of 85 - 90 °C with a 2 - 4% solution of water glass, which provides hydrophilization and depression of calcite and fluorite [5].

It has been shown that it is possible to increase the efficiency of main scheelite flotation by using synthetic collectors of stable composition of alpha-substituted chlorocarboxylic or alpha-branched carboxylic acids [6, 7]. Nevertheless, studies to increase the contrast of flotation properties of calcium minerals through a combination of collectors continue to be conducted. It is proposed to use a mixture of a complexing collector of hydroxamate and sodium oleate in a ratio of 1 to 2,
the pulp pH is 8.3 [8, 9].

In the flotation of scheelite ores, liquid glass with module 2 is used as the main depressor of rock minerals. Liquid glass micelles or chains of silicon-oxygen tetrahedrons are able to neutralize the hydrophobizing effect of the collector. The depressing effect of water glass can be powered by combining with polyvalent metals salts or addition of oxalic acid. Another important group of inorganic depressants for scheelite-containing ores are polyphosphates derived from carbohydrates, such as starch, dextrin, carboxymethyl cellulose.

This paper presents the findings of a study of the effects of high-power electromagnetic pulses (HPEMP) parameters on the structural and chemical state of the surfaces of calcite, fluorite, and scheelite and their sorption and flotation properties.

2. Materials and methods

In this study, natural calciferous minerals were used – calcite (CaCO$_3$ > 98%), scheelite (CaWO$_4$ > 98%), and fluorite (CaF$_2$ > 98%), size fraction - 80 + 50 µm. Distilled water (MeS/H$_2$O = 10/1) was added to the samples before they were treated with high-power electromagnetic pulses (HPEMP–treatment).

HPEMP–treatment of the samples was performed on a lab scale in air by a series of pulses between 5 to 10 ns long. The electrical field component was 30 kV with a pulse frequency of 100 Hz. The pulse energy was approximately 0.1 J and the integral treatment pulse number varied from 5 x 10$^2$ to 1.5 x 10$^4$ pulses. Pulse parameters were kept constant and the number of pulses was controlled by varying the treatment duration between 5 s and 150 s.

The microhardness of minerals before and after electromagnetic treatment was determined by the Vickers method (HV, MPa) according to GOST-2999-75 (ISO 6507-1: 2005) on polished sections using a PMT-3M microhardness meter (LOMO, Russia); the impact on indenter is 100 g, loading time 10 - 15 s.

To identify and study the acid - base properties of the mineral surfaces, the Hammett indicator adsorption method from aqueous media was adopted [10].

The adsorption of oleate was determined using UV spectroscopy according to the residual concentration of oleate ion in the pulp filtrate after contact of the mineral sample with the reagent solution (SU 1322130). Sorption of sodium silicate was determined by the residual concentration of silicic (C$_3$H$_4$O) in the pulp filtrate by the method (PND 14.1.: 2.215 - 06), based on the interaction of silicic acid with ammonium molybdate in an acidic medium.

Flotation test conditions: monomineral sample fraction (CaCO$_3$, CaF$_2$, CaWO$_4$) sized - 80 + 50 µm, 1.00 g; pH = 10.0; contact time of the mineral with water (S/L = 1:6) 1 min, with sodium silicate (100 g/t) 3 min, sodium oleate (300 g/t) 3 min, flotation time - 1.5 min. Reagent treatment was defined to achieve the highest possible recovery in the flotation of untreated scheelite sample.

3. Results and discussion

Micromechanical studies have shown a monotonic decrease in microhardness (HV, MPa) of mineral samples with an increasing of the treatment times (10 - 150 s). The maximum relative drop in microhardness was determiner for treatment during 150 s (1.5 x 10$^4$ pulses) and it was about 66% for calcite (from 790 MPa to 265 MPa), for fluorite it is 53% (from 912 MPa to 410 MPa), for scheelite 49% (from 2087 MPa to 1017 MPa) (Figure 1).

Acid-base properties of the surface: As seen in Figure 2a, the surface of calcite (CaCO$_3$) in the native state is mainly composed of Broensted acidic sites (BAS) with pKa = 1.3 (55.4 x 10$^{-3}$ µmol/g); neutral sites (pKa = 7.3, n = 15.4 x 10$^{-3}$ µmol/g), base Lewis (pKa = - 4.4, n = 13.4 x 10$^{-3}$ µmol/g), and Broensted (pKa = 12.8, n = 16.6 x 10$^{-3}$ µmol/g) sites also constitute a major share. The number of sites with pKa = 4.1 and pKa = 6.4 is 0.5 x 10$^{-3}$ and 3.3 x 10$^{-3}$ µmol/g, respectively.

