Quality of Anode. Overview of Problems and Some Methods of their Solution
Part 2. Improving the quality of the anode

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Abstract

Laboratory studies have shown that charging the components of the anode mass increases the rate of wetting and impregnation of dust by the pitch in 2-3 times, as evidenced by the shorter time interval during which the moistened coke was immersed in the pitch melt.

The algorithm for obtaining anode mass according to the proposed technological scheme is the following. The components of coke batch - bunches of different fractional composition from bunkers 2 by metering devices 3 and transport screws 5 are fed to preheater 6 where they are heated to a temperature of 200-220 °C. Further, the heated components of the coke batch are fed into the mixer 11. Coke dust before entering the mixer 11 by the dispenser 4 from the bunker 1 is fed into the dust preheater 7 where it is also heated to a temperature of 200-220 °C. Pitch, preheated to a temperature of 200-2400°C, is fed into the dispenser 8. Further heated coke dust and pitch enter high-voltage charging units 9, where they acquire electrical charges for 1-3 seconds, pitch-negative, and coke dust-positive. At the same time, the voltage at the electrodes of high-voltage charging stations is in the range of 24000-50000 V DC. Charged in this way, pitch and coke dust also enter mixer 11, where they are mixed for 3-4 minutes with each other and with coke batch. From the mixer 11, the finished anode mass is fed into the extruder 10 to form briquettes. In general, the process of anode preparation takes 4-5 minutes, which is 2 times lower than with "traditional" technology.

The presented technology allows essentially to raise productivity of the manufacture equipment of an anode mass at the expense of time reduction of mixing more than in 2 times. Reducing the mixing time of components reduces the specific electricity consumption for anode production by 20-25 kW / t, taking into account the energy consumption of the high-voltage charging unit. An increase in the degree of impregnation of coke dust with pitch reduces the anode fall-off under the conditions of the active electrolyzer and reduces the yield of coal foam by 5-7 kg / ton of Al. Reducing the yield of coal foam reduces the time of the electrolyzer in the depressurized state by an average of 0.2-0.25 hours during the day, which reduces the specific fluoride emissions by 0.1-0.2 kg / ton Al.

Keywords: aluminum electrolyzer, anode, carbon, coal foam, consumption, oxidation, protection, method.

INTRODUCTION

Dissolved CO_2 in the electrolyte, moving to the surface of the electrolyte, reacts with both carbon in reaction (1) and with dissolved aluminum and sodium. In the case of a rapid chemical reaction, the rate of the gas dissolution is determined by the step of its removal from the surface of the emerging bubble. This case was considered by V.G. Levich for a single bubble, suspended in a turbulent flow of a liquid, and he also proposed an equation for the diffusion flux [1]:

\[
C + CO_2 = 2 CO
\]

\[
j = D^{0.5} \left( \frac{\rho}{\rho_l} \right)^{0.5} \cdot \frac{a^{0.75}}{L^{0.25}} \cdot \frac{C_0 - C}{\nu}
\]

Where: \(D\) is the diffusion coefficient of the substance in the liquid; \(U\) is the average velocity of the fluid flow; \(a\) is the initial radius of the bubble; \(C_0\) is the saturation concentration; \(L\) is the characteristic size of the bubble; \(\nu\) is the kinematic viscosity coefficient/

With decreasing particle size of coal foam, their apparent density \(\rho_p\) increases from 1600 kg / m^3 (d≤10 μm) and becomes higher than the electrolyte density (2000 kg / m^3). The floating velocity \(V_{fp}\) of a spherical particle of diameter (d≤ 1 μm) in the electrolyte is determined by the Stokes equation:

\[
V_{fp} = \frac{4}{18} \cdot \frac{\rho_l - \rho_p}{\mu} \cdot a^2
\]

