1	Assessment of chemical inhibitor addition to improve the gas production from
2	biowaste
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12 Abstract

13 The coexistence of sulphate-reducing bacteria and methanogenic archaea in the reactors during the anaerobic digestion from sulphate-containing waste could favor the 14 accumulation of sulfide on the biogas, and therefore reduce its quality. In this study, the 15 effect of sulphate-reducing bacteria inhibitor (MoO₄⁻²) addition in a two phase system 16 17 from sulphate-containing municipal solid waste to improve the quality of the biogas has been investigated. The results showed that although SRB and sulphide production 18 19 decreased, the use of inhibitor was not effective to improve the anaerobic digestion in a two phase system from sulphate-containing waste, since a significant decrease on 20 biogas and organic matter removal were observed. Before MoO_4^{-2} addition the average 21 values of volatile solid were around 12 g/kg, after 5 days of inhibitor use, those values 22 did exceed to 28 g/kg. Molybdate caused acidification in the reactor and it was 23 24 according to decrease in the pH values. In relation to microbial consortia, the effect of inhibitor was a decrease in Bacteria (44 %; 60% in sulphate-reducing bacteria) and 25 Archaea (38%) populations. 26

Keywords: biomethanization; inhibition; sulphate-containing solid waste; microbial
community structure.

30 **1. Introduction**

Conventional bioconversion of waste in anaerobic digestion (AD) systems is widely 31 recognized (Cuetos et al. 2008; Martín-González et al. 2013; Xing et al. 2014) and is 32 characterized by four steps: hydrolysis, acidogenesis, acetogenesis and methanogenesis. 33 The first three steps are carried out by various bacteria species while the fourth step 34 (methanogenesis) is usually dominated by special microorganisms belonging to the 35 Archaea domain (Zahedi et al. 2016). In the first and second steps, hydrolysis and 36 37 acidification take place by hydrolytic-acidogenic bacteria (HAB), and intermediate products such as volatile fatty acids (VFA), hydrogen (H₂) and carbon dioxide (CO₂) 38 are generated. In the third step, VFA are transformed into acetate, H₂ and CO₂ by 39 acetogenic bacteria. Usually, Propionate-utilizing acetogens (PUA) and butyrate-40 utilizing acetogens (BUA) are the majority of the acetogens in the anaerobic reactors 41 (Mara and Horan 2003). The methanogens occupy the terminal position in the anaerobic 42 food chain and are normally divided into two main groups based on their substrate 43 conversion capabilities. Acetoclastic methanogens (AUM) are able to convert acetate 44 45 into methane and carbon dioxide (Montero et al. 2008). Hydrogenotrophic methanogens (HUM) convert H_2/CO_2 to methane. These species play a key role in the overall process 46 by maintaining the very low partial pressures of H_2 (<10 Pa) necessary for the 47 functioning of the intermediate trophic group, the acetogens, which are responsible for 48 49 the conversion of acids organic and alcohol intermediates to direct methane precursors 50 (Montero et al. 2008).

51 During anaerobic treatment of sulphate-containing wastewaters, sulphate-reducing 52 bacteria (SRB) compete for substrate with other anaerobic bacteria or methanogens. 53 Sulphate competes against organic carbon as an electron acceptor and it leads to the 54 undesirable production of hydrogen sulphide (H₂S).

55 H₂S is a corrosive gas and its presence reduces the potency of biogas as a fuel for 56 boilers or electricity generation in a biogas engine. Besides, it greatly affects the flammability of biogas when used directly in burners (Rasi et al. 2007; Muñoz et al. 57 58 2015). In addition, H₂S causes malodor and health hazards due to the well-known toxicity and material corrosion tendencies (Auguet et al. 2015). The nutritional 59 requirements of SRB include an inorganic electron acceptor which is usually provided 60 61 by sulphate ion and an electron donor which essentially consist of VFA or H₂ and occasionally sugars and long chain fatty acids. 62

Two stages of inhibition exist as a result of sulphate reduction (Chen et al. 2008): (i) 63 64 primary inhibition due to competition for common organic and inorganic substrates from SRB and (ii) secondary inhibition which results from the toxicity of sulphide to 65 various microbial groups. In this regard, many researchers have used molybdate (MoO₄⁻ 66 ²) as sulphide inhibitor where different substrates were utilized (Newport and Nedwell 67 1988; Tucker et al. 1998; Ranade et al. 1999; Isa and Anderson 2005; Predicala et al. 68 2008; Rincon et al. 2008; Biswas et al. 2009; Jesus et al. 2015) and all of them show of 69 70 molybdate successful in inhibiting SRB activity.

