Mineralization of LCFA Associated With Anaerobic Sludge: Kinetics, Enhancement of Methanogenic Activity, and Effect of VFA

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Abstract: Long-chain fatty acids (LCFA) associated with anaerobic sludge by mechanisms of precipitation, adsorption, or entrapment can be biodegraded to methane. The mineralization kinetics of biomass-associated LCFA were established according to an inhibition model based on Haldane’s enzymatic inhibition kinetics. A value around 1,000 mg COD-LCFA/g VSS was obtained for the optimal specific LCFA content that allowed the maximal mineralization rate. For sludge with specific LCFA contents of 2,838 ± 63 and 4,571 ± 257 mg COD-LCFA/g VSS, the specific methanogenic activities in the presence of acetate, butyrate, and H₂/CO₂ were significantly enhanced after the mineralization of the biomass-associated LCFA. For sludge with a specific LCFA content near the optimal value defined by the kinetic model, the effect of adding VFA to the medium was studied during the mineralization of the biomass-associated LCFA. Different patterns were obtained for each individual substrate. Acetate and butyrate were preferentially consumed by the consortium, but in the case of propionate no evidence of a sequential consumption pattern could be withdrawn. It was concluded that LCFA do not exert a bactericidal neither a permanent toxic effect toward the anaerobic consortia. A discussion is addressed to the relative roles of a reversible inhibitory effect and a transport limitation effect imposed by the LCFA surrounding the cells. © 2004 Wiley Periodicals, Inc.

Keywords: LCFA; oleic acid; methanogenic activity; kinetics

INTRODUCTION

Lipids, abundant in slaughterhouses, edible oil/fat refineries, and dairy wastewaters, are potentially attractive for biogas production due to their high theoretical methane yield. However, in practice, physicochemical pretreatment is usually applied before anaerobic treatment, because neutral fats are easily hydrolyzed to long-chain fatty acids (LCFA), which are reported to exert an acute toxic effect on the microorganisms involved in the β-oxidation and methanogenic pathways (Angelidaki and Ahring, 1992; Hwu, 1997; Koster and Cramer, 1987; Rinzema, 1988).

In general, a severe toxic effect is referred, although granular sludge seems to be more resistant to LCFA toxicity than suspended sludge (Hwu, 1997). Rinzema (1988) reported that LCFA exert a bactericidal effect and that acetoclastic bacteria do not adapt to LCFA either upon repeated exposure to toxic concentrations or after prolonged exposure to nontoxic concentrations. The recovery after a lag phase usually observed in batch assays was attributed by this author to the growth of few survivors. Angelidaki and Ahring (1992) reported a permanent toxic effect that underlines the conclusions of Rinzema (1988). According to some authors, the mechanism of LCFA toxicity seems to be related to the adsorption of the surface active acids onto the cell wall, which affects its transport and/or protective functions (Demeyer and Henderickx, 1967; Galbraith et al., 1971; Rinzema, 1988). More recently, Lalman and Bagley (2001) reported the inhibitory effects of oleic and stearic acids to acetoclastic and hydrogenophilic methanogens. In their study, diethyl ether was added at a concentration of 14.2 g L⁻¹, to promote LCFA dispersion in the cultures, being a questionable approach, since this compound is not expected to occur under real conditions.

More than inhibition drawbacks, flotation, and washout are reported to be important problems associated to the adsorption of LCFA onto anaerobic sludge, as granulation and/or granule maintenance are difficult tasks for lipid/LCFA-containing wastewaters. Hwu (1997) added some insights to this phenomenon by assessing the critical LCFA-specific organic load that induced flotation and complete washout during the operation of expanded granular sludge bed (EGSB) reactors, concluding that it corresponded to a concentration far below the toxicity limit.

