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Development of a Modified Plug-Flow Anaerobic Digester for Biogas Production from Animal Manures

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Abstract: Traditional plug-flow anaerobic reactors (PFRs) are characterized by lacking a mixing system and operating at high total solid concentrations, which limits their applicability for several kinds of manures. This paper studies the performance of a novel modified PFR for the treatment of pig manure, characterized by having an internal sludge mixing system by biogas recirculation in the range of 0.270–0.336 m³ m⁻³ h⁻¹. The influence on the methane yield of four operating parameters (recirculation rate, hydraulic retention time, organic loading rate, and total solids) was evaluated by running four modified PFRs at the pilot scale in mesophilic conditions. While the previous biodegradability of organic matter by biochemical methane potential tests were between 31% and 47% with a methane yield between 125 and 184 L CH₄ kgVS⁻¹, the PFRs showed a suitable performance with organic matter degradation between 25% and 51% and a methane yield of up to 374 L CH₄ kgVS⁻¹. Operational problems such as solid stratification, foaming, or scum generation were avoided.

Keywords: plug-flow reactor; anaerobic digestion; animal manures; biogas; unconfined gas injection mixing; mixing recirculation

1. Introduction

Anaerobic digestion (AD) is a biological process in which organic matter breaks down naturally in the absence of oxygen to produce biogas [1]. The most common forms of large-scale anaerobic digesters are the continuously stirred tank reactor (CSTR) and plug-flow reactor (PFR) [2,3]. In a CSTR system, microorganisms are suspended in the digester through intermittent or continuous mixing, which offers good substrate–sludge contact with slight mass transfer resistance, but higher energy is required [4]. In an ideal CSTR, the concentration in any point of the reactor is identical. In contrast, in an ideal PFR, there is no lengthwise mixing of the substrates under digestion as they move through the PFR. Therefore, the concentration distribution is not uniform throughout the reactor [5]. Actually, PFR is characterized by the fact that the flow or fluid through the reactor is orderly with no element of fluid overtaking or mixing with any other element ahead or behind [6]. A PFR system is simple, economical [7], and attractive in terms of efficiency and overall bioconversion compared to CSTR [4].

The expected volatile solid (VS) conversion to gas both in CSTR and PFR with high loads is in the range of 35–45% [8], depending on the substrate. For a similar hydraulic retention time (HRT), PFR usually reaches similar or higher removal efficiencies [9,10] and a better utilization of the volume requirements due to hydraulics and the high solid concentrations [11–14] with lower initial investment and running costs than CSTR systems [4]. Moreover, no short-circuiting can happen in PFRs, and the
operational energy demand is lower than CSTRs due to the mixing and heating requirements [15]. PFRs are increasingly being used, particularly in North America, for high-solid manure digestion systems [12], and these have become an industry standard in the USA for scraped-manure treatments [16]. Among this, there are different examples of efficient pilot and industrial-scale PFRs reported in the literature for AD and the co-digestion of organic wastes [10–13, 17–23]. The PFRs are designed for manure with a high solid content in the range of 11–14% total solids (TS). Typical operational parameters are an HRT between 20 and 50 days, an organic loading rate (OLR) of 1–6 kgCOD m$^{-3}$ d$^{-1}$ and biogas production between 0.4 and 0.8 m$^{3}$ m$^{-3}$ d$^{-1}$ [13,24].

However, PFRs’ drawbacks are a lower mass transfer due to lack of mixing, a lower efficiency when treating low TS content substrates (<10%), thermal stratification and solid sedimentation, or floating/scum-formation problems [13,15,25]. These problems have generated unsuccessful fermentations when treating manure feedstock in conventional biogas plants [26]. It must be noted that neither dung or manure are homogeneous liquids which contain floating solids; in the case of PFRs for such materials, a length to width ratio in the range of 2.0–2.5 has been suggested, although conventional liquid-based PFRs usually have a larger length to width ratio to avoid mixing.

The most common option to prevent solid stratification in PFRs is a partial mixing of the inner content, using mechanical mixers or biogas blower mixers, combined with a recirculation rate (RR) of the effluent [27–29]. Biogas blower mixers by biogas reinjection were found to be the best solution for treating diluted manures (TS of 5%, 10% and 15%) in AD for dairy cattle [27,30]. Appels et al. [30] recommended a mixing biogas flow range of 0.27–0.30 m$^{3}$ m$^{-3}$ min$^{-1}$ for unconfined systems.

Another aspect regarding mixing is the intensity and pattern, but significant differences of values can be found. In general, mixing with an intermediate intensity and an intermittent pattern was concluded to be the most optimal in terms of biogas production [27,31]. In [32], typically, mixing power inputs between 5 and 8 W m$^{-3}$ were reported; Wu [33] found that 0.5–4 W m$^{-3}$ may be sufficient; and both recommendations are below the proposed range of 40–100 W m$^{-3}$ by Couper et al. [34].

This work describes the assessment of a modified PFR equipped with a biogas reinjection system in order to avoid operational problems such as solid stratification, foaming, or scum generation during the anaerobic digestion of animal slurries. The effect of the effluent RR, the HRT, and the inlet TS on the methane yield (MY) was evaluated, four modified PFRs running in parallel at the pilot scale for biogas production from animal manure (pig slurry) in mesophilic conditions.

