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Insights into the local interaction mechanisms between fermenting broken maize and various binder materials for anaerobic digester structures

Marie Giroudon1,2, Cédric Perez3,4,5, Matthieu Peyre Lavigne2, Benjamin Erable3, Christine Lors4,5, Cédric Patapy1, Alexandra Bertron1,*

1. LMDC, Université de Toulouse, UPS, INSA Toulouse, France
2. TBI, Université de Toulouse, CNRS, INRA, INSA, Toulouse, France
3. Laboratoire de Génie Chimique, Université de Toulouse, CNRS, INPT, UPS, Toulouse, France
4. IMT Lille Douai, Institut Mines Télécom, Univ. Lille, Centre for Materials and Processes, F-59000 Lille, France
5. Univ. Lille, Institut Mines Télécom, Univ. Artois, Junia, ULR 4515 - LGCgE, Laboratoire de Génie Civil et géo-Environnement, F-5900 Lille, France

* Corresponding author: bertron@insa-toulouse.fr

Highlights

- Cement matrices face biodeterioration but have little effect on anaerobic digestion
- CH₄ production was similar in all BMP reactors (with/without binder materials)
- Alkali-activated metakaolin shows a thinner chemically modified layer
- Metakaolin paste impacts the digestion in terms of [NH₄⁺], biomass nature, and pH
- The sessile and planktonic microbial communities are different

Abstract

Concrete structures of anaerobic digestion plants face chemically aggressive conditions due to the contact with the complex liquid fraction of the fermenting biowaste. This paper aims to determine the biogeochemical dynamic interaction phenomena at play between the biowaste and cementitious matrices at the local scale, and to identify durable binders in such environments. Binder materials
likely to show increased durability – slag and calcium aluninate cement, and a metakaolin-based alkali-activated geopolymer – and a reference Portland cement were inserted into sealed bioreactors during 5 cycles (245 days) of broken maize anaerobic digestion. Cementitious pastes suffered chemical and mineralogical alteration related mainly to carbonation and leaching. However, they had no negative impact on the bioprocess in terms of pH, metabolic evolution of volatile fatty acids and NH₄⁺, planktonic microbial community composition or CH₄ production. In all reactors, the microbial community was able to perform the anaerobic digestion successfully. The MKAA was only slightly altered in its outermost layer. Its presence in the biowaste induced lower NH₄⁺ concentrations, a slightly higher pH and a marked shift in the microbial community, but CH₄ total production was not affected. Substantial enrichment of acid forming bacteria, especially members of the genus Clostridium, was observed in the biofilm formed on all materials.

Keywords: Anaerobic digestion, cementitious materials, geopolymer, durability, biodeterioration, biofilm

Graphical abstract
1 Introduction

Anaerobic digestion (AD) is the process of transforming organic matter into a methane (CH\textsubscript{4})-rich gas (the biogas) and a moist conditioner and fertilizer (the digestate) by anaerobic microorganisms. In the current context of sustainable transition towards low carbon renewable energies and reduction of greenhouse gas emissions into the atmosphere, the development of the sector is being encouraged in Europe and the number of industrial installations is growing. As an illustration, in France, the objective of the Pluriannual Energy Programme is to reach the target of including 7\% to 10\% of biogas in the total gas consumption by 2030 (Ministère de la transition écologique et solidaire, 2020).

On an industrial scale, the AD bioprocess is implemented in anaerobic digesters, i.e. fermentation plants, which are mainly made of concrete because this material is economical, easy to implement, waterproof, and has a good thermal inertia. While the upper part of the structure, in contact with the biogas, is often protected from the aggressiveness of the environment by liners (Nathalie Bachmann, 2013), in the lower part, the concrete is in direct contact with the biowaste undergoing digestion. Concrete structures that are thus directly exposed to the fermenting biowaste experience severe deterioration (Giroudon et al., 2021a; Koenig and Dehn, 2016; Perez et al., 2021; Voegel et al., 2019, 2016) which could affect the sustainability of the structures and their durability, with both economic and environmental consequences.

In more detail, the microbiologically driven AD bioprocess requires the succession of four steps of biowaste degradation under anaerobic conditions: hydrolysis, acidogenesis, acetogenesis and methanogenesis (Batstone et al., 2002; Evans and Furlong, 2003). During the first step, macromolecules of biowaste are broken into smaller molecules. These compounds of lower molecular weight are then fermented by acidogenic bacteria during the second step, leading to the specific production of volatile fatty acids (VFA) (mainly propionic, acetic and butyric acids). During the following step of acetogenesis, the products of the previous step are converted into acetate, H\textsubscript{2} and
Finally, CO$_2$ and CH$_4$ are produced through two distinct pathways during methanogenesis: the hydrogenotrophic pathway produces CH$_4$ from H$_2$ and CO$_2$; and the acetoclastic pathway uses acetate as a precursor substrate (Krakat et al., 2010). All the AD reaction steps are carried out by microorganisms that can adopt clustered structures in the form of sludge, aggregates, or biofilms, which allow close interaction and cooperation between the microbial populations involved in the four reaction stages. Some of the microbial metabolites produced in the AD process, especially VFA, dissolved CO$_2$, and NH$_4^+$ ions (contained directly in some waste and/or produced along with VFA during acidogenesis (Meegoda et al., 2018)) are known to be aggressive for concrete (Giroudon et al., 2021a; Koenig and Dehn, 2016; Perez et al., 2021; Voegel et al., 2019) and are present in varying proportions. Moreover, the action of microorganisms organised in biofilms on the concrete surface can increase the local concentration of metabolites and locally accentuate the aggressiveness of the environment and thus the degradation of the concrete (Magniont et al., 2011).

With a view to enhancing the durability of concrete in this expanding industrial sector, this study aims at an extensive understanding of the biogeochemical interactions between cement matrices and the liquid fraction of biowaste in AD. On the one hand, the impact of the presence of different binder materials on the performance of the AD bioprocess is evaluated considering (i) the CH$_4$ production, (ii) the chemical composition of the liquid fraction in terms of VFA concentrations, NH$_4^+$ ion concentration and pH, and (iii) the microbial diversity of sessile and planktonic biomasses. On the other hand, the chemical and mineralogical changes of the binder materials previously exposed to the fermenting biowaste are also assessed. Cement pastes made of blast-furnace slag cement (CEM III), calcium aluminate cement (CAC), a metakaolin-based alkali-activated geopolymer (MKAA) likely to show increased durability in this medium (Bertron et al., 2007a; Drugă et al., 2018; Duan et al., 2015; Grengg et al., 2020; Gruyaert et al., 2012; Oueslati and Duchesne, 2012; Singh et al., 2015) and an ordinary Portland cement (CEM I) were immersed for 245 days in airtight biochemical methane potential (BMP) reactors (Holliger et al., 2016) inoculated with broken maize. This substrate was selected because it offers a high methanogenic potential (Bruni et al., 2010; Gerin et al., 2008) and
also because it allows the microbial production of a large amount of organic acids during the acidogenesis step, generating aggressive chemical conditions for the cement matrices. A high solid/liquid ratio (surface area of the material sample/liquid volume) was used in order to reproduce the local conditions occurring in the vicinity of the digester concrete walls.

2 Materials and methods

The experimental protocol was similar to the one developed and validated by Giroudon et al. (2021a) and is presented in Figure 1.

