INFLUENCE OF COKE QUenchING METHOD ON ITS REACTIVITY

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Abstract. The results of investigation for the influence of the coke quenching method on the coke reactivity were shown. Using thermogravimetry and scanning electron microscopy significant differences of the porous structure and peculiarities of the structure of coke carbon, which forms the pores walls in both wet and dry quenching coke, were discovered. It was established that higher reactivity of wet quenching coke is due to its higher porosity (60% higher than that of dry quenching coke).

Keywords: coke, structure, wet quenching, dry quenching, TGA, SEM, pores.

1. Introduction

Reactive power and postreactive strength became one of the main quality measures of blast furnace coke quality in the world practice, as it reflects its behavior in blast furnace process most effectively, especially with the use of pulverized coal injection method (PCI). Injecting up to 200 kg of coal per ton of crude iron increases the time of charge in the blast furnace almost twice [1]. It results in low reactive power.

Most frequently reactive power is measured by the method of ("Nippon Steel Corporation") – NSC, diagnosing reactive index (CRI) and coke postreactive strength (CSR) or “hot” strength [2]. In this regard, Western European and Northern American metallurgical plants impose high requirements to the coke quality according to the following indices: CSR – more than 60% and CRI – less than 30%.

Coke reactive index level and coke postreactive strength is known to be mainly provided by the properties of carbon raw material source. Nevertheless, some other factors, including out-of-furnace processing [3] can influence CRI and CSR.

One of the technological operations in coke processing is coke quenching done in wet or dry way. It was found out that stamp blends coke which was dry quenched in dry quenching chambers of coke plant with other equal conditions (charge content, preparation and coking technological regime) is characterized by lower rate of CRI and higher rate of CSR than wet quenched coke [4].

In our work cokes from the usual coal blends are studied.

2. Experimental

For the study of the factors determining the above-mentioned difference, wet quenched and dry quenched coke was produced at the Public Company “Avdeevskiy Coking Plant” from the charge of the following content (Table 1).

This blend had the following characteristics: $A_d = 8.9$, $S_{t}^{d} = 1.34$, $V_{daf}^{d} = 32.7\%$, $y = 15$ mm.

Dry and wet quenched coke from this blend was tested for quality measures, strength index, actual and apparent density and reactive capacity according to all-Union state standard 10089-89 and Ukrainian national standardization system 4703:2006 (ISO 18894:2006).

Thermogravimetric analysis¹ was performed with the use of derivatograph “Paulic-Paulic-Erdey” mark Q-1500 with computer control and logging system. Temperature and weight loss was registered with the computer once a second. The software allows printing derivatograms or exporting them to hard disc drive in .png file format.

Dynamic thermogravimetric analysis was performed in inert medium (with Ar purity 99.9%). Oxidation thermogravimetry was performed in air medium. The research was done in the temperature range of 293–1273 K and with the heating rate of 10 K/min. Accuracy of temperature regulation is ±1 K. Sample weight is 350–400 mg. Reference substance is incinerated aluminum oxide. Gas purging speed is 0.5 l/min. Air purging is done with microprocessor through U-tube with CaCl₂.

Statistical analysis of experimental data was performed with the help of ORIGIN 8.0.

¹Thermogravimetric analysis was performed by senior researcher Tolmachev N.V.
### Table 1

**Rank and component composition of coal blend**

<table>
<thead>
<tr>
<th>Name of the supplier</th>
<th>Coal rank</th>
<th>Content in charge</th>
<th>Total for rank</th>
</tr>
</thead>
<tbody>
<tr>
<td>Processing plant «Samsonovskaya» + Processing plant «Selidovskaya»</td>
<td>Gas coals</td>
<td>14</td>
<td>36</td>
</tr>
<tr>
<td>Processing plant «Zarechnaya»</td>
<td>16</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Promugolservis</td>
<td>6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Processing plant «Mikhailovskaya»</td>
<td>Fat coals</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>Processing plant «Dzerzhinskaya» + Processing plant «Mikhailovskaya»</td>
<td>8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Processing plant «Samsonovskaya» + Processing plant «Dzerzhinskaya»</td>
<td>9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Processing plant «Duvanskaya» + Processing plant «Kievskaya»</td>
<td>9</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Kolosnikovskaya</td>
<td>Close-burning coals</td>
<td>16</td>
<td></td>
</tr>
<tr>
<td>Open-pit mine «Neryungrinskaya» (No.1)</td>
<td>5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Processing plant «Sholokhovskaya»</td>
<td>Semi-lean coals</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>Processing plant «Uzlovskaya»</td>
<td>4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Table 2