Nevertheless, data on the outcome of competition between SRB and the microorganisms mentioned above are contradictory in literature (Chen et al. 2008). The considerable variation in the inhibition/toxicity levels reported for sulphate is due to the complexity of AD process where mechanisms such as antagonisms, synergism, acclimation and competition could significantly affect the phenomenon of inhibition.

Studies involving use of inhibitors to suppress activity of SRB and, consequently,
promote growth of methanogens have been reported in literature (Ranade et al. 1999;
Isa and Anderson 2005; Patidar and Tare 2005; Chen et al. 2008). However, other

79 studies documented that SRB and methanogens could have a symbiosis between them
80 (Vossoughi et al. 2003; Zahedi et al. 2013a; Auguet et al. 2015), although these
81 organisms are constantly competing as electron acceptors. Therefore, the feasibility of
82 using SRB inhibitor for the control of sulphate reduction and the improvement of
83 methane production in biological reactor are not established.

In view of this, the present study was undertaken to investigate the effect of molybdate supplementation on the two-phase dry-thermophilic AD process of sulphate-containing municipal solid waste. Structure and dynamics of the anaerobic consortia developed along the experiment were analyzed by fluorescent in situ hybridization (FISH) employing different oligonucleotide probes.

89 2. Methods

90 2.1 Experimental equipment

Two laboratory-scale continuously stirred tank reactors were employed (Figure 1). The 91 92 first reactor, dedicated to the hydrogen production (HP) (first phase), had a 5.5 liters 93 working volume, while the second reactor (second phase) dedicated to the methane production (MP) had a 5 liters working volume, both heated by recirculating water 94 through a thermostatic jacket. PRECISTERM 6000142/6000389 (SELECTA S.A.) 95 96 baths were used, with a maximum capacity of 7 liters of water. The stainless steel 97 reactors lid have a diameter of 200 mm and contain three openings, one for the biogas outlet, a feed inlet and another opening for the stirring system. The bottoms of the 98 reactors had a discharge valve with a 40 mm i.d., used for sampling. The biogas was 99 100 collected in 40 liter capacity Tedlar (a polyvinyl fluoride plastic polymer) bags. The stirring systems consisted of an IKA EUROSTAR Power Control visc-P4 overhead 101 102 stirrer coupled to a stainless steel blade with scrapers which allows homogenization of the waste at a speed of 23 rpm. The system was fed semi-continuously, once per day, and the hydraulic retention times (HRT) were 1.5 (organic loading rate (OLR) = 57 g/l/d) and 5 d (OLR = 8 g/l/d) for the first and second phase, respectively.

106 **2.2 Inoculum, substrate and feeding**

107 The seed used as acidogenic and methanogenic inoculum were collected from a two 108 phase dry-thermophilic system of urban wastes. The total solid (TS) and volatile solid 109 (VS) concentrations in the second phase (methanogenic inoculum) were 67 g/kg and 33 110 g/kg as against their concentrations in the first phase (acidogenic inoculum) which were 111 82 g/kg and 50 g/kg, respectively.

112 The tested substrate in the first phase was the urban wastes from the 30 mm trommel of 113 the municipal solid waste treatment plant in Cadiz, Spain. The urban wastes was stored in 25 kg drums at - 4° C to avoid AD by the microorganisms found in the solid waste 114 itself (Zahedi et al. 2013b). The TS concentration of the feed of the first reactor was 115 116 adjusted to 20 % (which is characteristic of dry AD) by adding tap water. Composition of the substrate (urban solid wastes and water; 20 % of TS) employed in the first phase 117 118 is shown in Table 1. The substrate used in the second phase was the effluent of the first 119 phase (Table 2). Both reactors were fed once a day (semi-continuous).