![Figure 1: Schematic representation of the experimental protocol describing the samples, the experiments, and the temporal management of the solid, liquid and microbial fraction analyses, adapted from Giroudon et al. (2021a)](image)

It consisted of immersing pastes made of ordinary Portland cement CEM I 52.5R (CEM I), CEM III/B 42.5N (CEM III), calcium aluminate cement, Calcoat ® RG (CAC), and a metakaolin-based alkali-activated (MKAA) paste, in reactors inoculated with broken maize at 35 °C, in order to reproduce anaerobic digestion during 5 cycles of digestion.
2.1 Binders

The cement pastes (CEM I, CEM III, CAC), called cementitious materials or cement pastes below, were poured with a water/binder ratio of 0.30. The MKAA geopolymer was made according to the procedure of Pouhet (2015) by using metakaolin, liquid sodium silicate (molar ratio $\frac{SiO_2}{Na_2O} = 1.7$) and water. Geopolymers are aluminosilicate materials formed by the activation of an aluminosilicate source, such as metakaolin, by a strongly basic alkaline solution (Pouhet et al., 2019).

The samples were made according to the procedure described in the study by Giroudon et al. (2021a), i.e., the paste specimens were mixed using the French standard NF EN 196-1 (2016) and cast in cylindrical moulds 75 mm high and 25 mm in diameter. They were cured in sealed plastic bags and were then exposed to the biowaste in AD.

The water porosities of the pastes were measured according to the NF P18-459 standard (AFNOR, 2010). The measurements show a significantly higher porosity of the MKAA paste (48.6 %) and a lower porosity of the CAC paste (24.0 %) in comparison with the CEM I and CEM III pastes, whose water porosity values were intermediate - about 32.0 % and 35.6 %, respectively.

2.2 Immersion of cement pastes in laboratory BMP reactors

The protocol of immersion consisted of immersing cement paste specimens in airtight BMP reactors (Holliger et al., 2016) containing 80 mL of microbial inoculum from an industrial biogas plant located in Haute-Garonne (France) and broken maize from a farm in Barcelonne-du-Gers (Gers, France) as the biowaste. For each kind of material, the experiment was carried out in duplicate with two BMP reactors each containing a sample. In addition, two control BMP reactors without material were also operated and monitored over the same period of time. The detailed protocol is presented in Giroudon et al. (2021a) (Section 2.1).

The mass of added broken maize increased with the cycles of AD (Table 1), in order to increase the aggressiveness of the medium with time. Each cycle of AD was considered complete when the gas...
production in the BMP reactors stopped (Table 1). During the third cycle, once the biogas production was complete, the BMP reactors were kept at room temperature for 5.5 weeks because the laboratory closed for the summer break.

| Table 1: Mass of broken maize added per cycle and duration of each cycle |
|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|-----------------------------|
|                             | 1\textsuperscript{st} cycle | 2\textsuperscript{nd} cycle | 3\textsuperscript{rd} cycle | 4\textsuperscript{th} cycle | 5\textsuperscript{th} cycle |
| Temperature                 | 35°C                       | 35°C                       | 35°C                       | Room                       | 35°C                       | 35°C                       |
| Mass of broken maize added  | 3                          | 3                          | 4                          | 5                          | 5                          |
| (g)                         |                             |                             |                             |                             |                             |
| Duration of a cycle (weeks) | 10                         | 4.5                        | 5                          | 5.5                        | 5                          | 5                          |

2.3 Analyses of liquid and gas fractions

Several times a week, the BMP reactors were shaken manually, the gas pressure was measured with a manometer, and gas and liquid were sampled using syringes. Gas samples were analysed using gas chromatography (GC Trace 1300 Thermofisher, Mobile phase Helium, Separation Column Hayssep N 60-80 1.0 m x 1/16, oven temperature 110°C, detector TCD) (O2-N2, H2, CH4, CO2 and H2S). pH was immediately measured in the liquid samples before they were centrifuged for 5 min (Eppendorf, Centrifuge 5430R, 4 °C, RCF 7197) and the supernatant was filtered through a 0.2 μm filter for further analysis. The NH4\textsuperscript{+} concentration was analysed by ion chromatography (Dionex Thermofisher ICS2000, Guard column AG19, IonPac CS12 3x250 mm, at 30 °C with eluent generator cartridge EGC III MSA for methanesulfonic acid) and the concentrations of some VFA (acetic, propionic, butyric, isobutyric, valeric, and isovaleric acids) were analysed by gas chromatography (Varian 3900/430, WAX column: length 15 m, external diameter 0.53 mm, thickness of the internal phase 1 μm, detector FID). In addition, gas and liquid compositions were used to calculate the aqueous inorganic CO2 concentration in the liquid fraction using Henry’s law considering a temperature of 35 °C and atmospheric pressure (values of the Henry’s law constants from Batstone et al. (2002)).

2.4 Analyses of the changes in binder materials

The binder materials were removed from the BMP reactors after the 3\textsuperscript{rd} and the 5\textsuperscript{th} cycles and slices were sawn from their ends with a diamond saw. The remaining cylinders were re-immersed in the
BMP reactors as quickly as possible to continue the experiments, while the slices were used in analyses of the mineralogical and chemical changes following exposure to the fermenting biowaste. The mineralogical degradation in depth was assessed by qualitative X-Ray Diffraction (XRD) (Brucker Avance, Co cathode, 40 kV, 40 nA) by successively analysing and abrading the plane side of the slice until the core of the material was reached (initial mineralogical composition) (Bertron et al., 2005a). The chemical composition was quantitatively characterized by Electron Probe Micro-Analysis (EPMA) ( Cameca XFive, 15 kV, 20 nA) on polished sections: series of chemical punctual analyses (Ca, Si, Al, Fe, Mg, P and S) were performed from the surface in contact with the liquid fraction to the core of the specimen, avoiding the anhydrous grains. Calibrations were performed on synthetic natural controls before each run. The graphs are the combination of two chemical profiles in mass percentage of oxides from the same sample (each chemical profile consisting of a series of punctual chemical analyses, from the surface in contact with the biowaste to the sound core) and smoothed over 3 points for better readability, using a moving average (Bertron et al., 2009). Trend and smoothed curves were then established from these graphs.

### 2.5 Analysis of microbial populations

The protocol for collecting sessile microbial communities was adapted from Perez et al. (2021) in order to capture two distinct layers in the thickness of the biofilm successively. The "weakly adhered" biofilm was removed from the cementitious material surface by an immersion in phosphate buffered saline solution (PBS, 0.1 M, pH 7.4) for 15 minutes. The "strongly adhered" biofilm was removed by 3 minutes of sonication treatment (FB 15061 Fisher Scientific, ultrasonic frequency 37 Hz) in PBS. For both treatments, the entire PBS volume containing the detached biomass was then recovered. A 2 mL sample was also collected from the liquid fraction of each BMP reactor (Figure 1). DNA was extracted from the three types of samples ("weakly adhered" biomass, "strongly adhered" biomass and planktonic biomass from the liquid fraction) with the Qiagen DNeasy power biofilm DNA extraction kit, according to the protocol described by the manufacturer. The extracted DNA was sent to RTL Genomics (Texas, USA), where the 16s rDNA was amplified using the 515F and 806R primers,
targeting both bacteria and archaea. Sequencing of the amplified DNA was also performed by RTL Genomics (USA) using a MiSeq Illumina platform (details of the amplification and sequencing protocols are available in the supplementary data). Data analysis for DNA quality, DNA sequence alignment, and clustering in operational taxonomic units and taxonomic assignment were also performed by RTL Genomics according to their protocol (available at https://rtlgenomics.com/amplicon-bioinformatics-pipeline). The sequencing raw sequence reads were published in the NCBI SRA database (accession: PRJNA758226; direct link: https://www.ncbi.nlm.nih.gov/Traces/study/?acc=PRJNA758226). The number of OTUs (Operational Taxonomic Units) identified at the taxonomic levels of species and genera were analysed statistically using the R software, (R Core Team, 2017). A heatmap was produced with the "Marray" package using Ward's classification method and the calculation of distance based on the correlation between the standardized abundance scores of each OTU. Principal component analyses (PCA) were performed with the "Factominer" and "Factoextra" packages.