**Proximate analysis, coke strength and density indices**

<table>
<thead>
<tr>
<th>Coke</th>
<th>Technical analysis, %</th>
<th>Strength index</th>
<th>Structural strength according to Gryaznov, %</th>
<th>Abrasive strength according to Ginsburg, mg</th>
<th>Actual density, g/cm³</th>
<th>Apparent density, g/cm³</th>
<th>Porosity, %</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$A'$</td>
<td>$S_i'$</td>
<td>$V_{ad}$</td>
<td>$M_{25}$</td>
<td>$M_{10}$</td>
<td>$S_r$</td>
<td>$S_a$</td>
</tr>
<tr>
<td>No.1</td>
<td>13</td>
<td>1.27</td>
<td>0.9</td>
<td>86.2</td>
<td>8.6</td>
<td>87</td>
<td>102</td>
</tr>
<tr>
<td>No.2</td>
<td>12.9</td>
<td>1.20</td>
<td>0.1</td>
<td>87.6</td>
<td>7.9</td>
<td>87</td>
<td>104</td>
</tr>
</tbody>
</table>

### Table 3

**Reactive index of wet quenching and dry quenching coke**

<table>
<thead>
<tr>
<th>Coke</th>
<th>Coke reactive index</th>
<th>According to all-Union state standard 10089-89, cm³/g·s</th>
<th>According to Ukrainian national standardization system 4703:2006</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CRI %</td>
<td>CSR, %</td>
<td></td>
</tr>
<tr>
<td>No.1 (wet quenching)</td>
<td>0.28</td>
<td>54.6</td>
<td>25.0</td>
</tr>
<tr>
<td>No.2 (dry quenching)</td>
<td>0.24</td>
<td>50.9</td>
<td>28.2</td>
</tr>
</tbody>
</table>

### 3. Results and Discussions

Results of coke No.1 (wet quenched) and coke No.2 (dry quenched) analysis are shown in Tables 2 and 3.

Less devolatization from dry quenched coke can be explained by its long holding in dry quenching coke plant prechamber at the high temperature, which was close to the one at which coke was poked out from the coking chamber.

Coking proved the known fact of lower mechanic strength ($M_{25}$ and $M_{10}$) of wet quenched coke in comparison with dry quenched coke.

Less mechanical strength ($M_{25}$) for wet quenched coke can be explained by heat shock at its quenching with water, leading to additional crack formation, which takes place in a Micum-drum. Besides, it can be the result of higher porosity and probably of the Rebinder effect, which, as is widely known, results in solid body strength decrease due to the presence of absorbed water in pores.

Other indices of coke properties, given in the Table 2, do not show any significant differences between wet quenched and dry quenched coke.

At the same time reactive index of wet quenched and dry quenched coke from the same coal blend differ a
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lot according to all-Union state standard 10089-89 and CRI index (Table 3). It can happen because of the difference of porous structure or because of the difference in chemical activity of the carbon, forming pores walls.

However, as it can be seen in Table 2, porosity, estimated as difference quotient of actual and apparent actual density for both types of coke has almost the same rate. Probably, ethanol molecule used for picnometric analysis does not penetrate all pores formed in the process of wet coke quenching and makes it necessary to measure its porosity in some other way, for example, by the water absorbed in pores. Thus, according to the data [5], estimated value of ethanol and water absorption platform is 23.1 and 10.5 nm², respectively.

Therefore, matching data from Table 2 and 3 allows to suppose that the data received from measuring actual and apparent density do not reflect real picture of wet and dry quenched coke porosity and other more informative methods are required.

Thermogravimetry experiments turned out to be more informative with coke samples being heated. Thermogravimetry allowed defining the following indices that characterize structural and thermochemical properties of cokes studied. These are pores capacity with respect to moisture $W$ (mg $H_2O$/g coke) and coke ignition initiation temperature (according to oxidation thermogravimetry).

For thermoplastic records of coke samples, see Fig. 1. Coke porous structure was evaluated by moisture volume, kept by coke, i.e. assuming that coke pores volume is equal to the volume of absorbed moisture.

For porosity tests coke samples dried to constant weight at 423 K were prepared. After cooling down to room temperature, the samples were put into an exicator with a vessel filled with water on the bottom. Coke samples were held in moist environment at room temperature for 15 h. After that part of the samples were taken (after thorough mixing) for tests on derivatograph.