2. Materials and Methods

2.1. Biomass Sources: Animal Manure and Inoculum

Fresh pig manure (FM), selected as an example of animal slurries, was collected in an intensively rearing pig farm (Girona, Spain) five times for the experiment. FM samples were characterized and used as the influent of four PFR digesters by applying different dilutions with tap water to adjust the inlet total solid content, depending on the experimental conditions (see next section). Both fresh and diluted manure were periodically characterized; Table 1 shows the characterization of FM samples. The net methane yield (L$_{CH_4}$ kgVS$^{-1}$), at 273 K and 1013 hPa, and the biodegradability (% COD) were calculated through biochemical methane potential assays at 35 °C on FM samples [35,36]. Inoculum (5 gVS L$^{-1}$), bicarbonate (1g gCOD$^{-1}$), deionized water (to accomplish a 0.5 L of medium per vial), and FM (5 gCOD L$^{-1}$) were added to 1.2 L glass vials for 30 days at 35 °C. In parallel, controls were prepared to determine the residual biogas production of the inoculum. In order to calculate the methane production rate I and the lag phase ($\lambda$) of the fresh manure (see Table 1), the modified Gompertz Equation (1) was used [37]:

$$R(t) = R_0 \exp\left\{-\exp\left(\frac{R_{max}e}{R_0} \left[\lambda - t\right] + 1\right)\right\}$$

(1)
The inoculum used for the start-up of the digesters and for the biochemical methane potential assays was sampled in a mesophilic CSTR digester of a municipal wastewater treatment plant (WWTP) in Barcelona (Spain).

Table 1. Characterization of fresh pig manure. (FM, fresh manure. COD, chemical oxygen demand. TS, VS, total and volatile solids. TKN, TAN, total Kjeldahl and ammonia nitrogen. R, methane production rate. λ, lag phase. Nd, not determined).

| Fresh Manure (FM) | FM1  | FM2  | FM3  | FM4  | FM5  | Averages |
|------------------|------|------|------|------|------|----------|
| COD (gO₂ kg⁻¹)   | 120.5| 96.1 | 90.4 | 83.1 | 113.5| 100.7 ± 14.1 |
| TS (g kg⁻¹)      | 105.1| 105.6| 99.3 | 104.4| 113.0| 105.5 ± 4.4  |
| VS (g kg⁻¹)      | 75.2 | 84.3 | 79.3 | 75.7 | 80.0 | 78.9 ± 3.3   |
| TKN (gN kg⁻¹)    | 6.0  | 6.1  | 6.0  | 6.8  | 6.3  | 6.2 ± 0.3    |
| TAN (gN kg⁻¹)    | 3.7  | 3.8  | 4.0  | 4.5  | 4.2  | 4.0 ± 0.3    |
| Biodegradability (%COD) | 33% | nd   | 31%  | 47%  | nd   | 37 ± 7.0     |
| CH₄ Yield (m³CH₄ kgVS⁻¹) | 0.184 | nd   | 0.125 | 0.181 | nd   | 0.163 ± 0.027 |
| R (m³CH₄ kg₆COD⁻¹ d⁻¹)  | 5.8  | -    | 6.9  | 14.0 | -    | 8.9 ± 4.1    |
| λ (d)             | 1.9  | -    | 14.0 | -    | -    | -           |

2.2. Pilot-Scale Biogas Plant

A pilot-scale plant comprised of four horizontal PFRs (R1, R2, R3, and R4), a feeding system, a heating system, a biogas flow meter, and a programmable logic controller (PLC) to automate the plant equipment (timing and data acquisition system). All recorded data (temperature profiles of each reactor, biogas flows, inlet flows, etc.) were downloaded periodically as Excel files directly from the panel. Figure 1 shows a scheme of the pilot plant.

Once collected, FM samples were stored at ambient temperature (10–40 °C) in a 1 m³ tank that was diluted with tap water daily just before the feeding of each digester. The weight of tap water and FM added to the dilution tank were registered by a weight cell BL-7 (Sensocar S.A., Terrassa, Spain). The diluted FM was stored in another 1 m³ tank, also at ambient temperature, which was periodically stirred with a waterproof pump GR BluePRO (Zenit Europe, Bascharage, Luxembourg) or a vertical rotor (dilute manure tank) during the storage.

Each PFR (2830 mm length, 646 mm width) consisted of a horizontally oriented U-shaped container of stainless steel and a methacrylate cover, which allowed periodic visual revisions of the inner material, with a working volume of 160 L (total volume of 235 L). Each PFR had its own external gas holder (flexible balloon of 100 L) located on the digester’s cover that, along with the gas headspace of the stainless tank, led to a total volume of 175 L per PFR for gas storage. The biogas composition was measured off-line daily with a gas analyzer, equipped with electrochemical (H₂S, O₂) and dual-beam infrared (CH₄, CO₂) sensors BIOGAS5000 (Geotech Ltd., Conventry, UK). Each digester was heated with individual electric blankets For-Flex Super (Electricfor S.A., Rubí, Spain) and insulated with polyurethane boards. The working temperature was set up at 34 °C and monitored with three temperature probes Pt-100 PR-24-3-100-A-G1/4-6-150 (OMEG, Connecticut, USA) that were distributed regularly along the length of each reactor, controlled by PLC.