Where: \(\mu\) is the dynamic viscosity of the electrolyte, equal to 3 mPa·s

The bubble rise rate \(V_{br}\) (cm / s) depends on their radius \(r\) and is determined by three equations [2]:

\[
V_{br} = \frac{1}{9} \cdot \frac{\rho_l a^2}{\mu} \quad (r < 0.8 \text{ mm})
\]
\[ V_{br} = 1.35 \left( \frac{\sigma}{\rho l q} \right)^{0.5} \quad (0.8 < r < 2.5) \] (5)

\[ V_{br} = 1.18 \left( \frac{\sigma q}{\rho l} \right)^{0.5} \quad (r > 2.5 \text{mm}) \] (6)

The rate of bubble uplift, depending on their size, is 20 - 40 cm/s, which is an order of magnitude higher than the rate of emergence of carbon particles. Approximately at the same speed, carbon particles adhering to the bubble will also surface, because the main amount of foam is on the surface of the electrolyte, then, taking into account the low velocities of the emergence of particles with a diameter of \( \leq 100 \mu \text{m} \), it can be concluded that they rise due to flotation. The floating foam, the temperature of which is about 950 °C, reacts with the oxygen of the air. The burning of carbon in air is limited by mass transfer [3] and the burning rate of carbon is \( \sim 1 \text{ g/cm}^2 \cdot \text{h} \). The rate of oxygen delivery to the carbon particles is affected by the integrity of the cryolite-alumina crust, the wetting of the foam by electrolyte and the speed of the gas pump.

**ANODE GASES**

Anode gases are heavily diluted with air (up to 1 - 3% by volume of \( \text{CO}_2 \)), however, cryolite-alumina crust prevents air penetration into the space side-anode. This leads to a reduction in the burnout of the foam and the anode. Even in the presence of cracks and holes in the crust (for example, under the feeders), the intake of air under the crust is difficult due to the difference in pressure. The pressure under the crust is approximately 100 Pa higher than under the shelter shields of the electrolyzer.

In the case of air delivery under the crust, the influence on the access of oxygen to carbon is exerted by wetting of the foam with electrolyte. Depending on the angle of wetting, there are three possible options for the behavior of a carbon particle on the surface of the electrolyte (Fig. 1) [4]:

A) the wetting angle \( \theta \) is less, b) is equal and c) is greater than 90 °.

It is known that near the vertical surfaces of the anode immersed in the melt and under the crust \( \theta = 100-1100\), so the foam is poorly wetted by the electrolyte and part of its surface is accessible to the gas (Figure 1c). It should be noted that in cases a) and b) it is also possible to form a thin film of electrolyte on the surface of carbon particles [5].

Coal foam accumulates beneath the feed points of the bath with alumina, and can adversely affect the solubility of alumina, since instead of dispersing alumina, clumps are formed in the melt that are difficult to penetrate into the melt. The poor solubility of alumina is the reason for the formation of sediments on the bottom, the occurrence of anode effects, and an increase in the temperature of the electrolyte. The concentration of alumina in the bath plays a decisive role. Local depletion of the concentration may lead to a local anode effect.

The situation of increasing the resistance of the electrolyte due to the accumulating foam is equivalent to the bubble resistance in the bath, which is proportional to the volume of the gas-containing layer of the electrolyte. The increase in the resistance of the bath is proportional to the volume fraction of carbon in the melt. If the content of the foam in the electrolyte is 5% by weight, in order to compensate for its resistance, a 2 mm reduction in IPD (inter-pole distance) is necessary. By experimental measurements it was established that, with a highly-foamed electrolyte, the baths work with IPD, reduced by 4-6 mm. Many automatic process control systems (ACSs) in this case reduce the voltage on the bath, which is accompanied by a further decrease in IPD. Decrease in IPD affects the stability and productivity of the bath, which increases the formation of foam, forming a closed circle, shown in Fig. 2 [6].