3 Results

3.1 Characterization of the AD bioprocess with respect to the immersed binder

Table 2 gives the total production of CH₄ in each AD cycle for all BMP tests, with and without binder materials, and the cumulative total production at the end of the experiment.

Table 2: Total production of CH₄ (NmL.g⁻¹ of broken maize) at the end of each digestion cycle and cumulative total production at the end of the experiment (Total). Results are expressed as mean value of the two duplicates of BMP reactors ± the standard deviation.

<table>
<thead>
<tr>
<th></th>
<th>Control BMP reactors</th>
<th>CEM I</th>
<th>CEM III</th>
<th>CAC</th>
<th>MKAA</th>
<th>Mean values per cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Production of CH₄ (NmL.g⁻¹ of broken maize)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1st cycle</td>
<td>1530 ± 232</td>
<td>1439 ± 13</td>
<td>1579 ± 67</td>
<td>1379 ± 252</td>
<td>1336 ± 145</td>
<td>1453 ± 101</td>
</tr>
<tr>
<td>2nd cycle</td>
<td>1298 ± 88</td>
<td>1271 ± 91</td>
<td>1234 ± 48</td>
<td>1186 ± 136</td>
<td>1136 ± 131</td>
<td>1225 ± 65</td>
</tr>
<tr>
<td>3rd cycle</td>
<td>1308 ± 82</td>
<td>1214 ± 23</td>
<td>1270 ± 5</td>
<td>1283 ± 32</td>
<td>1178 ± 123</td>
<td>1251 ± 53</td>
</tr>
<tr>
<td>4th cycle</td>
<td>1005 ± 380</td>
<td>996 ± 290</td>
<td>1236 ± 81</td>
<td>1620 ± 556</td>
<td>1129 ± 39</td>
<td>1197 ± 256</td>
</tr>
<tr>
<td>5th cycle</td>
<td>1011 ± 174</td>
<td>1102 ± 45</td>
<td>1124 ± 3</td>
<td>1247 ± 46</td>
<td>1222 ± 35</td>
<td>1141 ± 96</td>
</tr>
<tr>
<td>Total</td>
<td>6152 ± 100</td>
<td>6022 ± 144</td>
<td>6392 ± 3</td>
<td>6715 ± 175</td>
<td>6001 ± 395</td>
<td>6267 ± 306</td>
</tr>
</tbody>
</table>
Despite the presence of the binder materials, the production of CH$_4$ was similar in all the BMP reactors during the whole experiment (Table 2). Even though the amount of broken maize biowaste was increased between the first and the last cycle, there was a slight decrease in the amount of CH$_4$ produced, probably due to the evolution, during storage, of the biowaste, the methanogenic potential of which decreased with time.

Figure 2 gives the evolution in time of the pH, the total concentration of VFA and the ammonium concentration in the liquid fraction of the BMP reactors.

The production of the different VFA in the different BMP reactors as a function of time is available in the supplementary materials (Appendix A). It shows that the production of acetic acid was predominant, and that the production and consumption of the different VFA was simultaneous in each type of BMP reactor.
Figure 2: Evolution of the pH, of the total VFA concentration and of the ammonium concentration during the five cycles of AD in the BMP reactors, with or without binder materials. Mean values of the two duplicate BMP reactors are presented with the standard deviations.
At the beginning of the experiment (day 0), the pH values were very close in all the BMP reactors (8.2 ± 0.1). After three days, the pH had significantly decreased, reaching 5.8 ± 0.23, 6.2 ± 0.12, 6.4 ± 0.14, 6.2 ± 0.01, and 6.7 ± 0.06 in the control BMP reactors and the BMP reactors containing CEM I, CEM III, CAC and MKAA pastes, respectively. On the same day (day 3), the VFA concentration increased to 1.8 ± 0.1 g.L⁻¹. Thereafter, the VFA were consumed and the pH increased in the BMP reactors containing the binder materials. In the meantime, the pH continued to decrease in the control BMP reactors with no decrease of the acid concentration. Until day 13, the pH in these reactors continued to drop and reached 5.4 ± 0.20, with an extremely high total VFA concentration of 3.3 ± 0.23 g.L⁻¹. In order to avoid acidosis (acidic inhibition) in the BMP reactors, 30 mL of a NaHCO₃ solution (80 g.L⁻¹) was injected into each control reactor to increase the pH and promote favourable conditions for the AD bioprocess. The natural rise in pH in the BMP reactors containing the binder materials was due to the strong buffering effect of the cementitious and alkali-activated materials. The addition of the NaHCO₃ solution in the control reactors led to an increase of the pH and therefore allowed methanogenic microbial activity to take place (confirmed by the methane production rate measured, data not shown), leading to the consumption of the VFA and to the production of biogas.

Subsequently, the variations in pH were typical of a fed-batch AD bioprocess: the pH was between 7.0 and 7.5 in the steady state (suitable for AD microbial populations) and a decrease in pH occurred after each new addition of biowaste, due to the fast production of VFA. Starting from cycle 2, the microbial communities in the control BMP reactors adapted to the AD of broken maize, and the pH naturally rose after each addition of broken maize. The lower buffering effect of CAC material in comparison with the other materials induced a slightly lower pH during the experiment, which may have had an impact on the balance of microbial populations.

During the first cycle, among the BMP reactors containing binder materials, the pH was slightly higher in those containing the MKAA (about +0.5) while the VFA concentrations were similar. The pH
in these reactors remained that much higher during the whole experiment. However, with the increase of the biowaste load from 3 to 5 g, the presence of MKAA induced a higher VFA concentration and a slower consumption of acids. This could reflect a poorer efficiency of the methanogenesis step. All VFA were, however, consumed at the end of each cycle and in all BMP reactors.

The presence of the cementitious materials CEM I, CEM III and CAC did not significantly influence the NH$_4^+$ concentration in the liquid fraction compared to that in the control BMP reactors. At the beginning of each cycle, the evolution of the NH$_4^+$ concentration resulted from two opposite effects: the addition of water led to a decrease in the NH$_4^+$ concentration by dilution, whereas the addition of biowaste generated a gradual increase in NH$_4^+$ concentration. This increase was much more significant during cycles 4 and 5, probably because the biowaste load was increased from 4 to 5 g. The NH$_4^+$ concentrations in the BMP reactors containing the cementitious materials and the controls were of the order of 150 to 300 mg.L$^{-1}$ during this experiment, which is much lower than the values measured with bovine manure (approximately 750 mg.L$^{-1}$ at the end of the first cycle) (Giroudon et al., 2021a). As highlighted in the study by Giroudon et al. (2021a), the presence of MKAA induced significantly lower NH$_4^+$ concentrations (between 0 and 50 mg.L$^{-1}$), probably due to the NH$_4^+$ adsorption capacity of this material.