The biogas reinjection system was used between the feeding and effluent withdrawal operations. This system includes for each PFR, a silica filter, a gasholder, a compressor V-DTN16 (Elmo Rietschle, Gardner Denver Iberia S.L., Madrid, Spain), and inner 45 polyethylene gas diffusers of 8 mm diameter Tee quick connection (Ningmao Hydraulic Pneumatic Components Factory, Zhejiang, China), which were distributed along the digester floor, near the dividing wall. The gas pipes were made of polyamide. The diffuser distribution fitted well with recommended configurations reported in the literature [38]. The reinjected biogas, introduced perpendicularly to the PFRs, generated turbulence to mix the reactor’s content. The stored biogas was compressed to attain a specific flow range of 0.270–0.336 m³ m⁻³ h⁻¹ and intermittently released (2 min h⁻¹ and 8 times d⁻¹), with the gas flow range being 4–5 m³ h⁻¹.
Safety valves were located along the cover of each digester in order to keep the internal pressure of each PFR ≤20 mbar (differential pressure).

Feeding, effluent recirculation and effluent withdrawal were done manually once per day, from Monday to Friday, dividing the loading equally over the 5 days. Daily flows were recorded directly in the PLC.

![Scheme of the pilot plant](image)

**Figure 1.** Scheme of the pilot plant: Numbers: (1) influent tank; (2) effluent tanks; (3) gas holder; (4) biogas flow-meter. Nomenclature: IN, influent; EF, effluent; B, biogas.

Sampling ports were located in the cover and in the bottom of the tank, allowing the collection of sludge samples from the initial, intermediate, and final points of the digesters. Influen ts and effluents of each PFR were characterized once per week by their content of total chemical oxygen demand (COD), total solids (TS), volatile solids (VS), total Kjeldahl nitrogen (TKN), and total ammonium nitrogen (TAN). Effluents were also characterized by pH, total and partial alkalinities (TA, PA), and volatile fatty acid (VFA) concentration once a week. All parameters were determined following standard methods [39], except for COD, which was determined as per Noguerol-Arias et al. [40] and VFA profile (acetic, propionic, i-butyric, n-butyric, i-valeric, n-valeric) which was determined by gas chromatography determined as per Rodriguez-Abalde et al. [35].

2.3. Experimental Conditions

Table 2 shows the recorded experimental conditions, which are also shown in Figure 2, in order to improve the understanding of the different conditions in each period and the dependence between the
different experimental parameters (inlet-TS, HRT, RR and OLR) shown. The experiment was developed in four periods (P1, P2, P3, and P4), being evaluated in terms of MY (LCH4 kgVS⁻¹, expressed at 273 K and 1013 hPa) and organic matter removal efficiency (VS removal). The length of each period was at least 2.5 × HRT or the minimum amount of time to achieve a “stable condition”, which was defined as that moment in which the biogas production and COD concentration in the effluent were inside 15% of the average value [27].

Table 2. Operational conditions. Average values ± Standard Error. Nomenclature: inlet total solids (inlet-TS), hydraulic retention time (HRT), recirculation rate (RR), organic loading rate (OLR). Abbreviations: period (P), average (Av), volatile solids (VS).

| Parameters     | P1           | P2           | P3           | P4           |
|----------------|--------------|--------------|--------------|--------------|
| Time Interval (Days) | 1–56        | 57–120       | 121–186      | 187–255      |
| Digester R1     |              |              |              |              |
| Inlet-TS (%)    | 3.58 ± 0.01  | 5.30 ± 0.54  | 5.85 ± 0.68  | 5.57 ± 1.07  |
| HRT (d)         | 21.08 ± 4.26 | 24.64 ± 2.84 | 25.37 ± 3.60 | 19.94 ± 2.68 |
| RR (%influent)  | 31.17 ± 6.23 | 30.52 ± 4.15 | 33.55 ± 5.51 | 39.74 ± 4.59 |
| OLR (kgVS m⁻³ d⁻¹) | 1.12 ± 0.22  | 1.39 ± 0.15  | 1.50 ± 0.24  | 1.92 ± 0.29  |
| Digester R2     |              |              |              |              |
| Inlet-TS (%)    | 3.68 ± 0.05  | 5.64 ± 0.57  | 5.45 ± 0.99  | 6.21 ± 0.33  |
| HRT (d)         | 20.35 ± 4.08 | 24.54 ± 3.00 | 30.15 ± 3.82 | 31.44 ± 2.54 |
| RR (%influent)  | 31.03 ± 6.21 | 30.53 ± 3.94 | 26.37 ± 4.38 | 32.71 ± 4.93 |
| OLR (kgVS m⁻³ d⁻¹) | 1.14 ± 0.23  | 2.40 ± 0.13  | 2.30 ± 0.60  | 2.90 ± 0.50  |
| Digester R3     |              |              |              |              |
| Inlet-TS (%)    | 3.57 ± 0.00  | 5.60 ± 0.86  | 6.81 ± 1.35  | 3.43 ± 0.18  |
| HRT (d)         | 20.39 ± 4.09 | 28.51 ± 4.41 | 27.06 ± 4.37 | 22.41 ± 2.24 |
| RR (%influent)  | 31.21 ± 6.24 | 15.17 ± 16.15| 40.57 ± 7.99 | 38.73 ± 4.89 |
| OLR (kgVS m⁻³ d⁻¹) | 1.11 ± 0.22  | 1.21 ± 0.18  | 1.61 ± 0.33  | 1.41 ± 0.17  |
| Digester R4     |              |              |              |              |
| Inlet-TS (%)    | 3.10 ± 0.00  | 5.10 ± 0.40  | 5.70 ± 0.69  | 6.12 ± 0.31  |
| HRT (d)         | 20.39 ± 4.09 | 25.76 ± 3.57 | 32.01 ± 4.07 | 33.19 ± 3.07 |
| RR (%influent)  | 31.23 ± 6.25 | 24.46 ± 4.17 | 46.08 ± 9.81 | 26.50 ± 4.78 |
| OLR (kgVS m⁻³ d⁻¹) | 1.60 ± 0.20  | 1.60 ± 0.30  | 1.60 ± 0.10  | 2.00 ± 0.20  |

