



# Investigations of geopolymeric mixtures based on phosphate washing waste

R.Dabbebi<sup>a,b\*,</sup> S.Baklouti<sup>a</sup>, J.L. Barroso de Aguiar<sup>b</sup>, F.Pacheco-Torgal<sup>b</sup>, B.Samet<sup>a</sup>

<sup>a</sup>University of Sfax, Laboratory of Industrial Chemistry, Sfax 3038, Tunisia <sup>b</sup>University of Minho, C-TAC Research Centre, Guimarães, 4800-058, Portugal

## Abstract

The extraction of the phosphate ore produces a high amount of waste causing serious environmental problems. This waste, termed as phosphate washing waste, was filtered and dried at 105°C for 24 h to remove the water. The dried waste was milled and then sieved in a 100 µm sieve. The resulting phosphates washing waste (PWW) particles size are below 70 µm. The phosphate washing waste was calcined at 700°C and 900°C.n Both calcined and uncalcined waste were investigated with X-ray fluorescence (XRF), X-ray powder diffraction (DRX), Fourier transform infrared (FTIR), simultaneous differential thermal and thermogravimetric analyses (DTA-TG) and particle size analysis. This waste was activated with sodium hydroxide (NaOH) and sodium silicate in order to produce geopolymeric materials. The influence of replacing PWW by 15% of metakaolin was also study. The results show that the highest compressive strength is obtained with metakaolin. The results also showed that compressive strength decreased with the increase of NaOH concentration.

Keywords: Phosphate washing waste, sodium hydroxide, sodium silicate, geopolymer.

#### 1. Introduction

Waste recycling is an important feature in the context of sustainable development and also important to the circular economy. The natural Gafsa phosphate is extracted by CPG (Gafsa Phosphate Company) from sedimentary deposits of marine origin, their geological origin dates from about 50 million years. The most used techniques by CPG to extract the phosphate are washing and flotation to eliminate mineral and organic impurities. Tunisia is an international pioneer in the field of natural phosphate and mineral fertilizers and it is the second country in the world to develop a large sterile matrix like clay, quartz to increase the percentage of  $P_2O_5$  which can reach 37%. The ore was washed with a flocculent. Afterwards, it was sorted according to the particle size. The biggest particles exceeding 2 mm were stored in heaps and the finest particles with a dimension below 70µm are the phosphate washing waste. This product was placed in a basin with around a dozen hectares [1, 2]. This waste is hazardous in nature constituting a potential source of pollution and its storage creates environmental problems related to land pollution. Fig 1 shows a view of the sludge pond.

נעראפין אראשאנגע אראשער אראשער אראשער אראשער אראשער געראראין אראשער געראפע אראשער געראפע אראשער געראפע אראשער דעראפע אראשער אראשער געראפע געראפע געראפע געראפע געראפע געראין געראפע געראפע געראפע געראפע געראפע געראפע געראפע דעראפע געראפע געראפע

I he treatment of bhosbhate ore consists in removing the View metadata, citation and similar papers at <u>core.ac.uk</u>

\* R.Dabbebi

rawiabali@gmail.com

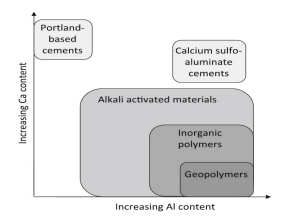
binding phase is almost exclusively aluminosilicate and very highly coordinated.

The available calcium content of the reacting components is usually low; to enable formation of a pseudo-zeolitic network structure rather than the chains characteristic of calcium silicate hydrates.



Fig 1.Tunisian phosphate sludge pond

Fig 2 shows the distinction between geopolymers and other binders. The activation of aluminosilicates can also be performed with an acid solution such as in the case of phosphate geopolymer or silico phosphate. Geopolymers have the ability to reuse several wastes [4,5]. They still present some shortcomings that need to be addressed such as a high cost and some durability problems [6]. Large number of researchers worked with a variety of raw materials to synthesis geopolymers. However, no work has addressed the use of phosphate washing waste as a raw material. Therefore, this study discloses preliminary results regarding not only the characterization of that raw material but also its geopolymerization.