![Figure 2: A vicious circle created by coal foam](image)

During the replacement of the anode, the foam floats to the surface of the melt. Because of intense radiation of heat by a black body, the electrolyte particles in the pocket crystallize, and the cold anode completes their freezing. At the same time, the thickness of the frozen electrolyte layer adhered to the anode bottom reaches 2 to 3 cm. This causes the effect of reducing IPD and reducing the melting rate of the adhered layer, due to the following reasons:

- the need for a higher temperature to melt the electrolyte containing coal particles;
- reduction of the contact area of the anode with the electrolyte;
- low overheating of the electrolyte under the anode, which suppresses melting of the adhered layer.

Thus, there is a high risk of further sticking of carbon particles and electrolyte and formation of a cone at the anode (Figure 3), causing a short circuit with cathode aluminum. More than 60% of the current of the entire anode flows through the cone, and the current loss can reach 3%, depending on the number of anodes and the duration of the short circuit. For example, for electrolytic cells equipped with 20 anodes, 10% of which are short-circuited with an average duration of 6 days, the permanent current loss is 1.5%.

With stable parameters of electrolysis, the formation of foam is a consequence of the low quality of the anode. In combination with technological disruptions, a high level of foam can lead to a crisis that reduces production performance for a long period.

**Effect of additives on the reactivity of the anode**

The effect of additions of Na$_2$CO$_3$, ZnS and Al$_2$O$_3$ on the reactivity and consumption of the anode in conditions of increasing the content of sulfur cokes and metallic impurities was considered in [7].

The reactivity of anodes with additions of Na$_2$CO$_3$, ZnS and Al$_2$O$_3$ was studied in the following way. The anode carbon materials used in this study were obtained by drilling out cores from full-scale fired anodes, or from small-scale laboratory blocks sealed with vibration and fired at 1265 °C. Samples drilled from industrial anodes were used to study the effect of Na$_2$CO$_3$ and ZnS additives, while samples from laboratory blocks were used to study the effect of the Al$_2$O$_3$ additive.

Na$_2$CO$_3$ was introduced into the sample by immersion in soda solutions of various concentrations for a certain time. After soaking, the samples were dried at 150 °C for 15 hours, cooled in a desiccator and weighed. This drying period was enough to get rid of all the water. The Na$_2$CO$_3$ content in the sample was determined by increasing its weight.

Due to the fact that ZnS is insoluble in water, its addition to the sample by the impregnation method is impossible. Therefore, for the tests, samples of anodes were prepared with the addition of specified amounts of finely divided Na$_2$CO$_3$ and ZnS.

The reactivity of carbon materials was determined by the usual method described in [8; 9; 10]. Samples were exposed to a continuous stream of CO or air for 2 to 4 hours at a constant temperature. Samples with a diameter of 20 mm and a height of 40 mm were suspended in an oven equipped with electronic scales, which allowed a continuous recording of the sample weight. The reaction capacity of the sample in CO$_2$ was studied at a temperature of 948 ± 3 °C, in air at 530 ± 2 °C.

The consumption of the anode material with the addition of Al$_2$O$_3$ was studied in a laboratory electrolytic cell equipped with an anode gas collection system (CO + CO$_2$). Anodes with a diameter of 42 mm for 3 hours were subjected to electrolysis at a constant current of 13 A and a temperature of 980 ± 2 °C in a cryolite containing 7% by weight of AlF$_3$ and 5% by weight of CaF$_2$, which was saturated with alumina and used as an electrolyte. The experimental conditions were strictly the same for all experiments.

The effect of the addition of Na$_2$CO$_3$ on the oxidation of anode samples in air and in CO$_2$ is shown in Figure 4.

![Figure 4: Effect of Na$_2$CO$_3$ on the reactivity of the anode in air and CO$_2$](image_url)

It can be seen from the graph that Na$_2$CO$_3$ catalyzes both reactions. While the effect of the additive on oxidation in air is small, in a CO$_2$ environment the catalytic effect is almost 7 times stronger.

