

## Lecture 3 Monitoring and lab analysis - assays

David Bolzonella



#### **Contents**



- 1. Why monitoring
- 2. Different monitoring approaches
- 3. Basic and sophisticated parameters for process monitoring
- 4. Evaluation of the biogas potential (BMP) of a substrate



## First of all, answer the question .... Why monitoring ?!





(credits: prof Juan Lema)



### .... to really control the process !!!



If...



...then...



(credits: prof Juan Lema)



### Anaerobic digestion: a complex combination of biochemical and physical-chemical processes





#### "Primitive" monitoring



We can have a VERY basic monitoring, based on the determination of the typical engineering parameters like the organic loading rate (OLR, kgVS per m<sup>3</sup> of reactor per day) or the hydraulic retention time (HRT, days).....

In this case, all we need to know is the volume of the reactor, the flowrate fed and the total and volatile solids concentration of the feed ..... The biogas flowrate ....

Uncreadibly, sometimes these are unknown ...



However, to really "drive" the process, beside engineering (operational) parameters, like temperature, hydraulic retention time, organic loading rate ... physical-chemical parameters need to be regularly monitored to check the state of the process (stability).

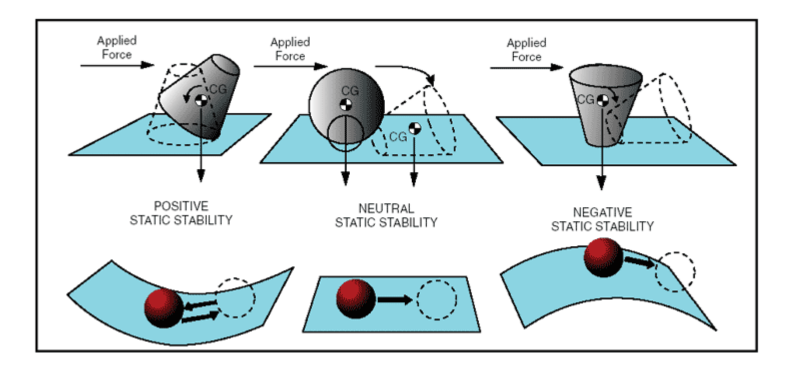
Among the most important we can remember here:

- ✓ pH
- ✓ alkalinity
- ✓ volatile fatty acids concentration and speciation
- ✓ biogas flowrate and composition
- √ ammonia concentration



These are also called "stability parameters" as they are helpful to define the "stable" operation of the anaerobic biological process ....

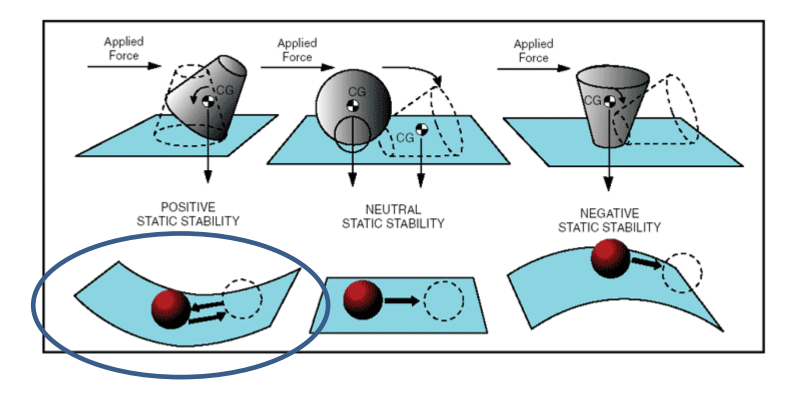






These are also called "stability parameters" as they are helpful to define the "stable" operation of the anaerobic biological process ....









Which level of monitoring ?!

Keep it simple ... but do not exsagerate ....



#### Simple monitoring (... but with some evident drawbakcs ...)



"Do you still have my rectal thermometer?"

(credits: prof Juan Lema)



生活習慣病+血管状態 Lifestyle & Blood Analysis

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0.00-0.11

6.5-8.3

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5-37

123-220

30-150

41-110

2.3-7.0

6.0-22.6

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1163

Complete Analysis

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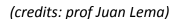
Weight(kg)

BMI Analysis

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#### Sophisticated monitoring ( ... but not that much ... )