120 **2.3 Inhibitor treatment methodology**

The effect of continuous dosing molybdate (MoO4⁻²) (2.5 mM) to improve the performance in two-phase dry-thermophilic AD of sulphate-containing urban waste was realized. No molybdate was added to the first phase, because under acid conditions the biogas was sulphide-free (Zahedi et al. 2013b). The whole experiment length was 50 d. During the first 45 d no inhibitor was used. In these 45 days two different period were considered: startup (0-20 d) and control/stationary phase (20-45 d). On day 46, sodium molybdate (MW=205.92 g/mol), 2.6 g, was added to the digester so as to have 2.5 mM concentration of the inhibitor in the reactor (V= 5 1) (Isa and Anderson 2005). From next day onwards, i.e. from day 47, based on the daily wash out, 0.52 g per 1000 ml of daily feed was added every day, so as to maintain the 2.5 mM inhibitor concentration in the digester.

132 **2.4 Analytical methods**

Total and soluble chemical oxygen demand (TCOD, SCOD), alkalinity, sulphate, VS, 133 134 pH and VFA were performed according to previous studies (Zahedi et al. 2013c; 135 Dahunsi et al. 2016a,b). Determination of total and partial alkalinity and ammonia (NH₄⁺–N) were carried out daily. Fluctuation in the concentration of volatile fatty acids 136 137 (VFA) was determined using a gas chromatography (GC2010) to which was attached a 138 Fused Silica Capillary Column (Supelco NUKOLTM, 15 x 0.53 x 0.5 µm film thickness) and with a flame ionization detector (200° C) with H₂ as the carrier gas. An 139 initial temperature of 80° C was used and was subsequently increased to 140° C, then 140 160° C and finally to 200° C at a rate of 10° C/min. The analyzed samples were 141 142 centrifuged and filtered through a 0.45 µm membrane.

Production of gas was continuously measured using a gas flow meter (Ritter Company, 143 144 drum-type wet-test volumetric gas meters), and the composition of the produced gas was determined by gas chromatography separation (SHIMADZU GC-2010). H₂, CH₄, 145 CO₂, O₂ and N₂ were analyzed by means of a thermal conductivity detector (TCD) 146 147 employing a Supelco Carboxen 1010 Plot column. Samples were taken using a 1 ml Dynatech Gastight gas syringe under the following operating conditions: split = 100; 148 149 constant pressure in the injection port (70 kPa); 2 min at 40 °C; ramped at 40 °C/min until 200° C; 1.5 min at 200° C; detector temperature: 250° C; and injector temperature: 150

151 200° C. Helium was used as carrier gas (266.2 ml/min) (Zahedi et al. 2017b) .
152 Commercial mixtures of H₂, CH₄, CO₂, O₂, N₂ and H₂S (Abelló Linde S.A.) were used
153 to calibrate the system.

154 2.5 Microbiological analysis and biochemical activity

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The cellular concentration and percentages of Bacteria and Archaea were quantified by epifluorescence method (FISH) according to the method of Zahedi et al. (Zahedi et al. 2013c; Zahedi et al. 2017a). The main steps of FISH of whole cells using 16S rRNAtargeted oligonucleotide probes are cell fixation, permeabilisation and hybridisation with the desired probe(s).

The cellular concentration and percentages of *Eubacteria*, *Archaea*, BUA, PUA, SRB, HUM and AUM were obtained by FISH according to Zahedi et al.(Zahedi et al. 2013c; Zahedi et al. 2014). The total population was calculated as the sum of the relative amounts of *Eubacteria* and *Archaea*, because the main anaerobic groups in the anaerobic reactors are contained within these two domains (Griffin et al. 1998). Acetogens were calculated as the sum of the relative amounts of PUA and BUA. HAB were calculated as the difference in the relative amounts of *Eubacteria* and acetogens.

168 **3. Results and discussion**

169 The process performances and the functional *Bacteria* and *Archaea* community 170 structures of the two-phase anaerobic reactors for HP and MP were investigated and 171 analyzed together. The section has been structured into two parts: hydrogenic phase 172 performance and methanogenic phase performance.