Problems of sludge flotation or unsuccessful granulation were ascribed to treatment failure of industrial- and pilot-scale UASB reactors treating lipids/LCFA-containing...
wastewater (Hawkes et al., 1995; Hwu, 1997; Rinzema, 1988; Samson et al., 1985; Sam-soon et al., 1991). Reasonable solutions to overcome flotation problems were searched in some reported works: sieve drums, biomass recirculation, or partial phase separation were tentatively applied for that purpose (Beccari et al., 1998; Hamdi et al., 1992; Hwu, 1997; Rinzema, 1988), without convincing reliability.

In recent studies conducted by our research group, it was observed that anaerobic sludge was able to mineralize LCFA in amounts up to 3 g COD·g VSS⁻¹, provided that it was associated with the sludge, not the bulk medium, evidencing that the anaerobic consortium remained active in such conditions (Pereira et al., 2002, 2003). This clearly contradicts the generally accepted idea that LCFA adsorption onto anaerobic sludge is closely linked to LCFA inhibition through mechanisms of cell membrane/wall damage, eventually inducing cytolysis and bactericidal effects (Demeyer and Henderickx, 1967; Galbraith et al., 1971). Salminen et al. (2001) observed the batch biodegradation of accumulating materials (mainly LCFA) in anaerobic digesters that previously failed the operation, during the semicontinuous treatment of poultry slaughterhouse waste.

These previous results prompted a renewed interest for a more deep study on the anaerobic biodegradation of LCFA that are mainly associated to the biomass, aiming to optimize its degradation rate. In this regard, the objectives of the present work are:

1. to establish the kinetics of anaerobic mineralization of the biomass-associated LCFA;
2. to evaluate and define the limits of methanogenic activity enhancement after the mineralization of the biomass-associated LCFA; and
3. to evaluate the effect of individual Volatile Fatty Acids (VFA) on the mineralization of the biomass-associated LCFA.

Oleic acid was used as a model for LCFA to obtain the loaded sludge in the continuous reactor operation because it is, in general, the most abundant of all LCFA present in wastewater (Komatsu et al., 1991), has a good solubility, and is one of the more toxic LCFA (Galbraith et al., 1971).

**MATERIALS AND METHODS**

**Sludge Sources**

Anaerobic sludge, in a total of 38 samples, was obtained from continuous laboratory-scale reactors operated at mesophilic conditions (37°C) with oleic acid as substrate. Table I presents the operating conditions prevailing at the moments of sample collection. The fixed bed reactor configuration was designed to allow the sampling of entrapped biomass along the operation (Alves et al., 1998). Prior to characterization, all the sludge samples collected were washed and centrifuged (1681 g, 10 min) twice, with the same anaerobic basal medium used in the batch experiments. The volatile suspended solids content (VSS) of each sample was determined according to Standard Methods (1989).

**Sludge Characterization**

**Microscopic Observations**

Phase contrast microscopy was performed with a Zeiss (Oberkochen, Germany) Axioscop microscope. The same microscope was used to visualize the autofluorescence of the methanogenic consortium (420 nm).

Staining of lipid compounds was performed using Sudan black B, according to the procedure described by Seviour and Blackall (1999). An aliquot of a sludge sample was placed on a gelatin-covered slide, air dried, and immersed on a Sudan black B alcoholic solution (0.3% wt/vol in ethanol 60%) for 10 min, followed with a 1-s rinse with water. After the excess water was removed, the slide was immersed in a safranin O aqueous solution (0.5% wt/vol) for 10 s. It was then well rinsed with water and allowed to dry prior to examination under bright field, using a Zeiss Axioscop microscope. A control procedure was performed to remove the lipid matter by keeping the samples in con-