In the control BMP reactors, the sodium concentration increased sharply between the start of the first cycle (64.7 ± 2.7 mg.L$^{-1}$) and the start of the second cycle (867.7 ± 22.0 mg.L$^{-1}$) due to the addition of the NaHCO$_3$ solution on the 13$^{th}$ day (Table 3). Subsequently, the sodium concentration decreased slightly during the experiment, from 867.7 ± 22.0 to 721.3 ± 13.2 mg.L$^{-1}$. This decrease was the consequence of a dilution effect due to the addition of water at the beginning of each cycle, in order to balance the volume losses related to liquid fraction sampling. Similar salinity values were observed for the BMP reactors containing the CEM I, CEM III and CAC: the sodium concentrations were between 74.0 ± 5.7 and 93.2 ± 2.1 mg.L$^{-1}$. The MKAA BMP reactors showed the highest salinity.
values (about 1500 mg.L\(^{-1}\) of sodium in solution) because MKAA were activated with sodium silicate, and thus released a larger quantity of sodium.

<table>
<thead>
<tr>
<th>[Na(^+)] mg.L(^{-1})</th>
<th>Start of the 1(^{st}) cycle</th>
<th>Start of the 2(^{nd}) cycle</th>
<th>Start of the 3(^{rd}) cycle</th>
<th>Start of the 4(^{th}) cycle</th>
<th>Start of the 5(^{th}) cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>64.7 ± 2.7</td>
<td>867.7 ± 22.0</td>
<td>818.4 ± 4.8</td>
<td>763.1 ± 3.3</td>
<td>721.3 ± 13.2</td>
</tr>
<tr>
<td>CEM I</td>
<td>74.0 ± 5.7</td>
<td>90.2 ± 8.7</td>
<td>90.5 ± 0.6</td>
<td>93.2 ± 2.1</td>
<td>86.1 ± 5.7</td>
</tr>
<tr>
<td>CEM III</td>
<td>77.4 ± 6.3</td>
<td>86.9 ± 4.0</td>
<td>85.9 ± 0.9</td>
<td>87.1 ± 4.6</td>
<td>79.5 ± 0.8</td>
</tr>
<tr>
<td>CAC</td>
<td>73.2 ± 2.8</td>
<td>89.4 ± 3.6</td>
<td>93.2 ± 2.2</td>
<td>89.6 ± 3.7</td>
<td>78.6 ± 2.2</td>
</tr>
<tr>
<td>MKAA</td>
<td>100.1 ± 4.4</td>
<td>1431.1 ± 26.7</td>
<td>1533.4 ± 1.8</td>
<td>1554.7 ± 26.3</td>
<td>1519.3 ± 3.1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Salinity mg.L(^{-1})</th>
<th>Start of the 1(^{st}) cycle</th>
<th>Start of the 2(^{nd}) cycle</th>
<th>Start of the 3(^{rd}) cycle</th>
<th>Start of the 4(^{th}) cycle</th>
<th>Start of the 5(^{th}) cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>475.8 ± 13.4</td>
<td>1230.5 ± 314.9</td>
<td>1462.7 ± 91.5</td>
<td>1132.4 ± 31.2</td>
<td>1115.5 ± 34.5</td>
</tr>
<tr>
<td>CEM I</td>
<td>537.1 ± 38.8</td>
<td>957.0 ± 35.5</td>
<td>1048.9 ± 32.4</td>
<td>848.1 ± 85.3</td>
<td>829.8 ± 147.7</td>
</tr>
<tr>
<td>CEM III</td>
<td>572.1 ± 52.3</td>
<td>859.6 ± 37.5</td>
<td>936.7 ± 38.7</td>
<td>627.4 ± 14.6</td>
<td>523.0 ± 178.0</td>
</tr>
<tr>
<td>CAC</td>
<td>520.4 ± 10.3</td>
<td>828.6 ± 30.3</td>
<td>904.8 ± 31.0</td>
<td>679.8 ± 33.4</td>
<td>699.6 ± 62.4</td>
</tr>
<tr>
<td>MKAA</td>
<td>532.3 ± 32.3</td>
<td>1579.3 ± 57.8</td>
<td>1703.8 ± 14.8</td>
<td>1656.3 ± 31.4</td>
<td>1609.1 ± 7.8</td>
</tr>
</tbody>
</table>

### Table 3: Concentrations of Na\(^+\) and salinity (\([\text{Na}^+] + [\text{K}^+] + [\text{PO}_4^{3-}] + [\text{Mg}^{2+}] + [\text{Ca}^{2+}]\)) in the liquid fraction of the BMP reactors at the start of each cycle (after the addition of water and maize), with or without the binder materials. Results are expressed as mean value of the two duplicates of BMP reactors ± the standard deviation.

#### 3.2 Microbial composition of sessile and planktonic biomasses

The clustering by similarity shown on Figure 3 allows different groups and subgroups to be distinguished within the samples on the X axis and the OTUs grouped by genera on the Y axis. A grouping according to the type of material used in the BMP reactor was observed: the MKAA group is clustered on the right of the Figure 3 and all the other materials and the negative sample, without material, are mostly grouped on the left side of this figure. Subgroups formed in accordance with the origin of the microorganisms samplings. Groups of planktonic and sessile samples were clearly divided. Concerning the grouping of the identified OTUs, there were also groups and subgroups. However, no strong relation was observed between the groups of samples and the group of OTUs. This classification method does not highlight the relationships between the OTUs, the type of samples and the type of material used.
Figure 3: Heatmap with colour scale of the next generation sequencing of the 16S DNA from planktonic biomass samples and “poorly” and “strongly” adhered biomass taken from the surface of cementitious material samples, and after 5 cycles of anaerobic digestion – L: weakly adhered biofilm; S: Strongly adhered biofilm; P: Planktonic; only OTU genera with an abundance of at least 1% in at least one sample are shown.

In order to assess these relationships, a principal component analysis (PCA) was carried out (Figure 4). Microbial samples are categorized according to the binder material used and the origin of the samples. The samples found on the left side of dimension 1 axis of Figure 4 all belong to the BMP reactors where MKAA (in blue) was immersed, while all the other samples are grouped on the right side of dimension 1 axis. The first PC, which explains 33.7% of the global variability, is thus linked to the presence/absence of MKAA.

The second PC, explaining 25.6% of the variability, is correlated with the origin of samples, since the strongly adhered biofilm samples (squares) are on the bottom part of dimension 2 axis, the liquid
samples (circles) on the top part, and the weakly adhered biofilm samples (triangles) are situated between these two groups. So, the second PC is highly correlated with the origin of the microbial samples and samples positioned towards the top of the dimension 2 axis should be the microbial populations that were situated the closest to the cementitious material surfaces. Also, the populations from the weakly adhered biomass should include a mix between the microbial populations of the other two groups.