173 3.1 Hydrogenic phase performance

As commented before, no molybdate was added to the first phase, because the biogaswas sulphide-free. The characterization physical-chemical and microbiological in the

effluent of the first phase are shown in the Table 2. The performance in the first phase 176 177 was according to Zahedi et al. (Zahedi et al. 2013c) study to 1.5 d HRT. This pH value was close to the ideal pH conditions for HP of 5.5 (Bolzonella et al. 2012). VFA 178 179 composition was composed mainly of butyric acid $(5.1\pm0. g/l)$. The dominant fermentation products were butyric acid and acetic acid with small amounts of propionic 180 181 also detected (<0.1 g/l). The sulphate values in the effluent were the same with those 182 measured in the feeding; therefore no sulphate consumption was detected (no SBR 183 activity was detected). A high solubilization (increase in SCOD and VFA and decrease in VS) was detected. The butyrate acid concentration was higher than acetic acid 184 185 concentration and it was in line to other researches of hydrogen production from similar 186 wastes (Cadiz-Spain urban wastes) (Romero Aguilar et al. 2013; Zahedi et al. 2013b; Angeriz-Campoy et al. 2017; Zahedi et al. 2017a). The ratio of butyrate:acetate (g 187 188 butyrate/g acetate) was 2.5 and it according to the ratios reported by these previous studies. The biogas produced was composed of H₂ and CO₂ without CH₄ and H₂S 189 190 detection. In terms of yields, biohydrogen in the first reactor was $47 \pm 4\%$ and the HP 191 was $2.0 \pm 0.3 L H_2/l/d$).

- 192 3.2 Methanogenic phase performance
- 193 *3.2.1 Process stability*

The stability of the process was evaluated based on the evolution of pH and the VFA/alkalinity ratio (VFA/Alk) before (1-45 d) and after (45-50 d) SRB inhibitor addition (Siles Lopez et al. 2009).

Fig. 2.a shows the evolution of pH throughout the test. At the beginning (before
inhibitor addition) the systems were able to self-regulate and reach a pH of between 7.0
and 8.5, the optimal pH for the activity of methanogens (De La Rubia et al. 2009;

Dahunsi et al. 2016). After inhibitor addition, the pH of the reactor decreased below 7.0
showing microbial inhibition of H₂ and acid consumer organisms.

Alkalinity is the capacity to neutralise acids, the total volatile fatty acids (TVFA)/Alk 202 203 ratio being typically used as a measurement to evaluate anaerobic system stability (Balaguer et al. 1992; Rincón et al. 2008; Siles Lopez et al. 2009). Values between 0.1 204 205 and 0.4 (equiv. acetic acid/equiv. CaCO₃) indicate favourable operating conditions without the risk of acidification. The evolution of these ratios is shown in Fig 2.b. 206 Before inhibitor addition stability was observed, however from the day after the 207 molybdate was added this ratio decline sharply, demonstrating the non-stability of the 208 second phase (ratios were higher than 0.4). Therefore, the pH values and the acids 209 gathered, thus preventing the activity of the methanogens, as will be explained later. 210

211 *3.2.2 Process performances*

The highest values for the removal of VS $(81\pm7\%)$ and SCOD $(58\pm2\%)$ were obtained 212 before inhibitor addition. These values were similar to those obtained by Brownie 213 (Browne et al. 2014) in biomethane production from the organic fraction of municipal 214 solids waste in semi-continuous systems. Before MoO₄⁻² addition the average values of 215 SCOD and VS in the effluent were around 9 g/l and 12 g/kg respectively, after MoO_4^{-2} 216 inhibition on day 50 those values did exceed to 25 g/l and 28 g/kg respectively (Figure 217 3). The reduction in the consumption of the organic matter means inhibition of the 218 anaerobic digestion process. 219

In relation to VFA, the effect of molybdate was an acidification in the reactor (increase in acid content, Figure 4) according to decrease in the pH values. In the present research, before molybdate addition, the concentrations of VFA were in the order propionic >acetic > butyric. Addition of molybdate caused butyric and acetic acids to

dominate over propionic acid at the end of the trial. The presence of acetic acid in the 224 225 effluent shows that non-availability of AUM substrate was not the underlying problem for the inhibition of CH₄ production in these studies, but chronic inhibition of AUM by 226 MoO₄⁻². In addition, butyric increment in the effluent was resulted of the non-227 availability to the BUA to consume the butyric. The amount of propionic acid generated 228 after molybdate addition was similar to the amount produced before of inhibitor 229 230 addition, suggesting that acetogenic activity of PUA was sufficient to achieve the normal propionic acid concentrations(Zahedi et al. 2013a). 231

