Chemical and Engineering Properties of Blended Cement Containing Micro- and Nano-silica

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Abstract: Nano-technology (NT) and nano-materials (NMs) has attracted considerable scientific attention, due to the new potential uses of nano-particles (NPs), which can change the concrete world. There is a great interest in replacing a long time used materials in concrete structure by NMs to improve the different properties of concrete. In this study, the hydration characteristics and microstructure of cement pastes and mortars with and without nano-silica (NS) were studied. Blended cement mix containing 10 mass, % silica fume (SF) was prepared, then the cement mix was partially replaced by NS up to 1.5 mass, %. The results show that, NS increases the water of consistency and final setting time. The values of chemically combined water, free lime, bulk density and compressive strength prove that, NS up to 1% seems to be an effective mineral additive for blending with OPC to improve its chemical and engineering properties. This is mainly attributed to that; NS behaves not only as filler to improve microstructure, but also as pozzolanic material (PZM). The formation of more hydrated silicates in presence of NS was confirmed by XRD, DTA, IR and SEM examinations. At 1.5 mass, % NS, the density and strength are reduced but still higher than those of 0.0 and 0.5% NS. Also, the results indicate that, the substitution of blended cement mix (OPC+10% SF) with 1 mass, % NS gives a reasonable hydration characteristics and improved engineering properties more than those of 0.0 and 0.5% NS at all hydration ages up to 90 days. Therefore, this level of NS is suggested to be the most suitable replacement level.

Keywords: Microsilica, Nanosilica, Setting Times, OPC, Chemical Properties, Engineering Properties and Microstructure

1. Introduction

Nanotechnology (NT) is advanced technology, which deals with the synthesis of nanoparticles (NPs), and their applications. More recently, NT has attracted considerable scientific interest, due to the new potential uses of particles in nanometer (10⁻⁹m) scale; the advent of NT has brought in various types of nano-materials (NMs), which may have good applications in civil engineering, especially concrete production, due to their major effects: size effect, surface effect, and interface effect [1-4]. It was stated that [5], “the fineness of cement affects it’s reactivity with water and generally the finer the cement the more rapidly it will react”. The role of NT in conceiving of innovative infrastructure systems has the potential to transform the civil engineering practice and dilate the vision of civil engineering [6]. NMs show unique chemical, physical and engineering properties different from those of the conventional materials. Indeed, NMs can change the concrete world, due to their properties at ultra-fine level [7]. There is a great interest in replacing a long time used materials in concrete structure by NMs to improve the engineering properties of concrete [8, 9]. Also, If NMs are combined with traditional building materials; this may lead to production of building materials with unique
properties and be very useful for construction industry.

NMs were used either to replace part of cement, producing ecological profile concrete or as admixtures in cement pastes [3]. In both cases, the addition of them improves the performance of cement; in fresh mixtures an improvement of rheological properties has been recorded [10] and in the hardened state, the compressive strength was increased [11, 12]. The nano-scale observations revealed the nano-crystallized structure of CSH [13, 14] and also NPs have been found to act as nuclei for cement phases, promoting their hydration [15, 16].

Nano-particles exhibit unique properties that develop a stronger, more durable concrete with favorable characteristics when added to concrete. Addition of NMs at the time of production of concrete can lead to significant improvements in the strength and life performance of concrete. One of the active areas of research dealing with cement and concrete is strength and durability of concrete containing NPs [17].

The role of NPs can be summarized as follows: i) NPs act as fillers in the empty spaces, ii) well distributed NPs act as crystallization centers (nuclei) of hydration products, increasing hydration rate, iii) NPs assist towards the formation of small sized Ca(OH)₂ crystals and homogeneous clusters of CSH composition, and iv) NPs improve the structure of the transition zone between aggregates and paste [7]. Commonly used NMs are nano-silica (NS), nano-alumina (NA), nano-titania and nano-ferrit (NF). The most NMs, which has a positive impact on engineering properties of cement concrete is NS. The performance of amorphous NMs is mainly influenced by chemical reaction at interface, enhancing the properties of concrete [18-20]. Nanosilica (NS) is typically a highly effective PZM. It is usually about 1000 times higher than the average cement particle. This has proven to be a superior mixture to improve and reduce the permeability of cement strength and durability. NS is in relation to the other components of silica and reduces the setting times of cement [21].

