IMPROVEMENT OF CEMENT PASTES COMPOSITE PROPERTIES CONTAINING CLAY NANOPARTICLES

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The present work aims to investigate the influence of clay nanoparticles (CNP) on the mechanical, physical and microstructure cement pastes composites. The cement paste composites were formulated by adding 20-60 mass % fly ash (FA), fine blast furnace slag (FBFS) and/or 6 mass % clay nanoparticles. The physico-chemical properties of the cement paste composites (CPC) were improved by the substitution of 6 % CNP in instead of OPC (ordinary Portland cement) in the consistence of the superplasticiser (SP). The physico-mechanical properties and hydration characteristics were investigated by the determination of the setting times (STs), the consistency (W/C, %), the free lime (FL), the combined water (Wn), the gel/space ratio (X), the total porosity (TP), the compressive strength (CS) and the bulk density (BD). The compressive strength values of the cement-CNP hybrid were higher than those of the cement paste composites without CNP. The STs of CPC-pastes containing CNP were accelerated. The CNP showed synergetic effect with the FA and FBFS to enhance the performance of pozzolanic reaction to form supplementary CSH, CAH and CASH, these phases are responsible for bridging, producing a rigid closed compact structure; hence the compressive strength, gel/space ratio, bulk density and chemically combined water increased, whereas the porosity decreased.

INTRODUCTION

Nowadays, the construction industry is mainly based on advanced concrete and modified cement paste composites, this is attributed to their, good performance, versatile applications easiness to fabricate, and low expenditure. However, the disadvantages of these cement-based materials, such as their low tensile strength, have led to multiple pathways of degradation in the technical properties of cement-based materials and the high costs of repairing them [1]. Recently, nano-materials were used to enhance all the properties and durability performances of modern concrete. It was shown that adding nano-materials may enhance the mechanical properties due to its surface effect, size effect, and interface effect. Nano-materials (NM) affect the performance of the physico-mechanical, chemical and microstructure of the cement composite by nucleation, seeding and filler effect. These materials facilitate the pozzolanic reaction to form C–S–H, CAH, and CASH phases within the cementitious matrices [2-6]. The most commonly used nanomaterials in cementitious products are nanoscale particles like silica-nanoparticles (SNP), Al2O3 (NA), TiO2 (NT), ZnO (NZ), Fe2O3 (NF), clay nanoparticles (CNP) and carbon nano-tubes [7-9]. Clay nanoparticles are inexpensive raw materials that improve the microstructure of the cement composite [10-12]. A feature of CNP is its platelet form having a thickness of 1 nm and width of 70 to 150 nm. The uses of CNP in concrete may enhance the mechanical properties in cementitious composites by decreasing the porosity and pore size of the cement matrix [13, 14]. It was found that pastes containing 4, 6 and 10 % CNP showed higher compressive strength values [15, 16]. It has been postulated that the filling effect of CNP contributes the strength development and the pozzolanic reaction starting within 3 - 7 days [17].

The previous study also showed that the CH content of a cement paste with 10 % CNP decreased up to 42 %, this was attributed to the pozzolanicity of CNP, so that CNP improves the texture of the hydration products of the cement pastes and results in a quite dense, compact and uniform microstructure [15, 18].

Mineral admixtures such as fly-ash (FA), silica fume (SF) and fine blast furnace slag (FBFS) are waste inorganic materials that have pozzolanic and/or latent hydraulic properties, improve the properties of concrete [19]. The technological and economic benefit use of these waste inorganic materials, as a partial replacement
in concrete include the improvement of the impermeability, chemical durability, thermal behaviour and enhance the strength. The partial replacement of waste inorganic materials may reduce the early strength, later strength, microstructure and durability of the concrete [20].

Bohá, et al., [21] stated that fine slag offers various technological, economical, and environmental advantages, where it has significantly lower CO₂ emissions in comparison with OPC. Subsequently, it is more actively used as an admixture for concrete. Fly ash is also utilised as a by-product of industrial manufacturing processes. Fly ash is the residue from burning coal in power plants, which has glassy spherical particles.

The present research is aimed at studying the effect of clay nanoparticles on the mechanical properties and microstructure of blended cement-pastes.

**EXPERIMENTAL**

The starting materials used in this study were OPC, FBFS, FA and clay nanoparticles (CNP). The chemical analyses of these materials were investigated by XRF spectrometry as shown in Table 1. The clay nanoparticles were calcined at 750 °C/2 hrs. The XRD pattern of the raw CNP is shown in Figure 1. The superplasticiser-based polycarboxylate (SP) is an opaque light-yellow liquid with a density of 1.08 g∙ml⁻¹.