Therefore, the data presented can mainly be categorised according to two main criteria, which are affiliated with the two CPs highlighted in Figure 4: firstly, the location of the sample, i.e. liquid or weakly/strongly attached biomass and, secondly, the material exposed, i.e. MKAA binder or all other binder materials tested.

It is interesting to look at Figure 3 while looking for the specific OTU genera found in these groups. Several microbial genera appear more likely to be detected in the MKAA samples. These are the acetogen genera *Petrimonas* and *Syntrophomonas* (T. E. Board, 2015; Sekiguchi, 2015). As for the samples without MKAA: the genera *Methanobacterium* (methanogen), *Pelospora*, *Bacteroides*, *Ruminoclostridium*, *Lachnocostridium*, *Aminomonas*, *Flexilinea* (acidogens) and *Acetobacteroides* (acetogen) are more abundant in those samples (Baena et al., 2015; Schink, 2015; Su et al., 2014; Tourlousse and Sekiguchi, 2018; Venkiteswaran et al., 2015).

When considering the other main criteria categorising our data, some kinds of OTUs were found more predominantly in the biofilm samples. These genera, identified as *Clostridium*, *Peptoclostridium*, *Pseudomonas* and *Turicibacter* (acidogen) (Bosshard, 2015), are the members of the A-1 group (Figure 3). The *Pseudomonas* genus mainly contains CO₂-producing respiring bacteria, known for their ability to form biofilms (Yoon et al., 2002). Also, among these four cited genera, the acidogen genera *Clostridium*, *Peptoclostridium* and *Turicibacter* were more abundant in the strongly adhered biofilm than in the loosely adhered one. As for the genera associated with the planktonic
lifestyle, the *Aminobacterium* genus and the *Synergistes* genus are both amino acid fermenters (E. Board, 2015).

The same clustering by similarities, initially presented at the genus level in Figure 3, was also performed at the level of the microbial species. The results are available as supplementary data (Appendix B). This clustering based on comparison at the microbial species level resulted in findings almost identical to those already shown in Figure 3. The genus *Clostridium* was highly represented, 14 OTUs being identified as belonging to this specific bacterial genus. The presence of each of these OTUs was significantly higher in the biofilm samples regardless of the type of binder material. This confirms the predisposition of this genus to widely form biofilms on surface materials. Among the methanogenic population, 4 genera were identified: *Methanobacterium*, *Methanosarcina*, *Methanoculleus* and *Methanosaeta*. *Methanosarcina*, *Methanoculleus* and *Methanosaeta* were present in all the microbial samples in both the planktonic and biofilm samples. However, the hydrogenotrophic methanogens, represented by the *Methanobacterium* genus, were almost absent in the MKAA samples. The amino acid fermenter genera, *Synergistes*, *Aminobacterium* and *Aminomonas*, leading to the release of NH$_4^+$ ions, were more often detected in the liquid fraction than in the biofilm.
Figure 4: PCA of microbial communities found in anaerobic digestion biomass samples and representation according to either the type of material or the location of the biomass sample – Distribution of all the samples according to the two main Principal Components (PC).

3.3 Chemical and mineralogical changes in the binder materials

The mineralogical and chemical changes of CAC pastes after 3 and 5 cycles of immersion in the fermenting broken maize are available in the Supplementary materials (Appendix C) since the effect of biodeterioration only marginally modified this material throughout the experiment. The mechanisms explaining these moderate mineralogical and chemical evolutions under AD conditions have already been described in detail in the work of Voegel et al. (2019).

For better readability, the results for the CEM I, CEM III and MKAA are presented in the form of trend graphs. Original data (CEM I and CEM III after 5 cycles of AD) are provided as Supplementary materials (Appendix D, Appendix E, Appendix F and Appendix G).

3.3.1 CEM I

Figure 5 shows the chemical and mineralogical features of CEM I pastes resulting from 3 and 5 cycles of immersion in the fermenting broken maize. The measurements were carried out respectively by EPMA and XRD, as a function of the distance to the surface of the CEM I paste in contact with the fermenting biowaste. The EPMA analysis highlighted three main zones after 3 cycles and 4 main zones after the 5th cycle. The sound paste was mainly made of calcium (50 %) and silicon (17 %) with minor amounts of aluminium, sulfur and iron. The mineralogical analyses showed the presence of typical hydrated (portlandite, ettringite) and anhydrous (C\textsubscript{2}S, C\textsubscript{3}S, brownmillerite, C\textsubscript{3}A) crystallized phases of a Portland cement-based hydrated matrix. The material mainly underwent decalcification with a progressive decrease in the CaO content together with some dissolution (decrease of the...
silicon and aluminium contents and dissolution of the Ca-bearing phases, mainly CH\(^1\)) associated with the carbonation of the outer layer of the CEM I paste (zones 3 and 2) with the precipitation of calcite and vaterite. Additionally, an enrichment in exogenous phosphorus, probably supplied by the fermenting biowaste, was spotted in the external zone. After 3 and 5 cycles, the sound zone was identified at about 600 µm deep. The main difference between these two limits was the presence of an intermediate zone spotted after 5 cycles (zone 3), between 100 and 300 µm deep, where calcium and silicon contents remained stable while the sulfur content fell to 0 %.

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Figure 5: Schematic representation of chemical composition of oxides and mineralogical composition (analysed by EPMA and XRD, respectively) of the CEM I pastes after 3 and 5 cycles of exposure to fermenting broken maize – Ett: ettringite; Ca:

\(^1\) Cement chemistry shorthand notations: A = Al\(_2\)O\(_3\), C = CaO, F = Fe\(_2\)O\(_3\), H = H\(_2\)O, M = MgO, S = SiO\(_2\)
calcite; Va: vaterite; Br: brownmillerite – **Bold characters = intensification of the XRD signal in comparison with the deeper zone; Parentheses = significantly lower intensity of the XRD signal in comparison with the main phase**

### 3.3.2 CEM III

The sound CEM III paste was mainly made of calcium, silicon and aluminium with lower amounts of sulfur and magnesium (Figure 6). The mineralogical analyses showed the same main crystalline phases as for the CEM I paste. After 3 cycles, an enrichment in calcium was observed between 400 µm and 1200 µm, together with a slight enrichment in sulfur, even though no mineralogical changes were observed. The decalcification of the paste stopped at about 400 µm deep, and the sulfur content dropped to a value close to 0 % (zone 3). The decrease of all the oxide contents (zone 4) and the phosphorus enrichment (zone 5) were observed in the external zones. The mineralogical analyses showed the carbonation of the CEM III paste with the presence of calcium carbonates calcite and vaterite.