#### Fig 2.Classification of different subsets of alkali activated materials with comparisons to OPC and calcium sulfoaluminate binder chemistry

# 2. Experimental works

The phosphate washing waste used in this study was taken from Metlawi's storage ponds (laverie VI) a region in the south of Tunisia. The phosphate washing waste was slurry with or without flocculent. Particle size of the slurry with flocculant was characterized by laser diffraction (Fritsch analysette 22 micritec plus). Then, It was filtered and dried at 105°C for 24 hours to remove the water. The dried phosphate washing waste was crushed, and then sieved to 100µm. It was calcined in a muffle furnace LAB TechDaihan LAB TECH CO.LTD for 2 hours with heating rate 5°/min at 700°C and 900°C. After calcination the samples were characterized with XRF, XRD and FTIR. The XRF were conducted by X ray fluorescence (Philips X'unique II), the X-ray diffraction (XRD) was conducted on a Bruker D8 Discover with Cu-Ka radiation ( $\lambda$ =1,54060Å) at 40 kV and 40 mA. Each sample was scanned from 5° to  $60^{\circ}$  at a speed of  $0.02^{\circ}$ s<sup>-</sup> <sup>1</sup>. The FTIR were scanned from 400 to 4000 cm<sup>-1</sup> wavenumbers using the Fourier Transform infrared model perkin Elmer spectrum BX spectrometer. The thermal behaviour of the phosphate washing waste was performed under argon atmosphere with heating rate 10°C/min from ambient to 1000°C with TA instruments SDT 2960 Simultaneous. The compressive strength of the consolidate material was evaluated with three cylindrical specimens with 35 mm in diameter and 70 mm in height. The compression tests were carried out using a LLOYD Instruments LR50K apparatus equipped with a 50 kN sensor. The descent rate of the upper platen was set at 0.2 mm/min.

### 3. Results and discussion

The chemical analysis of phosphate washing waste calcined and uncalcined was done by XRF and the results of this analysis are reported in the table 1. The table shows that the PWW has an important quantity of SiO<sub>2</sub>, CaO, Al<sub>2</sub>O<sub>3</sub> and P<sub>2</sub>O<sub>5</sub>. Figs 3 and 4 show the distribution of the particles size of the phosphate washing waste with and without flocculent. The Fig 5 shows some samples of phosphate washing waste powder before and after calcination.

Oxides	25°C	700°C	900°C
SiO <sub>2</sub>	42	44.3	42.7
Al <sub>2</sub> O <sub>3</sub>	9.77	10	9.45
CaO	26.5	24.9	26.7
P <sub>2</sub> O <sub>5</sub>	10	8.49	8.75
Na <sub>2</sub> O	1.12	1.64	1.50
K <sub>2</sub> O	0.673	0.798	0.779
MgO	3.09	2.75	2.91
SO <sub>3</sub>	3.39	3.96	4.10
Fe <sub>2</sub> O <sub>3</sub>	2.31	2.32	2.39
LOI	18	14.7	17

Table 1.Chemical composition (%) of PWW according to calcination temperature

PS: Phosphate sludge

Moisture of uncalcined PS = 0.1851% Melting temperature of PWW Tf°C =1150°C

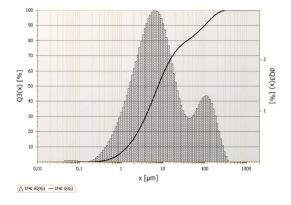


Fig 3. Particle size analysis of phosphate washing waste with flocculent

With flocculent the distribution is bimodal. The most dominant is between 0.9 and 60  $\mu$ m, and the majority of particles have a diameter of 7.5  $\mu$ m. The less dominant population is between 60 and 250  $\mu$ m, with particles majority having an approximate diameter of 100  $\mu$ m. The difference of the size particles can be explained by the presence of the flocculent used by the

company to recover the maximum of water. Fig.4 shows the distribution of particles size of the phosphate washing waste without flocculent.

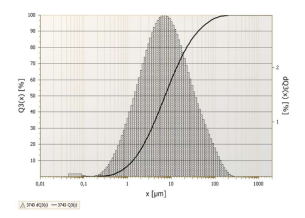


Fig 4. Particle size analysis of phosphate washing waste without flocculent

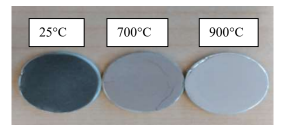


Fig 5. Phosphate washing waste powder before and after calcinations

The patterns of the XDR show the presence of amorphous and crystalline phases (Fig. 6).

The Ca/P ratio determined by the chemical composition of uncalcined material is 2.65>1.67: this indicates the presence of calcite or dolomite, or that there is a substitution of PO<sub>4</sub> by CO<sub>3</sub> [7], and this was confirmed by the XRD. The diffractogram shows the presence of calcite, fluorapatite, palygorskite, heulandites, quartz and gypsum.