Period P1 consisted in the start-up of all four digesters (R1, R2, R3, and R4), which were inoculated with 150 L inoculum obtained from a CSTR operating at a WWTP in Barcelona (Spain). Conservative conditions were established for 25 days. Different conditions were applied for each reactor in the next periods, P2 to P4, by changing inlet-TS, HRT, and RR values. The inlet-TS content was in the range of 3.0% and 7.0%, as the representative TS range of FMs. The HRT was between 10 and 40 days, as these values were reported as the minimum for anaerobic digesters in the literature [10,12]. The RR was fixed between 20% and 50% of the influent flow to ensure a stable process through the anaerobic biomass recirculation into the reactors.

Digesters R1 and R2 were used to evaluate the effect of changing the inlet flow and/or the inlet TS concentration. In this way, in R1 a progressive increase of the inlet flow and inlet-TS content through the periods were done, which increased the OLR and decreased the HRT from period P2 to P4.

In R2, the OLR was promptly doubled by pulse additions in each period. Two additions of dilute FM were done in period P2 reaching 2.5 kgVS m⁻³ d⁻¹ these days. Three pulses of different VFA (acetic acid 30 g pulse⁻¹, propionic acid 20 g pulse⁻¹, or butyric acid 16 g pulse⁻¹) were done in period P3 reaching 2.5 kgVS m⁻³ d⁻¹ these days. Finally, three glucose additions (30 g pulse⁻¹) in period P4 were done reaching 2.5 kgVS m⁻³ d⁻¹ these days. These increases in OLR were performed to evaluate resilience of the different group of microorganisms involved in the anaerobic digestion process: FM pulses help to evaluate reactor response focusing mainly on hydrolytic and acidogenic bacteria, VFA pulses help to evaluate reactor response focusing on methanogenic bacteria, whereas glucose pulses help to evaluate reactor response focusing on acidogenic and acetogenic bacteria.
Figure 2. Operational parameter: Hydraulic retention time (HRT), recirculation rate (RR), inlet total solids content (inlet-TS), and organic loading rate (OLR) per reactor and period. (a) R1; (b) R2; (c) R3; (d) R4. Note: Vertical grey lines denote periods; black lines denote average HRT per period; green lines denote average RR per period; blue lines denote average inlet-TS per period; red lines denote average OLR per period.

Digester R3 was used to evaluate possible problems at full-scale plants, such as pumping failures or feed blockages. In this way, during period P2, both feeding and recirculation of the effluent were stopped for 1 week and RR was stopped for another week at the end of this period.
Finally, digester R4 was used as a control by maintaining stable conditions, with an OLR of 1.6 kgVS m$^{-3}$ d$^{-1}$ and a HRT of 24–36 days. The only parameter changed was the effluent RR, varying from 50% to 25% depending on the experimental period (see Table 2).

Periodically, a mass balance calculation was done in terms of total N and COD. For that purpose, an accumulation term was estimated based on total N balance (since total N concentration is conservative in AD processes assuming a constant growth of the involved microorganisms). This accumulation was included in the COD mass balance in order to calculate both methane and biogas production. In this regard, Kinyua et al. [41] included N related compounds, such as struvite, and the presence of a dead volume inside tubular anaerobic digesters. Similarly, Jagadish et al. [26] discovered a floating layer that remained during the entire fermentation period inside horizontal PFRs when digesting a chopped blend of herbaceous terrestrial weeds and leaf biomass.

