

Blast-furnace slag cement and metakaolin based geopolymer as construction materials for liquid anaerobic digestion structures: Interactions and biodeterioration mechanisms

Marie Giroudon, Matthieu Peyre Lavigne, Cédric Patapy, Alexandra Bertron

▶ To cite this version:

Marie Giroudon, Matthieu Peyre Lavigne, Cédric Patapy, Alexandra Bertron. Blast-furnace slag cement and metakaolin based geopolymer as construction materials for liquid anaerobic digestion structures: Interactions and biodeterioration mechanisms. Science of the Total Environment, 2021, 750, 10.1016/j.scitotenv.2020.141518 . hal-02937600

HAL Id: hal-02937600 https://hal.insa-toulouse.fr/hal-02937600

Submitted on 14 Sep 2020 $\,$

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

- 1 Blast-furnace slag cement and metakaolin based geopolymer as construction materials for liquid
- 2 anaerobic digestion structures: interactions and biodeterioration mechanisms
- 3 Marie Giroudon^{1. 2.} *, Matthieu Peyre Lavigne^{2.}, Cédric Patapy^{1.}, and Alexandra Bertron^{1.}

4 ^{1.} LMDC, Université de Toulouse, UPS, INSA Toulouse, France

5 ^{2.} TBI, Université de Toulouse, CNRS, INRA, INSA, Toulouse, France

6 *Corresponding author

7 Keywords: Anaerobic digestion, slag cement, geopolymer, biodeterioration, ammonium

8 Abstract

9 In order to promote the development of the biogas industry, solutions are needed to improve 10 concrete structures durability in this environment. This multiphysics study aims to analyse the 11 multiphases interactions between the liquid phase of an anaerobic digestion system and 12 cementitious matrices, focusing on (i) the impacts of the binder nature on the anaerobic digestion 13 process at local scale, and (ii) the deterioration mechanisms of the materials. Cementitious pastes 14 made of slag cement (CEM III), innovative metakaolin-based alkali-activated material (MKAA), with 15 compositions presumed to resist chemically aggressive media, and a reference binder, ordinary 16 Portland cement (CEM I), were tested by immersion in inoculated cattle manure in bioreactors for a 17 long period of five digestion cycles. For the first time it was shown that the digestion process was 18 disturbed in the short term by the presence of the materials that increased the pH of the liquid phase 19 and slowed the acids consumption, with much more impact of the MKAA. However, the final total 20 production of biogas was similar in all bioreactors. Material analyses showed that, in this moderately 21 aggressive medium, the biodeterioration of the CEM I and CEM III pastes mainly led to cement matrix 22 leaching (decalcification) and carbonation. MKAA showed a good behaviour with very low degraded depths. In addition, the material was found to have interesting ammonium adsorption properties in 23 24 the chemical conditions (notably the pH range) of anaerobic digestion.

25 1 Introduction

26 Biogas production is based on the degradation of organic matter by microorganisms under controlled 27 anaerobic conditions (Chen et al., 2020; Huang et al., 2016; Yun et al., 2018). The process called 28 anaerobic digestion consists of four consecutive degradation reactions: hydrolysis, acidogenesis, acetogenesis and methanogenesis (Batstone et al., 2002; Evans and Furlong, 2003; Wang et al., 2018) 29 30 and has been evaluated as being one of the most energy-efficient and environmentally friendly 31 technologies for bioenergy production (Fehrenbach et al., 2008; Yun et al., 2019, 2018). In the 32 context of climate change and depletion of fossil resources, the development of anaerobic digestion 33 structures is encouraged by the European Union, in particular through the European Directives 2001/77/EC and 2009/28/EC (2001, 2009). The process leads to the production of (i) a renewable 34 35 energy source, biogas, which is mainly composed of methane (CH_4) and carbon dioxide (CO_2) (Lastella et al., 2002; Lesteur et al., 2010; Weiland, 2010) and (ii) a digestate that can be used as a fertilizer, 36 37 due to nitrogen, phosphorous and potassium content and as the bioavailability of nitrogen is greater 38 in the digestate than in the initial organic waste (Huang et al., 2016; Weiland, 2010). Optimized 39 productivity is achieved in mesophilic conditions (usually 35°C) (Kothari et al., 2014) and at pH 40 between 6.5 and 7.5 (Braun, 2007; Chandra et al., 2012; Liu et al., 2008). Although the anaerobic 41 digestion technology is widely used, biogas production and digestate reutilization still need to be 42 improved, in order to achieve maximum energy output. To this end, new strategies are being 43 developed such as the co-digestion or the addition of accelerants, leading to a better degradation 44 efficiency (Huang et al., 2016; Wang et al., 2019; Xu et al., 2020; Zhang et al., 2018). Anaerobic 45 digestion plants can be set up with different types of organic waste and the biogas recovered can be 46 used in many ways: production of heat, electricity, and vehicle fuel, or can even be injected after 47 purification, into the network of natural gas (Frigon and Guiot, 2010; Holm-Nielsen et al., 2009; Yun 48 et al., 2018).

49 The implementation of anaerobic digestion on an industrial scale requires airtight, isolated production structures. In this context, reinforced concrete, which is also an economical building 50 material, is the most widely used material for the construction of digesters. Inside the digester, the 51 cover concrete is in direct contact with the anaerobic digestion medium in the liquid phase, while 52 53 polymeric liners or plastic layers are often applied to protect concrete in the gas phase (Nathalie 54 Bachmann, 2013). At every stage of the digestion, the liquid medium contains species that are 55 particularly aggressive to concrete (Koenig and Dehn, 2016; Voegel, 2017; Voegel et al., 2016, 2015), 56 such as volatile fatty acids (VFA) (several g/L), ammonium (about 1 g/L) and dissolved CO_2 (several 57 g/L). These species are responsible for concrete leaching (for VFA and ammonium) and/or carbonation (for dissolved CO₂) of the matrix (Bertron et al., 2005a; Magniont et al., 2011; Voegel et 58 59 al., 2016). Moreover, the microorganisms colonize the concrete surface as a biofilm (Voegel et al., 60 2019a). Biofilms allow enhance the metabolic capacity of cells (Flemming et al., 2016) and seem to 61 amplify the material deterioration, in terms of kinetics and intensity of alteration (Magniont et al., 62 2011). So far, few studies have investigated the fate of concrete exposed to fermenting biowaste. 63 Koenig and Dehn (2016) studied the effects of anaerobic digestion in the liquid and gas phases of an 64 industrial scale pilot digester. After 1 ½ years in the liquid phase, the authors observed erosion of the cover concrete and a degraded depth of 12.8 mm, with significant decalcification in the outer layer, 65 features that are typical of an attack by acids, and precipitation of calcium carbonates. Voegel et al. 66 67 (2019a, 2019b, 2016) identified deterioration mechanisms corresponding to a combination of 68 decalcification and carbonation on cement paste specimens immersed in a synthetic biowaste for 4 69 weeks. The presence of alkaline cementitious materials slightly increased the pH (Voegel et al., 70 2015). Nevertheless, the liquid composition varies greatly over time and according to the substrate 71 (Fisgativa et al., 2016; Li et al., 2015; Xu et al., 2019). Thus, depending on the type of biowaste, the 72 composition in terms of VFA, ammonium and CO₂ varies and may impact the material differently. 73 Moreover, these studies considered only classic cement matrices (based on CEM I, CEM II, CEM III or 74 calcium aluminate cement). Thus, the degradation mechanisms of materials in anaerobic digestion systems remain unknown for various liquid compositions and on innovative matrices. Furthermore,
the impact of the materials on the process efficiency (biogas production) has never been studied.

77 The purpose of this laboratory study was to thoroughly evaluate the bio-geochemical interaction 78 mechanisms between the anaerobic digestion medium and the materials at the local scale and their 79 impact (i) on the quantitative and temporal production of methane and (ii) on the mineralogical and 80 chemical alteration of the material that may compromise its long-term durability and thus the 81 sustainability of the structure. In order to address these interface issues, the exploration strategy was to study the impact of different materials on anaerobic digestion by monitoring the liquid 82 environment and the biogas production on the one hand, and to evaluate changes in the solid 83 84 samples exposed to the medium on the other hand. Cement paste specimens made of blast-furnace 85 slag cement (CEM III), a metakaolin-based alkali-activated geopolymer, and an ordinary Portland 86 cement (OPC) paste (reference cement CEM I) were exposed to the liquid phase of inoculated cattle 87 manure for more than 36 weeks. Ground Granulated Blast furnace Slag (GGBS) is a by-product of the 88 iron industry and metakaolin is a powder obtained by calcination (600 to 800°C) and grinding of a clay mainly composed of kaolinite (AFNOR, 2012a). They may be used alone or as supplementary 89 90 cementitious materials (SCM) with ordinary Portland cement (EN 206/CN, NF EN 197-1) (AFNOR, 91 2014, 2012b) to reduce the environmental impact of the cementitious material manufacturing by 92 reducing the content of OPC, which is CO_2 emitter and energy-consuming (Juenger et al., 2019). 93 Moreover, CEM III cement pastes had already shown increased durability to acid attacks compared to 94 ordinary Portland cement (CEM I) due to the addition of aluminium-rich ground granulated blast-95 furnace slag (Bertron et al., 2005b; Gruyaert et al., 2012; Oueslati and Duchesne, 2012). The slag addition increases the chemical resistance to material leaching by decreasing the calcium content of 96 the binder and leads to a reduced portlandite content, a lower Ca/Si ratio of $C-S-H^1$ – the main 97 98 hydrated phase of the material – and the formation of C-A-S-H (Elakneswaran et al., 2016;

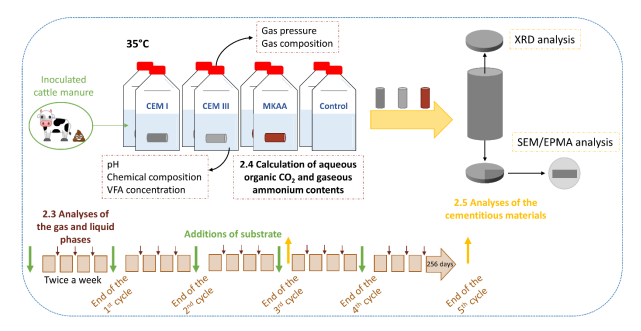
¹ Cement chemistry shorthand notations: $A = AI_2O_3$, C = CaO, $F = Fe_2O_3$, $H = H_2O$, M = MgO, $S = SiO_2$

99 Lothenbach et al., 2011), with higher chemical stability than C-S-H (Roosz et al., 2018). Furthermore, 100 the addition of slag to cement helps to densify the paste by filling the large pores with hydration 101 products and the resulting finer pore structure improves the transfer properties of the paste (Perlot 102 et al., 2006). The French prescription guide T47 (CIMbéton, 2009) recommends slag cements for use 103 in aggressive chemical media and they are actually used in the design of concrete for digester 104 structures. On the other hand, geopolymers have shown encouraging behaviour when exposed to 105 acidic abiotic (Drugă et al., 2018; Duan et al., 2015; Singh et al., 2015) and biotic (Grengg et al., 2020; 106 Khan et al., 2017) environments. However, the chemical and mineralogical compositions of 107 geopolymers are very variable and some of them, such as fly ash geopolymers (Khan et al., 2018), 108 and pure white metakaolin geopolymer (Grengg et al., 2020), showed poor resistance to microbially 109 induced deterioration (MID). The behaviour of these matrices therefore needs to be studied in order 110 to assess their suitability for use in an anaerobic digestion environment.

111 Cement paste specimens were exposed to the fermenting biowaste in biochemical methane 112 potential reactors with a high solid/liquid ratio in order to reproduce local chemical conditions in the 113 vicinity of the concrete walls of the digester. The liquid and gas phases were monitored over time. 114 Scanning Electronic Microscopy (SEM) observations coupled with Electron Probe Micro-Analysis 115 (EPMA), and X-Ray Diffraction (XRD) analyses were used to investigate the chemical and 116 mineralogical changes of the specimen.

117 2 Materials and methods

The experimental protocol implemented in this study is presented in Figure 1. It consisted of a series of bioreactors where construction materials samples were immersed in inoculated cattle manure at mesophilic temperature (35°C) in order to reproduce anaerobic digestion conditions. During the 256 days experiment (5 cycles of anaerobic digestion), both gas and liquid phases were monitored (see section 2.3) while the chemical and mineralogical changes occurred in the cementitious material samples were analysed at the end of the third and fifth cycles (see section 2.5).



124

Figure 1: Schematic representation of the experimental protocol to study the interactions between the anaerobic digestion environment and the cementitious materials

127 2.1 Binder materials

Cement pastes made of CEM III/B 42.5N (CEM III), a mixture of Portland cement and GGBS, and an 128 129 ordinary Portland cement CEM I 52.5R (CEM I), were poured with a water/binder ratio of 0.30. The 130 metakaolin-based alkali-activated (MKAA) paste was made with the following molar ratio - $\frac{SiO_2}{Al_2O_3} = 3.6$; $\frac{Na_2O}{Al_2O_3} = 0.9$; $\frac{H_2O}{Na_2O} = 14.5$ – by using metakaolin (Argeco), liquid sodium silicate (molar 131 ratio $\frac{SiO_2}{Na_2O} = 1.7$) and water. The metakaolin used in this study came from the calcination of a kaolin 132 with a relatively high level of impurity and included quartz (45%), mullite (2%), calcite (1%), anatase 133 134 (1%) and kaolinite (1%), as well as a considerable amount of iron oxide $(3.7\% \text{ of } Fe_2O_3)$ (Pouhet, 2015; 135 San Nicolas et al., 2013).

136 A procedure adapted from the French standard NF EN 196-1 (AFNOR, 2016) was used to mix the 137 paste specimens, i.e., for each cement paste, the binder and the water were mixed at low speed (140 \pm 5 rpm) for 60 seconds and at high speed (285 \pm 10 rpm) for 90 seconds. The pastes were cast in 138 139 cylindrical moulds 75 mm high and 25 mm in diameter. Each specimen was protected from the 140 atmosphere by a plastic coating after pouring. 24h after pouring, the CEM III and CEM I pastes were 141 removed from their moulds, sealed in plastic bags and kept at 20°C for a 27-day endothermic cure. 142 MKAA pastes were cured for 7 days in their closed moulds and then stored in sealed plastic bags. At 143 28 days of maturation, the paste specimens were exposed to the fermenting biowaste.