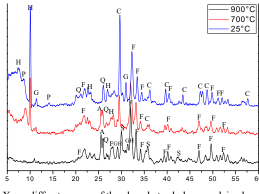
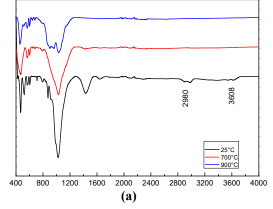


Fig 6. X ray diffractograms of the phosphate sludge uncalcined and calcined at 700°C and 900°C. C: Calcite (PDF: 00-002-0629), H:
Heulandites (PDF#01-076-2214), G: Gypsum (PDF#:00-021-0816)
, Q: Quartz (PDF#:01-070-2535), F: Fluorapatite (PDF#:00-002-0845), P: Palygorskite (PDF#00-029-0855), A: Anhydrite
(PDF#:00-003-0368), S: Calcium Iron Silicate (PDF#:00-003-1134), GH: Gehlenite (PDF#: 00-001-0982)

After calcination at 700°C, the curve shows the disappearance of palygorskite, and the presence of a new phase anhydrite  $CaSO_4$  a decrease in the intensity of the peaks of heulandites and of calcite, this was supported by the FTIR. At 900°C the curve shows a total disappearance of heulandites and calcite and the appearance of new phase calcium iron silicate and gehlenite.



The fig.7 shows the FTIR spectra of phosphate washing waste. The examination of the spectral shows bands at 1636-3543 cm-1 confirming the presence of the stretching hydroxyl group (OH<sup>-</sup>) and these bands disappeared at 700°C and 900°C. The peak at 3616 cm<sup>-1</sup> which characterizes the OH stretching vibration of the palygorskite mineral confirms the results present in the XRD patterns. The organic matter was shown by the presence of band at 2980 cm<sup>-1</sup> and it disappeared also at 700°C and 900°C.

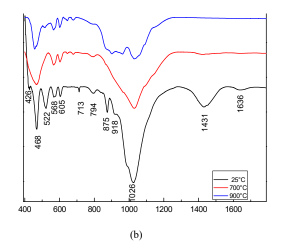


Fig 7. FTIR spectra of phosphate washing waste

(a) 400-4000cm<sup>-1</sup>; (b) 400-1600 cm<sup>-1</sup>)

The bands at 713, 875 and 1431 cm<sup>-1</sup> confirmed the presence of carbonate  $CO_3^{2-}$  [8]. These 3 bands were vanished at 900°C and the weight loss was attributed to the decomposition of carbonates. Asymmetric stretch silicate bands were observed at 1025 cm<sup>-1</sup>, a deformation band of Si-O-Al at 522 cm<sup>-1</sup>, a trace of quartz at 794 cm<sup>-1</sup> and a deformation band at 426 cm<sup>-1</sup> of Si-O. Two bands at 468 and 568 correspond to  $PO_4^{3-}$ . The FTIR spectrum shows also a deformation band at 918 cm<sup>-1</sup> of Al-OH-Al [9] of the palygorskite.

Fig. 8 shows the thermal behavior of phosphate washing waste, the first mass loss is an endothermic effect around 100°C corresponds to the removal of adsorbed water (moisture). The second mass loss is between 200-550°C, an endothermic peak corresponding to structural water of the natural zeolite [10], and an exothermic peak corresponding to the combustion organic matter. The third mass loss is an endothermic peak reflecting the decomposition of carbonates calcite CaCO<sub>3</sub> with release of CO<sub>2</sub>.

Mixtures with phosphate washing waste (uncalcined and calcined at 700°C and 900°C) activated with sodium silicate and sodium hydroxide were tested.

The chemical composition of the sodium silicate solution by mass was Na<sub>2</sub>O: 10.6%, SiO<sub>2</sub>: 26.5% and water: 45% with bulk density 1390 kg/m3. The sodium hydroxide solution was obtained by dissolving different amount of sodium hydroxide pellets (98% purity) in distilled water to have different concentrations 7, 9, 10, 12 and 14M before 24 hours of the mixing.

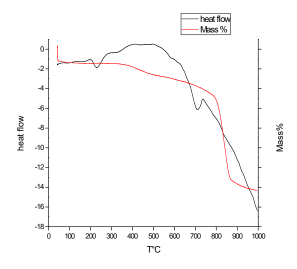


Fig 8. Thermal behavior of phosphate washing waste

concentrations 7, 9, 10, 12 and 14M before 24 hours of the mixing. The liquid/solid ratio of the mixture was  $\sim$ 0.75. The obtained mixtures were placed in a closed polystyrene mold and compacted with vibrating table to remove entrapped air bubbles. After they were cured at room temperature for 7 days. A summary of the different mixtures is presented in table 2.