**Table I.** Sources of the anaerobic suspended sludge used in this work.

<table>
<thead>
<tr>
<th>Reactor type</th>
<th>Operation mode</th>
<th>Influent COD (mg L⁻¹)</th>
<th>HRT (day)</th>
<th>Type of substrate⁴</th>
<th>Oleate loading rate (kg COD m⁻³·day⁻¹)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fixed bed</td>
<td>Steady state</td>
<td>12,000</td>
<td>1.4</td>
<td>SM (50% COD) + SO (50% COD)</td>
<td>4.3</td>
<td>Alves et al., 2001</td>
</tr>
<tr>
<td>Fixed bed</td>
<td>Steady state</td>
<td>12,000</td>
<td>1.4</td>
<td>SO (100% COD)</td>
<td>8.6</td>
<td>Alves et al., 2001</td>
</tr>
<tr>
<td>Fixed bed</td>
<td>Organic shock⁵</td>
<td>20,000</td>
<td>0.67</td>
<td>SM (50% COD) + SO (50% COD)</td>
<td>15</td>
<td>Cavaleiro et al., 2001</td>
</tr>
<tr>
<td>Fixed bed</td>
<td>Hydraulic shock⁶</td>
<td>4,000</td>
<td>0.13</td>
<td>SM (50% COD) + SO (50% COD)</td>
<td>15</td>
<td>Cavaleiro et al., 2001</td>
</tr>
<tr>
<td>EGSB</td>
<td>Steady state</td>
<td>4,000</td>
<td>1.0</td>
<td>SM (50% COD) + SO (50% COD)</td>
<td>2</td>
<td>Pereira et al., 2002</td>
</tr>
<tr>
<td>EGSB</td>
<td>Steady state</td>
<td>4,000</td>
<td>1.0</td>
<td>SO (100% COD)</td>
<td>4</td>
<td>Pereira et al., 2002</td>
</tr>
<tr>
<td>EGSB</td>
<td>Steady state</td>
<td>6,000</td>
<td>1.0</td>
<td>SO (100% COD)</td>
<td>6</td>
<td>Pereira et al., 2002</td>
</tr>
<tr>
<td>EGSB</td>
<td>Steady state</td>
<td>8,000</td>
<td>1.0</td>
<td>SO (100% COD)</td>
<td>8</td>
<td>Pereira et al., 2002</td>
</tr>
</tbody>
</table>

⁴Abbreviations: SM, skim milk; SO, sodium oleate.

⁵Seven samples were taken during 4 days.

⁶Ten samples were taken during 4 days.
tact, for 1 h at room temperature, with a solution consisting of chloroform and methanol (2:1, vol/vol) with addition of 1% HCl and 4% H$_2$O (High, 1984).

Scanning electron microphotographs were obtained in a Leica S360 microscope (Leica, Cambridge, England). Samples were fixed with 3% (wt/vol) glutaraldehyde solution in cacodylate buffer, pH 7.2, and dehydrated with ascending concentration ethanol solutions (10%, 25%, 50%, 75%, 90%, and 100%). Prior to microscopic examination, samples were sputter-coated with gold.