144 **2.2** Preparation of laboratory reactors and immersion of cement pastes

To start, 100 mL of microbial inoculum collected from an industrial biogas plant in Haute-Garonne (France) and cattle manure from the experimental farm of the Engineering School of Purpan (France) (the substrate) were poured into airtight biochemical methane potential (BMP) reactors (Holliger et al., 2016). For each kind of material, the experiment was carried out in duplicate with two bioreactors each containing a sample. In addition, two control bioreactors without material were also taken into account.

151 Immediately after the first feed with substrate, the paste specimens were immersed in bioreactors. 152 The solid/liquid ratio (cement paste surface area/total liquid volume) was approximately 85 cm².L⁻¹, 153 much higher than the in situ ratio in a standard industrial digester of approximately 4 cm².L⁻¹ (Voegel 154 et al., 2015). This ratio was chosen high in order to reproduce and emphasize local phenomena in the 155 vicinity of the concrete wall, where the chemical conditions are strongly influenced by the 156 interactions between the biowaste and the cement matrix.

After the immersion of the samples, water was added to reach a volume of 800 mL and the gas phase was flushed with N_2 in order to remove O_2 . The bioreactors were maintained at 35°C in an oven for 5 cycles. At each new cycle, the liquid medium from the previous cycle was kept in the reactors, supplemented with water (to reach the initial volume of 800 mL) and small amounts of new cattle

161 manure were incorporated (Table 1). The mass of cattle manure added increased with the cycles in 162 order to offset a decrease in the substrate's methanogenic potential over time. Each cycle was 163 considered complete when the gas production in the control reactors stopped, except for the last 164 cycle which lasted until the biogas production was over in all the bioreactors (see durations in Table 165 1). During the fourth cycle, once the biogas production was complete, the reactors were kept at 166 ambient temperature for 6 weeks because the laboratory closed for the summer break.

167	Table 1: Approximate mass of	f cattle manure added	per cycle and duration of each cycle

	1 st cycle	2 nd cycle	3 rd cycle	4 th cycle		5 th cycle
Temperature	35°C	35°C	35°C	35°C Room		35°C
Mass of cattle manure added (g)	6	6	8	10		13
Duration of cycle (weeks)	7	6.5	5	4	6	8

168

169 **2.3** Sampling and analysis of the anaerobic digestion medium

170 Twice a week, the gas pressure was measured and liquid was sampled. If the gas pressure was above 171 0.250 bar, some gas was sampled for analysis (O₂/N₂, H₂, CH₄ and CO₂) by gas chromatography (GC Trace 1300 Thermofisher) and the gas pressure was then brought back to ambient pressure. Samples 172 173 of 5 mL of the liquid medium were collected and the pH of each sample was measured. The liquid 174 samples were centrifuged (Eppendorf, Centrifuge 5430R) for 5 minutes (4°C, RCF 7197). In one hand, the concentrations of ionic species PO_4^{3-} , Na^+ , NH_4^+ , K^+ , Cl^- , Ca^{2+} and Mg^{2+} were analysed by Ion 175 176 Chromatography (Dionex Thermofisher DX320). In the other hand, 1 mL of the supernatant part was sampled and mixed with 0.5 mL of an internal standard (solution of 2-ethyl-2-butyric acid at 1 g.L⁻¹ in 177 demineralized water and containing 5% orthophosphoric acid H₃PO₄) to be analysed by gas 178 chromatography (Varian SERIE 3900/430) in order to provide the concentrations of some VFA (acetic, 179 180 propionic, isobutyric, butyric, isovaleric, and valeric acids).

181 **2.4** Calculation method for aqueous inorganic CO₂ and gaseous ammonium contents

As the analyses carried out did not allow the aqueous inorganic CO_2 and gaseous ammonium contents to be measured, gas and liquid chromatography results, together with thermochemical equilibria associated with Henry's law, were used to calculate these missing data.

185 2.4.1 Theoretical background: Henry's Law

Henry's law states that the partial pressure of the dilute solute in a solution is directly proportional to its liquid mole fraction. The proportionality factor is called the "Henry's law constant" (H_i) (Equation A).

$$Py_i = H_i x_i \tag{A}$$

189 In Equation A, P, y_i , H_i and x_i are the total pressure, the composition of solute in vapour gas (molar 190 fraction of the gas phase), Henry's law constant of solute i, and the molar fraction of the liquid 191 phase, respectively.

The constants were calculated considering a temperature of 35°C, the atmospheric pressure and the
 pH after each gas sampling (values of the Henry's law constants from Batstone et al. (2002)).

194 2.4.2 Aqueous inorganic CO₂ content

195 The inorganic CO₂ content corresponds to the molar fraction of the liquid phase x_{CO_2} divided by the 196 water concentration, 55.41 mol.L⁻¹ (Equation B).

$$[CO_{2 aq}] = \frac{x_{CO_2}}{55.41} \text{ where } x_{CO_2} = \frac{P_{atm}y_{CO_2}}{H_{CO_2}}$$
(B)

Using the thermochemical equilibria (Reactions 1 and 2), and the gas composition, the total inorganic CO₂ content can be calculated by successively calculating the $[HCO_3^-]$ and $[CO_3^-]$ concentrations.

$$H_2CO_3 + H_2O \rightleftharpoons HCO_3^- + H_3O^+, K_{C_1}$$
 (1)

$$HCO_3^- + H_2O \rightleftharpoons CO_3^- + H_3O^+, K_{C_2}$$
 (2)

where K_{C_1} and K_{C_2} are the thermodynamic equilibrium constants, values from Batstone et al. (2002). $K_{C_1} = 4.73E^{-7}$ and $K_{C_2} = 5.60E^{-11}$ at 35°C.

201 2.4.3 Gaseous ammonium content

Using the thermochemical equilibria (Reaction 3) and the total ammonium concentration in the liquid phase, the $[NH_4^+]$ and $[NH_3]_{aq}$ concentrations were calculated.

$$NH_4^+ + H_2 0 \rightleftharpoons NH_3 + H_3 0^+, K_N$$
 (3)

where K_N is the thermodynamic equilibrium constant, value from Batstone et al. (2002). $K_N = 6.16E^{-10}$ 205 ¹⁰ at 35°C.

Henry's law then allowed the gaseous ammonium content to be calculated, assuming equilibriumand no limitation of liquid/gas transfer (Equation C):

$$y_{NH_3} = \frac{x_{NH_3H_{NH_3}}}{P_{atm}}$$
 where $x_{NH_3} = \frac{[NH_3]_{aq}}{55.41}$ (C)

208 2.5 Analysis of cementitious materials

209 At the end of each cycle, the cylindrical cement paste specimens were removed from the bioreactors 210 and a diamond saw was used to collect slices for solid analyses. Quarter sections were intended for 211 SEM and EPMA analysis. They were embedded in an epoxy resin (Mecaprex Ma2+ from Presi), and 212 dry polished using silicon carbide polishing disks (Presi) (Bertron et al., 2009). Then, the polished 213 sections were coated with carbon. Analyses by SEM (JEOL JSM-LV, 15 kV) combined with EPMA 214 analyses (Cameca SXFive, 15 kV, 20 nA) enabled the microstructural and chemical characterization of 215 the specimens. For EPMA, the points to be analysed were chosen to avoid residual anhydrous grains. 216 Chemical profiles were established from the surface in contact with the liquid phase to the core of 217 the specimen. For each point, the following elements were analysed: Ca, Si, Al, Fe, Mg, S and P. 218 Element mass percentages were expressed as mass percentages of the associated oxides.

The mineralogical composition changes of the specimens were characterized by XRD (Brucker D8 Advance, Co cathode, 40 kV, 40 nA). The plane side of the slice, which was directly in contact with the liquid medium, was first analysed. Then, the sample was abraded in order to characterize the degradation in depth, and this until reaching the sound material (Bertron et al., 2005a).

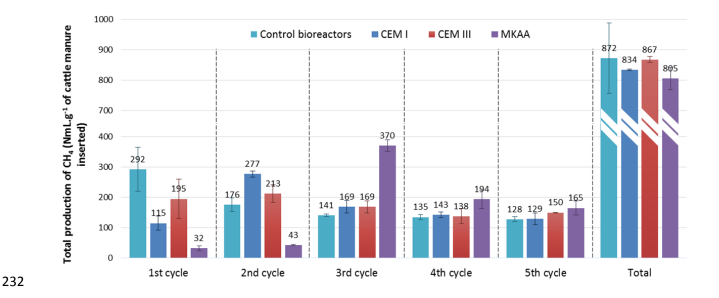
XRD and EPMA analyses were performed after the third cycle. SEM observations were carried outafter each cycle.

225 **3 Results**

3.1 Characteristics of the liquid phase according to the material during the anaerobic
 digestion

228 3.1.1 Biogas production and composition

Figure 2 gives the total production of CH₄ per gram of cattle manure in each type of bioreactor and the cumulative total production at the end of the experiment. Mean values of the two duplicate bioreactors are presented with the standard deviations.



233 Figure 2: Total production of CH_4 (NmL.g⁻¹ of cattle manure) at the end of each digestion cycle and cumulative total **234** production at the end of the experiment (Total)

In the control bioreactors, the production of CH_4 reached between 292 and 128 NmL of CH_4 per gram of cattle manure per cycle. The decrease of the methane production as the number of cycles increased expresses a decrease in the methanogenic potential of the substrate over time, which was probably due to the evolution of the cattle manure during the storage (4°C) time of the experiment.

The presence of the cementitious materials CEM I and CEM III, induced a significantly lower methane production during the first cycle than that in the control bioreactors (292 NmL of CH₄/g of cattle manure) with 115 and 195 NmL, respectively, of CH₄ per gram of cattle manure. During the second cycle, the bioreactors containing the CEM I and CEM III samples showed methane production rates (respectively 277 and 213 NmL of CH₄/g of cattle manure) that were higher than those of the control bioreactors (176 NmL of CH₄/g of cattle manure). Thereafter, production in the bioreactors containing cementitious materials was equivalent to those of the control bioreactors.

The biogas production in the bioreactors containing alkaline MKAA was strongly delayed compared to other bioreactors. About ten times less methane (32 NmL of CH_4 per gram of cattle manure) was produced in the MKAA bioreactors than in the control bioreactors during the first cycle and the production remained very low (43 NmL of CH_4 per gram of cattle manure) during the second cycle. The third cycle was marked by a sharp increase of methane production in these bioreactors (370 NmL of CH_4 per gram of cattle manure). During the fourth and fifth cycles, methane production in the MKAA reactors remained slightly higher than in the control bioreactors.

Even if the presence of the materials delayed the biogas production, at the end of the experiment the cumulative amount of methane produced was equivalent for all the bioreactors (-4.4%, -0.6% and -7.7% relative to the control for CEM I, CEM III and MKAA, respectively).

Table 2 gives the composition of the gas phase (N_2 , CH_4 , CO_2) of the bioreactors at the end of each cycle in volume percent. Mean values of the two duplicate bioreactors are presented with the standard deviations.

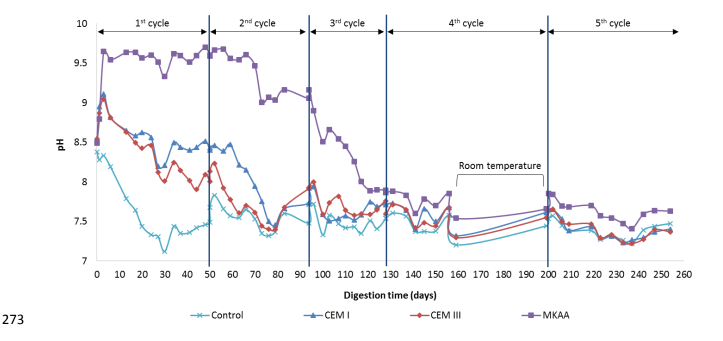
	%	End of 1 st cycle	End of 2 nd cycle	End of 3 rd cycle	End of 4 th cycle	End of 5 th cycle
	N ₂	24.5 ± 3.2	32.5 ± 3.8	31.7 ± 1.1	33.9 ± 1.4	27.9 ± 1.8
Control	CH ₄	54.5 ± 1.7	42.5 ± 1.5	42.8 ± 0.4	44.7 ± 0.4	46.8 ± 1.5
	CO2	22.6 ± 1.6	23.0 ± 2.4	24.0 ± 0.8	19.4 ± 1.2	25.3 ± 0.1
	N ₂	64.2 ± 7.1	33.7 ± 1.1	34.5 ± 5.0	31.4 ± 2.1	23.8 ± 2.6
CEM I	CH ₄	34.8 ± 6.4	48.8 ± 1.2	43.9 ± 3.1	44.0 ± 2.7	48.4 ± 1.0
	CO2	2.6 ± 0.8	15.1 ± 2.4	20.1 ± 1.3	21.2 ± 1.8	27.6 ± 2.0
	N ₂	49.2 ± 13.3	34.2 ± 6.4	32.6 ± 4.8	35.8 ± 5.7	24.7 ± 1.6
CEM III	CH ₄	45.3 ± 8.7	46.9 ± 5.8	45.4 ± 2.6	44.8 ± 3.4	49.6 ± 0.3
	CO2	7.1 ± 4.5	17.3 ± 0.7	20.2 ± 1.9	17.6 ± 1.8	25.1 ± 1.7
	N ₂	87.5 ± 2.9	81.4 ± 1.1	16.6 ± 1.2	26.7 ± 4.7	21.6 ± 4.5
МКАА	CH ₄	12.3 ± 3.2	15.1 ± 0.9	62.7 ± 1.3	52.9 ± 1.2	49.9 ± 3.4
	CO2	0.1 ± 0.0	0.6 ± 0.0	15.5 ± 2.7	17.2 ± 3.4	28.0 ± 0.4

260

261 Table 2 shows that the composition of the gas phase in the control bioreactors varied little over time, 262 with about twice as much CH₄ as CO₂, and approached the composition of an industrial biogas (55 to 263 65% of CH₄ and 30 to 40% of CO₂ (Rasi, 2009)). In contrast, in the bioreactors containing the 264 materials, the proportion of N₂ decreased through the cycles since more and more biogas was 265 produced. Moreover, the proportion CO_2/CH_4 varied depending on materials and time. While it was 266 almost constant in the control bioreactors (about 1/2), this ratio was very low in the bioreactors 267 containing materials during the first cycle. It then increased (more slowly in the bioreactors containing MKAA due to the delay of the efficient biogas production) and tended to reach 1/2 268 thereafter. 269

270 3.1.2 pH and volatile fatty acids production

Figure 3 gives the evolution of the pH during the five cycles of anaerobic digestion in the liquidfraction of the bioreactors.