2.4. Statistical Analysis

The statistical analysis was done with the SPSS Statistics v.20.0.0 (IBM) software (Armonk, N.Y., USA). An ANOVA and HSD Tukey post-hoc analyses were performed to determine significant differences ($p < 0.05$) for variables whose data were normally distributed and with equal variances. For data of variables with heterogeneous variances, the Welch statistic and Games–Howell post hoc analyses were used to determine which groups were significantly different ($p < 0.05$). Linear correlations between control parameters and operation conditions during the assays were studied using the Pearson correlation coefficient ($r$). Correlations were considered significant at $p < 0.05$.

3. Results and Discussion

3.1. Substrate Characterization

The collected FM (Table 1) was representative of typical pig slurry or liquid manure in terms of TS content, as reported in [42]. The TS (9.9–11.3% wet weight) and TKN (6.0–6.8 gN kg$^{-1}$) contents were almost constant among all experiments. The biodegradability of FM ranged between 31–47% COD with a MY between 125–184 L$_{CH4}$ kgVS$^{-1}$. These values were lower than the reported by Møller et al. [43] and Hansen et al. [44] when treating slurries from fattening pigs, with ranges of 47–78% COD and MY of 300–356 L$_{CH4}$ kgVS$^{-1}$. However, the VS and TAN contents of fresh FM varied, which could explain the different qualities regarding the biodegradability. This result is coherent with other references in the literature where the storage time in the farm facility was related with the quality and methane potential of animal slurries [45].

3.2. General Plant Performance

Pilot plant operation showed several challenges due to the complex automation and the operation of four reactors simultaneously.

During P2, P3, and P4, some problems regarding the recorded biogas flow with the flowmeter were observed, and the biogas production was lower than the calculated values by COD balance (Figure 3). Two main reasons for this were identified. The first one was caused by the compressor operation for the biogas reinjection. Because of it, a regular overpressure in the digesters was observed, causing the opening of the safety gas valves in order to keep the internal pressure below 20 mbar and losing some biogas without being recorded. Secondly, the pipe length between the gas extraction points and the gas flowmeter changed depending on the digester, leading also to some biogas losses and condensate generation inside of these pipes. For these reasons, the recorded biogas flows in the beginning (P1) were well measured in all reactors, but immediately lower in the next periods.

Reactor R4 had an extra time operation of 30 days. Before this period, a maintenance in gas piping was performed in order to avoid problems regarding the recorded biogas flow. After this maintenance, the MYs in both reactors suddenly increased (Figure 3), following a similar trend to the calculated values by COD balance. This showed the difficulty of operating an automated pilot biogas plant.
comprised of four different reactors, but also confirmed that the COD balance performed during the periods in which the biogas flow meter was not working appropriately, was correct and exact enough to accept methane yields obtained through this method.

Figure 3. Evolution of organic matter removal (VS removal) and methane yield (MY) per reactor and period: (a) R1, (b) R2, (c) R3, (d) R4. Note: Vertical grey lines denote periods; green lines denote average values of VS removal efficiency for each whole period; red and blue lines denote average values of MY registered by flowmeter and COD balance, respectively, for each whole period.

The biogas reinjection flow was well adapted to the literature [38] in the range of 0.270–0.336 m$^3$ m$^{-3}$ h$^{-1}$. Some of the 45 valve diffusers per reactor were clogged (10–15%), as reported in the literature [32].
but no foaming or scum problems were observed in the digesters. No corrosion of gas piping and equipment was observed.

In general, the reactors showed appropriate performance, avoiding common problems associated with the operation of plug flow reactors at low inlet-TS concentrations, such as solid stratification (sedimentation or crust formation), foaming or scum generation.

3.3. Pilot Plant Monitoring

3.3.1. Biogas Composition and Organic Matter Removal

Table 3 shows the results obtained throughout the experiment regarding methane yield and methane volumetric yield, organic matter removal (VS removal) and biogas composition.

Table 3. Control parameters (methane productivity and yield, and organic matter removal). Nomenclature: methane yield (MY), volumetric methane yield (VMY), period (P), average (Av).