It can clearly be observed that the addition of sodium molybdate on day 45 caused 232 immediate inhibition of sulphate reduction, thereby resulting in increase of sulphate 233 234 content in the effluent (Figure 5) and total absence of H₂S in the biogas (Figure 6a) on the following day. Before MoO_4^{-2} addition the average values of sulphate were around 235 0.7 g/l and on day 50 those values increased to 2.0 g/l. The values of H₂S decreased 236 237 from 45±5 mL H₂S/l/d on day 45 to 0 mL H₂S/l/d on day 46. Regarding to the MP, the values of the MP also involved a decrease due to inhibitor use (Figure 6b). Before the 238 use of the SRB inhibitor, MP was around 3.5 L CH₄/l/d and after the supplement it was 239 240 modestly decreased, until the end, where MP did not exceed 0.2 L CH₄/l/d (day 50). It 241 technically shows that adding this amount of inhibitor for this amount of time does not improve biogas production. Future efforts could incorporate dosing periods with 242 smaller amounts of inhibitor added in increasing increments and the observed return of 243 the system to the previous rate of biogas production and quality. 244

245 3.2.3 Microbial community

The evolution of the main microbial group involved in the methanogenic process is described in the Table 3. In the present research, concentrations of different microbial

groups were evaluated before and after inhibitor addition. All the results shown are 248 249 average values. Before inhibitor addition, the ratio of Eubacteria: Archaea was 55:45. These results are in accordance to those obtained by Zahedi et al. (Zahedi et al. 2013c) in 250 251 the second phase reactor of dry-thermophilic anaerobic digestion process of sulphatecontaining municipal solid waste and logically, lower than those obtained by Griffin et 252 253 al.(Griffin et al. 1998) and McMahon et al.(McMahon et al. 2001) in single-phase 254 reactors of organic waste. SRB population values were in line with those (20-28%) obtained by Zahedi et al. (Zahedi et al. 2013c) and Zhang et al. (Zhang et al. 2011) lower 255 256 than those (14%) obtained by Mohan et al. (Mohan et al. 2005).

It should be noted, that after inhibitor addition the microbial proportion in the reactor had not very altered, the microbial consortia and microbial activity were hardly altered. It is necessary to emphasize that although the proportion of microorganisms in the reactor is a key, not only the stability, but also the adequate dynamics ("flexibility") of the microbial community structure and high values of microbial activity are important for the stable performance of the reactors treating urban wastes¹³.

At the end of the trial, the microbial consortia were decreased in 42%. The removal 263 264 rates of Bacteria, Archaea, HAB, acetogens, AUM, HUM, SRB, BUA and PUA were 44 %, 38%, 48%, 39%, 35%, 41%, 60%, 63% and 15% respectively. The most affected 265 266 were SRB and BUA and the most resistant group was PUA which is in line with the 267 constant values of propionic acid in the effluent. The reductions in the microbial 268 consortia, the decrease in the MP, H₂S production, pH value, and increase in the sulphate, VS, SCOD and VFA contents all reveal a decrease in the microbial activity, 269 270 except for PUA.

271 In short, the use of the toxic to improve the biogas of the two-phase dry-thermophilic 272 anaerobic digestion process of sulphate-containing municipal solid is not effective. However, it is to be noted that the harmful effect of molybdate supplementation on the 273 274 SRB and methanogens and therefore, on the H₂S and CH₄ generation makes this treatment an interesting option on other fields, as sewerage system. Since, anaerobic 275 276 conditions in sewer pipes favor the accumulation of both H_2S and CH_4 and these 277 compounds have detrimental effects on the sewer system, with different consequences for both the installation and its surroundings (Auguet et al. 2015) (such as malodor, 278 279 health hazards due to the well-known toxicity of H₂S, and corrosion of both the inner 280 surface of pipes and the inlet zones of waste water treatment plants, etc)

281 Conclusion

282 Inhibitor addition has proven successful to remove the undesirable production of hydrogen sulphide (H₂S). However it too resulted in an increase in TVFA/Alk ratio, as 283 well as decrease in pH, organic removal organic matter and biogas generation. All this 284 indicated inhibition of all the steps to AD (except to propionic degradation). Therefore, 285 although molybdate is an effective bactericide for SRB, the use of the toxic would be 286 287 avoided, since molybdate supplementation did not improve the quality of the biogas, under the circumstances of this experiment. Microbial consortia were decreased in 288 42%. The removal rates of Bacteria, Archaea, HAB, acetogens, AUM, HUM, SRB, 289 BUA and PUA were 44 %, 38%, 48%, 39%, 35%, 41%, 60%, 63% and 15% 290 291 respectively.

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Table 1. Physical-chemical and microbiological characterization of the substrateemployed in the first phase.