Addition of NS in cement paste and in concrete can lead to completely different effects. One is size effect, i.e. primarily based on their particle nature, which makes it useful as filling material and the second is the commerce pozzolanic activity of the cluster compounds [22]. NS in cement mortar not only increases the compressive strength, but also results in a finer pore structure, and then reduces overall permeability of hardened concrete [23-25].

Pozzolanic activity of NS and SF was previously studied [26], and the results indicated that, the compressive strength development of pastes made with Ca(OH)₂ and NS, the pozzolanic reaction rate of NS with Ca(OH)₂ and the velocity of C-S-H formation showed marked increase over those of Ca(OH)₂ with SF. The bond strength as well as the bending strength of concrete incorporated with 3 mass,% NS increased more than those with SF, especially at early hydration ages.

Pozzolanic reactivity of NS was much greater than that of SF. The results suggest that, with a small amount of NS, the Ca(OH)₂ crystal at the interface between hardened cement paste and aggregate at early ages may be effectively absorbed in High Performance Concrete (HPC).

In a previous work [27] it was reported that, NS powder caused pronounced improvements in the mechanical properties of mortars and concretes, most notably in compressive strength. Senff et al. [28] found that, fluidity; spreading and hardening time of the cement paste and mortar were reduced as the proportion of NS powder was increased.

The aim of this work is to investigate the chemical and physico-mechanical properties of blended cement containing micro- and nano-sized silica. The water of consistency, initial and final setting times were determined for each cement paste. Also, the values of chemically combined water, free lime, bulk density and compressive strength of hardened cement specimens were measured as a function of curing time up to 90-days. To follow up the hydration rate of the prepared samples, some selected hardened samples were investigated using the XRD, DTA, IR and SEM techniques.

2. Materials and Experimental Details

2.1. Materials

The starting materials used in this study were the ASTM Type (I) ordinary Portland cement (OPC), silica fume (SF) and nanosilica (NS). OPC with Blain surface area of 3000±50 cm²/g was provided from El-Masryah Cement Company, Egypt. The oxide analyses of OPC and SF obtained by X-ray fluorescence (XRF) spectrometry is summarized in Table 1. The mineralogical composition of OPC is listed in Table 2. SF with specific surface area of about 30.774±2 m²/g was supplied from Sika Chemical Company, Egypt. NS with average particle size, Blain surface area and purity percentage of about 15 nm, 50 m²/g and 99.9%, respectively.

Table 1. Chemical Oxide Analyses of OPC and SF (mass, %).

| Oxides | SiO₂ | Al₂O₃ | Fe₂O₃ | CaO | MgO | SO₃ | Na₂O | K₂O | L. O. I | Total |
|--------|------|-------|-------|-----|-----|-----|------|-----|--------|-------|
| OPC    | 19.30| 3.94  | 3.80  | 62.67| 1.90| 3.22| 0.44 | 0.39 | 3.04   | 99.70 |
| SF     | 94.81| 0.16  | 0.84  | 0.89 | 0.49| 0.08| 0.05 | 0.20 | 2.43   | 99.95 |

Table 2. Mineralogical Composition of OPC.

| Compound         | Chemical formula | Abbreviation | Content, % |
|------------------|------------------|--------------|------------|
| Tri-calcium silicate | 3CaO·SiO₂       | C3S          | 66.08      |
| Di-calcium silicate   | 2CaO·SiO₂       | C2S          | 5.5        |
| Tri-calcium aluminate         | 3CaO·Al₂O₃      | C3A          | 4.02       |
| Tetra-calcium aluminoferrite     | 4CaO·Al₂O₃·FeO₃ | C4AF         | 11.55      |