### Methanogenic Activity Measurements

Specific methanogenic activity (SMA) tests were performed using a pressure transducer technique (Coates et al., 1996; Colleran et al., 1992). The pressure increase developed in the batch vials fed with nongaseous substrates (30 mM acetate, 30 mM propionate, and 15 mM butyrate) or pressure decrease in vials previously pressurized (1 bar) with gaseous substrates (H$_2$/CO$_2$, 80:20 vol/vol) was monitored, using a hand-held pressure transducer capable of measuring a pressure variation of 2 bar (0 to ±202.6 kPa) over a device range of −200 to +200 mV, with a minimum detectable variation of 0.005 bar, corresponding to 0.05 mL of biogas in a 10-mL headspace. The basal medium used in all batch experiments was made up with demineralized water, was composed of cysteine-HCl (0.5 g L$^{-1}$) and sodium bicarbonate (3 g L$^{-1}$), the pH was adjusted to 7.0–7.2 with 8 N NaOH, and was prepared under strict anaerobic conditions. Methane content of the biogas was measured by gas chromatography using a Chrompack Haysep Q (80–100 mesh) column (Chrompack, Les Ulis, France), with N$_2$ as carrier gas at 30 mL min$^{-1}$ and a flame-ionization detector. Temperatures of the injection port, column, and flame-ionization detector were 120, 40, and 130°C, respectively. The values of methane production were corrected for the standard temperature and pressure conditions (STP). Blank controls were used for liquid substrates (no added substrate) and for gaseous substrates (pressurized with N$_2$/CO$_2$, 80:20 vol/vol at 1 bar). All batch experiments were performed in triplicate. SMA values were determined dividing the initial slope of the methane production curve by the VSS content of each vial at the end of the experiment and were expressed in mL CH$_4$ g VSS$^{-1}$ day$^{-1}$, which were then converted to the equivalent chemical oxygen demand (COD). Background methane production due to the residual substrate was discounted. Ideally, the VSS values used to express the specific methanogenic activity should be the VSS initially present in the vials. However, those values account for all organic matter, including not only the cells but also the biomass-associated LCFA. To obtain the VSS content that accounts only for the cells, the methane production in the test vials was followed until stabilization in order to minimize the LCFA contribution to the VSS measured at the end of the test. These VSS values were considered for the specific methanogenic activity measurements and ideally they should be between 2 and 5 g L$^{-1}$.

This procedure may be questionable as some biomass growth can occur, underestimating the initial specific methanogenic activity. However, the error associated to this factor is certainly lower due to the slow growth rate of anaerobic biomass when compared to the huge amount of LCFA present in the sludge.

It should be noted that, concerning the SMA in the presence of indirect methanogenic substrates, the methane production rate will only be a valid measurement of the syntrophic activity on those substrates when the acetoclastic and hydrogenophilic activities are not rate limiting (Dolfing and Bloemen, 1985).

### Mineralization of LCFA Accumulated Onto the Sludge

These batch assays were similar to the blank control assays described above for the measurement of SMA in the presence of liquid substrates. Each sludge sample was incubated in sealed batch vials of 25 mL at 37°C, 150 rpm, under strict anaerobic conditions, without any added organic carbon source, calcium, or nutrients. Only anaerobic basal medium was added. The initial methane production rate and the maximum plateau achieved were determined for each test, according to the example presented in Figure 1. In all cases, the methane production rate refers to the initial slope, even when a lag phase was observed, indicating a null value. The cumulative methane production at the end of the batch assays was considered an indirect measurement of the amount of biomass-associated LCFA. The specific LCFA content was then determined by dividing the maximum plateau achieved in the methane production curve by the VSS content of each vial at the end of the experiment and was expressed as mg COD-CH$_4$ g VSS$^{-1}$ (the direct measurement), as mg COD-LCFA g VSS$^{-1}$ (the indirect measurement) or just as mg COD g VSS$^{-1}$. The specific methane production rate was obtained by dividing the initial slope of each curve of methane production by the VSS content of each vial at the end of the experiment, and was expressed as mg COD-CH$_4$ g VSS$^{-1}$ day$^{-1}$.

![Figure 1. Example of a methane production curve in a batch assay, during the mineralization of the biomass-associated LCFA.](image-url)
Biomass-Associated LCFA Mineralization

The SMA of the sludge before and after degrading the biomass-associated LCFA accumulated during the continuous operation was compared by performing parallel assays. After it had been washed with anaerobic basal medium and centrifuged (1681g, 10 min) twice, the sludge was incubated in 24 vials of 25 mL and 12 vials of 70 mL, at 37°C, 150 rpm under strict anaerobic conditions. Two sets of 18 vials (12 + 6) were separated for parallel assays. In the first set, the methanogenic activities in acetate, propionate, butyrate and H2/CO2 were determined in triplicate as described before. Six vials were used as blank controls: three for liquid substrates were no substrate was added, and three for gaseous substrates where N2/CO2 (80:20 vol/vol) was pressurized (1 bar). In the other set of 18 vials, no substrate was added, but the degradation of the biomass-associated LCFA was followed until stabilization. The vials were then depressurized, vented with N2/CO2 (80:20 vol/vol), and methanogenic activities in acetate, propionate, butyrate, and H2/CO2 were measured in triplicate, as described for the first set of vials. The VSS values taken into account to calculate the specific methanogenic activity before and after the depletion of the biomass-associated LCFA were the values measured at the end of the whole experiment. Even if some error was introduced due to the possible biomass growth, this was needed to eliminate the contribution of the LCFA to the initial VSS content in the vials.