As a result of HPEMP–treatment of the duration $t_{treatment}$ ~ 10 - 30 s (N ~ 10$^3$ - 3 x 10$^3$ pulses), a
reduction was identified in Broensted base sites from $19.3 \times 10^{-3}$ to $2.8 \times 10^{-3}$ µmol/g, an increase in the concentration of a protonic electron donor Lewis sites with pKa = -4.4 by ~1.2 times (from $13.4 \times 10^{-3}$ µmol/g to $16.7 \times 10^{-3}$ µmol/g) and proton donor Broensted sites with pKa = 1.3 and pKa = 4.1 by ~1.2 and ~5.2 - 8.6 times, respectively.

Increasing the treatment duration to $t_{\text{treatment}} \geq 50$ s ($N \geq 5 \times 10^3$) resulted in a consistent increase in the number of Broensted base sites with pKa = 12.8 by a factor of 10 - 19.5 (surface hydroxylation) and a decrease in the number of Lewis base (pKa = -4.4) and Broensted acid sites with pKa = 1.3 and 4.1 (Fig. 2b). Concentration of neutral sites (pKa = 7.3 and pKa = 6.4) did not change during the treatment.

Analysis of the calcite data reveals two key findings. The first one is the antechat change in the concentration (µmol/g) of sites with pKa = 12.8 and pKa = 1.3 depending on the duration of the pulse treatment, which may be interpreted as the conversion of Broensted base and acid sites indicating a change in the donor-acceptor equilibrium and stronger acceptor properties of the surface at $t_{\text{treatment}} \leq 30$ s ($3 \times 10^3$ pulses), followed by weakening with increasing treatment time. The second one is that the increase in the number of sites with pKa = 4.1 and pKa = -4.4 may be the...
result of accumulation and annihilation of structural defects.

Distribution of the adsorption sites on the original sample surface of fluorite (CaF$_2$) is shown in Figure 3a. The original mineral surface is dominated by Brønsted (pKa = 12.8, n = 195.6 x 10$^{-3}$ µmol/g) and Lewis (pKa = -4.4, n = 79.4 x 10$^{-3}$ µmol/g) base sites. Besides, Brønsted acid (pKa = 4.1, n = 48.2 x 10$^{-3}$ µmol/g; pKa = 1.3, n = 21.9 x 10$^{-3}$ µmol/g), neutral, and base (pKa = 7.3, n = 0.79 x 10$^{-3}$ µmol/g, pKa = 8.8, n = 0.87 x 10$^{-3}$) sites have been identified.

As can be seen in Figure 3, treatment of the mineral for t$_{treatment}$ ~ 10 s (N ~ 10$^3$) reduced the number of sites with pKa = -4.4 by more than an order of magnitude (from 83.21 x 10$^{-3}$ µmol/g to 8.19 x 10$^{-3}$ µmol/g) and increased the number of sites with pKa = 1.3 (from 21.9 x 10$^{-3}$ µmol/g to 118.59 x 10$^{-3}$ µmol/g) and 4.1 (from 48.2 x 10$^{-3}$ µmol/g to 58.2 µmol/g), i.e. by ~ 5.5 and 1.2 times, respectively.

Further increase in the duration of pulse treatment to t$_{treatment}$ ~ 30 s had the opposite effect, resulting in a significant increase in the sites with pKa = -4.4 and a decrease in the sites with pKa = 1.3; after the treatment, the concentration of the sites with pKa = -4.4 was nearly three times that in the original sample surface.

HPEMP--treatment for 50 seconds (5 x 10$^3$ pulses) resulted in a decrease by a factor of at least 11 in the concentration of base sites with pKa = -4.4. Importantly, the increase in Brønsted acid sites with pKa = 1.3 was not proportional (from 87.57 x 10$^{-3}$ µmol/g to 96.10 x 10$^{-3}$ µmol/g); sites with pKa = 4.1, to the contrary, decreased by a factor of 1.2 and the concentration of sites with pKa = 7.3, 8.8, and 12.8 did not change over the entire examined range of HPEMP values.

Based on the data from the fluorite samples, one can conclude that HPEMP--treatment over 10 s (10$^3$ pulses) results in a decrease of Lewis basicity with the conversion of Lewis base sites (pKa = -4.4) to Brønsted acid sites (pKa = 4.1 and 1.3), which indicates stronger electron-donor properties of the surface. At t$_{treatment}$ ~ 30 s, an explosive growth in Lewis base sites combined with a parallel decrease in Brønsted acidity were observed, which apparently is a result of both defect formation and deprotonation and / or dehydroxylation of the sites with pKa = 4.1 and 1.3.