274 Figure 3: Evolution of the pH during the five cycles of anaerobic digestion in the bioreactors

At the start of the first digestion cycle (day 0) the pH was identical in all the bioreactors (pH = $8.5 \pm$ 0.1). The subsequent measurements showed very different behaviours among the bioreactors. The control bioreactors first underwent a decrease of pH during the first 30 days. From the second cycle, the pH reached suitable values for anaerobic digestion and remained stable between 7.3 and 7.8 for the rest of the experiment.

The alkalinity of the cementitious materials (CEM I and CEM III) impacted the pH which increased significantly in the first few days, reaching a maximum of about 9 on day 3. Thereafter, the negative effect of the cementitious materials seems to have been overcome and the pH decreased, until values similar to those of the control bioreactors were observed at day 76. At the same time, biogas production accelerated in these bioreactors. Singular behaviour was observed in the bioreactor containing the MKAA. As for the bioreactors containing cementitious materials, the pH first increased due to the high alkalinity of the material. However, it reached a very high value, of 9.6, on the third day. The pH remained at this high value during the first cycle and started to decrease halfway through the second cycle, on day 70. During the third cycle, a rapid decrease of pH was observed in the MKAA bioreactors, and pH conditions approached those of production in the other bioreactors. This pH drop occurred at the same time as biogas production started.

During the fourth and fifth cycles, the pH was stable for all types of bioreactors. However, it was observed that the pH in the MKAA bioreactors remained about 0.3 higher than in the others.

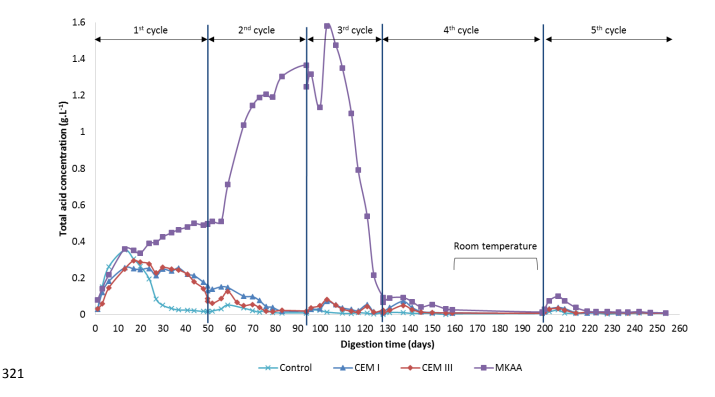
Figure 4 gives the total acid concentrations in the liquid phase of the bioreactors during the experiment. Among the VFA typically metabolized by microorganisms in anaerobic digestion (acetic, propionic, butyric, isobutyric and isovaleric acids) (Cibis et al., 2016), acetic acid was predominant far ahead of the others, as in several previous studies (Koenig and Dehn, 2016; Voegel et al., 2019b, 2016). The choice was thus made to present the total acid concentration for each material.

The first cycle showed a significant increase in the VFA concentration at the start of the digestion.Thereafter, two different types of behaviour occurred.

301 For the control bioreactors and the bioreactors containing CEM I and CEM III, the production reached a maximum during the second week, with concentrations of 0.35 g.L⁻¹, 0.26 g.L⁻¹ and 0.29 g.L⁻¹ 302 303 respectively. This initial increase corresponded to the acidogenesis step and was followed by the 304 slow consumption of the VFA marking the continuation of the digestion of the cattle manure. 305 Although the CEM I, CEM III and control bioreactors shared similar behaviour, it was observed that 306 the materials had an impact (i) on the quantity of VFA produced, the CEM I bioreactors producing 0.09 g.L⁻¹ less acid than the control ones, and (ii) on the kinetics of production: the decrease in the 307 308 total acid concentration was slower in the bioreactors containing materials and some VFA remained 309 at the end of the first cycle, highlighting a less efficient consumption of VFA. This induced a new, smaller increase in the VFA concentration at the start of the second cycle. Thereafter, the VFA werewell consumed and their concentration increased only very little at the start of subsequent cycles.

During the first 17 days, the evolution of the VFA concentration in the MKAA bioreactors followed the same trend as in the other bioreactors. Afterwards, the VFA concentration in the MKAA bioreactors presented a different evolution: instead of decreasing, it continued to increase during the first cycle and the accumulation of VFA accelerated with the start of the second cycle. The VFA concentration reached 1.58 g.L⁻¹ on day 103 and suddenly dropped to only 0.09 g.L⁻¹ at the end of the third cycle (day 128). This behaviour was strongly linked to the pH since the drop in the VFA concentration occurred just after the pH decrease.

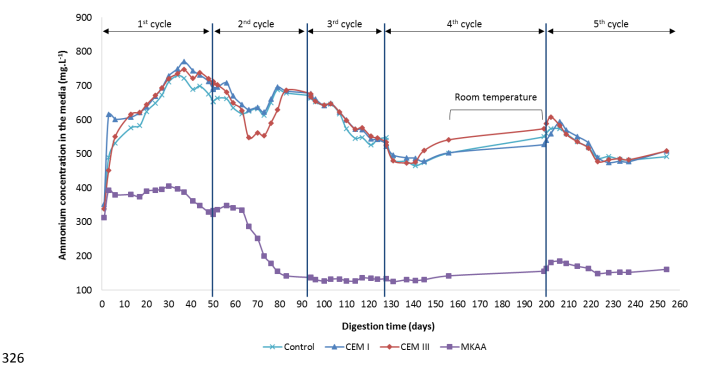
In the fourth and fifth cycles, all the bioreactors had low concentrations of VFA, showing their goodconsumption during the anaerobic digestion.



322 Figure 4: Evolution of the total acid concentration during the five cycles of anaerobic digestion in the bioreactors

323 3.1.3 Ammonium concentration

Figure 5 shows the evolution of the ammonium concentration during the anaerobic digestion of cattle manure in the bioreactors.





In anaerobic digestion, ammonium is present in the substrate and is also produced by the degradation of some proteins (Yenigün and Demirel, 2013), depending on the biowaste or the inoculum. In our case, it can be observed that the initial addition of inoculum and substrate induced a significant increase of the ammonium concentration while the following additions of substrate did not lead to such an increase. Thus, ammonium was provided by the inoculum and not by the biowaste. Once again, the singular behaviour of the bioreactor containing the MKAA can be noted.

In the bioreactors containing the cementitious materials and in the control bioreactors, the ammonium content reached about 750 mg.L⁻¹ after 30 days of digestion. After this initial increase, the ammonium concentrations tended to fall slowly through cycles, probably due to the consumption of ammonium for the growth of the microbial communities (Kuypers et al., 2018). The ammonium concentrations were much lower (about 400 mg.L⁻¹) in the bioreactors containing MKAA during the first cycle and dropped during the second cycle. The ammonium concentration remained around 130 mg.L⁻¹ for the rest of the experiment. This will be investigated further in the discussion section.

342 3.1.4 Release of ions in solution

Table 3 gives the Na⁺, K⁺, PO_4^{3-} and Mg^{2+} concentrations (mg.L⁻¹) in the liquid phase of the bioreactors at the start of each cycle. Mean values of the two duplicate bioreactors are presented with the standard deviations.

346 Table 3: Na^+ , K^+ , PO_4^{3-} and Mg^{2+} concentration (mg.L⁻¹) in the liquid phase of the bioreactors at the start of each cycle

	1	Start of the					
	mg.L ⁻¹	1 st cycle	2 nd cycle	3 rd cycle	4 th cycle	5 th cycle	
	Control	117.5 ± 13.4	167.0 ± 1.7	125.6 ± 0.9	149.8 ± 120.8	121.3 ± 0.9	
[Na ⁺]	CEM I	130.5 ± 13.4	195.5 ± 9.8	155.6 ± 2.2	154.4 ± 3.4	138.4 ± 2.7	
	CEM III	123.5 ± 0.7	197.6 ± 0.4	145.9 ± 1.6	140.4 ± 1.9	142.6 ± 1.2	
	ΜΚΑΑ	636.0 ± 77.4	2222.2 ± 80.6	1749.3 ± 61.2	1908.1 ± 2.2	2007.9 ± 13.1	
	Control	231.0 ± 19.8	379.1 ± 11.6	292.3 ± 9.9	390.7 ± 18.8	469.5 ± 33.7	
[K⁺]	CEM I	324.5 ± 30.4	655.6 ± 162.8	534.6 ± 16.0	681.5 ± 13.0	703.5 ± 7.3	
	CEM III	247.0 ± 1.4	436.4 ± 2.6	338.8 ± 3.0	425.1 ± 1.3	518.1 ± 6.5	
	ΜΚΑΑ	168.0 ± 18.4	100.0 ± 0.1	68.5 ± 0.8	119.3 ± 13.7	166.2 ± 24.2	
	Control	26.7 ± 9.0	6.2 ± 0.3	15.2 ± 0.5	11.7 ± 8.5	14.3 ± 6.6	
[PO4 ³⁻]	CEM I	25.1 ± 3.7	4.6 ± 0.9	13.9 ± 0.8	14.3 ± 2.6	16.6 ± 2.0	
	CEM III	22.5 ± 2.1	3.6 ± 1.1	11.9 ± 1.0	8.2 ± 5.7	16.7 ± 1.9	
	ΜΚΑΑ	21.0 ± 3.5	4.2 ± 2.7	21.6 ± 2.0	24.6 ± 1.0	26.7 ± 1.9	

	Control	2.2 ± 0.6	56.3 ± 3.5	32.6 ± 1.4	38.6 ± 4.2	38.3 ± 5.9
[Mg ²⁺]	CEM I	2.9 ± 1.3	56.7 ± 2.4	50.1 ± 7.8	55.6 ± 6.1	58.4 ± 3.5
	CEM III	2.8 ± 0.6	67.2 ± 2.5	65.3 ± 2.0	68.9 ± 4.8	65.7 ± 1.7
	ΜΚΑΑ	27.4 ± 2.9	45.3 ± 3.6	19.3 ± 1.6	19.9 ± 1.5	30.7 ± 2.6

347

In the bioreactors containing the cementitious materials and in the controls, the sodium concentration remained between 100 and 200 mg.L⁻¹ throughout the experiment. Because of its sodium activation, the MKAA released much more Na⁺: its concentration in the bioreactor liquid phase increased until day 60 and remained around 2000 mg.L⁻¹ thereafter.

352 The potassium concentration followed the same trend in the bioreactors containing the cementitious materials and in the control bioreactors: a first increase until day 35 and slight variations thereafter. 353 The potassium concentration in the control bioreactor was between 300 mg.L⁻¹ and 400 mg.L⁻¹ during 354 the experiment. Bioreactors containing the cementitious materials showed higher potassium 355 concentrations, between 300 mg.L⁻¹ and 700 mg.L⁻¹ in the bioreactor containing CEM I and between 356 250 mg.L⁻¹ and 500 mg.L⁻¹ in the bioreactors containing CEM III. The potassium concentration in the 357 bioreactors containing MKAA remained between 100 and 200 mg.L⁻¹ throughout the experiment. 358 359 Moreover, the K^{+} concentration for the MKAA bioreactors was found to be lower than that of the 360 control bioreactor during the whole experiment: the release of Na $^{+}$ could have led to an exchange of 361 the Na⁺ and the K⁺ ions in the material (O'Connor et al., 2010).

The phosphate concentrations followed a similar trend in the control bioreactors and in the bioreactors containing CEM I and CEM III: a first decrease from $25 \pm 2 \text{ mg.L}^{-1}$ to $5 \pm 1 \text{ mg.L}^{-1}$ and then an increase leading to stable concentrations between 8 and 15 mg.L⁻¹. In the bioreactors containing MKAA, the phosphate concentration was similar to that in the other bioreactors during two cycles, but rose significantly afterwards ($24 \pm 3 \text{ mg.L}^{-1}$). 367 Magnesium is provided by the inoculum and by the leaching of the materials. In the control bioreactors and in the bioreactors containing CEM I and CEM III, the magnesium concentration 368 started at about 3 mg.L⁻¹ and reached 60 ± 6 mg.L⁻¹ at the start of the second cycle. The magnesium 369 370 concentration was then stable at about 36 mg.L⁻¹, 55 mg.L⁻¹ and 67 mg.L⁻¹ in the control bioreactor, the bioreactor containing CEM I and the bioreactor containing CEM III, respectively. The initial 371 372 magnesium concentration (day 1) in the bioreactor containing MKAA was higher than in the others $(27.4 \pm 2.9 \text{ mg}.\text{L}^{-1})$ but the following increase was less significant than in the other bioreactors. The 373 374 magnesium concentration then remained between 20 and 30 mg. L^{-1} .