Effect of Added VFA in the Mineralization of the Biomass-Associated LCFA

In a set of vials with 500-mL working volume, the sludge containing accumulated LCFA was incubated in batch mode, keeping the environmental conditions applied in the SMA assays. Acetate, propionate, and butyrate were individually added and their depletion in the liquid phase was monitored. Possible VFA release to the medium during the mineralization of the biomass-associated LCFA was also assessed by setting blank controls where no VFA were added. VFA concentrations were determined by high-performance liquid chromatography using a Chrompack column (300 × 6.5 mm) and a mobile phase of 5 mM sulfuric acid (H2SO4) at 0.7 mL·min⁻¹. The column was set at 40°C, and the detection was spectrophotometric at 220 nm. Duplicate assays were performed.

RESULTS AND DISCUSSION

Microscopic Examination of Sludge With Associated LCFA

The ability of suspended sludge to accumulate LCFA was confirmed by microscopic observation of a sludge that had been in prolonged contact with oleic acid under continuous operation mode (Figs. 2 and 3).

In this sludge, clear whitish zones can be identified, likely representing the LCFA associated with the biomass (Fig. 2a,b). These white zones seem to act as a light emission barrier, decreasing the visible 420-nm autofluorescence of the methanogenic population (Fig. 2c,d). Scanning electron micrographs allowed observing a net difference on the sludge aspect before and after being loaded with oleic acid (Fig. 2e,f). In the later case, the presence of a nonbiologic matrix entrapping the microorganisms is suggested, and a crystalline structure can also be clearly identified (Fig. 2a,f,g,h).

Sudan black staining confirmed the presence of lipid matter in the sludge after it been loaded with oleic acid (Fig. 3a,b). This lipid material, which is stained as a blue/black color, was not detected in the sludge after the mineralization of the biomass-associated LCFA (Fig. 3c) or when the sludge was submitted to the control procedure for lipid removal (Fig. 3d). It is also interesting to note that the counterstaining of the cells with safranin was significantly more evident in the sludge without LCFA than in the sludge loaded with LCFA. Adsorption of LCFA onto the cell surface may impair cell staining, justifying this observation.

The anaerobic sludge structure was previously found to be important for the phenomenon of LCFA mineralization, suspended sludge, small fluffy and filamentous flocs, being more efficient for LCFA accumulation and degradation than granular sludge (Pereira et al., 2002). The localization of lipid matter inside the loose floc structure may justify the large amounts of LCFA that can be entrapped by this type of sludge compared to the dense granular sludge. From these microscopic observations is evidenced the important contribution of precipitation and entrapment on LCFA accumulation.

Kinetics of Biomass-Associated LCFA Degradation

LCFA accumulation onto the anaerobic sludge was previously found to be important as a way to promote its mineralization, but the conditions that allow the optimal conversion to methane were not yet defined. When the 38 sludge samples were individually incubated in batch vials without any added organic carbon source, calcium, or nutrients, methane was produced due to the mineralization of the LCFA that had accumulated onto the sludge. The specific methane production rate and the amount of the biomass-associated LCFA per VSS unit (specific LCFA content) were calculated for each sample, as exemplified in Figure 1, and are plotted in Figure 4. This graph represents the influence of the available substrate on the kinetics of its own mineralization. It is not a conventional kinetic plot, as the substrate is not dissolved in the medium but instead is associated with the sludge. In spite of that, experimental data were fitted by Eq. (1), which represents an adaptation of an inhibition model used before by Andrews (1971) to describe the inhibition of volatile fatty acids in the
anaerobic digestion process. This model is based on the substrate inhibition enzymatic kinetics developed by Haldane (1930).