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2.2. Experimental Details

2.2.1. Preparation of Cement Pastes and Mortars
Nano-silica was synthesized according to the following procedure: a desired amount of Na$_2$SiO$_3$ solution was diluted with distilled water; the solution stirred for 15 min, and then precipitated using diluted HCl. The precipitate was filtered, washed several times with distilled water until complete removal of NaCl from the precipitate, and then the precipitate was dried overnight. To decrease the particle size of the prepared powder, it was milled by ball mill (600 rpm) for 10 hours [29]. The amorphous nature of NS was verified using XRD and TEM techniques (Figures 1 & 2). OPC was partially substituted with NS up to 1.5 mass, %. Each dry mix was blended in a steel ball mill using five balls for 1 hour in order to attain complete homogeneity. The cement blends were mixed in a rotary mixer. NS-particles are not easy to disperse uniformly in water, due to their high surface energy. Accordingly, the mixing was performed as follows: a) NS was stirred with 25.0% of the required water for standard consistency at high speed of 120 rpm for 2 min., b) the cement and the residual amount of mixing water were added to the mixer and homogenized at medium speed (80 rpm) for another 2 min., c) the mixture was allowed to rest for 90 second, and then mixed for 1 min at 120 rpm and d) the paste was manually placed, pressed and homogenized in stainless steel moulds. After the top layer was compacted, the top surface of the mould was smoothened by the aid of thin edged trowel. For preparation of mortars, the sand was added gradually in step b) and mixed at medium speed for additional 30 sec. The mortars were prepared according to ASTM (C109-93) by mixing 1 part of cement and 2.75 parts of sand proportion by weighing with water content sufficient to obtain a flow of 110±5 with 25 drops of the flowing table [30-32]. Freshly prepared cement mortars were placed in 50×50×50mm cubic moulds into two approximately equal layers manually compacted and pressed until a homogeneous specimen was obtained. The moulds were vibrated for a few minutes to remove any air bubbles and to give a better compaction. The mix compositions of the prepared blends are shown in Table 3.

2.2.2. Determination of Water of Consistency and Setting Times
The required water of standard consistency gives a paste which permitted the settlement of the Vicat plunger (10 mm in diameter) to a point 5-7 mm from the bottom of the Vicat moulds. It was measured to get all specimens having the same workability. The required water of standard consistency and setting times for each mix were determined according to ASTM specification [33].

2.2.3. Curing and Stopping of Hydration
The specimens were cured in a humidifier (100% R. H) at room temperature 23 ± 2°C for 24 hours, then immersed in tap water until the time of testing. After the predetermined curing time, the hydration of cement pastes was stopped by pulverizing 10 grams of representative sample in a beaker containing methanol-acetone mixture (1:1), and then mechanically stirred for 1 hour. The mixture was filtered through a gouch crucible, G4 and washed several times with the stopping solution, then with ether. The solid was dried at 70°C for 1 hour to complete evaporation of alcohol, then collected in polyethylene bags; sealed and stored in desiccators for analysis [34].

2.2.4. Determination of Combined Water
The chemically combined water content (Wn, %) is used as an indication for the degree of cement hydration. Wn is that portion retained in the sample after the free water is removed. The Wn, % is considered as the percent of ignition loss of the dried sample (on the ignited weight basis). Approximately 2 grams of the pre-dried sample were ignited at 1000°C for 1 hour. The results of Wn, % were corrected for the water of free lime present in each sample [35].

Table 3. Mix Composition of Blended Cements, (mass, %).

| Mix No. | (OPC+10%SF) | N. S |
|---------|-------------|------|
| M0      | 100         | 0    |
| M1      | 99.5        | 0.5  |
| M2      | 99          | 1    |
| M3      | 98.5        | 1.5  |

Figure 1. XRD Pattern of Nano-silica.

Figure 2. TEM Photograph of Nano-silica.
2.2.5. Free Lime Determination
The free lime content of each hydrated cement paste was estimated by the following method, the sample (0.5 g) was poured in 40 ml of a glycerol-ethanol mixture (1:5 v/v), together with a small amount of anhydrous BaCl$_2$ (0.5g) as a catalyst, and phenolphthalein (Ph Ph) as an indicator. This mixture was kept in a conical flask, fitted with an air reflux, heated on a hot plate for 30 minutes (the color becomes pink). The contents of the flask were titrated with a standardized alcoholic ammonium acetate solution until the pink color just disappeared. Heating was again affected, and if the pink color reappears, the titration was completed with ammonium acetate solution until no further appearance of pink color occurs up on heating [36]. The free lime percentage was calculated as follow:

\[
\text{CaO, } \% = \left[ \frac{(W1 \times V)}{W} \right] \times 100, \quad W = \text{original weight}
\]

\[
W1 = \text{weight of CaO equivalent to amount of added alcoholic ammonium acetate, } V = \text{volume of ammonium acetate per ml.}
\]

2.2.6. Bulk Density and Compressive Strength Measurements
The bulk density (BD) was carried out on cement pastes. Samples were suspended weighed in water and in air (saturated surface dry). Each measurement was conducted on at least three similar cubes of the same mix composition and curing time. Then, the density was calculated as described elsewhere [37].