Based on the derivatogramme processing results the following data were received (Table 4).

As it is seen from the data in Table 4, pores volume of wet quenching coke (coke No.1) is by 66 % higher than that of dry quenched coke. Coke No.1 has lower burning point that equals to 775 K and is by 26 degrees lower than that of dry quenched coke (coke No.2).

![Fig. 1. Thermoplastic records of wet quenching (above) and dry quenching (below) coke samples](image)

<table>
<thead>
<tr>
<th>Coke</th>
<th>Coke thermochemical properties</th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Dehydration temperature ($T_{d,120}$), K</td>
<td>Inflaming temperature ($T_i$), K</td>
<td>Pores volume according to water capacity ($W$), ml/g</td>
<td>Furnace loss speed ($V_l$), g·s⁻¹·10⁻⁵</td>
</tr>
<tr>
<td>№1 (wet quenched)</td>
<td>354</td>
<td>502</td>
<td>16.0</td>
<td>2.6</td>
</tr>
<tr>
<td>№2 (dry quenched)</td>
<td>358</td>
<td>528</td>
<td>10.6</td>
<td>1.8</td>
</tr>
</tbody>
</table>
For characterizing coke porous structure peculiarities and pores wall structure the method of scanning electron microscopy (SEM) was used. Scanning electron microscope Jeol JSM 840 gave pictures of coke cleavage surface with different level of magnification: 100x, 500x, 2500x, 5000x, and 10000x (Figs. 2 and 3).

It was established that wet quenched coke has the minimum pores size of 0.6 micron and the maximum of about 30 micron. Minimum wall thickness is 0.8 micron and maximum thickness is more than 10 microns.

Dry quenched coke has the minimum pores size of about 0.7 micron and the maximum of about 30 microns. The minimum pores wall thickness is 1.6 micron, which is twice as much as for wet quenched coke, and the maximum thickness is more than 10 microns.

![Fig. 2. Pores walls of wet quenching coke (SEM, 100x); clarified area is represented rightward with 500x, 2500x and 5000x magnification](image)

![Fig. 3. Pores walls of dry quenching coke (SEM, 100x); clarified area is represented rightward with 500x, 2500x and 5000x magnification](image)
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For quantitative assessment of coke porosity pores distribution according to size was performed using pictures with 100x magnification (Fig. 4).

Quantitative assessment of pores distribution in coke samples enabled to define that wet quenched coke has less 0–2 micron pores (3.0 %, Table 4a) in comparison with dry quenched coke. Weighted average size value for wet quenched coke is 7.43 microns and for dry quenched coke it is 6.57 microns. Pores wall structure of coke samples was studied with 2500x, 5000x and 10000x magnification using SEM.

In the pictures with 5000x magnification sizes of coke structural elements were measured and calculated. In wet quenching coke elongated supramolecular formations have an average size of 0.337 micron. Ridge thickness of such formation is 0.05–0.07 micron.

Ridge length of supramolecular formations in dry quenched coke structure of the same charge is on the average 0.67 micron. Ridge thickness is 0.5–0.8 micron.

Figs. 5 and 6 represent pores wall surface of wet quenched and dry quenched coke with 10000x magnification.

As seen in Figs. 5 and 6 10000x magnification shows significant difference in coke sample structure. It can be marked that wet quenched coke pores walls have more developed surface owing to small structural elements. Dry quenched coke pores walls are characterized by bigger structural elements composed of elongated formations.

4. Conclusions

Application of thermogravimetry and scanning electron microscopy methods allowed to discover considerable differences in porous structure and pores
walls forming substance structure peculiarities in wet and dry quenched coke, which could not be detected using the conventional methods.

Coke wet quenching leads to coke pores volume increase up to 60% and their average size increase up to 11.6% in comparison with dry quenched coke.

Minimum pores wall thickness of wet quenched coke is twice as little as that of dry quenched coke.

Lower reactive index of dry quenched coke (according to CRI and inflaming temperature) is resulting, in our opinion, from the fact that with dry coke quenching the most reactive part of coke structure is burned out while it remains in case of wet quenching and is additionally activated by formation of micro porous structure.

Chemical activity of wet quenched coke pores surface is much higher than that of dry quenched coke; which is confirmed by inflammation temperature and furnace loss speed data.

References