In Figure 5, the reactivity data are expressed as a function of the total sodium concentration, by adding it with the introduced Na$_2$CO$_3$. This figure shows the data obtained in...
where the anode samples were made from raw materials with different sodium content. The data obtained in both cases show an extremely high sensitivity of the anode to oxidation in CO$_2$ in the presence of sodium, regardless of the content of other metallic impurities. It is believed that the large difference in the catalytic effect of Na$_2$CO$_3$ on the reactivity of the anode in CO$_2$ and in air is due to the different behavior of sodium carbonate at these temperatures during both reactions. In the works [11; 12] Such a behavior of Na$_2$CO$_3$ is due to its thermal dissociation.

Due to the high temperature, the following reaction is possible:

$$\text{Na}_2\text{CO}_3(s,l) + 2\text{C}(s) = 2\text{Na}(q) + 3\text{CO}(q)$$

(7)

The equilibrium constant for this reaction at 927 °C is ~ 4 · 10$^{-7}$, which can cause a significant increase in the vapor pressure of Na, with decreasing CO pressure. The separation of sodium vapor during the catalytic oxidation of graphite in CO$_2$ at 1000 °C was indeed noted in [17]. Since sodium vapor has a high reactivity, their formation can lead to the following reactions:

$$2\text{Na}(q) + \text{CO}_2(q) = \text{Na}_2\text{O}(s) + \text{CO}(q)$$

(8)

$$\text{Na}_2\text{O}(s) + \text{CO}_2(q) = \text{Na}_2\text{CO}_3(s,l)$$

(9)

Moreover, at the temperature under consideration, both reactions are thermodynamically possible.

In the case of the oxidation reaction of the anode in air, due to a decrease in temperature <600 °C, the sodium vapor pressure, in accordance with reaction (7), will be insignificant, since the equilibrium constant of reaction (7) is ~4 · 10$^{-33}$ at 527 °C, and at this temperature, the dissociation of Na$_2$CO$_3$, according to the reaction (4), will be very weak:

$$\text{Na}_2\text{CO}_3(s) = \text{Na}_2\text{O}(s) + \text{CO}_2(q) \quad \Delta G > 0$$

(10)

Nevertheless, according to [17], the dissociation of sodium carbonate in the presence of carbon and oxygen is enhanced as a result of the following successive reactions:

$$2 \text{Na}_2\text{O}(s) + \text{O}_2(q) = \text{Na}_2\text{O}_2(l)$$

(11)

$$2 \text{Na}_2\text{O}_2(l) + \text{C}(s) = 2 \text{Na}_2\text{O}(s) + \text{CO}_2(q)$$

(12)

Which can explain the weak catalytic effect.

The effect of the addition of ZnS with and without the use of Na$_2$CO$_3$ is shown in Figure 6.

![Figure 5: Reactivity of the anode in a current with CO$_2$ as a function of the sodium content](image)

**Figure 5:** Reactivity of the anode in a current with CO$_2$ as a function of the sodium content

Figure 6 shows that the addition of ZnS only to 0.40% by weight does not cause significant changes in reactivity in the current of CO$_2$, and the loss of reactivity of carbon is 10-11%, which can be attributed to the experimental error. Addition of 0.45% by weight of Na$_2$CO$_3$, as expected, increases the reactivity loss to 25%, in the absence of ZnS. An interesting feature of the experiment was the fact that the amount of Na$_2$CO$_3$ in the samples was maintained at a constant level of 0.45% by weight, while the content of the ZnS additive was increased from 0 to 0.47% by weight, which resulted in a decrease in the reactivity of the anode in CO$_2$ from 25 to 18%. In other words, the catalytic effect of Na$_2$CO$_3$ decreases with increasing S / Na ratio in the sample. The reason for this behavior is probably the presence of additional chemical equilibrium, which can be expressed as follows:
**ZnS(s) + 2Na(q) = Na₂S(s) + Zn(q)**  

(13)

As indicated earlier, the sodium vapor pressure is again formed by reaction (7) due to the dissociation of Na₂CO₃. However, in the second stage of the catalytic cycle, reaction (8) competes with reaction (13). From the thermodynamic point of view, the Gibbs energy of reaction (13) is ΔG = -136.8 kJ, and the reaction (7) ΔG = -67.8 kJ at 927°C. Therefore, it is reasonable to assume that at least a portion of the sodium vapor is converted to Na₂S, and thus the amount of free sodium for the catalytic cycle decreases.