| Parameters                  | Time Interval (Days) | P1        | P2        | P3        | P4        |
|-----------------------------|----------------------|-----------|-----------|-----------|-----------|
|                             | 1–56                 | 57–120    | 121–186   | 187–255   |
| Digester R1                 |                      |           |           |           |           |
| MY (m³CH₄kgVS⁻¹)            | 0.15 ± 0.02          | 0.16 ± 0.10| 0.21 ± 0.10| 0.10 ± 0.02| 0.02 ± 0.02|
| VMY (m³CH₄m⁻³d⁻¹)          | 0.19 ± 0.01          | 0.21 ± 0.12| 0.30 ± 0.14| 0.04 ± 0.04|
| CH₄ (%)                    | 63.01 ± 1.52         | 62.97 ± 3.34| 61.52 ± 1.62| 57.22 ± 2.02|
| O₂ (%)                     | 0.60 ± 1.12          | 0.26 ± 0.32| 0.71 ± 0.71| 1.14 ± 1.15|
| VS removal (%VS inlet)     | 48.86 ± 16.53        | 50.77 ± 18.08| 49.68 ± 16.70| 25.12 ± 30.01|
| Digester R2                 |                      |           |           |           |           |
| MY (m³CH₄kgVS⁻¹)            | 0.16 ± 0.02          | 0.17 ± 0.04| 0.20 ± 0.06| 0.19 ± 0.04|
| VMY (m³CH₄m⁻³d⁻¹)          | 0.20 ± 0.01          | 0.22 ± 0.06| 0.25 ± 0.10| 0.23 ± 0.06|
| CH₄ (%)                    | 61.32 ± 2.10         | 62.06 ± 3.75| 62.90 ± 1.49| 58.81 ± 3.32|
| O₂ (%)                     | 0.80 ± 0.54          | 0.31 ± 0.37| 0.43 ± 0.42| 1.04 ± 1.40|
| VS removal (%VS inlet)     | 45.34 ± 9.44         | 50.72 ± 10.05| 48.78 ± 13.83| 48.91 ± 14.08|
| Digester R3                 |                      |           |           |           |           |
| MY (m³CH₄kgVS⁻¹)            | 0.24 ± 0.03          | 0.18 ± 0.12| 0.37 ± 0.05| 0.22 ± 0.11|
| VMY (m³CH₄m⁻³d⁻¹)          | 0.30 ± 0.02          | 0.21 ± 0.13| 0.57 ± 0.04| 0.42 ± 0.13|
| CH₄ (%)                    | 60.12 ± 2.21         | 59.26 ± 3.29| 62.51 ± 2.33| 59.38 ± 4.89|
| O₂ (%)                     | 0.41 ± 0.76          | 0.45 ± 0.57| 0.23 ± 0.32| 1.06 ± 2.50|
| VS removal (%VS inlet)     | 46.38 ± 20.11        | 46.08 ± 29.33| 46.47 ± 20.03| 51.38 ± 26.01|
| Digester R4                 |                      |           |           |           |           |
| MY (m³CH₄kgVS⁻¹)            | 0.29 ± 0.03          | 0.14 ± 0.12| 0.22 ± 0.04| 0.12 ± 0.04|
| VMY (m³CH₄m⁻³d⁻¹)          | 0.23 ± 0.01          | 0.16 ± 0.26| 0.05 ± 0.16| 0.05 ± 0.05|
| CH₄ (%)                    | 56.49 ± 1.76         | 57.54 ± 1.95| 55.01 ± 5.01| 57.23 ± 4.34|
| O₂ (%)                     | 1.52 ± 0.81          | 0.43 ± 0.44| 3.13 ± 3.34| 0.63 ± 1.24|
| VS removal (%VS inlet)     | 43.62 ± 20.75        | 42.36 ± 32.29| 38.57 ± 14.66| 37.01 ± 18.53|

Biogas composition was similar for each period and reactor, with the highest methane content of 63.0 ± 3.3% in R1 during the P2 period and a lowest methane content of 55.0 ± 5.0% in R4 during the P3 period (Table 2). Biogas produced from animal slurries typically has a methane content of 60% [46,47] and it was shown that R1, R2, and R3 accomplish that during the P2 and P3 periods, but not during the P4 period, which is also true for the R4 reactor in any period. The unconfined gas injection mixing introduced O₂ as a consequence of air inlet in the digesters. The maximum and minimum average amounts of O₂ detected were 3.13 ± 3.34% in R4 during the P3 period and 0.23 ± 0.32% in R3 during the P3 period, respectively. In general, the O₂ concentration was higher during the previous hours at 10:00 a.m., in accordance with the reinjection mixing cycles, with the exception of R3, in which a higher H₂S concentration was evaluated during P3 and P4. It was assumed that R3 mixing was lower than other reactors, due to several mixing stoppings for different reasons such as leakage verifications, O₂ concentration determination, or compressor maintenance. Although there is no correlation between O₂ and H₂S concentrations with experimental data (see Table 4), a decrease in H₂S concentration is observed in the reactors after an increase of O₂ concentration. Figure 4 shows
O$_2$ and H$_2$S concentrations in each reactor before and after 10:00 a.m., the time in which mixing was stopped and the feeding and discharge operations were performed. This shows an average difference decrease before and after 10:00 a.m. of 602, 263, 473, and 589 ppm of H$_2$S in reactors R1, R2, R3, and R4, respectively.

**Figure 4.** Evolution of H$_2$S and O$_2$ daily concentrations per reactor: (a) R1; (b) R2; (c) R3; (d) R4. Note: Vertical grey lines denote measurements done before and after 10:00 am; blue squares denote H$_2$S concentrations; red circles denote O$_2$ concentrations.
Table 4. Correlation coefficients (Pearson coefficients) for parameters. Note: * Significant correlation at 0.05 levels (dark grey). ** Significant correlation at 0.01 levels (light grey). Notes: (1) OLR in gVS m\(^{-3}\) d\(^{-1}\). (2) Methane yield (MY) in L\(_{CH4}\) gVS\(^{-1}\). (3) Volumetric methane yield (VMY) in m\(^3\)CH\(_4\) m\(^{-3}\) d\(^{-1}\). (4) Total Kjeldahl and ammonia nitrogen (TKN, TAN) in mgN L\(^{-1}\). (5) Total volatile fatty acids (VFA) in g\_{eq-acetic acid} L\(^{-1}\). (6) Intermediate alkalinity (IA) in gCaCO\(_3\) L\(^{-1}\).