For compressive strength determination according to (ASTM Designation: C-150, 2007) [38], a set of three cubes was tested on a compressive strength machine of SEIDNER, Riedinger, Germany, with maximum capacity of 2000 KN force.

2.2.7. X-ray Diffraction
The powder method of X-ray diffraction (XRD) was adopted in the present study. For this, a Philips diffractometer PW 1730 with X-ray source of Cu ka radiation (\(\lambda = 1.5418 \text{ Å}\)) was used. The scan step size was 2θ, the collection time of 1.0 cm in diameter. Transfer the KBr disk to a sample holder and place it in the spectrometer. The FTIR spectral analysis was recorded from KBr disks using Mattson Gensis FTIR spectrometer in the range from 400 up to 4000 cm$^{-1}$ [40].

2.2.9. Differential Thermal Analysis
The DTA was carried out in air using a DT-30-Thermal Analyzer Shimadzu Co., koyoto, Japan. Calcined alumina was used as an inert material, about 50 mg (76 µm) of each of the finely grounded hydrated cement pastes was housed in a small platinum-rhodium crucible. A uniform heating rate was adopted in all experiments at 20°C/min.

2.2.10. Microstructure Examination
The microstructures of some selected samples were investigated by FEI Inspect S ESEM Holland. Backscattered electron (BSE) imaging was used to study the samples without coating.

3. Results and Discussion

3.1. Water of Consistency and Setting Times

The variations of water of standard consistency (w/c,%) of the investigated cement pastes are graphically represented in Figure 3. It obvious that, the water demand increases with NS%. The increase in water demand is attributed to the acceleration of hydration process of cement clinker phases under the effect of high fineness and surface energy of NS particles [41- 44]. Thus the specimens containing NS require more water to rapid formation of hydration products [27].

The variations of the initial and final setting times (IST&FST) of the investigated cement pastes with NS% are plotted in Figure 4. It can be seen that, IST decreases and FST increases with NS content. The decrease of IST is mainly due the acceleration of cement hydration under the effect NS, leading to the formation of excessive amounts of CSH, which fill up some of open pores originally filled with water that accelerates the setting. The increase of FST is attributed to the high water demand for standard consistency in presence of NS [3].

3.2. Combined Water Contents

The degree of cement hydration can be measured from the combined water (Wn) and free lime (FL) contents. Figure 5 indicates the variations of combined water content of hydrated cement pastes up to 90 days as a function of NS%. It is apparent that, Wn % increases with curing time for all hydrated cement pastes. This is essentially due to the continuous hydration of cement clinker phases, leading to the formation of more hydrated products. The values of Wn% increase with NS content from 0.5 to 1.5% with two different rates, the first (0.5-1%) is faster than the second (>1%). After 1% NS, the Wn% decreases, but still more than those containing 0.0 and 0.5% NS. The increase of Wn% with NS% seems to be due to two factors work together, the high water demand and pozzolanic action of NS. Nano-silica reacts with the liberated lime from the hydration of cement clinker phases, leading to the formation of additional and excessive hydrated products with high water contents, such
as CSH, CAH and CASH. The results also show that, NS accelerates the hydration of cement clinker phases, especially at early ages of hydration [43], beginning from the time of mixing up to 28 days.

3.3. Free Lime

The pozzolanic reaction of NS can be kinetically followed by monitoring the free lime contents at different curing times. The variation of free lime contents (FL %) of cement pastes with different NS contents (0-1.5%) and hydrated up to 90 days is shown in Figure 6. The results indicate that, the values of FL decrease with curing time for all investigated cement pastes with and without NS. This is due to the pozzolanic reaction between the amorphous glassy silica (SF&NS) with the free Ca(OH)$_2$ liberated during calcium silicate hydration [3]. At a given hydration time, the FL content decreases with NS%. This is due to the higher pozzolanic activity of NS comparing with SF.