The batch composition based on the aforementioned starting materials are tabulated in Table 2. The CNP and SP were stirred with a suitable W/C ratio (standard consistency) as given in Table 2, at speed of 120 rpm for two min; the anhydrous blends (OPC, FBFS and/or FA) were mixed to full homogenisation at 90 rpm up to 2 min and homogenised in a rotary mixer at 120 rpm up to 30 s; then the cement blends were allowed to rest for 90 s; this was followed by mixing at 120 rpm for 1 min, then the blended pastes were cast into 2.5 × 2.5 × 2.5 cm³ moulds, and then stored in a humidity cabinet (100 % RH) at 23 ± 1 °C for 24 hrs and then stored under water until the required time of testing (1, 3, 7, 28, 90 and 360-days).

The consistency and STs of the fresh mixed blends were monitored according to ASTM: C191 [22]. The hydrated pastes were stopped using a methanol-acetone mixture, and diethyl ether, and then dried at 70 °C up to 1 h [23]. The chemically combined water content (Wn), free lime (FL), bulk density (BD) and total porosity (TP) were determined as described in previous research [24, 25]. The compressive strength was calculated after the determination of the BD and TP according to the ASTM specifications (C-150) [26]. Some selected hydrated samples were examined using XRD, DTG/TGA and SEM techniques to verify the mechanism predicted by the chemical and mechanical tests. For the XRD analysis, a Philips X-ray Diffractometer of Cu Kα radiation (λ = 1.5418 Å) was used (model PW 1730). The speed of the scan was 20/min between 5° - 65°. The X-ray tube voltage was 40 kV and the current was 25 mA. The XRD analysis was performed with computer software searching for the PDF diffraction data. The TGA/DTG analysis was performed by using a Shimadzu DSC-50 thermal analyser at a heating rate of 10 °C∙min⁻¹, the rate flow of nitrogen was 30 cm³∙min⁻¹. The microstructure of the OPC and some other selected cement pastes were investigated by Scanning Electron Microscope (SEM).

### Table 1. The chemical analyses of the OPC, FBFS, FA and CNP (mass %).

<table>
<thead>
<tr>
<th></th>
<th>SiO₂</th>
<th>Al₂O₃</th>
<th>Fe₂O₃</th>
<th>CaO</th>
<th>MgO</th>
<th>SO₃</th>
<th>Na₂O</th>
<th>K₂O</th>
<th>LOI</th>
<th>Total</th>
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<tbody>
<tr>
<td>CNP</td>
<td>61.24</td>
<td>20.89</td>
<td>5.38</td>
<td>0.16</td>
<td>0.38</td>
<td>0.17</td>
<td>0.71</td>
<td>0.61</td>
<td>10.62</td>
<td>99.99</td>
</tr>
<tr>
<td>FBFS</td>
<td>43.21</td>
<td>9.97</td>
<td>0.59</td>
<td>35.96</td>
<td>5.43</td>
<td>1.37</td>
<td>0.79</td>
<td>0.67</td>
<td>1.98</td>
<td>99.97</td>
</tr>
<tr>
<td>FA</td>
<td>63.10</td>
<td>26.54</td>
<td>5.4</td>
<td>2.33</td>
<td>0.01</td>
<td>0.09</td>
<td>0.85</td>
<td>0.52</td>
<td>0.8</td>
<td>99.64</td>
</tr>
<tr>
<td>OPC</td>
<td>21.3</td>
<td>3.58</td>
<td>5.05</td>
<td>63.48</td>
<td>1.39</td>
<td>2.05</td>
<td>0.26</td>
<td>0.22</td>
<td>2.57</td>
<td>99.90</td>
</tr>
</tbody>
</table>

### Table 2. The composition of the different mixes (mass %).

<table>
<thead>
<tr>
<th>Mix</th>
<th>OPC</th>
<th>FA</th>
<th>FBFS</th>
<th>CNP</th>
<th>SP</th>
</tr>
</thead>
<tbody>
<tr>
<td>C0</td>
<td>100</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>C1</td>
<td>94</td>
<td>0</td>
<td>0</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>C2</td>
<td>60</td>
<td>40</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>C3</td>
<td>54</td>
<td>40</td>
<td>0</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>C4</td>
<td>60</td>
<td>0</td>
<td>40</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>C5</td>
<td>54</td>
<td>0</td>
<td>40</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>C6</td>
<td>54</td>
<td>20</td>
<td>20</td>
<td>6</td>
<td>1</td>
</tr>
<tr>
<td>C7</td>
<td>34</td>
<td>20</td>
<td>40</td>
<td>6</td>
<td>1</td>
</tr>
</tbody>
</table>