After 5 cycles, an enrichment in sulfur was observed between 600 and 1200 µm deep, together with the decalcification of the cement matrix (zone 4). Between 270 and 600 µm (zone 3), the calcium content increased, probably due to the carbonation of the paste, whereas the total oxide content, as well as the aluminium and the silicon contents, decreased. Once again, an intermediate zone (zone 2) was marked by the stagnation of the calcium, aluminium and silicon contents whereas the sulfur content dropped to 0 %, this time with an enrichment in phosphorus. The external zone showed drops in the major oxide contents whereas a new enrichment in phosphorus was seen (zone 1). The main crystalline phases were calcite and vaterite. For both limits, the modified depths of the CEM III paste were significantly higher than for the CEM I paste (1200 µm vs. 600 µm respectively).
Figure 6: Schematic representation of chemical composition of oxides and mineralogical composition (analysed by EPMA and XRD, respectively) of the CEM III pastes after 3 and 5 cycles of exposure to fermenting broken maize–Ett: ettringite; Ca: calcite; Va: vaterite; Br: brownmillerite – Bold characters = intensification of the XRD signal in comparison with the deeper zone

3.3.3 MKAA

Figure 7 shows the chemical composition profiles of the MKAA paste together with the mineralogical phases identified. Because of the amorphous nature of the metakaolin and associated products of reaction, the XRD analyses did not show any mineralogical changes between the sound zone and the external part exposed to the aggressive environment. Only a thin outer fringe (100 and 180 µm wide respectively after the 3rd and 5th cycles) was significantly chemically modified, probably due to the dissolution of the paste. A drop in the potassium content was observed only in the degraded outer zone, around the outer 100 µm. Zone 1 bis showed the increase in the phosphorus content. After the
third cycle, this also corresponded to the increase in the sodium content just before its decrease in the outer zone, corresponding to its release into the liquid fraction. For the MKAA, it seems that the sodium contents in the solid varied greatly according to the depth, and the trends also varied depending on the samples. However, the decrease in the sodium content, like that of potassium, occurred only in the last micrometres (about 100 µm).

Figure 7: Schematic representation of chemical composition of oxides and mineralogical composition (analysed by EPMA and XRD, respectively) of the MKAA pastes after 3 and 5 cycles of exposure to fermenting broken maize
3.3.4 Relative performances of the binders

During the experiment (5 AD cycles, 245 days of exposition), the CEM I and CEM III pastes showed greater chemical and mineralogical changes than the MKAA pastes, in particular linked to the decalcification of the cement matrix and the dissolution of the initial mineralogical phases. In addition, the modified depths of the CEM III paste were much greater than in other materials at the end of the experiment. The MKAA samples appeared to show very stable behaviour against biochemical attack, with a very shallow modified depth. Thus, a trend of chemical stability of mineralogical phases to attack by digesting biowaste can be established on the basis of the results of this study: CEM III < CEM I < MKAA.

4 Discussion

4.1 Effect of the AD bioprocess on the binder materials

Cementitious materials are porous and strongly basic reactive media that experience a chemical attack by the liquid fraction of the biowaste in AD, thus releasing chemical compounds that could disrupt microbial activities. In this study, despite the high solid/liquid ratio used, it was observed that the presence of the binder materials did not decrease the efficiency of the AD, the CH₄ production being similar in all BMP reactors.

These observations are different from what had been observed by Giroudon et al. (2021a) who, in a slightly different way, studied the impact of cementitious materials on the AD of cattle manure. However, they are consistent with the studies of Voegel et al. (2016, 2019b), where the authors focused on the AD of collective food catering waste in laboratory conditions. With or without the presence of cementitious materials, they described similar values and trends for the pH and for the production kinetics of VFA, even though the CH₄ production was not measured. Broken maize has a much greater methanogenic potential than cattle manure (French Chamber of Agriculture, 2010) and, above all, faster hydrolysis. Thus, much higher VFA concentrations were observed at the start of
the cycles (maximum of 0.35 g.L\(^{-1}\) for cattle manure and 1.8 g.L\(^{-1}\) for broken maize in the first cycle),

linked to a strong acidification of the environment, as in the studies by Voegel et al. (2016, 2019b).

The transformation of this large amount of fermentable biowaste led to a production of CH\(_4\)

approximately 7.5 times greater than that obtained with cattle manure (data not shown).

For both biowastes (cattle manure and broken maize), the presence of the binder materials impacted

the AD bioprocess. In the case of AD of cattle manure, the alcalis of the materials (Na\(^+\), K\(^+\)), released
during the first cycle, raised the pH and inhibited some of microorganisms implicated in the AD

bioprocess (acidogenic, acetogenic, and methanogenic populations). In the case of AD of broken

maize, the leaching of these alcalis counterbalanced the strong acidity of the liquid fraction, thus

avoiding the phenomenon of acidosis that appeared in the control BMP reactors during the first

cycle, and enabling the AD sequence to progress well.

In AD, NH\(_4^+\) is classically produced by the degradation of proteins or urea (Kayhanian 1999; Yenigün

and Demirel 2013), which justifies the high concentrations of NH\(_4^+\) during the AD of cattle manure

(approximately 750 mg.L\(^{-1}\)) (Giroudon et al., 2021a), while the concentrations did not exceed 350

mg.L\(^{-1}\) with the broken maize. According to the pKa of the acid and conjugate base NH\(_4^+\)/NH\(_3\), i.e.

9.25, the ammonia was mainly in the form of NH\(_4^+\) ions in this experiment, with no possible stripping

in the gas phase. These low concentrations of NH\(_4^+\) are below the concentrations known to

inhibit/impact the microbial digestion activity, since the lowest inhibition values collected from the

literature are of the order of 600 mg.L\(^{-1}\) (Karthikeyan and Visvanathan, 2013).

In both experiments (broken maize vs. cattle manure), in pH conditions suitable for AD, the presence

of MKAA led to a significant decrease in the concentration of NH\(_4^+\) ions. The hypotheses and possible

mechanisms between NH\(_4^+\) ions and the MKAA are detailed in Giroudon et al. (2021b).
4.2 Microbial communities associated with binder materials

4.2.1 Microbial communities with sessile and planktonic lifestyles

In our study, a strong presence of the *Clostridium* genus was found in the biofilm and even more in the strongly adhered fraction of the biofilm, i.e. the layers closest to the solid material surface. A similar finding has already been documented for a microbial biofilm formed on cellulose particles during the anaerobic digestion of the particles in a batch process of particulate cellulose AD. Hydrolytic bacteria colonized the particles during the first 14 days, then their number decreased to be ultimately supplanted by methanogenic archaea (Song et al., 2005). In our study, the methanogenic OTUs were present in both sessile and planktonic communities. Our samples were exposed for almost 2.5 times as long as those of Perez et al. (2021), who recently compared the microbial diversity of microbial communities collected from a fermenting biowaste, inoculated early with activated sludge, and on CEM I cement pastes immersed in the same fermenting biowaste for up to 15 weeks at 37 °C. They found that acidogenic bacterial populations were enriched in the community colonizing the CEM I surface, especially members of the *Clostridium* genus. In the liquid fraction, methanogens and acetogens were predominant. The longer exposure time of binder materials in our study probably offered both planktonic and sessile communities the possibility to mix and form a thicker, hierarchically organized biofilm on binder surfaces, explaining why, in our case, methanogenic and acetogenic populations were present in the biofilm.

The presence of the genus *Clostridium* in the strongly adherent layers of the biofilm might be interpreted as being due to its rapid implantation occurring very early in the cycle of biofilm formation on the surface of the binders. Following its installation on the surface of the solid pastes, its acidogenic metabolism would then lower the pH on the binder surface and make the pH conditions more suitable for the integration of both methanogenic and acetogenic microorganisms in the biofilm community. Consequently, this scenario is particularly detrimental to the cementitious
material, as the high local production of microbial acids on the surface would promote the acid
tack of the binder matrices and significantly increase the biodeterioration kinetics.

4.2.2 Material influence on the microbial communities

For all samples, regardless of the type of material, the microbial community identified corresponded
to a community capable of performing the four steps of anaerobic digestion and is commonly found
in anaerobic digestion environments (Venkiteshwaran et al., 2015).