**Figure 3.** The distribution of the adsorption centers throughout the pKa values for the initial fluorite sample (a); the dependence of the active centers with pKa = -4.4; 1.3; 4.1 on the number of pulses (b)

Considering the ambiguity of the accumulation mechanism of Lewis base sites (deprotonation indicates stronger acceptor properties, while dehydroxylation indicates the opposite), the effects of this treatment stage on the donor and acceptor properties cannot be conclusively identified.
However, increasing the treatment time to 50 seconds caused a sharp decrease in the number of Lewis base sites with a minor share converting to proton donor Bronsted sites; the latter indicates some increase in the donor ability of the mineral surface.

The surface of the original sample of scheelite CaWO₄ is mainly composed of proton donor Bronsted sites with pKa = 1.3 (n = 45.72 x 10⁻³ µmol/g), with 13.16 x 10⁻³ µmol/g and 19.82 x 10⁻³ µmol/g sites with pKa = 4.1 and 6.4, respectively. The concentration of Lewis (pKa = - 4.4) and Bronsted base sites is not that high: 3.33 x 10⁻³ µmol/g and 5.52 x 10⁻³ µmol/g, respectively (Fig. 4a).

Using the indicator method, it was demonstrated that the concentration of acid sites with pKa = 1.3, 6.4, 7.3 and pKa = 8.8 did not change as a result of the HPEMP–treatment. At the same time, the number of Bronsted base sites with pKa = 12.8 increased by a factor of 4.8 - 6 (from 5.5 x 10⁻³ µmol/g in the initial surface to 33.11 x 10⁻³ µmol/g as a result of the HPEMP–treatment of tₑ ~ 150 s). However, at tₑ ≤ 100 s, there is a decrease in the number of proton donor Bronsted sites with pKa = 4.1 (Figure 4b), resulting in an increased number of Lewis base sites and the concentration of proton donor Bronsted sites falling from 13.16 x 10⁻³ µmol/g in the original sample to 6.07 x 10⁻³ µmol/g in the sample after tₑ treatment ~ 100 s.

In other words, in scheelite, the main changes in the acid and base surface properties are associated with the increase in the number of Bronsted alkaline sites.

Considering the fact that the concentration of acid and neutral (pKa = 1.3, 6.4, 7.3, 8.8) sites does not change, the increase in Bronsted basicity may be logically explained by the stronger acceptor properties of the surface - activation of sites with pKa ≥ 14 accompanied by dissociative adsorption of water leading to growth in the number of sites with pKa = 12.8. The conversion of sites with pKa = - 4.4 to sites with pKa = 4.1 indicates weaker donor properties of the surface.

![Figure 4. The distribution of the adsorption centers throughout the pKa values for the initial scheelite sample (a); the dependence of the active centers with pKa = - 4.4; 1.3; 4.1 on the number of pulses (b)](image-url)

The sorption activity of calcium minerals towards oleate was investigated by UV-spectroscopy based on the residual concentration of oleic acid in the pulp after the contact of the mineral with the reagent solution. The obtained dates are presented in Figure 5, as it can be seen the preliminary HPEMP–treatment activated the sorption of the collector on the surface of calciferous minerals.

For scheelite samples HPEMP–treatment results to the increasing of the adsorbed oleate quantity (µg/g) by a factor of 1.2 - 1.3 times: from
139 µg/g (the mineral in the initial state) to 173 - 185 µg/g (the sample after the HPEMP–treatment, \( N_{\text{pulses}} = 3 \times 10^3 - 10^4 \)). For fluorite and calcite, the increasing of the adsorbed reagent quantity is 10 - 17%: the reagent adsorption for fluorite increased from 209 µg/g (initial state) to 238 - 244 µg/g (\( N_{\text{pulses}} = 3 \times 10^3 - 10^4 \)). For calcite samples, the rising of the sorption quantity was about 15%: from 229 µg/g (initial sample) to 260 - 269 µg/g for the mineral after HPEMP–treatment (\( N_{\text{pulses}} = 3 \times 10^3 - 10^4 \)).

The study of the HPEMP–treatment influence on adsorption of sodium silicate at the minerals surface demonstrates the decreasing in the quantity (mg/g) of the adsorbed reagent by 10 - 25% (Figure 5). In the case of scheelite and fluorite, the decreasing in the amount of adsorbed sodium silicate quantity was 10 - 14%: for scheelite - 3.2 mg/g (mineral in the initial state), 2.1 - 2.2 mg/g for samples after the HPEMP–treatment (\( N_{\text{pulses}} = 10^3 - 3 \times 10^3 \)). The influence is more effective for calcite: the decreasing in the quantity of adsorbed sodium silicate was \( \sim 20 - 25\% \) as compared to the sample in the initial state (6.7 mg/g) and 4.8 - 5.1 mg/g for the sample after the HPEMP–treatment (\( N_{\text{pulses}} = (3 - 5) \times 10^3 \)).