375 **3.2** Microstructural, chemical and mineralogical changes in the material samples

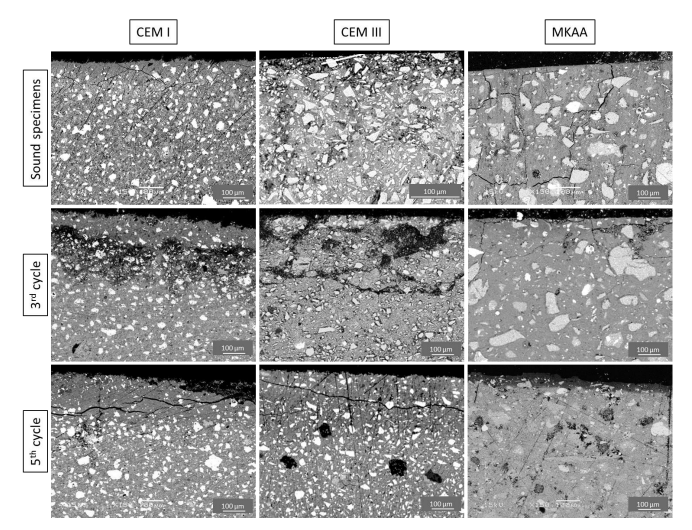
376 3.2.1 Microstructural changes

377 Figure 6 shows the images of the samples of materials obtained by SEM in back-scattered electron (BSE) mode before immersion and after the third and the fifth cycles. The surfaces in contact with 378 379 the fermenting biowaste are at the top of the images and the cores of the specimens at the bottom. 380 BSE images are characterized by a range of grey levels, which are proportional to the average atomic 381 weight of the analysed volume (or density). Thus, the grey level gives information about the different 382 phases, the heaviest elements being bright and the lightest elements being dark. The presence of a small quantity of solid (i.e. a high quantity of porosity) in an analysed area is characterized by a 383 384 darker contrast.

After the immersion, the cores of the specimens showed a high density of anhydrous residual grains (lighter grey grains). The outer layer (in contact with the biowaste) showed darker grey zones, corresponding to a decrease of mass density following the exposure to the fermenting biowaste. Most of the cracks observed were probably linked to shrinkage caused by sample preparation and to the more fragile nature of the external zones. The CEM I paste already showed a lower density zone after 3 cycles and a local dissolution was observed after the 5th cycle (top right of the image). This dissolution may have been associated with high local concentrations of acid.

After the 3rd cycle, the CEM III paste showed cracks, probably due to the way the sample was prepared. However, after the 5th cycle, no dissolution was observed but a zone of lower density appeared in the external part.

In contrast, the MKAA paste did not show any sign of deterioration after 3 cycles and only a thin layeron the external part presented a lower density after 5 cycles.



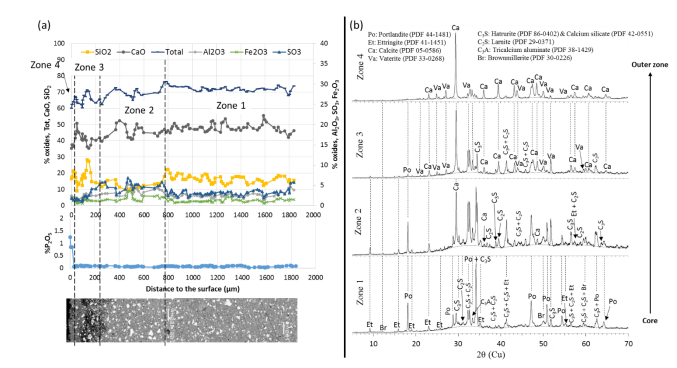
397

Figure 6: Observation of the paste by scanning electronic microscopy in back-scattered electron mode, before immersion and
after the third and fifth cycles. Surface in contact with the fermenting biowaste is at the top of each image and the core of

400 the specimen is at the bottom.

401 3.2.2 Chemical and mineralogical changes

402 Chemical composition profiles (EPMA), microscopic observations (SEM in BSE mode) and XRD 403 analyses were carried out to identify the structural, chemical and mineralogical changes after the 3rd 404 cycle for the CEM I (Figure 7) and CEM III pastes (Figure 8).



405 3.2.2.1 CEM I and CEM III



407 Figure 7: (a) Chemical composition profile, according to the distance to the surface, of a CEM I paste immersed in the liquid
408 phase in anaerobic digestion conditions (EPMA), and SEM observations of the polished sections in BSE mode. (b)
409 Mineralogical analyses of the same sample immersed for three consecutive cycles

Zonation, from the core to the outer layer, was identified for the sample and is represented in thechemical composition profile (zones 1 to 4):

Zone 1 corresponds to the sound zone. The SEM images show unaltered cement pastes where white anhydrous grains are surrounded by the hydrated paste. Pastes are predominantly composed of calcium and silicon, with minor amounts of aluminium, sulfur and iron. The mineralogical analyses show peaks of hydrated phases and anhydrous phases: portlandite, ettringite, C_2S , C_3S and C_4AF , and also merwinite C_3MS_2 for the CEM III paste.

Zone 2 is 400 µm thick and shows an enrichment in sulfur. The relative silica content drops in this zone whereas the aluminium content increases. The slight intensification of the ettringite peaks in the mineralogical patterns suggests that, as in the study by Voegel et al. (2016), the sulfur enrichment could come from the precipitation of secondary ettringite in this zone. This secondary ettringite precipitation is classically observed in transition zones of specimens exposed to leaching because of the diffusion of sulfates from the altered zone (Bertron et al., 2005a; Faucon et al., 1998). The total oxide content remains stable.

Zone 3, the thickness of which is about 400 μm in the CEM I paste and 180 μm in the CEM III paste, is
first marked by a decrease in the calcium and sulfur contents. The total oxides content decreases as a
result of the decrease of CaO. In this zone, the SEM image shows a very low density and a
heterogeneous layer. The mineralogical patterns show the precipitation of calcium carbonates,
calcite and vaterite, with a decrease of the peak intensities in the initial phases. This suggests
combined dissolution/precipitation phenomena.

Zone 4, about 30 μ m thick, is marked by a sudden increase of the phosphorus content - from 0 to 1.25% in the CEM I paste and from 0 to 1% in the CEM III paste. This zone also shows a decrease in the CaO, SiO₂ and total oxides contents. After the third cycle, the only remaining crystallized phases are the calcium carbonates, calcite and vaterite.

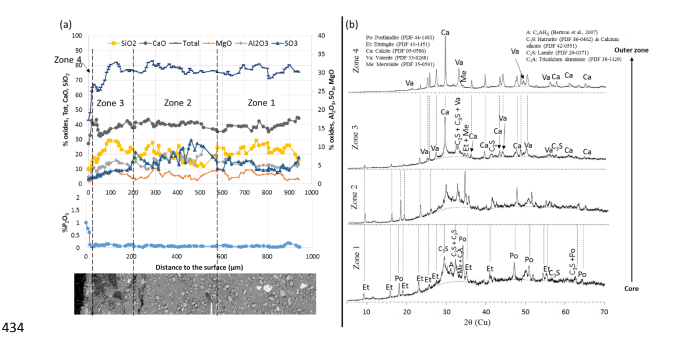


Figure 8: (a) Chemical composition profile, according to the distance to the surface, of a CEM III paste immersed in the liquid
 phase in anaerobic digestion conditions (EPMA), and SEM observations of the polished sections in BSE mode. (b)
 Mineralogical analyses of the same sample immersed for three consecutive cycles

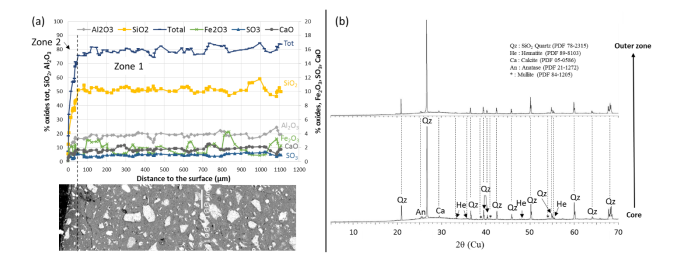
438 **3.2.2.2 MKAA**

The chemical composition of the MKAA specimen after the third cycle is presented in Figure 9. The

440 specimen did not seem to present degradation at any stage of the experiment, except on a thin outer

441 layer (50 μm after the third cycle), which was darker on the SEM image: the release of alkaline ions

- did not impact the major oxides over more than 50 μ m.
- The XRD analyses after the third cycle were identical to the initial XRD analyses of the sound paste.
- 444 The crystallized phases were mainly quartz but hematite, calcite, anatase and mullite were also
- 445 present. Their analyses through time did not highlight any mineralogical change between the core of
- the specimen and the outer zone. However, it should be noted that the mainly amorphous nature of
- the sound MKAA paste did not allow every mineralogical change to be identified.



448

449 Figure 9: (a) Chemical composition profile according to the distance to the surface (EPMA) of an MKAA paste immersed in 450 the liquid phase in anaerobic digestion conditions), and SEM observations of the polished sections in BSE mode. (b) 451 Mineralogical analyses of the same sample immersed for three consecutive cycles

Discussion 4 452

Characteristics of the digestate as a function of the material nature during anaerobic 453 4.1

digestion 454

Production of biogas and volatile fatty acids 455 4.1.1

Correlations of the biogas production, the VFA production and the pH of each type of bioreactor 456 457 were investigated to understand the effect of the nature of each sample on the anaerobic digestion process: the presence of materials had an impact on the digestion but it was limited in time. In the 458 short term, two types of behaviour were identified: 459

- In the bioreactors containing CEM I and CEM III, the VFA were not well consumed and 460 accumulated in the liquid environment during the first cycle but biogas was still produced. 461
- In the bioreactors containing MKAA, the material presence had a strong impact on the liquid 462 environment for three cycles since the VFA accumulated first and no biogas was produced 463 during two cycles.
 - 25

For the control bioreactors, the first cycle showed an acidification of the media from a pH of 8.38 up 465 466 to 7.12, corresponding to VFA production (hydrolysis and acidogenesis steps) (Han et al., 2019). It can be noticed that the initial value of the pH was much higher than the ones reported by Han et al. 467 (2019) and Yun et al. (2018), respectively 6.7 and 6.1. According to Yun et al. (2018), this low pH 468 469 comes from the partial acid peculiarity of the cattle manure. In the current study, since the cattle 470 manure was not fresh, the substrate acids were probably already consumed. Moreover, the inoculum 471 used was adapted to the substrate and was rich in bicarbonates, which explains the higher pH. A 472 maximum VFA concentration of 0.35 g.L-1 was reached and the VFA were then well consumed by the 473 microorganisms: after the thirteenth day, the total acid concentration decreased and biogas was 474 produced, meaning that the anaerobic digestion process was complete. This effective digestion of 475 the substrate continued through the following cycles with a pH that remained stable between 7.3 476 and 7.8, which is slightly lower but coherent with the pH values of the control check sample of Han et 477 al. (2019).

478 The bioreactors containing CEM I and CEM III pastes underwent a pH increase in the first three days 479 due to the alkaline nature of the materials, which released alkaline ions during the first cycle. 480 Afterwards, the pH decreased while the VFA concentration increased over a period of 10 days. The 481 increase in VFA concentration was slower and less significant in these bioreactors than in the control ones (maximum VFA concentration of 0.26 g.L⁻¹ and 0.29 g.L⁻¹ in the bioreactors containing CEM I and 482 483 CEM III respectively). From day 13, a slow consumption of the acids by the methanogenic archaea 484 occurred, highlighted by a decrease of the VFA concentration. The lack of efficiency in the VFA 485 production and consumption was associated with a low production of biogas during the first cycle. During the second cycle, the bioreactors containing cementitious materials reached a pH about 7.5, 486 487 which was similar to that of the control bioreactors and suitable for anaerobic digestion - especially 488 the methanogenic microbial activities, for which the optimal pH range is between 6.5 and 8.2 489 (Kothari et al., 2014). During this cycle, the bioreactors containing the cementitious materials 490 produced biogas in greater quantities than the control ones did, due to the undigested cattle manure

remaining after the first cycle. From the third cycle, these bioreactors behaved similarly to thecontrols and the presence of materials no longer disturbed the anaerobic digestion process.

493 The lower VFA production of the first cycle in the bioreactors containing cementitious materials was also observed by Voegel et al. (2016). In their study, CEM I pastes (water/cement = 0.4) were 494 495 immersed in an inoculated synthetic biowaste, representative of organic domestic waste, for a single 496 cycle of anaerobic digestion. The substrate induced significantly higher maximum acid concentrations than in the current study, about 2.90 g.L⁻¹ (43 mmol.L⁻¹) in the control bioreactors and a slightly lower 497 concentration, about 2.85 g.L⁻¹ (41 mmol.L⁻¹), in the bioreactors containing CEM I paste. Moreover, 498 499 the pH values between 7 and 8 found in the studies by Voegel et al. (2019b, 2016) were similar to 500 those encountered in the current study, whereas a slightly lower pH, around 7, was observed in the 501 liquid phase of a laboratory fermenter in a study by Koenig and Dehn (2016).

502 In the MKAA bioreactors, the pH was very high, between 9 and 9.5, during the first two cycles. This 503 can be explained by the strong release of alkalis that occurred during the first cycle (about 2 g/L of 504 Na⁺, Table 3). These high pH values were much higher than the ones tolerated by the methanogenic 505 archaea (optimal pH around 7 (Chae et al., 2010)). These are the most fragile microorganisms of the 506 anaerobic digestion process and are sensitive to pH variations (Chae et al., 2010), unsuitable pH 507 leading to very low biogas production. At the same time, the VFA accumulated in the medium and 508 accentuated the environment's aggressiveness. The formation of VFA in the medium showed that 509 the high pH did not disturb the first steps of the anaerobic digestion (hydrolysis, acidogenesis), 510 although the optimal operating pH of these bacteria is between 5.5 and 6.5 (Hagos et al., 2017; Han 511 et al., 2016; J. Kim et al., 2003; M. Kim et al., 2003; Yu and Fang, 2002). The third cycle showed a 512 decrease of the pH, probably due to the accumulation of the VFA in the medium, since most alkalis 513 were released during the first cycle. This decrease of pH allowed the methanogenic activities to take 514 place, and thus the rapid consumption of the VFA, leading to a significant production of biogas. The accumulation of methane produced during the third cycle indicated that not all the methane
potential was expressed earlier, due to the delay in the anaerobic digestion reactions.

According to the thermochemical equilibria (Reactions 1 and 2) associated with Henry's law at 35°C, the increase in pH during the first cycles in the bioreactors containing materials led to a reduction of the volatilization potential of CO_2 , which can explain the lower CO_2 amount in the gas phase during this period.