| Parameter | Period | OLR (1) | HRT (d) | VMY (2) | MY (3) | RR (%) | NTK (4) | TAN (4) | pH | IA (5) | VFA (6) | VS-Rem (%) | CH\(_4\) (%) | CO\(_2\) (%) | H\(_2\)S (ppm) | COD-Rem (%) |
|-----------|--------|---------|---------|---------|--------|--------|---------|---------|----|--------|---------|------------|-------------|-------------|--------------|-------------|
| Period    | 1      | 0.411 ** | -0.115 | -0.150 | -0.370 ** | 0.100 | 0.410 ** | 0.461 | 0.412 ** | -0.311 | -0.356 ** | 0.461 | 0.412 ** | -0.213 | -0.335 ** |
| OLR (1)   |        |         | -0.800 ** | -0.150 | -0.279 | 0.438 ** | 0.636 ** | 0.139 | 0.392 ** | 0.684 ** | -0.231 | 0.062 | -0.073 | 0.467 ** | -0.339 ** |
| HRT (d)   | 1      | 0.190 | -0.211 ** | -0.323 | -0.567 ** | -0.175 | -0.100 | -0.592 ** | -0.331 | -0.201 | 0.205 | -0.142 | 0.279 |            |              |
| VMY (2)   | 1      | 0.972 ** | 0.051 | -0.448 ** | -0.243 | -0.240 | -0.191 | -0.275 | 0.596 ** | -0.001 | -0.005 | 0.271 | 0.812 ** |        |              |
| MY (3)    | 1      | -0.053 | -0.502 ** | -0.299 ** | -0.247 | -0.233 | -0.329 ** | 0.658 ** | -0.073 | 0.070 | 0.168 | 0.866 ** |      |              |
| RR (%)    | 1      | 0.236 | -0.004 | 0.021 | 0.159 | 0.217 | -0.156 | 0.030 | -0.038 |      |              |
| NTK (4)   | 1      | -0.204 | 0.028 | 0.259 | 0.515 ** | -0.277 | 0.201 | -0.200 | -0.062 |      |              |
| TAN (4)   | 1      | 0.000 | 0.578 ** | 0.325 | -0.623 ** | -0.134 | 0.129 | 0.220 | -0.474 ** |      |              |
| pH        | 1      | -0.277 | 0.018 | -0.122 | 0.008 | -0.002 | -0.251 | -0.343 ** |      |              |
| IA (5)    | 1      | 0.682 ** | -0.367 | -0.283 | 0.277 | 0.219 | -0.266 |      |              |
| VFA (6)   | 1      | -0.456 ** | -0.217 | 0.217 | -0.342 ** |      |              |
| VS-Rem (%)| 1      | 0.112 | -0.113 | 0.037 | 0.818 ** |      |              |
| CH\(_4\) (%) | 1      | -1.000 ** | -0.045 | -0.019 | 0.023 | 0.018 |      |              |
| CO\(_2\) (%) | 1      | 0.034 |      |      |      |              |
Because of operational problems regarding biogas measurements, biogas production was estimated using the COD balance. This methodology has been widely used in many publications and textbooks [48–50], and we validated the use of COD balance in the present work by comparing the MY values obtained during period P1 from both methods, by flowmeter and by COD balance, as in this period the flowmeter ran correctly. MY averages values are shown in Figure 3, and how a good correlation between them during this period P1 (0.15 ± 0.02, 0.16 ± 0.02, 0.24 ± 0.05, 0.29 ± 0.03 m³CH₄ kgVS⁻¹ in reactors R1, R2, R3, and R4, respectively) can be observed. This is also shown during period P4, when gas piping maintenance was performed in R4 and flows of both methods (flowmeter and COD balance) were similar. Figure 5 shows the average MY values for all periods and reactors for the different operational conditions (HRT, RR, inlet-TS, and OLR). The maximum MY values obtained were in R3 during P3, with an average of 373.7 ± 49.7 LCH₄ kgVS⁻¹, together with the lowest O₂ concentrations and an organic matter removal of 46.5 ± 20.0% VS. The minimum MY values obtained were in R1 during P4, with an average of 23.5 ± 23.0 LCH₄ kgVS⁻¹, matching with the lowest organic matter removal of 25.1 ± 30.0% VS.

Figure 5. Comparison of methane yields (MY) and the hydraulic retention time (HRT), the recirculation rate (RR), the inlet total solids content (TS) or the organic loading rate (OLR). Note: Vertical grey lines denote reactors; black line denotes average values of MY registered in a whole period; blue, red, green and yellow bars denote average values of a whole period (P1, P2, P3, and P4, respectively) of the corresponding operational parameter per reactor.