\[ q_p = \frac{\hat{q}_p \times S_{ba}}{K_S + S_{ba} + \frac{S^2_{ba}}{K_i}} \]  

where \( q_p \) is the specific methane production rate \([\text{M}_{\text{substrate}} \times \text{M}_{\text{biomass}}^{-1} \times \text{t}^{-1}] \), \( S_{ba} \) is the biomass-associated substrate per VSS unit \([\text{M}_{\text{substrate}} \times \text{M}_{\text{biomass}}^{-1}] \), and \( \hat{q}_p \), \( K_S \), and \( K_i \) are model parameters.

Table II summarizes the results obtained by the fitting as well as the corresponding correlation coefficient, determined by the Levenberg–Marquardt method (Marquardt, 1963).

Based on this model, a value of 1,080 mg COD-LCFA·g VSS\(^{-1}\) was found to be the critical biomass-associated substrate per VSS unit \( S_{ba}^{\text{crit}} \), which leads to the maximal mineralization rate, found to be 250 mg COD-CH\(_4\)·g VSS\(^{-1}\)·day\(^{-1}\) (Eq. (2)).

\[ S_{ba}^{\text{crit}} = \sqrt{K_S K_i} \]  

The low correlation coefficient obtained for the fit (0.56) imposes a significant uncertainty on the value of \( S_{ba}^{\text{crit}} \). Therefore, a value around 1,000 mg COD-LCFA·g VSS\(^{-1}\) represents just a rough estimate of the specific LCFA load that allows the maximal mineralization rate of the biomass-associated LCFA. However, even with precision limitations, it has practical interest as far as the mineralization of LCFA is concerned.

The low correlation coefficient obtained by this fitting also suggests that the mechanism of inhibition is more complex than a simple substrate inhibition phenomenon, in
part because the β-oxidation process involves a sequence of substrates and products that can compete with each other. Furthermore, the method used to measure the amount of the accumulated substrate is based on the measurement of the final product (methane) and thus depends on the metabolic activity of different trophic groups in the consortium. On the other hand, as previously suggested by Pereira et al. (2003), transport limitations may have an important role on LCFA mineralization. As the LCFA seems to be mostly in intimate contact with the biomass, substrate diffusion limitations are not expected to limit the degradation rate, but product diffusion limitation, e.g., biogas release, can be hampered by the LCFA associated with the sludge and thus may induce a delay on the measured methane production. This is not taken into account in the inhibition model and may also contribute to the relative discrepancy between model prediction and experimental data.

Enhancement of Specific Methanogenic Activity After the Mineralization of the Biomass-Associated LCFA

The use of an inhibition model to fit the experimental data suggests a decrease in the consortium capacity to mineralize specific LCFA contents above about 1,000 mg COD-g VSS⁻¹. This decrease may be due to a metabolic or a physical inhibition (transport limitations), as previously suggested by Pereira et al. (2003). Whatever the phenomenon that is behind the inhibition (metabolic or physical), it is important to evaluate if it is reversible. In order to obtain

Table II. Kinetic constants for the applied inhibition model.*

<table>
<thead>
<tr>
<th></th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>( q ) (mg COD-CH₄·g VSS⁻¹·day⁻¹)</td>
<td>384,361</td>
</tr>
<tr>
<td>( K_S ) (mg COD-CH₄·g VSS⁻¹)</td>
<td>790,495</td>
</tr>
<tr>
<td>( K_I ) (mg COD-CH₄·g VSS⁻¹)</td>
<td>1.48</td>
</tr>
<tr>
<td>Correlation coefficient</td>
<td>0.56</td>
</tr>
</tbody>
</table>

*Based on Haldane’s enzymatic kinetics (Haldane, 1930).
Samples A, B, and C were characterized in terms of the SMA with acetate, propionate, butyrate, and H$_2$/CO$_2$, before and after allowing the mineralization of the biomass-associated LCFA. Table III summarizes the obtained results.