3.4. Bulk Density

The bulk density (BD) values of cement pastes with different NS contents (0.0-1.5%) and hydrated up to 90 days are graphically represented in Figure 7. The results show that, the bulk density increases with curing time for all hydrated cement pastes, due to the continuous hydration of cement phases, leading to the formation and accumulation of successive amounts of hydrated silicates (CSH) and alumino-silicates (CASH), which tends to increase the gel/space ratio as well as the bulk density [34, 45], leading to the formation of uniform and compact microstructure. The BD value increases with NS percentage up to 1 mass%, and then decreases at 1.5%, but still higher than those containing 0.0 and 0.5% NS. This can be interpreted as follows [27]: Suppose that NS particles are uniformly dispersed in cement paste and each NS-particle is contained in a cube pattern, therefore the distance between NS-particles can be determined. After the hydration begins, the hydration products diffuse and envelop NS particles as kernel [46]. If the NS content and the distance between them are appropriate, the crystallization will be controlled to be a suitable state through restricting the growth of Ca(OH)$_2$ crystals. Moreover, the NPs located in cement paste as kernel can further promote cement hydration, due to their high activity. This makes the size of Ca(OH)$_2$ crystals smaller and the cement matrix more homogeneous and compact. Consequently, the pore structure is improved. With increasing the NS content more than 1%, the improvement on the pore structure of cement paste is weakened, but still more better than those of 0.0 and 0.5%NS. This can be due to that, the distance between NPs decreases with increasing the content of nanoparticles, and Ca(OH)$_2$ crystals cannot grow up enough due to limited space and the crystal quantity is decreased, which leads to the decrease of crystal to strengthening gel ratio. Therefore, the shrinkage and creep of cement matrix are increased [47].
3.5. Compressive Strength

The effect of NS% on the compressive strength values of hydrated cement mortars up to 90 days is graphically represented in Figure 8. It is clear that, the compressive strength values increase with curing time for all hydrated cement pastes, due to the continuous hydration and formation of excessive amounts of hydration products. These products accumulate in water filled pores to form a more compact body [45]. The results also indicate that, the compressive strength of the investigated mortars increases with NS content up to 1%, and then decreases at 1.5%, but still more than those containing 0.0 and 0.5% NS. The improvement of compressive strength in presence of NS up to 1% is attributed to that, NS behaves not only as a filler to improve the microstructure of cement mortar, but also as an activator to promote pozzolanic reaction. Both the nucleation and pozzolanic effects of NS lead to more accumulation and precipitation of hydration products in the open pore volume, leading to the formation of uniform, dense and compact microstructure. Consequently the bulk density and compressive strength increase with NS up to 1%. The decrease of compressive strength at 1.5% NS seems to be due to the decrease of the distance between NS-particles with NS%. Therefore, Ca(OH)₂ crystals cannot grow up enough due to the limited space and the crystal quantity is decreased, which leads to the reduction of crystal/gel ratio [47], thus the bulk density and compressive strength are relatively decreased.

3.6. Interpretation of XRD Results

Figure 9 represents XRD patterns of cement pastes containing 1% NS and hydrated at different curing times up to 90 days. The results indicate that, the peaks of Ca(OH)₂ and anhydrous silicates decrease with curing time in contrast with the peak of Tobermorite like phase (CSH). This seems to be due to the hydration progress as well as the pozzolanic reaction of SF and NS with the liberated Portlandite, leading to the formation of additional amounts of hydrated silicates. Also, the calcite (CaCO₃) peak increases with curing time, due to the carbonation of hydration products. It is also clear that, the peaks related to formation of the hydration products shifts to appear at earlier curing times, indicating, the positive effect of NS-particles on the formation of both Portlandite and Tobermorite phases at early hydration ages [27].

The X-ray diffractograms of hydrated cement pastes with and without NS for 28 days are shown in Figure 10. It is clear that, the all hydrated cement mixes show Portlandite Ca(OH)₂, Tobermorite (CSH) and anhydrous silicates. Nano-modified cement mixes exhibit lower Portlandite peak intensity and higher Tobermorite peak intensity than the corresponding peaks in case of the control mix. The intensity of Portlandite peak decreases with NS content. The peaks of anhydrous silicate phases superpose with those of the hydrated silicates and decrease with NS content. But, the CSH has opposite manner to both Portlandite phase and anhydrous silicates. This is due to the pozzolanic reaction of NS with Ca(OH)₂ formed from cement clinker hydration.
3.7. IR Spectroscopy