Parameter	Value
рН	5.3 (0.6)
TS (g/l)	120 (15)
VS (g/l)	85 (7)
Sulphate (g/l)	1.9 (0.3)
VFA (g acetic acid/l)	1.8 (0.5)
Acetate (g/l)	1.87(0.5)
Propionate (g/l)	0.0 (0.0)
Butyrate (g/l)	0.4 (0.2)
Total population (10^8 cells/ml)	6 (2)
Eubacteria (%)	78 (3)
Archaea (%)	22 (2)

442 Average values are shown, with standard deviations in parentheses.

Table 2: Physicochemical and microbiological characterization of the first phase reactoreffluent

Physicochemical parameters								
 рН	SCOD (g/l)	VS (g/kg)	Alkalinity (gCaCO ₃ /l)	Sulphate (g/l)	TVFA (g acetic/l)	Acetic (g/l)	Propionic (g/l)	Butyric (g/l)
 5.3±0.3	33±2	42±5	4±0	1.9±0.2	10.4±0.9	2.5±0.4	0.1±0.1	5.1±0.5

Microbiological parameters

Total population (10 ⁸ cells/mL)	Eubacteria (%)	HAB (%)	Acetogens (%)	BUA (%)	PUA (%)	SRB ^a (%)	Archaea (%)	AUM (%)
9.5±0.6	88±2	70±2	18±2	8±1	10±1	14±1	12±1	5±0

448 ^aPercentages compared to total *Eubacteria*.

449

Period (Day)						
1-45*	46	47	48	49	50	
robiological j	paramete	ers				
21.8±2.5	16.3	14.5	14.9	13.9	12.8	
55±2	56	63	51	49	53	
28±1	34	37	27	25	25	
27±2	22	26	24	24	28	
14±1	6	9	8	7	9	
13±1	16	17	16	17	19	
26±2	18	18	16	17	18	
45±2	44	37	49	51	48	
24±1	24	21	29	28	27	
21±1	20	15	21	23	21	
	1-45* robiological p 21.8±2.5 55±2 28±1 27±2 14±1 13±1 26±2 45±2 24±1 21±1	1-45*46robiological parameter 21.8 ± 2.5 16.3 55 ± 2 56 28 ± 1 34 27 ± 2 22 14 ± 1 6 13 ± 1 16 26 ± 2 18 45 ± 2 44 24 ± 1 24 21 ± 1 20	Period (I1-45*4647robiological parameters 21.8 ± 2.5 16.314.5 55 ± 2 5663 28 ± 1 3437 27 ± 2 2226 14 ± 1 69 13 ± 1 1617 26 ± 2 1818 45 ± 2 4437 24 ± 1 2421 21 ± 1 2015	Period (Day)1-45*464748robiological parameters 21.8 ± 2.5 16.314.514.9 55 ± 2 566351 28 ± 1 343727 27 ± 2 222624 14 ± 1 698 13 ± 1 161716 26 ± 2 181816 45 ± 2 443749 24 ± 1 242129 21 ± 1 201521	Period (Day)1-45*46474849robiological parameters 21.8 ± 2.5 16.314.514.913.9 55 ± 2 56635149 28 ± 1 34372725 27 ± 2 22262424 14 ± 1 6987 13 ± 1 16171617 26 ± 2 18181617 45 ± 2 44374951 24 ± 1 24212928 21 ± 1 20152123	

451 Table 3: Microbiological characterization of the second phase reactor effluent.

452 ^aPercentages compared to total *Eubacteria*.

453 *Values corresponding to the analytical determinations in steady conditions (between day 21 and 45).

455 Figure Captions

- 456 Figure 1: The laboratory-scale reactors used in this study. Hydrogenic reactor to the left
- 457 and methanogenic reactor to the right.
- 458 Figure 2: (a) pH evolution. (b) TVFA/Alk evolution (g acetic/g CaCO₃)
- 459 Figure 3: (a) SCOD evolution (g/l). (b) Volatile Solid evolution (g/kg).
- 460 Figure 4: (a) VFA evolution (g/l).
- 461 Figure 5: (a) Sulphate evolution (g/l).
- 462 Figure 6: (a) Sulphide production (SP) evolution (ml $H_2S/l/d$). (b) Methane production
- 463 (MP) evolution ($l CH_4/l/d$).





470 Figure 2.b





















