Utilization trials of lignite solid byproducts of West Macedonia and Peloponnesus lignite fired power plants for the production of lightweight aggregates

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UTILIZATION TRIALS OF LIGNITE SOLID BYPRODUCTS OF WEST MACEDONIA AND PELOPONNESUS LIGNITE FIRED POWER PLANTS FOR THE PRODUCTION OF LIGHTWEIGHT AGGREGATES

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Abstract

The production of lightweight aggregates from the utilization of solid byproducts (fly ash (FA) and bottom ash (BA)), of the lignite fired power plants from West Macedonia (Kardia, Ptolemaida) and Peloponnesus (Megalopolis), has been investigated in this study. Sintering of FA and BA with the grate sinter band method was selected because it exploits the energy content of the raw materials for the increase of temperature and completion of the material’s sintering. These byproducts contain the necessary carbon content for temperature increase during the sintering process. Detailed XRD analyses show that Megalopolis’ FA contains high amounts of amorphous phase (25-54 wt%) while fly ashes from Kardia and Ptolemaida power units are more crystalline. Lime and anhydrite are the most abundant crystalline phases in Ptolemaida’s and Kardia’s fly ashes. The bottom ashes revealed high percentages of amorphous content (58-64 wt%) and anorthite and quartz as the major crystalline phases. Mixtures of different BA/FA ratios were prepared for sintering tests. The sintered mixtures from Kardia’s and Ptolemaida’s regions did not have sufficient mechanical properties for further treatment. The sintercake produced from Megalopolis’ ashes exhibited good mechanical properties and was selected for further study for use as light aggregate raw materials to the lightweight concrete production.

Key words: fly ash; bottom ash, use, grate sintering, mineralogical composition.

Περίληψη

Στην παρούσα εργασία μελετήθηκε η παραγωγή ελαφροαδρανών από τη χρησιμοποίηση επιμέμησης τύφρων (ΠΠ) και τύφρων πυθμένα (ΠΠΠ), που παράγονται από την καύση λιγνίτη στις περιοχές Καρδιάς, Πτολεμαίδας (Δ.Μακεδονία), Μεγαλόπολης (Πελοπόννησος). Επιλέχθηκε η μέθοδος πυροσυσσωμάτωσης με σχάρα, διότι εκμεταλλεύεται ως καύσιμο τον άκαυστο άνθρακα που περιέχουν οι τύφρες, για την αύξηση της θερμοκρασίας που απαιτείται κατά την πυροσυσσωμάτωση. Σύμφωνα με τις ορυκτολογικές αναλύσεις περιθλασιμετρίας ακτίνων-Χ, οι ΠΙΠ περιέχουν υψηλότερο ποσοστό υμέρης μόριος.
1. Introduction

Nowadays, the utilization of solid fuel combustion byproducts is rapidly increased all over the world, not only because of the lack of land disposal sites but for the global demand of new recycling raw materials, aiming to the combination of environmental and economic profits. The fly ash (FA) and bottom ash (BA) have been mainly found applications to the building construction, road construction or as additives in the cement or concrete (Nishigaki, 2000; Lin, 2006; Vegas et al., 2008; Anagnostopoulos et al., 2010). Many researchers have been aimed to the production of lightweight aggregates by sintering of several municipal solid wastes incineration (Bhattay and Reid, 1989; Cheeseman et al., 2005).

Lignite combustion in West Macedonia and Peloponnesus of Greece results in production of approximately 14Mt/year of FA and BA byproducts. Today, only 10-15% of FA produced in Greece is utilized as raw material mainly in cement production while the BA use is very restricted due to its high carbon content.

This study investigates the utilization of FA and BA from the lignite fired power plants from North Greece (Kardia, Ptolemaida - West Macedonia) and South Greece (Megalopolis-Peloponnesus) as raw materials in the production of lightweight aggregates, using the grate sinter band method. This method has been applied with a lot of advantages to the bottom ash utilization (Anagnostopoulos and Stivanakis, 2009). The later contains a high carbon content which is used as the main fuel for the temperature increase during the sintering process whilst its high humidity could be used as the main binder phase for the agglomeration.