Flotation test results: Figure 6 illustrates the relationship between the floatability of monomineral fractions of calcite, fluorite, and scheelite and the duration of HPEMP–treatment.

It can be seen that, in the studied minerals, there is an increase in the froth yield as a result of a short \( (t_r \leq 50 \text{ s}) \) HPEMP–treatment - the largest increase in recovery achieved was 8% in calcite, 6% in fluorite, 12% in scheelite.

To confirm the obtained effect, using the above parameters of HPEMP, flotation experiments were carried out with a mixture of pure minerals weighing 3 g, with a ratio of calcite : fluorite : scheelite = 1 : 1 : 1.
The obtained results showed that preliminary electromagnetic treatment (HPEMP–treatment) of the artificial mixture allows to increase recovery of tungsten in concentrate. It was experimentally determined that with the consumption of sodium oleate 200 g/t and sodium silicate 100 g/t, the rising in the W recovery is 6.1 - 7.2% : 44.4% (0 imp is the reference sample), 51.5% (3000 pulses) and 50.6% (5000 pulses), the content of W in the concentrate, at the same time, increases by 1.9 - 2.5%. An increase in the consumption of sodium oleate to 250 g/t eliminates the effect of electromagnetic pulsed exposure: W extraction increases by 3.0 - 3.3% from 95.5% (0 imp) to 98.5 - 98.8% (3000 imp and 5000 imps, respectively), and the quality (W content) of the concentrate is improved by 1.5 - 2.0%.

As applied to selectivity increasing of the flotation separation of calcite from fluorite under monomineral flotation conditions (consumption of sodium oleate - 100 g/t, aluminum sulfate and sodium silicate - 500 g/t) the possibility of the minerals selectivity separation was determined because of fluorite activation on 12 - 14% at 10 x 10^{4} pulses.

Thus, it can be concluded that as a result of the research conducted, the principal possibility of using pulsed energy effects to increase the efficiency of the flotation process of calcium-containing minerals has been experimentally substantiated.

4. Conclusion

1. Using the Hammett color indicator adsorption method, following key evolution mechanisms were identified of the acid-base sites in calcium mineral surfaces exposed to high-voltage nanosecond pulses: stronger electron-donor properties of fluorite surfaces and stronger acceptor properties of scheelite surfaces as a result of HPEMP–treatment; mutual conversion at $t_{\text{orb}} \leq 30$ s of Broensted base and acid sites in calcite surfaces resulting in weaker donor properties and stronger acceptor properties of the mineral surfaces, and then, at $t_{\text{orb}} > 30$ s, recovery (strengthening) of the donor properties of the surfaces.

2. It was determined that HPEMP–treatment of calciferous minerals causes the increasing in their sorption activity towards a fatty acid collector (sodium oleate) and a decrease in the quantity (mg/g) of adsorbed depressor (sodium silicate). The increasing of adsorbed sodium oleate quantity for scheelite was 20 - 30% as compared with the initial sample, for calcite and fluorite, the increasing of the adsorbed reagent quantity was about 10 - 20%. The decreasing in the quantity of adsorbed sodium silicate (mg/g) was 10 - 15% for scheelite and fluorite and 20 - 25% for calcite.

3. In monomineral flotation of calcite, fluorite, and scheelite, the optimal parameters of the HPEMP–treatment were identified ($t_{\text{treatment}}$ ~ 10 - 50 s) and the reagent treatment was optimized for scheelite recovery, resulting in an increase of recovery of 8% in calcite, 6% in fluorite, and 10 - 12% in scheelite.

Note

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5. References

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UTICAJ ELEKTROMAGNETNIH PULSEVA VELIKE SNAZE NA STANJE POVRŠINE I TEHNOLOŠKE KARAKTERISTIKE MINERALA GVOŽĐEVITOG KREČNJAKA

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Izvod

U ovom radu je ispitivana promena strukturnih hemijskih osobina prirodnog fluorita, šelita i kalcita pod uticajem elektromagnetnih pulseva velike snage (HPEMP postupak) na osnovu adsorpcije kiselo-baznih indikatora. Utvrđeno je da tokom HPEMP postupka u periodu od 30 sekundi (3 x 10⁷ pulseva) dolazi do povećanja sposobnosti doniranja elektrona na površini fluorita, kao i do povećanja akceptorskih svojstava na površini kalcita i šelita.

Flotacijski testovi potvrđili su da postupak elektromagnetnim pulsevima velike snage omogućava poboljšanje flotabilnosti minerala gvožđevitog krečnjaka. Povećanje iskorišćenja šelita iznosilo je 10-12%, fluorita 5-6%, a iskorišćenje kalcita povećalo se za 7-8%.

Ključne reči: Kalcit; Fluorit; Šelit; Flotabilnost; Elektromagnetni pulsevi velike snage.