3.3.2. VFA, pH, and Alkalinity Profiles

VFA, pH, and alkalinity are important control parameters during AD [47]. The pH average values in all reactors were slightly higher (8.1 ± 0.2, 8.0 ± 0.1, 8.0 ± 0.1 and 8.0 ± 0.2 in reactors R1, R2, R3, and R4, respectively) than the range for the normal operation of AD, between 7.5 and 7.8. This was probably due to the highest TA values registered during all periods, with averages of 11.1 ± 1.9, 10.1 ± 0.6, 10.6 ± 0.7, 10.6 ± 1.1 gCaCO₃ L⁻¹ in reactors R1, R2, R3, and R4, respectively. Moreover, changes in VFA were shown to have a significant effect on TA, with the biggest fall in R1 during P3, when a decrease of VFA from 1859 to 333 mg L⁻¹ caused a decrease of TA from 15.7 to 9.2 gCaCO₃ L⁻¹. The percentages of acetic acid were 91 ± 10.5, 98 ± 2.5, 94 ± 9.7, and 92 ± 16.6 in reactors R1, R2, R3, and R4, respectively. These were higher than other studies reported in the literature, with ranges of 35% to 75% [51] or 60% to 75% [52]. Angelidaki et al. [53] found a common value of 1.5 g L⁻¹
was considered to be limiting for a stable operation of AD reactors. It can be seen that total VFA concentrations in effluent (Figure 6) were lower than 2 g·L⁻¹ in all reactors with the exception of R1 when it was operated at high OLR during the P4 period. The average total VFA values were 1.0 ± 1.0, 0.4 ± 0.2, 0.4 ± 0.5, and 0.6 ± 0.4 g·L⁻¹ in reactors R1, R2, R3, and R4, respectively.

Figure 6. Evolution of volatile fatty acids (VFA) per reactor and period: (a) R1, (b) R2, (c) R3, (d) R4. Symbols: Acetic (●), propionic (■), i-butyric (▲), n-butyric (○), i-valeric (x), n-valeric (Δ), i-caproic (■), n-caproic (□).

3.3.3. Nitrogen (TKN, TAN) Profile

In general, the inhibition of the anaerobic digestion process has been reported to start at a TAN level of 1.5–2.0 gN·L⁻¹. However, in the case of pig manure, it has been reported to start at 3.1 gN·L⁻¹. The registered TAN averages values were 2.1 ± 0.2, 2.1 ± 0.1, 2.7 ± 0.2, 2.9 ± 0.3 gN·L⁻¹ in reactors R1, R2, R3, and R4, respectively. The maximum amount of TAN registered (3.8 gN·L⁻¹) was in reactor R1 during P4, where a lower MY value was obtained (23.5 ± 23.0 L₈CH₄ kgVS⁻¹). That would confirm the starting TAN level of 3.1 gN·L⁻¹ to be inhibitory for the process.

3.3.4. Effect of Recirculation

Figure 2 shows the operational parameter evolution for each period and reactor. The reported literature has confirmed positive effects by varying RR in AD, such as the liquefaction and inoculation of the fresh biogas substrate, the stabilization of biogas synthesis and the optimization of biogas production [54].

When RR decreased, VFA concentration increased. That can be seen clearly in R4, when changing from P2 to P3, the RR variation from 24.5 ± 4.2 to 46.1 ± 9.8% caused a maximum decrease of 90% of VFA. Increases in MY values were observed with higher RR values, as can be seen in R3 and R4 from P2 to P3 (increases of 105.5% and 57.1% in R3 and R4, respectively). Results suggest RR values up
to 40% should be used in order to keep stable conditions of the bacterial population inside the reactor, optimizing biogas production.

3.3.5. Effect of the Organic Loading Rate

In general, the expected trend for HRT and OLR was confirmed in all reactors, as the biogas production increased when increasing HRT or OLR, with some exceptions: R1 during P4, probably due to the ammonia inhibition effects mentioned before. Also increases of OLR in R3 during P2 and in R4 during P2 and P4 did not increase the MY, probably due to the decreased RR in these reactors and periods. An increase of acetic and propionic acid concentrations was also observed with OLR increases, such as in R1 and R2 during both P3 and P4 periods.

3.3.6. Effect of Total Solid Content

As the OLR depends on inlet-TS concentration, similar performances were evaluated. At higher inlet-TS concentrations, higher MY values were obtained. The maximum MY obtained was in R3 during P3 (373.7 ± 49.7 LCH4 kgVS−1) with 6.81 ± 1.35 % TS. H.M. El-Mashad et al. [51] concluded that a higher TS content resulted in higher VFA concentrations. Although there was not a clear correlation in our results between inlet-TS contents and VFA (see Table 4), a similar trend was exhibited as the reported study of Page et al. [55]; when evaluating dairy manure effluent at 2.4% TS, a range of 89–439 mg L−1 of VFA was obtained.

4. Conclusions

A novel modified plug-flow anaerobic reactor for treating animal slurries was developed and tested successfully with appropriate performance. The mixing power inputs and the intermittent mixing pattern allowed for the avoidance of operational problems such as solid stratification, foaming, or scum generation. In general, there was no inhibition due to ammonia concentration, except for reactor R1 during P4. VFA generation allowed stable conditions. The methane yields were optimized during some of the periods, showing how significant the operational parameters evaluated are. However, it seems that complex automation and equipment design involves facing the important problem of the scale factor.

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