The effect of curing time up to 90 days on the hydration characteristics of cement pastes containing 1% NS can be seen from the IR spectra in Figure 11. The observed peak at about 3645 cm\(^{-1}\) is due to the \(-\text{OH}\) group of Ca(OH)\(_2\). Its intensity decreases with hydration time. This is due to that, NS acts as activator to promote the hydration of cement clinker phases, liberating free lime. Also, NS is a highly pozzolanic material (PZM) to react with the liberated Portlandite from cement hydration. With curing time, the pozzolanic effect of NS exceeds its accelerating action. Therefore, the rate of lime consumption becomes more than that of its production. The appeared band in the range 950-985 cm\(^{-1}\) is assigned to the formation of Tobermorite like phase (CSH) \([50]\). The broad band centered at about 3400 cm\(^{-1}\) and that at 1640-1650 cm\(^{-1}\) are due to stretching and bending vibration of water bound in the hydrated silicates. The band around 1480 cm\(^{-1}\) is attributed to C\(\equiv\)O bond stretching of CO\(_3^{2-}\). The CSH peak has opposite pattern to that of Ca(OH)\(_2\).

Figure 11. IR Patterns of M2 at Different Curing Times.

Figure 12 represents the IR spectra of cement pastes with different NS% and hydrated for 28 days. It is well known that, the phases of non-ordered structures cause the increase of band width and decreases its intensity. The results show the presence of peak at 3637-3640 cm\(^{-1}\) that can be assigned to the \(-\text{OH}\) band from Portlandite phase. This band is useful for diagnosis of the initiation of hydration process \([48]\). The intensity of that peak reduces with NS%, due to the pozzolanic reaction of NS with the liberated Portlandite during cement phases hydration. The broad band at about 3448 cm\(^{-1}\) and that observed at the range 1640-1650 cm\(^{-1}\) are mainly due to the stretching and bending vibration of lattice water in the hydrated silicates and alumino-silicates (CSH and CASH)\([49]\). Their intensities increase and their widths decrease with NS%, due to the pozzolanic action of NS, which accelerates the formation of more hydrated silicates. Also, the appeared band at 950-980 cm\(^{-1}\) is due to the formation of CSH \([50]\). The band at 1480 cm\(^{-1}\) is due to C-O bond stretching of CO\(_3\), which result from the carbonation of hydrated silicates as well as the residual Ca(OH)\(_2\). The exposure of cement pastes to CO\(_2\) results very quick carbonation \([48]\). Its intensity increases with NS%. This may be due to that; NS behaves as activator to promote the hydration rate of cement phases. The observed band at 495 cm\(^{-1}\) is due to bending vibration of Si-O bond in SiO\(_4^{2-}\) tetrahedral \([51]\). Its intensity increase with NS content, due to the above-mentioned reasons.

Figure 12. IR Patterns of Hydrated M0, M2 and M3 at 28 Days.

3.8. Differential Thermal Analysis (DTA)

Figure 13 represents the DTA thermograms of M\(_2\) at different curing times up to 90 days. It can be seen that, the intensity of CH endotherm increases and shifted to higher dissociation temperature with curing time to 7 days, then decreases up to 90 days. Therefore, the area of Ca(OH)\(_2\) peak and its temperature range are reduced with curing time after 7 up to 90 days. The endothermic peak of CSH has opposite manner to that of Portlandite, because the more consumption of lime means more CSH formation.

The DTA results of cement mixes containing different NS percentages and hydrated at 28 days are graphically represented in Figure 14. Very small endothermic peaks appear at low temperatures up to 250°C, may be due to the decomposition of hydrated silicates as Tobermorite like phase (CSH). The endothermic peak located at 477-480°C may be assigned to the dehydration of free Ca(OH)\(_2\) whereas the broad endotherm at 700-750°C is mainly related to the decomposition of calcite phase (CaCO\(_3\)) or carbonated CSH. It is clear that, the Ca(OH)\(_2\) endothermic peak is occurred at lower temperature range than the normal. Also, its area decreases with NS%. This is mainly due to the pozzolanic reaction of NS particles with free lime. The results of DTA analysis are in a good harmony with those of XRD, IR and chemical analyses.

3.9. Microstructure Discussion

To verify the mechanism predicted by the experimental results obtained in this work, SEM examinations were performed. Figure 15 represents the microstructure
photographs of M2 at different curing times (3, 7, 28 and 90 days). The micrographs show increase of compaction and homogeneity of the internal microstructure with curing time for all hydrated mixes. This is principally due to the continuous hydration of cement clinker phases as well as the pozzolanic reaction of NS with free lime, leading to the formation of successive and additional amounts of hydrated products, which responsible for compaction and strength properties of cement paste.