2. Experimental Procedure and Analytical Techniques

The developed process for the production of lightweight aggregates from the solid lignite combustion by-products consists of two stages, pelletization and sintering, respectively (Anagnostopoulos and Stivanakis, 2009). The whole pilot-scale sintering installation at ELKEME is shown in Figure 1. Pre -designed ratios of the solid raw materials are first mixed into a mechanical mixer and then introduced into the pelletization disc with the addition of sprayed water, if necessary. The rotating disc is of 0.6m diameter and operates at a slope of 45° with a rotation speed of 40-50 rpm. The formed pellets are introduced into the sinterpot, which is of 0.16m surface. Five thermocouples are set up along the height of the sinterpot for the measuring of temperature inside the formed bed of pellets. A layer with charcoal of 3-4cm is placed at the top of the bed of pellets, in order to assist the ignition of the pellets. The ignition of the pellets occurs the same time with the start-up of the suction blower, connected at the bottom of the pot. The blower creates suction pressure inside the pot and causes movement of the combustion zone downwards. The process of sintering is terminated when the combustion zone reaches the bottom of the pot. The flue gases of sintering are passed through a filter-bag for the removal of the solid particles.
Chemical analyses of the studied samples (raw materials and final sintered products) was carried out in cooperation between UPATRAS (Dep. Chem. Engineering, METLAB)-ELKEME, using AAS and ICP, while C, S contents have been measured using a Carlo –Erba analyzer. XRD analyses, using a Bruker D8 Advance diffractometer with Ni filter and CuKα radiation, were also performed to identify the phases present in the samples, in the Laboratory of applied Mineralogy (MINTECHLAB), Department of Mineral Resources, Techn. University of Crete, Greece. In addition, quantitative analysis was performed by Rietveld Method using Rayflex Autoquan Software. Microstructures of samples have been studied in ELKEME using a FEI XL40 SFEG scanning electron microscope (SEM). The physical parameters of sintered products have been measured and evaluated according to ASTM 373-88, standards (ASTM C373-88, 2002) in the METLAB.

3. Results and Discussion

The raw materials BA, FA and ML (metallurgical lime) which are used for the sintering production of the final light aggregates materials are characterized below.

3.1. Raw Materials

3.1.1. Chemical Analyses

The %wt of major oxides, carbon and LOI in the raw materials are given in Table 1. Chemical analyses of BA revealed higher percentage of loss of ignition compared with this in FA samples, due to their higher carbon content. In addition, the bottom ashes are significantly poorer in CaO content against to FA samples. In BA samples, the SiO₂ content ranges from 37.46 (in BAP=BA Ptolemaidas) to 41.21wt% (in BAK=BA Kardias), while the Al₂O₃ ranges from 16.27 (in BAK) to 17.39 (in BAP=BA Ptolemaidas). Fly ashes from Kardia and Ptolemaida contain significantly higher amounts of calcium oxide compared with that of Megalopolis. The maximum silicon oxide and aluminium oxide occur in FAM (=FA Megalopolis), while the minimum are contained in FAP (FA Ptolemaidas) and FAK (FA Kardias) samples respectively.

![Figure 1 - Pilot-scale sintering installation.](http://epublishing.ekt.gr)
Table 1 - Chemical analyses (wt%) of major oxides, C and LOI in the raw materials.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Al₂O₃</th>
<th>CaO</th>
<th>Fe₂O₃</th>
<th>SiO₂</th>
<th>C</th>
<th>LOI</th>
</tr>
</thead>
<tbody>
<tr>
<td>BAK*</td>
<td>16.27</td>
<td>14.32</td>
<td>4.88</td>
<td>41.21</td>
<td>10.13</td>
<td>14.50</td>
</tr>
<tr>
<td>BAP</td>
<td>17.39</td>
<td>12.90</td>
<td>3.95</td>
<td>37.46</td>
<td>14.51</td>
<td>23.00</td>
</tr>
<tr>
<td>BAM</td>
<td>16.31</td>
<td>12.80</td>
<td>7.87</td>
<td>39.04</td>
<td>9.77</td>
<td>15.30</td>
</tr>
<tr>
<td>FAK</td>
<td>14.60</td>
<td>33.15</td>
<td>5.63</td>
<td>29.36</td>
<td>1.64</td>
<td>4.50</td>
</tr>
<tr>
<td>FAP</td>
<td>15.03</td>
<td>32.55</td>
<td>6.30</td>
<td>26.36</td>
<td>1.73</td>
<td>4.60</td>
</tr>
<tr>
<td>FAM</td>
<td>19.27</td>
<td>14.90</td>
<td>10.68</td>
<td>46.71</td>
<td>1.22</td>
<td>1.10</td>
</tr>
<tr>
<td>ML</td>
<td>88.37</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>11.03</td>
</tr>
</tbody>
</table>