In all the bioreactors containing materials, when conditions suited to anaerobic digestion were well established, the process remained efficient over time. Moreover, the final total production of methane was equivalent in all the bioreactors (Figure 2), so the presence of the materials nevertheless allowed the expression of the methane potential of the entire amount of substrate. It can be assumed that the retarding effect of the MKAA on the biogas production is only expressed in the short term, as geopolymers rapidly lose their alkaline charge, which is mainly present in the liquid contained inside pores (Khan et al., 2020).

528 4.1.2 Ammonium production

A significant difference of ammonium concentration was observed between, on the one hand, the control bioreactors and the bioreactors containing the cementitious materials and, on the other hand, the bioreactors containing MKAA. This suggests a specific behaviour of the MKAA towards ammonium, which needs to be studied more thoroughly.

533 4.1.2.1 Control bioreactors and bioreactors containing CEM I and CEM III

The ammonium concentrations in the CEM I, CEM III and control bioreactors increased until a maximum of about 750 mg.L⁻¹ was reached on the 35th day. Similar concentrations were found in Voegel et al.'s study (2016) (800 mg.L⁻¹), where the presence of CEM I cement paste also did not influence the ammonium production. It could be expected that each addition of cattle manure would lead to an increase in the potential of hydrolysable nitrogen but the ammonium concentrations 539 actually decreased during the experiment, meaning that this element was mainly brought by the 540 inoculum. In Voegel et al. (2019b), the ammonium concentration did not increase with a new substrate addition either, since the measured concentrations were 988 mg.L⁻¹ (54.9 mmol.L⁻¹) in the 541 542 first cycle and 940 mg.L⁻¹ (52.24 mmol.L⁻¹) in the second. The decrease of the ammonium 543 concentration could be explained by several phenomena: nitrogen consumption by bacteria for 544 growth (Kuypers et al., 2018), stripping of the ammonium in the gas phase in the form of ammonia 545 (Limoli et al., 2016), and/or formation of precipitates containing ammonium (struvite, for example) 546 (Cerrillo et al., 2015; Escudero et al., 2015).

Using the chemical oxygen demand (COD) of the CH₄ production with the biomass production yields proposed by the Anaerobic Digestion Model n° 1 (ADM1) (Batstone et al., 2002), the quantity of biomass produced per cycle was estimated and its growth only required 61 mg of nitrogen, which is insufficient to justify the difference of 253 mg of nitrogen lost between the maximum concentrations (day 37) and the end of the fifth cycle.

The method used for gas chromatography did not allow identifying and measuring ammonia in the gas phase. Thus, the thermochemical equilibrium (Reaction 4) associated with Henry's law at 35°C and atmospheric pressure, were used and it was found that the amount of volatile ammonia passing into the gas phase did not exceed 0.15 mg for the whole experiment in these bioreactors.

$$NH_4^+ + H_2O \rightleftharpoons NH_3 + H_3O^+$$
: pKa = 9.25 (4)

Moreover, thermodynamic calculations using the geochemical code PHREEQC and considering the temperature, the pH and the concentrations in the solution (N as ammonium, P as phosphate, K, Mg, Cl, Ca, Na, inorganic carbon) (Table 3) were carried out to assess the possibility of struvite precipitation during the digestion. The MINTEQ database was used together with the struvite solubility value at 35°C; $pK_{sp \ struvite}$ = -13.20 (Bhuiyan et al., 2007). Figure 10 shows the variation of the struvite saturation index of the liquid phase of the bioreactors containing CEM I and MKAA with time. Precipitation conditions are met for a positive index, while a negative index corresponds to dissolution if struvite is present. The bioreactors containing CEM I were chosen to represent the bioreactors containing CEM I and CEM III and the control bioreactors. In these bioreactors, struvite precipitation was continuously possible during the experiment, with higher indices in the first cycle and values closer to 0 at the end of the experiment. The ammonium loss during the experiment could thus be explained by the precipitation of struvite.

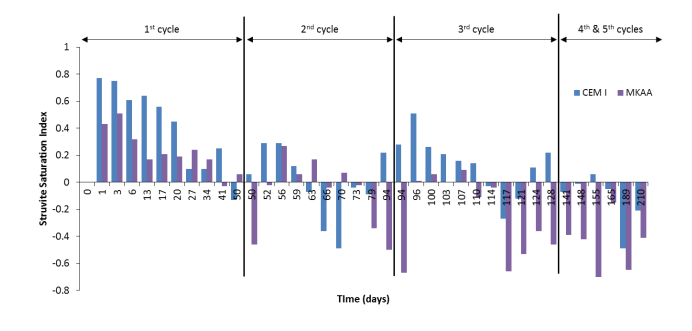


Figure 10: Variation with time of the Struvite Saturation Index of the liquid phase of the bioreactors containing CEM I and
MKAA

571 4.1.2.2 Bioreactors containing MKAA

In comparison with the other bioreactors, the bioreactors containing MKAA reached a lower 572 maximum ammonium concentration, of about 400 mg.L⁻¹, on day 30. At this stage of the process, the 573 pH was about 9.5. According to the pKa of the acid and conjugate base NH_4^+/NH_3 , i.e. 9.25 (Reaction 574 575 4), the ammonium was thus present in the form of both ammonium and ammonia. The volatile 576 ammonia form could have passed into the gas phase, leading to a weaker ammonium concentration 577 measured in the liquid phase of the MKAA bioreactors. Nevertheless, this hypothesis implies that a 578 fall in pH should lead to an increase in the ammonium concentration in the liquid medium, whereas 579 the concentration actually dropped quickly during the second cycle, at the same time as the pH

started decreasing. Moreover, the thermochemical equilibrium (Reaction 4) associated with Henry's law at 35°C and atmospheric pressure allowed the amount of ammonia in the gas phase to be calculated: this amount was very small throughout the experiment and did not exceed 0.5 mg of NH₃, which does not explain the lower ammonium concentration found in the liquid phase of the MKAA bioreactors.

585 The thermodynamic calculations using the geochemical code PHREEQC allowed us to follow the 586 variation of the Struvite Saturation Index of the liquid phase of the bioreactors containing MKAA 587 (Figure 10). The index was positive during the first cycle, but with lower values than in the other 588 bioreactors, while the ammonium concentration was lower in the bioreactors containing MKAA. 589 Moreover, whereas the acidification of the environment (from day 66) led to negative or neutral 590 struvite saturation indices, the ammonium concentration decreased in the bioreactors containing 591 MKAA. Thus, even if struvite can precipitate at the beginning of the experiment, the trend of the 592 ammonium concentration in the bioreactors containing MKAA remains unexplained, and is probably 593 the result of a combination of phenomena.

594 According to the literature, natural zeolites (hydrated aluminosilicates) have been successfully used 595 in wastewater treatment to remove ammonium (Luukkonen et al., 2016; Wang et al., 2011). The ion 596 exchange properties of the material are attributed to the charge-balancing cation (Na⁺ or K⁺) being replaced by another cation, such as K⁺, Ag⁺, NH₄⁺ or Pb²⁺ (O'Connor et al., 2010). As synthetic zeolites, 597 598 metakaolin-based alkali-activated geopolymers have been studied for ammonium removal by Luukkonen et al. (2016). In this study, the maximum capacity of metakaolin geopolymer to remove 599 NH_4^+ was 46% higher than that of natural zeolite, showing its potential interest in such an 600 601 application. The adsorption process is influenced by the pH: for pH higher than 9.25, NH4⁺ 602 deprotonates to NH_3 (Reaction 4) which is not favourable to adsorption. Actually, the ammonium 603 adsorption capacity seems to decrease even from $pH \ge 7$ (Marañón et al., 2006).

604 According to the former results, it is thus possible that ammonium was adsorbed on the surface of 605 the MKAA pastes through an exchange mechanism between a cation of the sound paste and the NH₄⁺ 606 cation in the liquid phase. The liquid medium analyses showed high concentrations of Na⁺ (2000 mg.L⁻¹ on average), this element being leached from the MKAA matrix. The drop of ammonium 607 608 concentration during the second cycle might be explained by the sudden drop of pH (due to the VFA 609 accumulation) leading to more suitable adsorption conditions. Then, although the ammonium 610 concentration appeared to be lower in the bioreactors containing MKAA, the production of 611 ammonium in the liquid medium due to digestion was probably similar in all the bioreactors, the 612 difference being adsorbed by the MKAA.

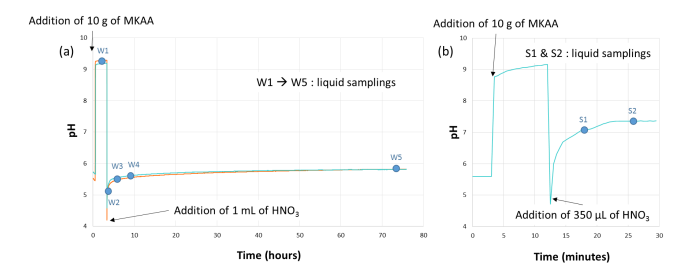
613 **4.2** Exchange phenomena between the MKAA and NH₄⁺

In order to investigate the assumption concerning the adsorption of NH_4^+ by the MKAA, based on the literature, an additional test was carried out to study the interaction between the MKAA and the NH_4^+ cation. The test design was adapted from the experimental protocols implemented by Lin et al. (2013) and Luukkonen et al. (2016).

This experiment aimed (i) to investigate the exchange phenomena between the MKAA and NH₄⁺ for different pH values and (ii) to understand how the ammonium cation interacted with the material. To this end, MKAA powder was immersed in an ammonium nitrate solution and the ammonium concentration was monitored for various pH values. The remaining MKAA powder was also analysed.

622 4.2.1 Experimental protocol

An 800 mg.L⁻¹ (0.0444 mol.L⁻¹) ammonium nitrate solution was prepared with demineralized water (according to the ammonium concentration found in the control bioreactors) and a cylindrical specimen of MKAA paste (75 mm high and 35 mm in diameter) was ground to 80 μm using a disc mill. 627 500 mL of solution was poured into a glass reactor thermostatically controlled at 35°C with magnetic stirring (Annexe A). The pH value was measured continuously and recorded every 15 seconds. After 628 629 about 30 minutes (the time for the temperature of the ammonium nitrate solution to reach 35°C), 10 g of MKAA powder was poured into the reactor. After 3 hours, 1 mL of concentrated HNO₃ acid (69%) 630 631 was added in order to lower the pH. Samples of the liquid were then taken at different stages of the 632 experiment to be analysed by Ion Chromatography (Thermo Electron ICS 300, CS16 column). The pH 633 evolution and the distribution of the liquid samples in time (W1 to W5) are presented in Figure 11 634 (a). The experiment was performed in duplicate (reactor 1 and reactor 2). In order to reach another 635 final pH, a supplementary experiment was run with the addition of $350 \ \mu L$ of HNO₃ only. The pH 636 evolution and the distribution of the liquid samples S1 & S2 in time are presented in Figure 11 (b).



637

638 Figure 11: pH evolution according to time and distribution in time of the liquid sampling: (a) in the two duplicate reactors for **639** 1 mL of added HNO₃ and (b) in the solution of the supplementary experiment for 350 μ L of added HNO₃

After the end of the main experiment, the solution was centrifuged in order to collect the MKAA powder. It was then immersed in isopropanol for 15 minutes to dry by solvent exchange (Scrivener et al., 2016; Snellings et al., 2018). After another centrifugation, the powder was collected and placed in a vacuum desiccator for 1 hour to finish drying. This powder and a control MKAA powder were analysed in several ways: thermogravimetric analysis (NETZSCH STA 449F3); ²⁷Al, ²⁹Si and ¹H nuclear magnetic resonance (NMR) (Brucker Avance 4000 III HD MAS); and infrared spectrometry (FTIR) 646 (Perkin-Elmer Spectrum). No significant differences were observed in the ²⁷Al and ²⁹Si NMR spectra 647 between the control MKAA powder and the powder from the experiment (called MKAA – NH_4^+ 648 powder), and these results are not presented.

649 4.2.2 Liquid analysis

650 The ammonium concentration $[NH_4^{\dagger}]$ was measured by Ion Chromatography and the results from 651 reactor 1 and the supplementary experiment are presented in Table 4. These analyses highlighted an 652 exchange mechanism between the MKAA and the ammonium solution since the measured concentrations were different from the initial ammonium concentration of the solution (800 mg. L^{-1}). 653 654 When the MKAA powder was added, the pH increased to 9.29 and the ammonium concentration decreased from 800 mg.L⁻¹ to 437.9 mg.L⁻¹. The pH decrease from 9.29 to 7.06 led to a decrease of 655 the ammonium concentration to 351.1 mg.L⁻¹. From then on, the ammonium concentration 656 increased when the pH was decreased, but still remained below 800 mg.L⁻¹. 657

658 Table 4: Ammonium concentration of the liquid withdrawals according to the pH, in mg.L⁻¹

Samples	W1	S2	S1	W5	W4	W3	W2
Decreasing pH	9.29	7.36	7.06	5.82	5.52	5.49	5.29
[NH4 ⁺]	437.9	357.9	351.1	520.4	525.3	530.1	530.0

659 4.2.3 MKAA powder analyses

660 Differential Thermo Gravimetry (DTG), ¹H NMR and FTIR analyses of the control MKAA powder and

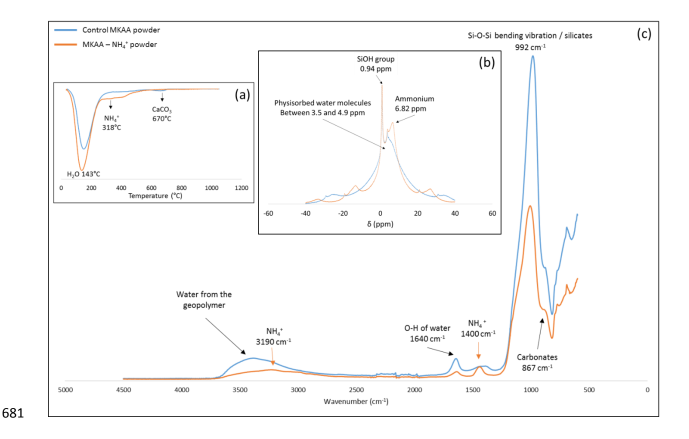
the powder from the reactor 1 are presented in Figure 12 (a), (b) and (c) respectively.