Addition of NS was found to affect the hydration behavior of cement paste and leads to differences in the microstructure of the hardened cement paste. Figure 16 represents the microstructure photographs of M0, M2 and M3 at 28 days. The CSH gel existed in the form of ‘stand-alone’ clusters, lapped and jointed together by many needle hydrates. At the same time, Ca(OH)₂ crystals were distributed among the cement paste in the microstructure of M0. But, the microstructures of the pastes containing NS, which are of higher strength than M0 revealed a pores filling with dense and compact structure, i.e., the texture of hydrated products was more dense and compact. Big crystals such as Ca(OH)₂ are nearly absent, due to the pozzolanic reaction of NS with the lime liberated during cement clinker hydration, leading to the formation of more hydration products. Thus the number and size of calcium hydroxide (CH) crystals are reduced. It is clear that, 1% NS gives cement paste with more uniform and denser microstructure than 1.5%. The suggested mechanism that, the NS can improve the microstructure and consequently the strength of cement paste is illustrated as follows: When a small quantity of NS is uniformly dispersed in the cement paste, the hydrated products of cement will deposit on the nano-particles (NPs), due to their great surface area and high surface energy. The hydration products will grow to form conglomeration containing the NS-particles as ‘nuclei’. Due to its high pozzolanic activity, the NS-particles located in the cement paste as nuclei will further activate and accelerate the hydration of cement clinker phases. The uniformly disperse situation of NS gives a good microstructure with uniformly distributed conglomeration. According to Wu’s ‘centroplasm’ hypothesis, the aggregates, sands and other particles are considered as centroplasm that acts as skeleton, and gel as transmitter substance. The binding force between centroplasm and transmitter substance has an important effect on the strength of concrete [52]. The NS-particles distributing in cement paste as ‘sub-centroplasm’ can tightly bond with the hydrated products around the interfacial transition zone (ITZ) between the NPs and hydrated products. On the other hand, the NPs among the hydrated products will prevent the crystal from growing, such as Ca(OH)₂ and AFm, and such fine crystals are favorable for the strengthening properties of cement paste [53–55]. Also, the NS-particles will fill pores to increase the strength. However, when NS cannot be well dispersed, as the case of its extensive content, the aggregating NS-particles will create weak zone, in form of voids. Consequently, the homogeneous microstructure cannot be formed, and low strength will be expected.
4. Conclusions

Based on the experimental results of this work, the following conclusions were drawn:

1. Nano-silica (NS) significantly increases the water demand for standard consistency, and consequently final setting time (FST). This seems to be controlled by the particle size distribution and high specific surface area of NS.

2. The free lime decreases with NS%, due the high pozzolanic action of NS.

3. Chemically combined water contents increase with NS%. This is attributed to the pozzolanic reaction of NS with the lime liberated from cement hydration, leading to the formation of more hydrated products with higher water contents than that of Ca(OH)$_2$.

4. The bulk density (BD), microstructure and consequently the mechanical properties of the investigated cement mortars are improved with NS up to 1%, and then decreases at 1.5%, but still higher than those containing 0.0 and 0.5% NS. With increasing the NS content more than 1%, the improvement on the pore structure of cement paste is weakened, but still better than those of 0.0 and 0.5% NS. This can be due to that, the distance between nano-particles decreases with increasing the content of nanoparticles, and Ca(OH)$_2$ crystals cannot grow up enough due to limited space and the crystal quantity is decreased, which leads to the decrease of crystal to strengthening gel ratio.

5. The compressive strength values increase with curing time for all hydrated cement pastes, due to the continuous
hydration and formation of excessive amounts of hydration products. These products accumulate in water filled pores to form a more compact body. More rapid formation of hydrated products in presence of nano-SiO\textsubscript{2} was confirmed by XRD, DTA, IR and SEM examinations. Their results are in a good harmony with those of chemical and physico-mechanical of the investigated materials.

6. The NS up to 1% proved to be an effective mineral additive for blending with Portland cement to improve its chemical and engineering properties.

7. The substitution of blended cement mix (OPC+10% SF) with 1wt.% NS gives a reasonable hydration characteristics and improved engineering properties more than those of 0.0 and 0.5% NS at all hydration ages up to 90 days. Therefore, this level of NS is suggested to be the most suitable replacement level.

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