In the experiment before allowing the mineralization of the biomass-associated LCFA, the less loaded sludge, i.e., sludge A, exhibited a measurable specific activity in acetate, butyrate, H$_2$/CO$_2$, but no activity with propionate as substrate. Sludge B and C presented no initial SMA, except with H$_2$/CO$_2$ as substrate. After the mineralization of the biomass-associated LCFA, an increase on the SMA in the range of specific LCFA contents between 1,000 and 5,000 mg COD·g VSS$^{-1}$ was observed for all the selected substrates. With the single exception of the SMA with hydrogen as substrate, the difference between the activities measured before and after the mineralization of the accumulated substrate was maximal for the sludge B, which has a specific LCFA content of 2,838 mg COD·g VSS$^{-1}$.

This result proves that the effect of LCFA in the decrease of the SMA, is a reversible phenomenon in the range of specific LCFA content between 1,000 and 5,000 mg COD·g VSS$^{-1}$. This is a remarkable result that clearly contradicts the bactericidal or permanent toxic effects of LCFA, previously reported by Rinzema (1988) and by Angelidaki and Ahring (1992), and creates a new challenge for the anaerobic treatment of wastes/wastewater with high lipid contents.

The low or null SMA measured before the depletion of the biomass-associated LCFA may result from a strong effect of transport (diffusion) limitations imposed by the LCFA layer surrounding the cells, which could hamper the access of the added substrates, as well as the subsequent biogas release. This is reinforced by the fact that H$_2$, the smallest substrate used, was, in all cases, easily mineralized to methane, suggesting a fast transport of this molecule through the LCFA layer. Transport limitations phenomena may be responsible by the observed lag phases that have been ascribed to mechanisms of cell wall damage and bactericidal effects, widely accepted for years. The recovery after a lag phase, attributed to the growth of few survivors (Rinzema, 1988), could be a delay imposed by transport limitations. However, although the hypothesis of

![Figure 5. Cumulative methane production due to the mineralization of the biomass-associated LCFA. Each curve represents an average of three replicates. Sludges A, B, and C have specific LCFA contents of 1,221 ± 144, 2,838 ± 63, and 4,571 ± 257 mg COD-CH$_4$·g VSS$^{-1}$, respectively.](image-url)
transport limitations seems to be reasonable, there is no absolute sureness of it, because if LCFA exerted a reversible inhibitory effect, the result could be similar. Probably both phenomena are involved, its dominance depending on the specific LCFA content of the sludge.

Before the depletion of the biomass associated LCFA, sludges B and C presented null SMA values with the tested VFA because the initial methane production was lower than the one measured for the blank controls. Figure 6 shows the initial methane production pattern obtained during the SMA measurement in sludge B, exemplifying this behavior.

A similar effect was also previously observed when oleic acid was added, inducing a severe delay in the mineralization of the biomass-associated LCFA (Pereira et al., 2002).

These results suggest that optimal conditions for LCFA mineralization occur in the absence of organic carbon in the bulk medium. However, the presence of other organic carbon sources in the medium, for instance VFA, is highly expected in any real anaerobic digestion process.

Sludge A, which contains a specific LCFA content near the optimal value (determined according to the kinetic model), was selected to more deeply evaluate the interactions between individual VFA externally added (acetate, propionate, and butyrate) and the mineralization of the biomass-associated LCFA. The choice of this sludge was based on the practical interest of a better understanding of the role of VFA in the conditions that allow the optimal mineralization of the accumulated LCFA.