![Intensity (a.u.) vs. 2θ (°) graph](image-url)

Figure 1. The XRD pattern of the raw CNP.
RESULTS AND DISCUSSION

Water consistency, initial, and final setting times

The mixed and design patches of the starting materials are represented in Figure 2. The results show that the CPC pastes containing CNP need excess water for consistency and have shorter setting times when compared with the neat cement paste. The CPC containing CNP was homogenised by a considerable amount of consistency for good workability, in addition to the presence of SP, leading to increases in the fluidity of the fresh CPC; therefore, the CPC cement pastes possess high hydraulic characteristics, followed by an acceptable setting time when compared with the blank cement paste. This may be due to the acceleration of the ettringite formation and the CSH, CAH and CASH hydrated gel products by the added CNP [27-31].

The results showed that the consistency of CPC pastes containing FBFS are lower than those of the CPC-cement containing the FA-composite. It was found that the FA-composite cement paste containing FA was homogenised with a high amount of consistency and gives longer setting times, this is attributed to the lower activity of the FA than the FBFS. On the other hand, the cement containing FBFS showed lower consistency, while the setting times were reduced. It was concluded that mix C5 showed the lowest setting times when compared with all the cement mixes, this may be due to the formation as mentioned before. The STs of the CPC-pastes containing CNP accelerated in the existence of CNP [27]. The results also show that the consistency of mixes C6 and C7 decreases, whereas the STs elongated, this is due to the reduction of the OPC-content, as shown in Figure 2.

Free lime contents

Figures 3 represents the free lime content (FL) of the cement composite pastes containing the CNP after 360 days of curing. From Figure 3, it can be found that all the CPC-pastes mixes showed a decreasing trend, while the blank cement paste (OPC) showed an increasing trend from day 1 up to day 360, respectively. All the CNP mixes cured after 3 days showed a slight decrease in the FL content. For the blank OPC cement paste (mix C1), during all the curing ages of the hydration reaction, it liberates excess Ca(OH)₂ (CH) while the addition of CNP acts as a promotor, which accelerates the pozzolanic reaction which means decreasing the CH by formation of CSH, CAH and CASH gel hydrates [32, 33]. For the mixes containing FA, FBFS and CNP, which showed a synergetic effect to enhance the pozzolanic reaction performance, leads to a decrease in the free lime contents.

The results show that the FL content of the OPC paste increases with the age up to 360 days, which is attributed to the continued hydration reaction, liberating the CH through the hydration period. The presence of CNP tends to decrease the residual portlandite, due to the pozzolanic interaction of the CNP with the portlandite leading to the formation of CSH, CAH and CASH.
Improvement of cement pastes composite properties containing clay nanoparticles

The presence of CNP, leading to the raising of the FL content to 3 days due to the enhances in the hydration reaction OPC (lime production), and then decreases up to 360 days where the rate of consumption of the CH is higher than the rate of liberation due to higher pozzolanic activity of the CNP (lime consumption). The occurrence of SP tends to approach the CNP particles from the liberated lime to form further CSH, CAH and CASH, which precipitated in the vacant pore-system; hence, the porosity decreases as shown later. Mix C7 showed lower values than mixes C6 and C5.

Chemically combined water contents

The chemically combined water content ($W_n$) of the different CPC mixes is illustrated in Figure 4. All the mixes showed increasing trends all over the curing ages. The mixes containing 6 mass % CNP have higher $W_n$ values in comparison with the neat-OPC paste. The hydration reaction of the OPC cement phases and the pozzolanic reaction is considered to be the main reason for this increase, the presence of CNP promotes the pozzolanic reaction via the formation of many hydrated phases like CSH(I) and CSH(II) with a considerable quantity, which is responsible for increasing the chemically combined water content.

The results show that the pozzolanic action of the FA is lower than that of the FBFS, the FA-composite-cement-pastes have lower $W_n$ values than the FBFS-CPC-cement-pastes. The results also show that the chemically combined water content increases by increasing the content of the FBFS, where mix C7 has higher $W_n$ values than mix C6, especially at the early ages of hydration, this may due to the pozzolanic reaction of the FBFS, the CNP with the liberated CH from the OPC cement hydration, which enhances the later hydration times [34, 35].