*Abbreviations: BAK = BA Kardias; BAP = BA Ptolemaidas; BAM = BA Megalopolis; FAK = FA Kardias; FAP = FA Ptolemaidas; FAM = FA Megalopolis; ML: metallurgical lime

3.1.2. XRD Analyses

According to the XRD results, the FAM contains the higher amount of amorphous phase (~55%) while in FAK and FAP it ranges to lower percentages (~25 to 30% respectively). In FAK and FAP samples lime and anhydrite appear as the major crystalline phases. Moreover, anorthite, quartz and gehlenite have been detected in all FA samples.

In BA samples the amorphous phase ranges from ~57-64% while anorthite and quartz have been identified as the major crystalline phases. In the metallurgical lime (ML), which has been used as additive, approximately 24% of lime has been hydrated to portlandite.

3.2. Design of Different Raw Materials Mixtures

The ratios of the raw materials (BA, FA, ML) in the different designed mixtures prepared for sintering, are given in Table 2. Taking into account the chemical analyses of all BA samples, the mixtures of Table 2 contain the requisite carbon content used as fuel during the sintering process.

3.3. Characterization of Final Products (pellets)

3.3.1. Macroscopic Characterization

According to macroscopic observations of the sintered products those from Kardia’s region did not comprise sound sinter-cake. Formation of sinter-cake (sound bonding of the pellets) is considered to be a characteristic essential visual parameter reflecting the efficiency of the sintering process. Thus, it comprises a criterion for the product’s mechanical suitability for further treatment (Figure 2). Ptolemaida’s mixtures (Figure 2), although they obtained the base mechanical strength after sintering, autogeneous disintegration occurred after two-three weeks later. The solid sinter cake produced from Megalopolis’ ashes (Figure 2) was selected for further study for use as light aggregate raw material to the lightweight concrete production.

3.3.2. Sintering Behaviour of Pellets

A representative temperature-time diagram of sintering process for mixture MMEG is presented in Figure 3. Temperature T1 is recorded at the top of the bed of pellets and the temperature T5 at the bottom of the bed. The temperature T6 corresponds to the flue gases of sintering. The duration of the sintering process is approximately 50min. The temperature of sintering is between 1200 and 1250°C. The temperature increase rate is very fast, from 600°C/min until 1100°C/min.
Table 2: % of raw materials in the different prepared mixtures for sintering.

<table>
<thead>
<tr>
<th>Name of mixture</th>
<th>Origin of raw materials</th>
<th>BA (%)</th>
<th>FA (%)</th>
<th>ML (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMEG</td>
<td>Megalopolis</td>
<td>90</td>
<td>7</td>
<td>3</td>
</tr>
<tr>
<td>MP1</td>
<td>Ptolemaidas</td>
<td>55</td>
<td>42</td>
<td>3</td>
</tr>
<tr>
<td>MP2</td>
<td>Ptolemaidas</td>
<td>75</td>
<td>22</td>
<td>3</td>
</tr>
<tr>
<td>MK1</td>
<td>Kardias</td>
<td>55</td>
<td>42</td>
<td>3</td>
</tr>
<tr>
<td>MK2</td>
<td>Kardias</td>
<td>75</td>
<td>22</td>
<td>3</td>
</tr>
<tr>
<td>MK3</td>
<td>Kardias</td>
<td>90</td>
<td>7</td>
<td>3</td>
</tr>
</tbody>
</table>

*Abbreviations: M: mixture; MEG: Megalopolis; P: Ptolemaidas; K: Kardias

Figure 2 - Representative macroscopic images of sintered products.