Liquid/solid	N. C.P.S	C.P.S 900°C	C.P.S 700°C	C.P.S 700°C+MK
S.H 7,9,10,12,14M	NO	NO	NO	NO
S.S	NO	NO	YES	YES
S.H in S.S	NO	NO	NO	NO
S.S+7M	NO	NO	YES	YES
S.S+9M	NO	NO	YES	YES
S.S+10M	NO	NO	YES	YES
S.S+12M	NO	NO	YES	YES
S.S+14M	NO	NO	YES	YES

Table 2. Summary of the mixtures

N.C.P: uncalcined phosphate sludge C.P.S: calcined phosphate sludge Consolidate material "yes" refers to a solid state after 24 hours while non-consolidate material refers to mixtures that were still fresh after 24 hours. This are labelled NO in the table 2. S.H refers to NaOH solution. S.S + xM refers to sodium silicate solution with sodium hydroxide solution. Those mixtures were composed by 50% of sodium silicate solution and 50% of sodium hydroxide solution. S.H in S.S refers that the pellets of sodium hydroxide were dissolved in the sodium silicate solution. The uncalcined powder and calcined at 900°C show different results compared with powders calcined at 700°C which implies that the two powders have more stable phases.

Fig. 9 shows the compressive strength results for mixtures with 7 curing days with sodium silicate and sodium hydroxide. The maximum strength (6MPa) was obtained when using a sodium hydroxide concentration below 10M. Mixtures show lower compressive strength for higher sodium hydroxide concentration. Mixtures with 15% metakaolin showed a higher compressive strength with a maximum of 10 MPa for a sodium hydroxide concentration of 7M.

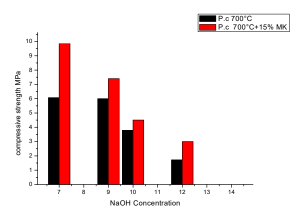


Fig 9. Compressive strength of the geopolymer with sodium silicate and different NaOH concentration

The concentration of sodium hydroxide affects the microstructure and the compressive strength of geopolymers [11]. The increase of NaOH concentration in the aqueous phase of geopolymeric system decreases the dissolution rate of Si and Si-Al phase of phosphate washing waste calcined at 700°C, which affects negatively the mechanical properties of the obtained materials. An excess of hydroxide ion OH<sup>-</sup> concentration caused a precipitation of aluminosilicate gel at a very early stages, and it will accelerate the dissolution, so the geopolymerisation will be stopped which leads to a lower compressive strength [12].

Moreover, in the condition of high NaOH concentration, the sodium cation consumes the surfaces species (T-OH and T-O-), so the chemical bonding between the insoluble solid particles and geopolymeric framework take place in the final stage of the geopolymerization [13].

Fig 10 shows the compressive strength results for mixtures with 7 curing days with only sodium silicate as activator.

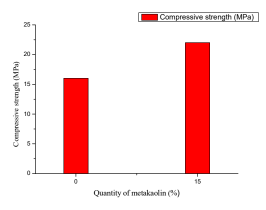


Fig 10. Compressive strength of the geopolymer with only sodium silicate

From the fig 10, the compressive strength of the samples activated with only sodium silicate solution S.S was equal to 16 MPa for the PS. This mixed with 15% of metakaolin shows a compressive equal to 22 MPa. These results were higher compared to the results mentioned in fig 9. The results show that the compressive strength depends necessarily on the type of the alkali activators. The results can be explained by the presence of more active silica provided by the sodium silicate which involve in the geopolymerization process and it improves the compressive strength [14]. these types of the sodium silicate [15]

### 4. Conclusions

In this research, the Tunisian phosphate washing waste was studied by fluorescence X, X-ray diffraction, FTIR, thermal and particle size analysis. The structural changes of phosphate washing waste after calcinations at 700 and 900°C were studied by fluorescence and Xray diffraction and FTIR. After the calcination, physical and chemical properties of the phosphate washing waste are affected, and the preliminary tests of geopolymerization confirmed the results. The phosphate sludge can be used as a source material for making geopolymers cured at ambient temperature. The geopolymer pastes and the compressive strength are dependent on the structure of the precursor and on the NaOH concentration. The sodium hydroxide concentration of 7M with sodium silicate and 15% of metakaolin shows the higher compressive strength after cured 7 days. The results also show that an increase of concentration sodium hydroxide decrease the compressive strength. We should mention also that the compressive strength of geopolymers based on phosphate washing waste and sodium silicate without and with 15% of metakaolin show the higher compressive strength after 7 days of curing: 16MPa and 22 MPa.