Figure 12 (a) shows that, despite the drying by solvent exchange, the MKAA – NH_4^+ powder contained more free water than the control powder (intense peak at 143°C). The DTG analysis differed slightly with a weak, broad peak centred on 318°C in the MKAA – NH_4^+ powder, which could be correlated with the thermal decomposition of the NH_4^+ cation identified in two different clays at 327°C and 666 330°C by Gautier et al. (2010) and Yariv (1985), respectively. Some calcium carbonates were 667 identified by a bump at 670°C (Wianglor et al., 2017) in the control MKAA powder.

Figure 12 (b) shows a fine, intense peak for both specimens at 0.94 ppm, which corresponds to the SiOH group usually found between 1 and 2.2 ppm in the literature (Hunger, 1996; O'Connor et al., 2010). All the peaks between 3.5 and 4.9 were associated with water and, more precisely for hydrated Na-form zeolites, with physisorbed water molecules (Hunger, 1996; O'Connor et al., 2010). Finally, a peak was detected in the MKAA – NH_4^+ powder only at 6.82 ppm and corresponded to the ammonium cation, whose signal appears between 6.5 and 7 ppm (Jacobs et al., 1993; O'Connor et al., 2010).

In addition to the peaks associated with water (3500 and 1640 cm⁻¹) (Ahmed et al., 2018; Aredes et al., 2015; Wang et al., 2005), the silicates (992 cm⁻¹) (O'Connor et al., 2010) and the carbonates (867 cm⁻¹), peaks centred on 3190 cm⁻¹ and 1400 cm⁻¹ can be seen in Figure 12 (c) and correspond to the ammonium cation (Mookherjee et al., 2005).

Thus, the analyses of the materials showed that ammonium from the solution was indeed exchangedtowards the MKAA.



682 Figure 12: (a) Differential thermogravimetry, (b) ¹H Nuclear Magnetic Resonance, and (c) Fourier Transform Infrared **683** Spectroscopy analyses of the control MKAA powder and the powder from the experiment (MKAA – NH_4^+). The main peaks **684** are identified.

685 4.2.4 Exchange phenomena conclusions

686 In light of the previous results, it can be concluded that exchange phenomena between MKAA and NH₄⁺ do exist. More precisely, the liquid analysis showed that the efficiency of this exchange 687 depended on the pH, with a much greater transfer of ammonium to the MKAA when the pH was 688 689 around 7. Moreover, the analyses of the MKAA powders confirmed the presence of the ammonium 690 cation in the material, but without changing the Si-Al structure of the geopolymer. The nature of this 691 exchange phenomenon would therefore be adsorption. Only 10 g of material was inserted in 500 mL of a 800 mg.L⁻¹ nitrate ammonium solution, whereas the cylinders immersed in the bioreactors are 692 about 100 g for 800 mL of solution (ammonium in similar concentration, about 750 mg.L⁻¹). Even 693 though the exchange surface is much higher in the case of a powdered material, the monolithic 694

- 695 samples of MKAA constituted, in theory, a sufficient source of exchange that could explain the lower
- ammonium concentrations found in the bioreactors containing MKAA.
- 697 **4.3 Deterioration of the material samples**

698 4.3.1 CEM I and CEM III

The chemical composition analysed in this study can be compared with results from previous studies. Voegel (2017) studied the biodeterioration phenomena of cementitious paste (water/cement = 0.40) in the submerged part of a digester with a laboratory test using typical food biowaste (solid/liquid ratio=224 cm².L⁻¹). After chemical and mineralogical tests, a zonation was established and phenomena were highlighted.

The chemical profiles were correlated with mineralogical changes and the CEM I and CEM III paste zonations were as follows:

- A sound zone at depth, corresponding to Zone 1, composed of the crystalline phases
 commonly found in the CEM I and CEM III matrices.
- A decrease in the CaO content and an enrichment in sulfur in Zone 2, probably correlated
 with the precipitation of secondary ettringite.
- Variations in the CaO content in Zone 3 and a slight decalcification, together with the
 precipitation of CaCO₃.

- A peak of phosphorus and decalcification of the matrix in Zone 4

From a mineralogical point of view, the X-Ray patterns obtained in the different zones of the CEM I and CEM III pastes after the third cycle showed leaching and carbonation phenomena: the initial phases (ettringite, portlandite, brownmillerite, C₃S and C₂S) were dissolved and calcium carbonates (calcite and vaterite) precipitated. These main findings are in accordance with the results obtained by Bertron et al. (2017), where the authors sum up results on CEM I and CEM III from Voegel's study (2017). In addition, a fifth zone was found in Voegel's studies, which presents almost complete decalcification and the complete dissolution of hydrates and anhydrous phases, leading to an amorphous gel of silicon and aluminium.

The values from Table 2 allowed the inorganic carbon contents (H_2CO_3 , HCO_3^- and CO_3^{-2} forms) in the 721 liquid medium to be calculated at the end of each cycle. Most of the values were between 1000 and 722 2000 mg.L⁻¹ in the control bioreactors and in the bioreactors containing CEM I and CEM III. They were 723 much higher than the values found by Voegel et al. (2019b, 2016) (about 140 mg.L⁻¹ or 2.3 mmol.L⁻¹) 724 725 who used a different digestate that initially contained less dissolved CO₂, but were consistent with the concentrations of between 2000 and 3000 mg.L⁻¹ found in industrial digesters (McCarty, 1964). 726 727 Moreover, the inorganic carbon content being strongly dependent on the pH and the rate of CO₂ in 728 the gas phase, higher concentrations could be encountered in the medium during the digestion cycle, when the pH is higher or when the proportion of methane is smaller and that of CO₂ is larger. These 729 730 high values accounted for the carbonation of the CEM I and CEM III samples. The presence of vaterite 731 in the external part could be explained by the presence of highly decalcified C-S-H with low Ca/Si 732 ratio (Morandeau et al., 2014). However, it could also be linked to the lower pH of the external zone 733 since the crystallization of vaterite is predominant for pH<10 (Oral and Ercan, 2018; Tai and Chen, 734 1998). Moreover the experiment temperature of 35°C (close to 30°C) promotes the precipitation of 735 both calcite and vaterite (Girou and Roques, 1971; Ševčík et al., 2015; Stepkowska et al., 2003; Wray 736 and Daniels, 1956).

737 In the study by Koenig and Dehn (2016), in addition to calcium carbonates, struvite and quartz were 738 detected in the surfaces of samples immersed in the liquid phase in a pilot scale fermenter. In the 739 same study, vivianite, calcium-aluminium-iron-chloride-silicates, calcium-aluminium-iron oxides, 740 ettringite and portlandite phases were measured on the near-surface area of samples immersed in

38

the laboratory. The presence of the initial phases portlandite and ettringite after 173 days ofimmersion demonstrated low leaching of the materials.

743 After the third cycle, the same phenomena as in Voegel's study (2017) were detected for the CEM I 744 and CEM III samples, with equivalent degraded depths between the two materials. However, it can 745 be noted that, even with a longer exposure time in the cattle manure, the degradation of the 746 samples was less since the altered zone seemed to stop at zone 4, where the phosphorous peak was 747 found. This may have been due to (i) the different substrate used, which led to lower concentration 748 of aggressive agents, and/or (ii) the lower water/cement ratio of our matrices, leading to a reduction 749 of the porosity, influencing the transport mechanisms and reducing the specific surface area (Koenig 750 and Dehn, 2015).

751 4.3.2 MKAA

752 Even though the amorphous part of the material could not be characterized, the MKAA samples 753 seemed to show very stable behaviour with a small apparent degraded depth. However, the MKAA 754 released a large amount of alkalis at the start of the immersion, leading to a greater buffering 755 capacity compared to other binders: more than 90% of the water introduced was "free water" 756 (Pouhet et al., 2019) and contained a large amount of alkaline ions, which are easily released from 757 the paste via the porous network. This behaviour could be linked to the observations of Khan et al. 758 (2020), who studied the degradation of a fly ash (FA) based geopolymer mortar in a naturally 759 aggressive sewer environment. A mixture of sodium hydroxide solution and sodium silicate solution 760 was used as the activator. Although the geopolymer showed less degradation in terms of mass loss, 761 dry bulk density, compressive strength and surface disintegration compared to a Sulfate Resistant 762 (SR) ordinary Portland cement mortar, the geopolymer was also found to have a much greater 763 neutralization depth with a loss of alkalinity. This shows a greater penetration of the aggressive 764 agents and could be a concern for steel reinforced concrete. Moreover, the same geopolymer was 765 compared to an alkali-activated slag mortar (AASm) in another study (Khan et al., 2018) (same

39

766 activator) and the FA geopolymer showed much higher overall matrix deterioration than the AASm. 767 Nevertheless, the very different chemistries of the FA-based and AAS mortars do not allow a 768 qualitative comparison with the current study. In the study by Grengg et al. (2020), a metakaolin 769 similar to the one used in this study, which was quartz-rich, rust brown iron-rich (GP-K2) with 770 potassium silicate, was placed in a power main outlet storage basin heavily affected by MID in 771 Austria, together with a geopolymer made with high purity white metakaolin (GP-K1), a calcium 772 aluminate cement and a CEM I based material. Among these materials, after 18 months, the degree 773 of surface cracking and the estimated degradation rates were the lowest for the GP-K2 but highest 774 for the GP-K1. Thus, as pointed out by Grengg et al. (2020) and Khan et al. (2018), depending on the 775 design and the type of geopolymer, its resistance to bacteriological attacks in acid medium can vary 776 widely.

777 **5 Conclusion**

This study presented the exposure of three types of binder paste specimens to five consecutive anaerobic digestion cycles of cattle manure in laboratory conditions, in an 8 months experiment. A high solid/liquid ratio used for this study was chosen to exacerbate and evaluate the local interaction mechanisms between the different materials and the anaerobic digestion medium.

782 The presence of the alkaline materials disrupted the efficiency of the anaerobic digestion process at 783 the local scale by slowing down the production and consumption of VFA and decreasing the 784 production of biogas. Whereas the Portland based materials, CEM I and CEM III pastes, impacted the 785 anaerobic digestion process for only one cycle, the alkali-activated materials (MKAA) released larger 786 amounts of alkalis at the start of the immersion, inducing a much higher pH, the VFA accumulation, 787 and almost no biogas production during the first two cycles. After 5 cycles, the final biogas 788 productions were equivalent for all the bioreactors and the anaerobic digestion process worked 789 under normal, efficient conditions. Thus, given the moderately aggressive conditions induced by the 790 liquid phase of anaerobic digestion, all materials can therefore be considered for construction791 application in such environment.

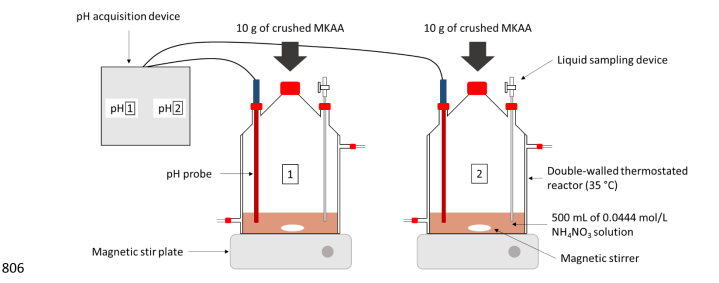
In terms of deterioration pattern the CEM I and CEM III biodeterioration were expressed by a combination of dissolution of the initial phases and decalcification on the surface, and precipitation of CaCO₃, in accordance with literature recent results. The original study of an alkali-activated geopolymer in an anaerobic digestion medium has revealed a very good behaviour of the material with very low degradations. In addition, the material was found to have interesting ammonium adsorption properties at the pH of the anaerobic digestion, significantly reducing the ammonium concentration in the liquid medium.

799 6 Acknowledgements

The authors gratefully thank the French National Research Agency (ANR) for funding the project BIBENdOM – ANR – 16 – CE22 – 001 DS0602. The authors also thank Evrard Mengelle, Simon Dubos, Mansour Bounouba and Chantha Kim for the pilot's design and the analytical support of the bioreactors.