Compressive strength

The compressive strength (CS) of the composite-pastes containing 6 % CNP up to 360-days is illustrated in Figure 5. The data obtained from Figure 5 represent an increase in the CS with an increase in the age-period for all the hardened pastes, qualified to the formation of a successive amount of CSH, CAH and CASH as the main source of strength, these hydrates precipitated in the accessible vacant pores to form a closed compact body [30]. The inclusion of 6 mass% CNP gives a higher CS than pastes without the CNP (Figure 5), ascribed to the formation of micro- and nano-sized hydrates in the cementitious matrix. The CS of the composite-cement-pastes showed better-quality in the presence of 6 % CNP for all the CPC pastes containing FA and/or FBFS; this is due to the pozzolanic reaction of the CNP and the free CH liberated during the OPC hydration production [36].

Figure 5 shows that the CS of mix C5 is higher than the FA-CPC pastes (mix C3) this is attributed to the pozzolanic activity of the hybrid CNP and FBFS, which worked to develop the hydration process of the CPC, especially at the later hydration times. The data also, showed that the CS of mix C7 had higher values than mix C6, due to the pozzolanicity of the FBFS and CNP.
show a higher gel/space ratio value in comparison with mix C3, due to the presence of 6 % CNP, which activates the reaction of the FBFS to give a higher CS and an X-ratio as shown in Figures 5 and 6. The represented results showed that the gel/space ratio increased by the FBFS content (mixes C7 and C6). Mix C7 showed higher gel/space ratio values than C6, this demonstrates that 40 % of the FBFS is more suitable than 20 % of the FBFS which gives more hydration products.

Figure 6. The gel/space ratio of the pastes containing 6 mass % CNP.

The values of the bulk density (BD) and the total porosity (TP) of the CPC pastes in the existence of 6 % CNP and the SP in the hybrid effect treated up to 360 days are tabulated in Table 3. The BD increases wherever the TP decreases with the treatment ages, this is accredited to the increases in the excessive hydration products. These products act as binding centres amongst the unhydrated and hydrated phase fractions of the CPC-grains through the progressive hydration, hence the TP decreased. The gel/space ratio and the bulk density increase in addition, the total porosity decreases. The BD cement composite pastes containing CNP represents higher values than the neat paste and/or the other CPC mixes lacking the CNP [30]. The BD of mix C5 has higher values than those of the neat-paste and mix C3, especially at later hydration ages. By increasing the content of the FBFS up to 40 %, the BD increases and the TP decreases. From the derived results, it can be concluded that the physico-mechanical properties, namely the CS, gel/space ratio, TP, and BD are in good agreement with each other and they represent an increasing trend, except for the TP, which showed a decreasing trend from 1 day up to 360-days.

Table 3. The bulk density and total porosity of the composite pastes in the presence of 6 mass % CNP.

<table>
<thead>
<tr>
<th>Time (days)</th>
<th>Bulk density (g·cm⁻³)</th>
<th>Total porosity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>C0</td>
<td>C1</td>
</tr>
<tr>
<td>1</td>
<td>2.146</td>
<td>2.267</td>
</tr>
<tr>
<td>3</td>
<td>2.178</td>
<td>2.301</td>
</tr>
<tr>
<td>7</td>
<td>2.197</td>
<td>2.324</td>
</tr>
<tr>
<td>28</td>
<td>2.226</td>
<td>2.349</td>
</tr>
<tr>
<td>90</td>
<td>2.252</td>
<td>2.367</td>
</tr>
<tr>
<td>360</td>
<td>2.288</td>
<td>2.410</td>
</tr>
</tbody>
</table>

Figure 7 represents the X-ray diffraction patterns of the hardened CPC pastes namely the C0, C3, C5, C6 and C7 mixes treated at 1 day and 90 days. Figure 7 represents the hydrated and anhydrate-phase diffraction lines of the CH, β-C₂S, C₃S, C, CSH and quartz as shown in Figure. 7. The diffraction lines are consistent to the C0 mix exhibiting higher intensities as represented for the portlandite diffraction lines.

As the hydration-proceeds, the intensity of the peak, illustrating the diffraction patterns of the CSH, increased, whereas the diffraction patterns corresponding to the CH decreased from one-day up to 90 days, as represented in the composite pastes containing CNP (mixes C3 and C5). This is due to the pozzolanic reaction between the CNP with the lime liberated during the hydration reaction of the cement, leading to the higher consumption of portlandite and the formation of further C–S–H. Increasing the content of the FBFS-FA-CPC pastes, the peaks characteristic to the portlandite and calcite phases decreased; this is due to lowering the content of the OPC.
This result is in good agreement with the results of the compressive strength and the gel/space ratio.