Figure 3 – Representative temperature -time diagram of sintering process (MMEG).

3.3.3. Chemical Analyses and Physical Parameters

The %wt of major oxides in the sintered products are given in Table 3. As expected, MMEG mixture is the most poor in CaO and the richer in SiO2, reflecting the chemistry of the raw materials. The maximum iron oxide has been analysed in MMEG sample. Minor oxides of K2O (<2.16wt%), MgO (<2.89wt%), Na2O (<0.51wt%), P2O5 (<0.05wt%), TiO2 (<1.06wt%), Cr2O3 (<0.04wt%), CuO (<0.29wt%), MnO (<1.19wt%) and ZnO (<4.29wt%) have been also determined. The low percentage of loss of ignition (max 1.6% in MP2) revealed that the incombustible C

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content of raw materials has been utilized as fuel well enough during the sintering process, promoting simultaneously the porous microstructure.

The results of physical parameter’s measurements (apparent porosity, apparent specific gravity and water absorption) of products are displayed in Table 4. All samples except of MMEG and MK2, were broken down during the measurement (boiling). MMEG exhibited low apparent specific gravity (0.89g/cm$^3$) while the MK2 (1.43g/cm$^3$) higher than a commercial light aggregate one (1g/cm$^3$). Additionally, MMEG exhibited higher percentage of water absorption and apparent porosity compared to MK2.

Table 4 – Physical parameters of sintered products.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Apparent porosity (%)</th>
<th>Apparent specific gravity (g/cm$^3$)</th>
<th>Water absorption (%)</th>
<th>Dry weight before sintering (kg)</th>
<th>Weight after sintering (kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MMEG</td>
<td>59.7</td>
<td>0.89</td>
<td>67.5</td>
<td>33</td>
<td>19.6</td>
</tr>
<tr>
<td>MP1</td>
<td>NM*</td>
<td>NM</td>
<td>NM</td>
<td>30</td>
<td>19.4</td>
</tr>
<tr>
<td>MP2</td>
<td>NM</td>
<td>NM</td>
<td>NM</td>
<td>27.5</td>
<td>21.25</td>
</tr>
<tr>
<td>MP3</td>
<td>NM</td>
<td>NM</td>
<td>NM</td>
<td>25</td>
<td>18</td>
</tr>
<tr>
<td>MK1</td>
<td>NM</td>
<td>NM</td>
<td>NM</td>
<td>35</td>
<td>22</td>
</tr>
<tr>
<td>MK2</td>
<td>42.8</td>
<td>1.43</td>
<td>30.2</td>
<td>32.5</td>
<td>12.7</td>
</tr>
<tr>
<td>MK3</td>
<td>NM</td>
<td>NM</td>
<td>NM</td>
<td>32.5</td>
<td>13.5</td>
</tr>
</tbody>
</table>

*Abbreviation: NM: not efficient for measurement

3.3.4. XRD Analyses

The semiquantitative mineralogical compositions of the sintered products are presented in Table 5. The amorphous phase ranges from ~36 (MK1) to 44 wt% (MP3) while anorthite, gehlenite and diopside (in two of them) are the major detected crystalline phases. It has been previously proved that in such materials, the mechanical strength is increased further by the anorthite appearance than the gehlenite (Anagnostopoulos, 2010). Samples MMEG, MP2 and MP3 contain the higher anorthite percentages (~34-38%) and the lower gehlenite (~3-9%), whilst the maximum diopside (~15%) is presented in MMEG, thus expected to improve further its mechanical strength (Bethanis et. al, 2002). Moreover in MMEG most of the hematite has been reducted to the magnetite. Maximum amount of free lime has been detected in the MK3 sintered product. Its
presence is undesired, since its subsequent transformation to portlandite by air humidity is accompanied by volume expansion resulting in disintegration of the solid matrix, making it unsuitable for further utilization. The different ratios of phases in the final products could be attributed to the combination of: 1) different geochemical/mineralogical compositions of BA and FA materials (as well as their lignite raw materials derived from), (Sakorafa et. al., 1996; Filippidis et. al., 1996; Siavalas et. al., 2009), 2) the sintering process, and 3) the different raw material ratios in the prepared mixtures. Moreover the chemically complicated systems of sintered products are reflected to their mineralogical compositions consisting of stable primary phases (eg quartz,) and new formed ones (eg diopside, magnetite) after different reactions mechanisms (decompositions, solid state reactions, reduction reactions, melting), not all in equilibrium conditions during the rapid sintering process (Anagnostopoulos, 2009).