**Effect of Acetate, Propionate, and Butyrate on the Mineralization of the Biomass-Associated LCFA**

Figure 7 presents the methane production and VFA depletion when sludge A is characterized in terms of SMA with acetate, propionate, or butyrate. Individual VFA were also monitored during the mineralization of the biomass-associated LCFA (“blank” assay), which revealed that, in these conditions, there was no detectable release of VFA to the medium. This was assumed to prevail also in the test vials. During propionate degradation, acetate and butyrate were detected in concentrations not exceeding 140 and 74 mg/L, respectively. Acetate was also detected during butyrate degradation, in concentrations lower than 170 mg/L.

In order to allow direct comparison with the corresponding methane production, VFA depletion rates were calculated taking into account the total VFA present in the medium and were expressed as COD. As different working
Table IV. Comparison between specific activities of a sludge with a LCFA content of 1.221 ± 144 mg COD·g VSS⁻¹, with measured VFA depletion and methane production (mean ± SD).

<table>
<thead>
<tr>
<th>Specific activity in the presence of</th>
<th>VFA depletion rate mg COD-VFA·g⁻¹ VSS⁻¹·day⁻¹</th>
<th>CH₄ production rate a mg COD-CH₄·g⁻¹ VSS⁻¹·day⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetate</td>
<td>362 ± 3</td>
<td>387 ± 8</td>
</tr>
<tr>
<td>Propionate</td>
<td>47 ± 5</td>
<td>172 ± 11</td>
</tr>
<tr>
<td>Butyrate</td>
<td>359 ± 13</td>
<td>345 ± 6</td>
</tr>
<tr>
<td>Blank controls (no added substrate)</td>
<td>(n.d)</td>
<td>244 ± 27</td>
</tr>
</tbody>
</table>

aWithout discounting the production of the blank controls.

b"Blank control" correspond to the mineralization of the biomass-associated LCFA.

cn.d. non-detectable VFA in the medium.

Volumes were used in the gas and liquid assays, all the values were expressed per volume unit.

Table IV summarizes the results obtained for the specific methanogenic activity of the sludge A, measured by VFA depletion and CH₄ production.

From the results obtained, it can be observed that, when acetate was added to the LCFA-loaded sludge (Fig. 7a), the initial methane production was due mainly to the consumption of the added acetate because the acetate depletion rate was similar to the corresponding methane production (Table IV). A similar pattern was observed when the added VFA was butyrate (Fig. 7c), but different behavior was observed for propionate (Fig. 7b). The VFA concentration decreased only 10% in the first 50 h, whereas in the same time period, 99.1% and 83.6% of total VFA disappeared from the medium when acetate and butyrate were added, respectively. Furthermore, in the presence of propionate, the initial specific methane production rate was significantly lower than in the situation where the mineralization of the biomass-associated LCFA was monitored (Fig. 7b; Table IV). In the former situation, methane production seems to be a result of the interaction between both substrates (propionate + LCFA) degradation, because there was no evidence of a sequential consumption pattern.

CONCLUSIONS

The results presented in this work point out new insights on the anaerobic biodegradation of biomass-associated LCFA. It is important to highlight the sludge ability to mineralize biomass-associated LCFA up to specific contents of 5,000 mg COD-LCFA·g VSS⁻¹ as well as the enhancement of sludge activity after the LCFA mineralization.

The mineralization kinetics of the LCFA associated to the sludge was established and a specific content of about 1,000 mg COD-LCFA·g VSS⁻¹ was found to be the optimal for an efficient methane production rate.

For sludge loaded with a LCFA specific content near the optimal value, the presence of acetate, propionate, or butyrate (with acetate and propionate at 30 mM and butyrate at 15 mM), did not impair the process as no delay was observed in terms of overall methane production.

From a practical viewpoint, the results presented herein clearly support the approach of using sequential accumulation and degradation steps for the treatment of lipid/LCFA-based effluents.

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