A previous work immersing CAC, CEM I and CEM III cement pastes in a lab scale AD bioprocess
system showed a short-term inhibition of the microbial colonization of CAC material that
disappeared after 10 weeks (Voegel et al., 2020). In the present study, no significant difference
between CAC samples and CEM I, CEM III and planktonic samples was observed. This tends to
confirm that there is no inhibition caused by the CAC, for exposure times longer than 10 weeks
(Voegel et al., 2020).

The specific change in the microbial community in relation to the presence of the MKAA material is
most probably related to the lower NH$_4^+$ concentration observed in the reactors containing MKAA
material. This drop in NH$_4^+$ concentration is the only environmental variable that differs when the AD
batches with MKAA or with the other binder materials are compared. Sodium concentration was also
different when the AD bioprocess evolution with MKAA and with the other materials were
compared. But the values were really close to the concentration measured in the control. Although
NH$_4^+$ is a very common inhibitor of the AD bioprocess at concentrations above 600 mg.L$^{-1}$
(Karthikeyan and Visvanathan, 2013), the effects of low NH$_4^+$ concentrations on AD microbial
communities are not documented in the literature. It is therefore not entirely possible at this stage to
state with absolute conviction that NH$_4^+$ is responsible for this marked shift in the microbial
community. A dedicated comparative study involving dynamic monitoring of an anaerobic digestion
microbial community exposed to low or high concentrations of NH$_4^+$ could help to confirm the key
role of NH$_4^+$. Nevertheless, CH$_4$ production was approximately the same in all BMP reactors,
indicating that, although the material had an influence on the microbial populations, it did not affect the main function of anaerobic digestion.

### 4.3 Deterioration mechanisms of the material samples

#### 4.3.1 CEM I and CEM III

Both CEM I and CEM III pastes showed mechanisms corresponding to a combination of leaching and carbonation, with a phosphorus enrichment in the external zone, as identified in other studies evaluating deterioration mechanisms of cementitious materials by biowaste in anaerobic digestion (Bertron et al., 2017; Giroudon et al., 2021a; Voegel et al., 2019, 2016). However, unlike the pastes immersed in inoculated cattle manure (Giroudon et al., 2021a), the CEM I and CEM III samples did not show identical degradation mechanisms, since particular chemical changes were observed on the CEM III pastes. Contrary to the findings of Koenig and Dehn’s study (2016), where the use of low-clinker binders with blast-furnace slag in the liquid fraction of a pilot scale fermenter led to a reduction in depths of modification, the modified depths of the CEM III paste were twice those of CEM I.

The CEM I paste showed deterioration mechanisms similar to the ones encountered for acids with soluble salts and for NH$_4^+$ attacks, with a decalcification and gradual dissolution of the initial mineralogical phases (Bertron et al., 2004; Bertron and Duchesne, 2013; Duchesne and Bertron, 2013; Escadeillas, 2013). However, despite the high concentrations of VFA (maximum VFA concentrations between 1 and 3 g.L$^{-1}$ at the beginning of the cycles), it appears that carbonation played an important role in the chemical and mineralogical changes of the paste. Calcium carbonates were detected in the outer layers, and the degraded depth did not increase as the experiment progressed. This could be explained by the clogging of the cementitious matrix by calcium carbonates (Baroghel-Bouny et al., 2008; Shah et al., 2018), leading to a slowing of the ingress of aggressive agents'.
Carbonation phenomena are linked to the microbial production of CO\textsubscript{2} in the liquid fraction of the fermenting biowaste. Dissolved CO\textsubscript{2} reacts with calcium released by the dissolution of cementitious matrix phases to form calcium carbonates. However, the analyses of the liquid fraction of the fermenting broken maize showed concentrations of total inorganic carbon similar to or even lower than those reported by Giroudon et al. (2021a), with values of about 1000 mg.L\textsuperscript{-1}. At the equilibrium, for the experiment with broken maize, the total inorganic carbon concentrations in the liquid fraction were lower due to a lower concentration of CO\textsubscript{2} in the gas phase and a lower pH in the liquid fraction. However, the quantity of CH\textsubscript{4} produced in the course of the experiment was significantly higher in the case of the broken maize. Thus, even for lower observed concentrations of CO\textsubscript{2}, its microbial production was actually much higher during the digestion of the broken maize, which induced a much greater flow of CO\textsubscript{2} in the liquid fraction in contact with the materials. This, in turn, could explain the predominant effect of carbonation in this environment. This was confirmed by experiments performed with chemical metabolites alone (Giroudon et al., 2021b) where the authors immersed Ordinary Portland Cement pastes in single-compound-based solutions.

Carbonation phenomena were also identified in the CEM III paste, but the modified depth increased significantly with time. The differences of behaviour between the CEM I and the CEM III pastes could be linked to the initial greater porosity of the CEM III paste or to the poorer performance of slag cements toward carbonation (Osborne, 1999).

In Portland based cements, portlandite is an abundant constituent of the paste and is the hydration product that reacts the most readily with carbon dioxide (Galan et al., 2015; Thiery et al., 2007). Its carbonation results in the precipitation of calcium carbonates in the pore network, decreasing the total pore volume and shifting the pore size distribution curve toward smaller pore diameters (Šavija and Luković, 2016). Moreover, the carbonation in Portland cement also causes a loss of pore connectivity (Han et al., 2015).
The C-S-H decomposition by carbonation consists of the decalcification of the C-S-H gel with the decrease of the Ca/Si ratio and can lead to a silicate polymerisation and to the formation of an amorphous silica gel and calcium carbonates (Li et al., 2017; Sanjuán et al., 2018; Šavija and Luković, 2016; Steiner et al., 2020). Furthermore, it appears that the C-S-H decomposition increases with decreasing Ca/Si ratio (Sevelsted and Skibsted, 2015). For extensive decalcification, the precipitation of calcium carbonate polymorphs and the formation of a silica gel result in significant shrinkage, loss of cohesion, increase in the number of pores and increased porosity (Li et al., 2017; Nedeljković et al., 2018; Puertas et al., 2006; Sanjuán et al., 2018).

The use of GGBS as a supplementary material leads to a reduced portlandite content and a lower Ca/Si ratio in the C-S-H gel (Lothenbach et al., 2011). Thus, the carbonation of slag cements leads to a decrease of their micro-mechanical properties (Nedeljković et al., 2018) and to the coarsening of their pore structure (Ngala and Page, 1997; Šavija and Luković, 2016). Several authors have reported extensive cracking and increases of the chloride diffusion coefficient and the oxygen permeability with carbonation on blended cements (Borges et al., 2010; Ngala and Page, 1997).