804 Annexe

805



807 Annexe A: Experimental setup for evaluating the interaction between the MKAA and the NH_4^+ cation

808 **References**

- AFNOR, 2016. NF EN 196-1. Methods of testing cement Part 1: Determination of strength. Paris,
 France.
- AFNOR, 2014. NF EN 206/CN. Concrete Specification, performance, production and conformity National addition to the standard NF EN 206. Paris, France.
- AFNOR, 2012a. NF P 18-513 Addition for concrete Metakaolin Specifications and conformity
 criteria. Paris, France.
- AFNOR, 2012b. NF EN 197-1. Cement Part 1 : composition, specifications and conformity criteria for
 common cements. Paris, France.
- Ahmed, A., Chaker, Y., Belarbi, E.H., Abbas, O., Chotard, J.N., Abassi, H.B., Van Nhien, A.N., El Hadri,
 M., Bresson, S., 2018. XRD and ATR/FTIR investigations of various montmorillonite clays
 modified by monocationic and dicationic imidazolium ionic liquids. J. Mol. Struct. 1173, 653–
 664. https://doi.org/10.1016/j.molstruc.2018.07.039
- Aredes, F.G.M., Campos, T.M.B., Machado, J.P.B., Sakane, K.K., Thim, G.P., Brunelli, D.D., 2015. Effect
 of cure temperature on the formation of metakaolinite-based geopolymer. Ceram. Int. 41,
 7302–7311. https://doi.org/10.1016/j.ceramint.2015.02.022
- Batstone, D.J., Keller, J., Angelidaki, I., Kalyuzhnyi, S.V., Pavlostathis, S.G., Rozzi, A., Sanders, W.T.M.,
 Siegrist, H., Vavilin, V.A., 2002. The IWA Anaerobic Digestion Model No 1 (ADM1). Water Sci.
 Technol. 45, 65–73.
- Bertron, A., Duchesne, J., Escadeillas, G., 2005a. Accelerated tests of hardened cement pastes
 alteration by organic acids: analysis of the pH effect. Cem. Concr. Res. 35, 155–166.
 https://doi.org/10.1016/j.cemconres.2004.09.009
- Bertron, A., Duchesne, J., Escadeillas, G., 2005b. Attack of cement pastes exposed to organic acids in
 manure.
 Cem.
 Concr.
 Compos.
 27,
 898–909.
 https://doi.org/10.1016/j.cemconcomp.2005.06.003
- Bertron, A., Escadeillas, G., de Parseval, P., Duchesne, J., 2009. Processing of electron microprobe
 data from the analysis of altered cementitious materials. Cem. Concr. Res. 39, 929–935.
 https://doi.org/10.1016/j.cemconres.2009.06.011
- Bertron, A., Peyre Lavigne, M., Patapy, C., Erable, B., 2017. Biodeterioration of concrete in agricultural, agro-food and biogas plants: state of the art and challenges. RILEM Tech. Lett. 2, 838
 83–89. https://doi.org/10.21809/rilemtechlett.2017.42

- Bhuiyan, M.I.H., Mavinic, D.S., Beckie, R.D., 2007. A Solubility and Thermodynamic Study of Struvite.
 Environ. Technol. 28, 1015–1026. https://doi.org/10.1080/09593332808618857
- Braun, R., 2007. Anaerobic digestion: a multi-faceted process for energy, environmental management and rural development, in: Improvement of Crop Plants for Industrial End Uses.
 Springer, Dordrecht, pp. 335–416. https://doi.org/10.1007/978-1-4020-5486-0_13
- Cerrillo, M., Palatsi, J., Comas, J., Vicens, J., Bonmatí, A., 2015. Struvite precipitation as a technology
 to be integrated in a manure anaerobic digestion treatment plant removal efficiency,
 crystal characterization and agricultural assessment. J. Chem. Technol. Biotechnol. 90, 1135–
 1143. https://doi.org/10.1002/jctb.4459
- Chae, K.-J., Choi, M.-J., Kim, K.-Y., Ajayi, F.F., Chang, I.-S., Kim, I.S., 2010. Selective inhibition of methanogens for the improvement of biohydrogen production in microbial electrolysis cells.
 Int. J. Hydrog. Energy, 3rd Asian Bio Hydrogen Symposium 35, 13379–13386.
 https://doi.org/10.1016/j.ijhydene.2009.11.114
- Chandra, R., Takeuchi, H., Hasegawa, T., 2012. Methane production from lignocellulosic agricultural
 crop wastes: A review in context to second generation of biofuel production. Renew. Sustain.
 Energy Rev. 16, 1462–1476. https://doi.org/10.1016/j.rser.2011.11.035
- Chen, J., Yun, S., Shi, J., Wang, Z., Abbas, Y., Wang, K., Han, F., Jia, B., Xu, H., Xing, T., Li, B., 2020. Role
 of biomass-derived carbon-based composite accelerants in enhanced anaerobic digestion:
 Focusing on biogas yield, fertilizer utilization, and density functional theory calculations.
 Bioresour. Technol. 307, 123204. https://doi.org/10.1016/j.biortech.2020.123204
- Cibis, K.G., Gneipel, A., König, H., 2016. Isolation of acetic, propionic and butyric acid-forming
 bacteria from biogas plants. J. Biotechnol. 220, 51–63.
 https://doi.org/10.1016/j.jbiotec.2016.01.008
- CIMbéton, 2009. Guide de prescription des ciments pour des constructions durables. Cas des bétons
 coulés en place, Collection Technique CIMbéton.
- Brugă, B., Ukrainczyk, N., Weise, K., Koenders, E., Lackner, S., 2018. Interaction between wastewater
 microorganisms and geopolymer or cementitious materials: Biofilm characterization and
 deterioration characteristics of mortars. Int. Biodeterior. Biodegrad. 134, 58–67.
 https://doi.org/10.1016/j.ibiod.2018.08.005
- Buan, P., Yan, C., Zhou, W., Luo, W., Shen, C., 2015. An investigation of the microstructure and durability of a fluidized bed fly ash-metakaolin geopolymer after heat and acid exposure.
 Mater. Des. 74, 125–137. https://doi.org/10.1016/j.matdes.2015.03.009
- Elakneswaran, Y., Owaki, E., Miyahara, S., Ogino, M., Maruya, T., Nawa, T., 2016. Hydration study of
 slag-blended cement based on thermodynamic considerations. Constr. Build. Mater. 124,
 615–625. https://doi.org/10.1016/j.conbuildmat.2016.07.138
- Escudero, A., Blanco, F., Lacalle, A., Pinto, M., 2015. Struvite precipitation for ammonium removal
 from anaerobically treated effluents. J. Environ. Chem. Eng. 3, 413–419.
 https://doi.org/10.1016/j.jece.2015.01.004
- Evans, G.M., Furlong, J.C., 2003. Environmental Biotechnology Theory and Application. John Wiley &
 Sons.
- Faucon, P., Adenot, F., Jacquinot, J.F., Petit, J.C., Cabrillac, R., Jorda, M., 1998. Long-term behaviour
 of cement pastes used for nuclear waste disposal: review of physico-chemical mechanisms of
 water degradation. Cem. Concr. Res. 28, 847–857. https://doi.org/10.1016/S00088846(98)00053-2
- Fehrenbach, H., Giegrich, J., Reinhardt, G., Sayer, U., Gretz, M., Lanje, K., Schmitz, J., 2008. Kriterien
 einer nachhaltigen Bioenergienutzung im globalen Ma\s sstab. UBA-Forschungsbericht 206,
 41–112.
- Fisgativa, H., Tremier, A., Dabert, P., 2016. Characterizing the variability of food waste quality: A need
 for efficient valorisation through anaerobic digestion. Waste Manag. 50, 264–274.
 https://doi.org/10.1016/j.wasman.2016.01.041

- Flemming, H.-C., Wingender, J., Szewzyk, U., Steinberg, P., Rice, S.A., Kjelleberg, S., 2016. Biofilms: an
 emergent form of bacterial life. Nat. Rev. Microbiol. 14, 563–575.
 https://doi.org/10.1038/nrmicro.2016.94
- Frigon, J.-C., Guiot, S.R., 2010. Biomethane production from starch and lignocellulosic crops: a
 comparative review. Biofuels Bioprod. Biorefining 4, 447–458.
 https://doi.org/10.1002/bbb.229
- Gautier, M., Muller, F., Le Forestier, L., Beny, J.-M., Guegan, R., 2010. NH4-smectite: Characterization,
 hydration properties and hydro mechanical behaviour. Appl. Clay Sci. 49, 247–254.
 https://doi.org/10.1016/j.clay.2010.05.013
- Girou, A., Roques, H., 1971. Etude des cinétiques de précipitation des carbonates de calcium. Bull.
 Assoc. Géographes Fr. 48, 227–233. https://doi.org/10.3406/bagf.1971.7651
- Grengg, C., Ukrainczyk, N., Koraimann, G., Mueller, B., Dietzel, M., Mittermayr, F., 2020. Long-term in situ performance of geopolymer, calcium aluminate and Portland cement-based materials exposed to microbially induced acid corrosion. Cem. Concr. Res. 131, 106034. https://doi.org/10.1016/j.cemconres.2020.106034
- 904 Gruyaert, E., Van den Heede, P., Maes, M., De Belie, N., 2012. Investigation of the influence of blastfurnace slag on the resistance of concrete against organic acid or sulphate attack by means 905 906 accelerated degradation tests. Cem. Concr. of Res. 42, 173-185. 907 https://doi.org/10.1016/j.cemconres.2011.09.009
- Hagos, K., Zong, J., Li, D., Liu, C., Lu, X., 2017. Anaerobic co-digestion process for biogas production:
 Progress, challenges and perspectives. Renew. Sustain. Energy Rev. 76, 1485–1496.
 https://doi.org/10.1016/j.rser.2016.11.184
- Han, F., Yun, S., Zhang, C., Xu, H., Wang, Z., 2019. Steel slag as accelerant in anaerobic digestion for
 nonhazardous treatment and digestate fertilizer utilization. Bioresour. Technol. 282, 331–
 338. https://doi.org/10.1016/j.biortech.2019.03.029
- Han, G., Shin, S.G., Lee, J., Lee, C., Jo, M., Hwang, S., 2016. Mesophilic Acidogenesis of Food Waste Recycling Wastewater: Effects of Hydraulic Retention Time, pH, and Temperature. Appl.
 Biochem. Biotechnol. 180, 980–999. https://doi.org/10.1007/s12010-016-2147-z
- 917 Holliger, C., Alves, M., Andrade, D., Angelidaki, I., Astals, S., Baier, U., Bougrier, C., Buffière, P., 918 Carballa, M., de Wilde, V., Ebertseder, F., Fernández, B., Ficara, E., Fotidis, I., Frigon, J.-C., de 919 Laclos, H.F., Ghasimi, D.S.M., Hack, G., Hartel, M., Heerenklage, J., Horvath, I.S., Jenicek, P., 920 Koch, K., Krautwald, J., Lizasoain, J., Liu, J., Mosberger, L., Nistor, M., Oechsner, H., Oliveira, 921 J.V., Paterson, M., Pauss, A., Pommier, S., Porqueddu, I., Raposo, F., Ribeiro, T., Rüsch Pfund, 922 F., Strömberg, S., Torrijos, M., van Eekert, M., van Lier, J., Wedwitschka, H., Wierinck, I., 923 2016. Towards a standardization of biomethane potential tests. Water Sci. Technol. 74, 924 2515-2522. https://doi.org/10.2166/wst.2016.336
- Holm-Nielsen, J.B., Al Seadi, T., Oleskowicz-Popiel, P., 2009. The future of anaerobic digestion and
 biogas utilization. Bioresour. Technol., OECD Workshop: Livestock Waste Treatment Systems
 of the Future: A Challenge to Environmental Quality, Food Safety, and Sustainability 100,
 5478–5484. https://doi.org/10.1016/j.biortech.2008.12.046
- Huang, X., Yun, S., Zhu, J., Du, T., Zhang, C., Li, X., 2016. Mesophilic anaerobic co-digestion of aloe
 peel waste with dairy manure in the batch digester: Focusing on mixing ratios and digestate
 stability. Bioresour. Technol. 218, 62–68. https://doi.org/10.1016/j.biortech.2016.06.070
- Hunger, M., 1996. Multinuclear solid-state NMR studies of acidic and non-acidic hydroxyl protons in
 zeolites. Solid State Nucl. Magn. Reson. 6, 1–29. https://doi.org/10.1016/09262040(95)01201-X
- Jacobs, W.P.J.H., de Haan, J.W., van de Ven, L.J.M., van Santen, R.A., 1993. Interaction of ammonia
 with Broensted acid sites in different cages of zeolite Y as studied by proton MAS NMR. J.
 Phys. Chem. 97, 10394–10402. https://doi.org/10.1021/j100142a022
- Juenger, M.C.G., Snellings, R., Bernal, S.A., 2019. Supplementary cementitious materials: New
 sources, characterization, and performance insights. Cem. Concr. Res. 122, 257–273.
 https://doi.org/10.1016/j.cemconres.2019.05.008

- Khan, H.A., Castel, A., Khan, M.S.H., 2020. Corrosion investigation of fly ash based geopolymer
 mortar in natural sewer environment and sulphuric acid solution. Corros. Sci. 108586.
 https://doi.org/10.1016/j.corsci.2020.108586
- Khan, H.A., Castel, A., Khan, M.S.H., 2017. Performance Evaluation of Geopolymer against In Situ
 Aggressive Sewer Environment, in: Concrete 2017. Presented at the Concrete 2017, Adelaïde,
 Australia.
- Khan, H.A., Khan, M.S.H., Castel, A., Sunarho, J., 2018. Deterioration of alkali-activated mortars
 exposed to natural aggressive sewer environment. Constr. Build. Mater. 186, 577–597.
 https://doi.org/10.1016/j.conbuildmat.2018.07.137
- Kim, J., Park, C., Kim, T.-H., Lee, M., Kim, S., Kim, S.-W., Lee, J., 2003. Effects of various pretreatments
 for enhanced anaerobic digestion with waste activated sludge. J. Biosci. Bioeng. 95, 271–275.
 https://doi.org/10.1016/S1389-1723(03)80028-2
- Kim, M., Gomec, C.Y., Ahn, Y., Speece, R.E., 2003. Hydrolysis and acidogenesis of particulate organic
 material in mesophilic and thermophilic anaerobic digestion. Environ. Technol. 24, 1183–
 1190. https://doi.org/10.1080/09593330309385659
- Koenig, A., Dehn, F., 2016. Biogenic acid attack on concretes in biogas plants. Biosyst. Eng. 147, 226–
 237. https://doi.org/10.1016/j.biosystemseng.2016.03.007
- Koenig, A., Dehn, F., 2015. Acid Resistance of Ultra High-Performance Concrete (UHPC), in: Sobolev,
 K., Shah, S.P. (Eds.), Nanotechnology in Construction. Springer International Publishing,
 Cham, pp. 317–323. https://doi.org/10.1007/978-3-319-17088-6_41
- Kothari, R., Pandey, A.K., Kumar, S., Tyagi, V.V., Tyagi, S.K., 2014. Different aspects of dry anaerobic
 digestion for bio-energy: An overview. Renew. Sustain. Energy Rev. 39, 174–195.
 https://doi.org/10.1016/j.rser.2014.07.011
- Kuypers, M.M.M., Marchant, H.K., Kartal, B., 2018. The microbial nitrogen-cycling network. Nat. Rev.
 Microbiol. 16, 263–276. https://doi.org/10.1038/nrmicro.2018.9
- Lastella, G., Testa, C., Cornacchia, G., Notornicola, M., Voltasio, F., Sharma, V.K., 2002. Anaerobic
 digestion of semi-solid organic waste: biogas production and its purification. Energy Convers.
 Manag. 43, 63–75. https://doi.org/10.1016/S0196-8904(01)00011-5
- Lesteur, M., Bellon-Maurel, V., Gonzalez, C., Latrille, E., Roger, J.M., Junqua, G., Steyer, J.P., 2010.
 Alternative methods for determining anaerobic biodegradability: A review. Process Biochem.
 45, 431–440. https://doi.org/10.1016/j.procbio.2009.11.018
- Li, K., Liu, R., Sun, C., 2015. Comparison of anaerobic digestion characteristics and kinetics of four
 livestock manures with different substrate concentrations. Bioresour. Technol. 198, 133–140.
 https://doi.org/10.1016/j.biortech.2015.08.151
- Limoli, A., Langone, M., Andreottola, G., 2016. Ammonia removal from raw manure digestate by
 means of a turbulent mixing stripping process. J. Environ. Manage. 176, 1–10.
 https://doi.org/10.1016/j.jenvman.2016.03.007
- Lin, L., Lei, Z., Wang, L., Liu, X., Zhang, Y., Wan, C., Lee, D.-J., Tay, J.H., 2013. Adsorption mechanisms
 of high-levels of ammonium onto natural and NaCl-modified zeolites. Sep. Purif. Technol.
 103, 15–20. https://doi.org/10.1016/j.seppur.2012.10.005
- Liu, C., Yuan, X., Zeng, G., Li, W., Li, J., 2008. Prediction of methane yield at optimum pH for anaerobic
 digestion of organic fraction of municipal solid waste. Bioresour. Technol. 99, 882–888.
 https://doi.org/10.1016/j.biortech.2007.01.013
- 984Lothenbach, B., Scrivener, K., Hooton, R.D., 2011. Supplementary cementitious materials. Cem.985Concr. Res., Conferences Special: Cement Hydration Kinetics and Modeling, Quebec City,9862009& CONMOD10, Lausanne, 2010987https://doi.org/10.1016/j.cemconres.2010.12.001
- Luukkonen, T., Sarkkinen, M., Kemppainen, K., Rämö, J., Lassi, U., 2016. Metakaolin geopolymer
 characterization and application for ammonium removal from model solutions and landfill
 leachate. Appl. Clay Sci. 119, 266–276. https://doi.org/10.1016/j.clay.2015.10.027