TGA/DTG thermograms

The TGA/DTG thermograms of the OPC and composite pastes containing 6 % CNP and 1 % SP cured at 1 day and 90 days are represented in Figure 8. The DTG/TGA curves show the typical temperature events up to 1000 °C. The peaks presented at 80 – 190 °C are attributed to the decomposition of the calcium silicate hydrate, tri-sulphoaluminate hydrate (Aft), and/or monosulphate-hydrated products (AFm). The peak situated at 120 - 190 °C is related to the dehydration of the CSH in the different crystalline states, which overlapped with the Aft, AFm and CASH hydrates [38]. However, the peak situated at 450 - 525 °C is interrelated to the dehydroxylation of CH [39, 40]. The endothermic peaks detected at 680 and 720 °C are accredited to the calcination of the amorphous and crystalline CaCO3 (C) [41]. The exothermic peak located at 850 - 950 °C is due to the recrystallisation of the monocalcium silicate, which is produced from the pozzolanic reaction [42]. Figure 8A illustrates the TGA/DTG thermograms of the C0 mix hydrated for 1 day and 90 days, it can be deduced that the areas under the peaks corresponding to the C-S-H and CH phases increase with the treatment ages. Whereas the addition of CNP to the FA and/or FBFS CPC-paste leads to an increase in the area under the peak identical to CSH and used up the CH in its pozzolanic action, hereafter, the area under the peak identical to the CH decreases when compared with the neat paste as shown in Figure 8B and 8C.

Microstructure and morphology of cement paste composites

The SEM micrographs of the selected cement composite mixes C3, C5, C6, and C7 are represented in Figure 9a-e. From Figure 9a-e, it was found that the sample C3 cured after 90 days of hydration showed...
a denser structure, which is mainly composed of the fully nano-crystalline hydration products and rod-like particles of the CSH (Figure 9a). The dense structure may be due to the effect of the CNP particles’ nucleus, which accelerates the pozzolanic activity leading it to form a homogenous compact CPC.

The SEM micrograph of mix C5 treated after 90 days depicts a porous microstructure with wider pores available for the crystallisation of the formed CSH, CAH, CASH. These microstructures exhibit a higher degree of crystallinity of a mineral hexagonal form of gehlenite like hydrate (C$_2$ASH$_8$), the pore size decreases due to the higher activity of the CNP as shown in Figure 9b. This microstructure reflects the higher compressive strength values, which promotes a decrease in the free lime by the formation of the CSH, CAH, CASH phases [43-45].

Figure 9. The SEM micrographs of the composite cement pastes; a) C3-90d; b) C5-90d; c) C6-90d with 50μm magnification; d) C6-90d with 10μm magnification; e) C7-90d.
Figure 9c and 9d represent the micrographs of the hardened cement paste composite C6 with different magnifications treated at 90 days. The SEM micrograph of mix C6 displayed the presence of C–S–H and CH as the main hydrated products in addition to the presence of C–A–H, C2ASH8 and C3ASH8. The microstructure of mix C6 is a heterogeneous structure where it contains different types and number of pores when compared with mix C5. This microstructure reflects the variation of both the compressive strength, bulk density and porosity.

Figure 9e represents the micrograph of C7, the microstructure represents the flaky plate-like morphology of C2ASH8. It also shows the presence of ill-crystalline and microcrystalline hydrates of C–S–H with the complete disappearance of Ca(OH)2. After 90 days of hydration, more hydrated products were placed in the pores where the denser structure is obtained. The micrograph also shows longer rod-like formations of C–S–H. This CSH phase is in a large quantity, which is responsible for bridging the cement particles, producing a rigid closed compact structure.

CONCLUSIONS

It can be concluded that:

- The results show that the CPC pastes containing CNP need excess water for consistency and shorter setting times when compared with the neat cement paste. The STs of the CPC-pastes containing CNP accelerated in the existence of the CNP. The results also show that the consistency of mixes C6 and C7 decreased, whereas the STs elongated.

- The free lime content of the CPC-pastes mixes showed a decreasing trend, while the OPC showed increasing trends from 1 day up to 360 days. The mixes containing CNP showed a synergetic effect with the FA and the FBFS to enhance the pozzolanic reaction performance forming further CSH, CAH and CASH, which precipitated in the vacant pore-system producing a compact matrix; hence, the compressive strength, gel/ space ratio, bulk density and chemically combined water increased, whereas the porosity decreased.

- The microstructure of the CPC represented the presence of the flaky plate-like morphology of C2ASH8 as well as the ill-crystalline and microcrystalline hydrates of C-S-H with the complete disappearance of Ca(OH)2. This CSH phase is responsible for bridging, producing a rigid closed compact structure.

REFERENCES