Under these CO$_2$-rich conditions, CEM III cement based matrices, well-known for their interesting performances against acid attacks (Bertron et al., 2005b; Gruyaert et al., 2012; Oueslati and Duchesne, 2014) or in mainly acidic environments (Koenig and Dehn, 2016), seem more sensitive to the aggressiveness of this environment than CEM I cement pastes do, as portlandite is present in lower quantities in the CEM III pastes and does not play its role of a sacrificial phase generating calcium carbonates via its reaction with CO$_2$. The C(-A)-S-H are thus attacked, which damages the paste. The carbonation may have increased the porosity of the CEM III pastes and induced a greater penetration of aggressive agents into the sample, which could explain the chemical and mineralogical changes highlighted above and also the deeper penetration of phosphorous from the liquid fraction into the material. Significant carbonation followed by acidification cycles could also have weakened the material.
4.3.2 MKAA

MKAA pastes showed only a very low degraded zone over the last hundreds of microns, as in the previous study by Giroudon et al. (2021a) with cattle manure, and despite more aggressive conditions in terms of acid concentrations and CO$_2$ flow. The chemical profiles highlighted the dissolution of the matrix (Si, Al, Fe, Na), which has already been identified during acid attacks in other studies (Bakharev, 2005; Burciaga-Diaz and Escalante-Garcia, 2012; Ukrainczyk and Vogt, 2020). Thus, in the case of MKAA, the acid attack seems to have a predominant effect. According to Burciaga-Diaz and Escalante-Garcia (2012), the deterioration of the paste in an acidic environment is due to the destruction of the geopolymeric structure and the release of Na, Al and Si into the solution. Several authors (Bakharev, 2005; Burciaga-Diaz and Escalante-Garcia, 2012) agree that it causes the breakdown of the aluminosilicate network of geopolymers. Nevertheless, in this environment, the MKAA did not show any sign of intense cracks or lack of mechanical strength and showed really interesting behaviour.

5 Conclusion

This study investigated the biogeochemical interactions between fermenting broken maize and binder materials intended for anaerobic digester structures. Four types of binder materials were inserted into reactors inoculated with broken maize during five consecutive cycles of anaerobic digestion (245 days). The presence of the cement pastes CEM I, CEM III and CAC did not influence the AD bioprocess in terms of pH, production of VFA and NH$_4^+$ ions, production of CH$_4$ nor did it influence microbial populations. CEM I and CEM III pastes nevertheless experienced biodegradation, with phenomena of decalcification, carbonation and enrichment in P$_2$O$_5$ on the surface. The CEM III paste generated a different zonation and a greater modified depth at the end of the experiment, which was probably linked to the high sensitivity of the slags when exposed to carbonation.

In contrast, alkali-activated metakaolin-based geopolymer showed better behaviour when faced with the aggressive environmental conditions, and exhibited a small modified depth, with a thickness of
several hundred micrometres only at the end of the experiment. The presence of this material led to very low NH$_4^+$ concentrations, probably due to these adsorption capacities, which could be beneficial for the treatment of nitrogen-rich biowaste. Moreover, the presence of the geopolymer resulted in a different microbial population (absence of the genera *Methanobacterium* and *Aminomonas*) and a slightly higher pH, although the CH$_4$ production was similar to that of other BMP reactors. Finally, differences were observed between sessile and planktonic populations in all BMP reactors, with more acidogens in the biofilm, mainly members of the genus *Clostridium*. The main difference observed was a greater presence of acidogens within the depth of the biofilm, especially members of the genus *Clostridium*, which were frequently encountered in the strongly adhered biofilm formed on all the materials.

**CRediT authorship contribution statement**

Marie Giroudon: Conceptualization; Methodology; Investigation; Validation; Writing - Original Draft; Writing - Review & Editing; Visualization

Cédric Perez: Conceptualization; Methodology; Investigation; Validation; Formal analysis; Writing - Original Draft; Writing - Review & Editing; Visualization

Matthieu Peyre Lavigne: Conceptualization; Methodology; Investigation; Resources; Writing - Review & Editing; Supervision

Benjamin Erable: Conceptualization; Methodology; Resources; Writing - Review & Editing; Supervision

Christine Lors: Conceptualization; Methodology; Writing - Review & Editing; Supervision

Cédric Patapy: Conceptualization; Methodology; Resources; Writing - Review & Editing; Supervision
Declaration of competing interest

The authors declare that they know of no competing financial interests or personal relationships that could have influenced the work reported in this paper.

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Appendix A: Evolution of the VFA concentrations during the five cycles of AD in the BMP reactors, with or without binder materials. Mean values of the two duplicate BMP reactors are presented with the standard deviations.
Appendix B: Heatmap with colour scale of the next generation sequencing of the 16S DNA from planktonic biomass samples and “poorly” and “strongly” adhered biomass taken from the surface of cementitious material samples and after 5 cycles of anaerobic digestion – L: weakly adhered biofilm; S: Strongly adhered biofilm; P: Planktonic; only OTUs with an abundance of at least 1% in at least one sample are shown.
Appendix C: Chemical (EPMA) and mineralogical (XRD) changes in the CAC pastes after 3 and 5 cycles of anaerobic digestion of broken maize – Ca: calcite; Ar: aragonite – Bold characters = intensification of the XRD signal in comparison with the deeper zone; Parentheses = significantly lower intensity of the XRD signal in comparison with the main phase
Appendix D: Chemical composition profile (EPMA), according to the distance from the surface, of the CEM I paste after 5 cycles in the broken maize in digestion.
Appendix E: Mineralogical analyses of the CEM I paste after 5 cycles in the fermenting broken maize
Appendix F: Chemical composition profile (EPMA), according to the distance from the surface, of the CEM III paste after 5 cycles in the fermenting broken maize
Appendix G: Mineralogical analyses of the CEM III paste after 5 cycles in the fermenting broken maize, \( \text{C}_4\text{AH}_x \) identification from Bertron et al. (2007b)

RT-LABS Amplification and Sequencing protocol

**Illumina 2-step**

Samples were amplified for sequencing in a two-step process. The forward primer was constructed (5’-3’) with the forward Illumina overhang adapter (TCGTCGGCAGCGTCAGATGTGTATAAGAGACAG) added to the 515F primer (GTGCCAGCMGCGCGGTAA). The reverse primer was constructed (5’-3’) with the reverse Illumina overhang adapter (GTCTCGTGGGCTCGGAGATGTGTATAAGAGACAG) added to the 806R primer (GGACTACHVGGGTWTCTAAT). Amplifications were performed in 25 µL reactions with Qiagen HotStar Taq master mix (Qiagen Inc., Valencia, California), 1 µL of each 5 µM primer and 1 µL of template. Reactions were performed on ABI Veriti thermocyclers (Applied Biosystems, Carlsbad, California) under the following thermal profile: 95 °C for 5 min, then 35 cycles of 94 °C for 30 s, 54 °C for 40 s, 72 °C for 1 min, followed by one cycle of 72 °C for 10 min and 4 °C hold.
Products from the first stage amplification were added to a second PCR based on qualitatively determined concentrations. Primers for the second PCR were designed based on the Illumina Nextera PCR primers as follows: Forward - AATGATACGCGACACGATCTACAC[i5index]TCGTCGGCAGCGTC and Reverse - CAAGCAGAAGACGGCATACGAT[i7index]GTCTCGTGGGCTCGG. The second stage amplification was run in the same way as the first stage except for 10 cycles.

Amplification products were visualized with eGels (Life Technologies, Grand Island, New York). Products were then pooled at equimolar ratios and each pool was size selected in two rounds using SPRiselect Reagent (BeckmanCoulter, Indianapolis, Indiana) in a 0.75 ratio for both rounds. Size selected pools were then quantified using the Qubit 4 Fluorometer (Life Technologies) and loaded on an Illumina MiSeq (Illumina, Inc. San Diego, California) 2x300 flow cell at 10 pM.