The vapor of zinc, formed as a result of reaction (13), is usually oxidized to ZnO:

**Zn(q) + CO₂(q) = ZnO(s) + CO(q)**  

(14)

As indicated above, ZnO is not active with respect to oxidation of the anode in CO₂, and Na₂S is a very stable compound, hardly active as a catalyst for reactivity. Thus, reaction (13) is considered to be an obstacle to the observed catalytic effect produced by Na₂CO₃.

In the light of the foregoing, it is possible to use a similar interaction of sodium and sulfur in the compositions of industrial anodes. Petroleum coke with a high sulfur content may have a positive effect on reducing the reactivity of the anode in a current of CO₂ by suppressing the catalytic ability of sodium. This may cause disagreement with previously published data on the effect of sulfur on the oxidation of the anode in CO₂.

The effect of the addition of Al₂O₃ on the physical properties of the anode is shown in Table 1.

<table>
<thead>
<tr>
<th>Al₂O₃ content (wt%)</th>
<th>Apparent Density (g cm⁻³)</th>
<th>Cumulative Pore Volume (cm³ g⁻¹)</th>
<th>Total Electrical Conductivity (S cm⁻¹)</th>
<th>Thermal Conductivity (W m⁻¹ K⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1.53</td>
<td>0.156</td>
<td>23.6</td>
<td>62.2 ± 1.4</td>
</tr>
<tr>
<td></td>
<td>1.50</td>
<td>-</td>
<td>-</td>
<td>66.4 ± 1.1</td>
</tr>
<tr>
<td>1</td>
<td>1.52</td>
<td>0.157</td>
<td>23.9</td>
<td>3.9 ± 0.04</td>
</tr>
<tr>
<td>2</td>
<td>1.51</td>
<td>-</td>
<td>60.3 ± 2.0</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>1.55</td>
<td>0.153</td>
<td>23.7</td>
<td>69.8 ± 0.6</td>
</tr>
<tr>
<td>4</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>3.8 ± 0.04</td>
</tr>
<tr>
<td>5</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

The apparent density and porosity of the anodes with the addition of Al₂O₃ remain unchanged, as is the pore size distribution, as illustrated in Figure 7 for "pure" anode samples and samples with an additive of 5% by weight of Al₂O₃.

**Figure 7:** Typical pore spectra of anode samples without the addition of Al₂O₃ and with the addition of 5% by weight of Al₂O₃

The correlation between the resistivity of the anode and the content of Al₂O₃ is not observed, and the thermal conductivity decreases almost in proportion to the increase in the Al₂O₃ addition.

Reactivity of anode samples with an additive of 5% by weight of Al₂O₃ is shown in Figure 8.

**Figure 8:** Effect of Al₂O₃ on the reactivity of the anode in air and CO₂

While the reactivity in air linearly decreases with increasing Al₂O₃ content, the reactivity in the current of CO₂ remains at the same level, at any content of aluminum oxide. This different behavior of the anode samples in air and in CO₂ is believed to be due to the temperature difference at which these reactions were studied.
From the technological point of view, the reduction in oxidizability in air is very useful, since the consumption of the anode decreases. Another useful aspect is a decrease in thermal conductivity with the addition of Al$_2$O$_3$, which will reduce the temperature of the anode in the industrial cell. According to [13], a decrease in thermal conductivity of 0.1 W·K$^{-1}$ reduces the anode surface temperature by 2.75 °C. Even such a small decrease in temperature in the upper part of the anode will lead to significant savings in the carbon material, since its reactivity in air depends strongly on temperature. This combined effect makes the addition of Al$_2$O$_3$ to industrial anodes attractive.