- Magniont, C., Coutand, M., Bertron, A., Cameleyre, X., Lafforgue, C., Beaufort, S., Escadeillas, G.,
 2011. A new test method to assess the bacterial deterioration of cementitious materials.
 Cem. Concr. Res. 41, 429–438. https://doi.org/10.1016/j.cemconres.2011.01.014
- Marañón, E., Ulmanu, M., Fernández, Y., Anger, I., Castrillón, L., 2006. Removal of ammonium from
 aqueous solutions with volcanic tuff. J. Hazard. Mater. 137, 1402–1409.
 https://doi.org/10.1016/j.jhazmat.2006.03.069
- 997 McCarty, P.L., 1964. Anaerobic Waste treatment Fundamentals. Public Works 95, 66.
- 998Mookherjee, M., Welch, M.D., Pollès, L.L., Redfern, S.A.T., Harlov, D.E., 2005. Ammonium ion999behaviour in feldspar: variable-temperature infrared and 2H NMR studies of synthetic1000buddingtonite, N(D,H)4AlSi3O8. Phys. Chem. Miner. 32, 126–131.1001https://doi.org/10.1007/s00269-005-0455-x
- Morandeau, A., Thiéry, M., Dangla, P., 2014. Investigation of the carbonation mechanism of CH and
 C-S-H in terms of kinetics, microstructure changes and moisture properties. Cem. Concr. Res.
 56, 153–170. https://doi.org/10.1016/j.cemconres.2013.11.015
- 1005 Nathalie Bachmann, E.S.A., 2013. 8 Design and engineering of biogas plants, in: Wellinger, A.,
 1006 Murphy, J., Baxter, D. (Eds.), The Biogas Handbook, Woodhead Publishing Series in Energy.
 1007 Woodhead Publishing, pp. 191–211. https://doi.org/10.1533/9780857097415.2.191
- 1008 O'Connor, S.J., MacKenzie, K.J.D., Smith, M.E., Hanna, J.V., 2010. Ion exchange in the charge balancing sites of aluminosilicate inorganic polymers. J. Mater. Chem. 20, 10234–10240.
 1010 https://doi.org/10.1039/C0JM01254H
- 1011 Oral, Ç.M., Ercan, B., 2018. Influence of pH on morphology, size and polymorph of room temperature
 1012 synthesized calcium carbonate particles. Powder Technol. 339, 781–788.
 1013 https://doi.org/10.1016/j.powtec.2018.08.066
- 1014Oueslati, O., Duchesne, J., 2012. The effect of SCMs and curing time on resistance of mortars1015subjected to organic acids. Cem. Concr. Res. 42, 205–214.1016https://doi.org/10.1016/j.cemconres.2011.09.017
- Perlot, C., Verdier, J., Carcassès, M., 2006. Influence of cement type on transport properties and chemical degradation: application to nuclear waste storage. Mater. Struct. 39, 511–523.
- Pouhet, R., 2015. Formulation and durability of metakaolin-based geopolymers (phd). Université de
 Toulouse, Université Toulouse III Paul Sabatier.
- Pouhet, R., Cyr, M., Bucher, R., 2019. Influence of the initial water content in flash calcined
 metakaolin-based geopolymer. Constr. Build. Mater. 201, 421–429.
 https://doi.org/10.1016/j.conbuildmat.2018.12.201
- 1024 Rasi, S., 2009. Biogas composition and upgrading to biomethane (Jyväskylä studies in Biological and
 1025 Environmental Science). University of Jyväskylä, Finland.
- Roosz, C., Vieillard, P., Blanc, P., Gaboreau, S., Gailhanou, H., Braithwaite, D., Montouillout, V.,
 Denoyel, R., Henocq, P., Madé, B., 2018. Thermodynamic properties of C-S-H, C-A-S-H and M S-H phases: Results from direct measurements and predictive modelling. Appl. Geochem. 92,
 140–156. https://doi.org/10.1016/j.apgeochem.2018.03.004
- San Nicolas, R., Cyr, M., Escadeillas, G., 2013. Characteristics and applications of flash metakaolins.
 Appl. Clay Sci. 83–84, 253–262. https://doi.org/10.1016/j.clay.2013.08.036
- Scrivener, K., Snellings, R., Lothenbach, B., 2016. A Practical Guide to Microstructural Analysis of
 Cementitious Materials. CRC Press.
- 1034Ševčík, R., Pérez-Estébanez, M., Viani, A., Šašek, P., Mácová, P., 2015. Characterization of vaterite1035synthesized at various temperatures and stirring velocities without use of additives. Powder1036Technol. 284, 265–271. https://doi.org/10.1016/j.powtec.2015.06.064
- 1037Singh, B., Ishwarya, G., Gupta, M., Bhattacharyya, S.K., 2015. Geopolymer concrete: A review of some1038recentdevelopments.Constr.Build.Mater.85,78–90.1039https://doi.org/10.1016/j.conbuildmat.2015.03.036
- Snellings, R., Chwast, J., Cizer, Ö., De Belie, N., Dhandapani, Y., Durdzinski, P., Elsen, J., Haufe, J.,
 Hooton, D., Patapy, C., Santhanam, M., Scrivener, K., Snoeck, D., Steger, L., Tongbo, S.,
 Vollpracht, A., Winnefeld, F., Lothenbach, B., 2018. RILEM TC-238 SCM recommendation on

- 1043hydration stoppage by solvent exchange for the study of hydrate assemblages. Mater. Struct.104451, 172. https://doi.org/10.1617/s11527-018-1298-5
- Stepkowska, E.T., Pérez-Rodríguez, J.L., Sayagués, M.J., Martínez-Blanes, J.M., 2003. Calcite, vaterite
 and aragonite forming on cement hydration from liquid and gaseous phase. J. Therm. Anal.
 Calorim. 73, 247–269. https://doi.org/10.1023/A:1025158213560
- 1048Tai, C.Y., Chen, F.-B., 1998. Polymorphism of CaCO3, precipitated in a constant-composition1049environment. AIChE J. 44, 1790–1798. https://doi.org/10.1002/aic.690440810
- 1050 The European Parliament, The European Union Council, 2009. Directive 2009/28/EC, 140.
- 1051 The European Parliament, The European Union Council, 2001. Directive 2001/77/EC.
- 1052 Voegel, C., 2017. Impact biochimique des effluents agricoles et agroindustriels sur les 1053 structures/ouvrages en béton dans la filière de valorisation par méthanisation (ou digestion 1054 anaérobie). Institut National Polytechnique de Toulouse (INP Toulouse), Toulouse, France. 1055 http://www.theses.fr/2017INPT0044.
- Voegel, C., Bertron, A., Erable, B., 2016. Mechanisms of cementitious material deterioration in biogas
 digester. Sci. Total Environ. 571, 892–901. https://doi.org/10.1016/j.scitotenv.2016.07.072
- 1058 Voegel, C., Bertron, A., Erable, B., 2015. Biodeterioration of cementitious materials in biogas digester.
 1059 Matér. Tech. 103, 202. https://doi.org/10.1051/mattech/2015023
- Voegel, C., Durban, N., Bertron, A., Landon, Y., Erable, B., 2019a. Evaluation of microbial proliferation
 on cementitious materials exposed to biogas systems. Environ. Technol. 1–11.
 https://doi.org/10.1080/09593330.2019.1567610
- 1063 Voegel, C., Giroudon, M., Bertron, A., Patapy, C., Peyre Lavigne, M., Verdier, T., Erable, B., 2019b. 1064 Cementitious materials in biogas systems: Biodeterioration mechanisms and kinetics in CEM I 1065 and CAC based materials. Cem. Concr. Res. 124, 105815. 1066 https://doi.org/10.1016/j.cemconres.2019.105815
- 1067Wang, H., Li, H., Yan, F., 2005. Synthesis and mechanical properties of metakaolinite-based1068geopolymer.ColloidsSurf.Physicochem.Eng.Asp.268,1–6.1069https://doi.org/10.1016/j.colsurfa.2005.01.016
- 1070 Wang, P., Wang, H., Qiu, Y., Ren, L., Jiang, B., 2018. Microbial characteristics in anaerobic digestion
 process of food waste for methane production–A review. Bioresour. Technol., Bioconversion
 of Food Wastes 248, 29–36. https://doi.org/10.1016/j.biortech.2017.06.152
- 1073 Wang, Q., Yang, Y., Yu, C., Huang, H., Kim, M., Feng, C., Zhang, Z., 2011. Study on a fixed zeolite
 1074 bioreactor for anaerobic digestion of ammonium-rich swine wastes. Bioresour. Technol. 102,
 1075 7064–7068. https://doi.org/10.1016/j.biortech.2011.04.085
- 1076 Wang, Z., Yun, S., Xu, H., Wang, C., Zhang, Y., Chen, J., Jia, B., 2019. Mesophilic anaerobic co-digestion
 1077 of acorn slag waste with dairy manure in a batch digester: Focusing on mixing ratios and bio1078 based carbon accelerants. Bioresour. Technol. 286, 121394.
 1079 https://doi.org/10.1016/j.biortech.2019.121394
- Weiland, P., 2010. Biogas production: current state and perspectives. Appl. Microbiol. Biotechnol. 85,
 849–860. https://doi.org/10.1007/s00253-009-2246-7
- Wianglor, K., Sinthupinyo, S., Piyaworapaiboon, M., Chaipanich, A., 2017. Effect of alkali-activated
 metakaolin cement on compressive strength of mortars. Appl. Clay Sci. 141, 272–279.
 https://doi.org/10.1016/j.clay.2017.01.025
- 1085 Wray, J.L., Daniels, F., 1956. Precipitation of Calcite and Aragonite. J. Am. Chem. Soc. 79, 2031–2034.
- 1086Xu, H., Li, Y., Hua, D., Mu, H., Zhao, Y., Chen, G., 2019. Methane production from the anaerobic1087digestion of substrates from corn stover: Differences between the stem bark, stem pith, and1088leaves. Sci. Total Environ. 694, 133641. https://doi.org/10.1016/j.scitotenv.2019.133641
- 1089Xu, H., Yun, S., Wang, C., Wang, Z., Han, F., Jia, B., Chen, J., Li, B., 2020. Improving performance and1090phosphorus content of anaerobic co-digestion of dairy manure with aloe peel waste using1091vermiculite.1092https://doi.org/10.1016/j.biortech.2020.122753
- 1093Yariv, S., 1985. Study of the adsorption of organic molecules on clay minerals by differential thermal1094analysis. Thermochim. Acta 88, 49–68. https://doi.org/10.1016/0040-6031(85)85414-9

- 1095Yenigün, O., Demirel, B., 2013. Ammonia inhibition in anaerobic digestion: A review. Process1096Biochem. 48, 901–911. https://doi.org/10.1016/j.procbio.2013.04.012
- Yu, H.-Q., Fang, H.H.P., 2002. Acidogenesis of dairy wastewater at various pH levels. Water Sci.
 Technol. 45, 201–206.
- Yun, S., Fang, W., Du, T., Hu, X., Huang, X., Li, X., Zhang, C., Lund, P.D., 2018. Use of bio-based carbon
 materials for improving biogas yield and digestate stability. Energy 164, 898–909.
 https://doi.org/10.1016/j.energy.2018.09.067
- Yun, S., Zhang, C., Wang, Y., Zhu, J., Huang, X., Du, T., Li, X., Wei, Y., 2019. Synergistic effects of Fe
 salts and composite additives on anaerobic digestion of dairy manure. Int. Biodeterior.
 Biodegrad. 136, 82–90. https://doi.org/10.1016/j.ibiod.2018.10.011
- 1105Zhang, C., Yun, S., Li, X., Wang, Z., Xu, H., Du, T., 2018. Low-cost composited accelerants for anaerobic1106digestion of dairy manure: Focusing on methane yield, digestate utilization and energy1107evaluation.Bioresour.Technol.263,517–524.1108https://doi.org/10.1016/j.biortech.2018.05.042
- 1109