3.3.5. Microstructure by SEM

SEM observations of final products revealed that all of them are characterized by a heterogeneous microstructure. Representative backscattered images are given in Figure 4. The MMEG sample exhibits the most compact structure because of the advanced sintering (via solid state reactions or melting phase bonding). Pores forming from outgassing are very often due to C, S devolatilization, the structural destruction of Ca sulphates/ carbonates and iron oxide reduction. Pore sizes vary not only among the different samples but and in the structure of the same sample, due mainly to the extend grain variation of BA (~2-1000 μm). Porosity forming is accompanied with boating effect when the simultaneous mechanisms of melting on the grains surface, trapping gas inner, out gassing at high temperature are taking place, leading to the final lightweight products (Anagnostopoulos, 2009).

Table 5 – Semiquantitative (%) analyses of major crystalline phases in the sintered products.

<table>
<thead>
<tr>
<th>Sample</th>
<th>MMEG</th>
<th>MP1</th>
<th>MP2</th>
<th>MP3</th>
<th>MK1</th>
<th>MK2</th>
<th>MK3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Phases</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>amorphous</td>
<td>37</td>
<td>36</td>
<td>37</td>
<td>44</td>
<td>36</td>
<td>32</td>
<td>41</td>
</tr>
<tr>
<td>anhydrite</td>
<td>tr</td>
<td>1.5</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1.5</td>
<td>1</td>
</tr>
<tr>
<td>gehlenite</td>
<td>3</td>
<td>20</td>
<td>9</td>
<td>4</td>
<td>27</td>
<td>17</td>
<td>16</td>
</tr>
<tr>
<td>Dicalcium silicate</td>
<td>1</td>
<td>7</td>
<td>7</td>
<td>2.5</td>
<td>5</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>diopside</td>
<td>15</td>
<td>5</td>
<td>5</td>
<td>5</td>
<td>6</td>
<td>10.5</td>
<td>5</td>
</tr>
<tr>
<td>anorthite</td>
<td>36</td>
<td>26</td>
<td>34</td>
<td>38</td>
<td>22</td>
<td>25</td>
<td>21.5</td>
</tr>
<tr>
<td>quartz</td>
<td>4.5</td>
<td>2.5</td>
<td>3.5</td>
<td>3</td>
<td>2.5</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>Hematite/ magnetite</td>
<td>1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>tr</td>
<td>-</td>
<td>2</td>
</tr>
<tr>
<td>Brawn millerite</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>-</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Lime/ Portlandite</td>
<td>-</td>
<td>tr</td>
<td>tr</td>
<td>tr</td>
<td>tr</td>
<td>-</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>-</td>
<td>tr</td>
<td>1</td>
<td>-</td>
<td>tr</td>
<td>-</td>
<td>1.5</td>
</tr>
</tbody>
</table>
4. Conclusions
The bottom ashes from Kardia, Ptolemaida and Megalopoli regions contain the demanded incombustible C for its utilization as fuel during the sintering process (till 1200°C) by the sinter band method. The final mineneralogical compositions of sintered products reflecting the complicated mechanisms have taken place during the rapid sintering, leading to the booming effect and porous structures. The higher SiO₂ content of MMEG is believed to be essential for the evolution of the sintering process by participation in the formation of crystalline phases which enhance bonding and lead to the production of the desired sinter-cake. Presence of free lime in the sintered products is undesired, since its subsequent transformation to portlandite by air humidity is accompanied by volume expansion resulting in disintegration of the solid matrix, making it unsuitable for further utilization. According to the macroscopic characterization, the physical properties as well as the mineral compositions and microstructures, the MMEG sintered mixture could be characterized as a light aggregate material and it is proposed as suitable for further treatment in the lightweight concrete production.

5. Acknowledgments
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