The effect of the addition of Al$_2$O$_3$ on the anode consumption

The effect of the addition of Al$_2$O$_3$ on the anode consumption in the laboratory cell is shown in Figure 9.

![Figure 9: Effect of Al$_2$O$_3$ on the anode consumption after electrolysis at 980 °C and 13 A, for 3 hours. The anode consumption is determined by the loss of its weight minus the Al$_2$O$_3$ dissolved in the melt.](image)

The theoretical anode consumption corresponds to its 100% oxidation in CO$_2$. It is seen that with an increase in the Al$_2$O$_3$ content, the anode is consumed at a faster rate. Nevertheless, some of these losses are caused by the dissolution of Al$_2$O$_3$ in the electrolyte.

Anode weight loss correction for the corresponding Al$_2$O$_3$ content gives a net carbon consumption, which also increases with the addition of alumina. These data are presented in Figure 10. Oxidability of the anode does not cause an increase in the anode flow rate and this agrees with the results of the tests for the determination of the reactivity in the CO$_2$ current.

To a higher consumption of carbon material leads the formation and removal of coal foam. In accordance with Figure 10, the most optimal is the addition of Al$_2$O$_3$ in an amount of 1.9% by weight.

Summarizing the results obtained, the authors of [14] draw the following conclusions:

- Na$_2$CO$_3$ acts as a strong catalyst for reactivity, while its effect on the rate of carbon oxidation in air is very modest;
- ZnS is not active for reactivity, nevertheless it has the ability to significantly reduce the catalytic ability of sodium, especially in the presence of Na$_2$CO$_3$;
- Al$_2$O$_3$ does not affect the reactivity in CO$_2$, but reduces the reactivity in air.

It was established in [15] that V and Ni are catalysts for oxidation of the anode in air. The same is true for Na and Fe if we consider the reactivity of the anode in CO$_2$. However, it is often difficult to separate the effect of different elements due to the joint presence of other impurities. An example of this is the presence of sulfur, about which contradictory results were reported in the work.

Additives AlF$_3$, B$_2$O$_3$, SiO$_2$ and phosphorus-containing compounds were tested as inhibitors of anode oxidation [9]. However, only a few published data about the effect of Al$_2$O$_3$ on the reactivity and anode carbon consumption. It was reported in [16] that Al2O3 catalyses the oxidation of graphite in air. A significant increase in the consumption of Soderberg test anodes containing Al$_2$O$_3$ as an additive was reported in [17]. In the works [18; 19] as an alternative to the traditional, composite anodes containing from 20 to 85% by weight of Al$_2$O$_3$ were proposed, while in [13] there is a minimum...
oxidation of such an anode by air, apparently due to a
decrease in the fraction of carbon in it.

Proposed measures to reduce the consumption of pure
carbon and the yield of coal foam.

According to [20], the surface area available for reaction (7)
depends on the properties of the pitch and the depth of its
penetration into the pores of the coke-filler. When coke is
impregnated, the maximum capillary rise height $h_{\text{max}}$ is
determined by the surface tension ($\sigma$) of the liquid phase, the
wetting contact angle ($\cos \theta$) and the density $\rho_l$ and can be
determined from the expression:

$$h_{\text{max}} = \frac{2\sigma \cos \theta}{\gamma\rho_l}$$

According to [21], the Maximum lifting height of the pitch
along the capillaries depends on its softening temperature and
reaches 7 mm. The most rapid lifting height of the pitch in the
capillaries varies in the temperature range 150 - 175 ° C. A
further increase in temperature to 200 ° C does not lead to a
noticeable change in the lifting height of the pitches in the
capillaries.