# Acknowledgment

The authors acknowledge the Gafsa Phosphate Company "CPG Tunisia" for providing us the phosphate washing waste sample and the Portuguese Foundation for Science and Technology (FCT) for the finance of the project UID/ECI/04047/2013.

# References

 Khaled Boughzela, N. F., Khaled Bouzouita, Habib Ben Hassine: Etude minéralogique et chimique du phosphate naturel d'Oum El Khecheb (Gafsa, Tunisie); Revue science des matériaux, 2015; Vol. 6. pp. 11-29.
 Loutou, M.; Hajjaji, M.; Mansori, M.; Favotto, C.; Hakkou, R. Phosphate sludge: Thermal transformation and use as lightweight aggregate material. Journal of Environmental Management 2013, 130, 354-360.

[3] Van Deventer, J.; Provis, J.; Duxson, P.; Brice, D.: Chemical Research and Climate Change as Drivers in the Commercial Adoption of Alkali Activated Materials. Waste Biomass , 2010 ;1:145–155

[4] Bernal, S.; Rodríguez, E.; Kirchheim, A.; Provis, J.: Management and valorisation of wastes through use in producing alkali-activated. Journal of Chemical Technology & Biotechnology (in press) 2016

[5] Payá, J.; Monzó, J.; Borrachero, M.; M. Tashima, M.: Reuse of aluminosilicate industrial waste materials in the production of alkali-activated concrete binders, in Handbook of Alkali-Activated Cements, Mortars and Concretes, F. Pacheco-Torgal, J. Labrincha, A. Palomo, C. Leonelli, P. Chindaprasirt, Eds, WoodHead Publishing, Cambridge, 2014; pp. 487-518

[6] Pacheco-Torgal, F.; Abdollahnejad, Z.; Miraldo, S.;Kheradmand,M. Alkali-activated cement-based binders (AACB) as durable and cost competitive low CO2 binders: Some shortcomings that need to be addressed, in Handbook of low carbon concrete, 1<sup>st</sup> A.

Nazari, J. Sanjayan, , Elsevier Science and Tech, Waltham, 2016; pp.195-216.

[7] Slansky, M.: Geology of sedimentary phosphates; Elsevier Science Pub. Co. Inc., New York, NY; None, 1986.

[8] Huang, C. K., and Paul F. Kerr. " Infrared study of the carbonate minerals." American Mineralogist 45.3-4 (1960): 311-324

[9] Neji, S. B.; Trabelsi, M.; Frikha, M. H. Activation d'une argile smectite Tunisienne à l'acide sulfurique: rôle catalytique de l'acide adsorbé par l'argile. Journal de la Société Chimique de Tunisie 2009, 11, 191-203.

[10] Gelves, J.; Gallego, G. S.; Marquez, M. Mineralogical characterization of zeolites present on basaltic rocks from Combia geological formation, La Pintada (Colombia). Microporous and Mesoporous Materials 2016, 235, 9-19.

[11] Somna, K.; Jaturapitakkul, C.; Kajitvichyanukul, P.; Chindaprasirt, P. NaOH-activated ground fly ash geopolymer cured at ambient temperature. Fuel 2011, 90, 2118-2124.

[12] Lee, W.; Van Deventer, J. The effects of inorganic salt contamination on the strength and durability of geopolymers. Colloids and Surfaces A: Physicochemical and Engineering Aspects 2002, 211, 115-126.

[13] Panias, D.; Giannopoulou, I. P.; Perraki, T. Effect of synthesis parameters on the mechanical properties of fly ash-based geopolymers. Colloids and Surfaces A: Physicochemical and Engineering Aspects 2007, 301, 246-254. [14]T. Phoo-ngernkham, S. Hanjitsuwan, N.

Damrongwiriyanupap, and P. Chindaprasirt, "Effect of sodium hydroxide and sodium silicate solutions on strengths of alkali activated high calcium fly ash containing Portland cement," KSCE Journal of Civil Engineering, vol. 21, pp. 2202-2210, 2016.

[15]Z. Abdollahnejad, F. Pacheco-Torgal, T. Félix, W. Tahri, and J. Barroso Aguiar, "Mix design, properties and cost analysis of fly ash-based geopolymer foam," Construction and Building Materials, vol. 80, pp. 18 30, 2015.