Increase lifting height of the pitch in coke capillaries, and thus
reduce the consumption of the binder is proposed through the
use of a sound capillary effect, when ultrasound causes an
abnormally deep penetration of liquid into capillaries and
narrow slits. In this case, the height of the lifting and the depth
of penetration significantly exceed the corresponding values,
caused only by the surface tension of the liquid pitch [22].
The use of a sound-capillary effect will ensure an
improvement in the quality of the anode mass with a lower
pitch consumption, which will have a favorable effect on the
environmental performance of both the anode production and
aluminum production in general.

Another way to increase the degree of impregnation by pitch
is to impart a multi-polar electrical charge of coke dust and
pitch using a high-voltage DC charging unit that provides a
voltage on the electrodes from 24 to 50 thousand volts. In this
case, the pitch is given a negative charge, the coke dust is
positive [23].

Laboratory studies have shown that charging the components
of the anode mass increases the rate of wetting and
impregnation of dust by the pitch in 2-3 times, as evidenced
by the shorter time interval during which the moistened coke
was immersed in the pitch melt (Figures 11-16) [24].
The technology of charging components of pitch and coke with multi-polar charges can be easily adapted to the existing technological process of obtaining anode mass. In this case, a high-voltage dust charging installation is located between the coke dust preheater and the mixer, pitch - between the pitch meter and the mixer (Figure 17).

The algorithm for obtaining anode mass according to the proposed technological scheme is the following. The components of coke batch - bunches of different fractional composition from bunkers 2 by metering devices 3 and transport screws 5 are fed to preheater 6 where they are heated to a temperature of 200-220 ° C. Further, the heated components of the coke batch are fed into the mixer 11. Coke dust before entering the mixer 11 by the dispenser 4 from the bunker 1 is fed into the dust preheater 9 where it is also heated to a temperature of 200-220 ° C. Pitch, preheated to a temperature of 200-2400C, is fed into the dispenser 8. Further heated coke dust and pitch enter high-voltage charging units 7, where they acquire electrical charges for 1-3 seconds, pitch-negative, and coke dust-positive. At the same time, the voltage at the electrodes of high-voltage charging stations is in the range of 24000-50000 V DC. Charged in this way, pitch and coke dust also enter mixer 11, where they are mixed for 3-4 minutes with each other and with coke batch. From the mixer 11, the finished anode mass is fed into the extruder 10 to form briquettes. In general, the process of anode preparation takes 4-5 minutes, which is 2 times lower than with "traditional" technology [25-33].

The presented technology allows essentially to raise productivity of the manufacture equipment of an anode mass at the expense of time reduction of mixing more than in 2...
times. Reducing the mixing time of components reduces the specific electricity consumption for anode production by 20-25 kW / t, taking into account the energy consumption of the high-voltage charging unit. An increase in the degree of impregnation of coke dust with pitch reduces the anode fall-off under the conditions of the active electrolyzer and reduces the yield of coal foam by 5-7 kg / ton of Al. Reducing the yield of coal foam reduces the time of the electrolyzer in the depressurized state by an average of 0.2-0.25 hours during the day, which reduces the specific fluoride emissions by 0.1-0.2 kg / ton Al.

According to [25], oxidation of the anode occurs over all its surfaces, as shown in Figure 18.

![Figure 18: Dimensions and contour of the anode and anode cinder](image)

To reduce the oxidation of the side surface of the anode is proposed by means of a device on its side, facing the flange sheet of the cathode device of the electrolyzer of the notch-shelf, as shown in Figure 19 [26].

![Figure 19: Annealed anode with a notch on the side surface](image)

ACKNOWLEDGEMENT

This paper has been prepared with financial support from the Ministry for Education and Science of Russian Federation (Order No.218 adopted on the 9th of April 2010 by the Government of the Russian Federation.) within the framework of the project 02.G25.31.0181 "Development of heavy duty energy efficient technology for aluminum RA-550" in the framework of realization of complex projects in high-tech production, approved by the RF Government decree № 218 from April 9, 2010.

REFERENCES