#### Off – line monitoring

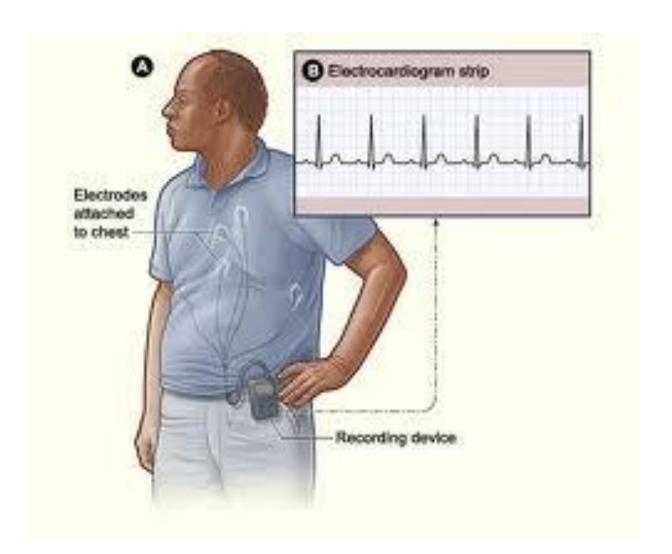




(credits: prof Juan Lema)



#### On – line monitoring







#### "BASIC" MONITORING

- √ temperature
- ✓ pH



#### "MILD" MONITORING

- √ temperature,
- **√**pH
- √ total and partial alkalinity

#### "HARD" MONITORING

- √ temperature
- ✓ pH
- ✓ total and partial alkalinity
- ✓ VFA conc and composition
- ✓ Biogas flowrate and composition
- ✓ Any other parameter.....



#### Monitoring of pH



pH is a measure of H<sup>+</sup> activity in an aqueous solution.

Because of its simplicity to be determined it is the most used parameter for AD monitoring: unfortunately, it is a "generic" parameter and almost useless in most of the case (see below)!

"Normal" pH values for AD processes follow in the range 6.5 – 8.5 but very different values can be found in stable AD processes perfectly working: this is because pH is the result of the presence several compounds in the bulk (ammonia, VFA, phosphates ....)



Lab pH-meters (off-line) or industrial on-line systems can be used for the purpose of pH measure









#### Please note that

In terms of routine monitoring, pH measurement cannot form the sole indication of imminent failure, because in medium or well-buffered solutions high VFA concentration would have to form in order to cause a detectable drop in pH, by which time failure would already occur.

Consequently, direct measurement of either (or both) VFA, partial and total alkalinity concentration is necessary.



#### Monitoring of alkalinity

Alkalinity measures the capability of a solution to neutralize acids and can be expressed using several different units (typically calcium carbonate concentration).

In AD processes it is strongly influenced by the presence of carbonate and bicarbonate ( $CO_2$  dissolved in water), ammonia, phosphate, silicate, volatile fatty acids .... And is generally very high (some g/L as  $CaCO_3$ ).

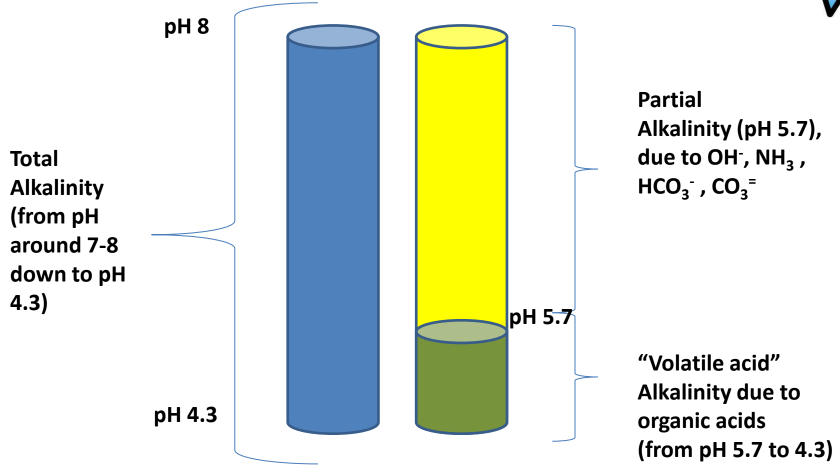
It is generally determined by tritation using 0.1 N HCl (or another strong acid like  $H_2SO_4$ ) at two equivalent points:

pH 5.75 partial alkalinity (PA) pH 4.3 or total alkalinity (TA) (or similar, see specific literature)



The difference between TA and PA is the so called IA (intermediate alkalinity) and is related to VFA presence







Partial to total alkalinity ratio (Ripley et al., 1986)

They suggested the tritation of the digestate to two final points: 5.75 and 4.3.

Then, the ratio ( $\alpha$ ) between the "VFA alkalinity" (determined as the difference between alkalinity at pH 4.5 and 5.75, intermediate alkalinity) and the partial alkalinity, determined at pH 5.75, is suggested to define the "state" of the anaerobic process.

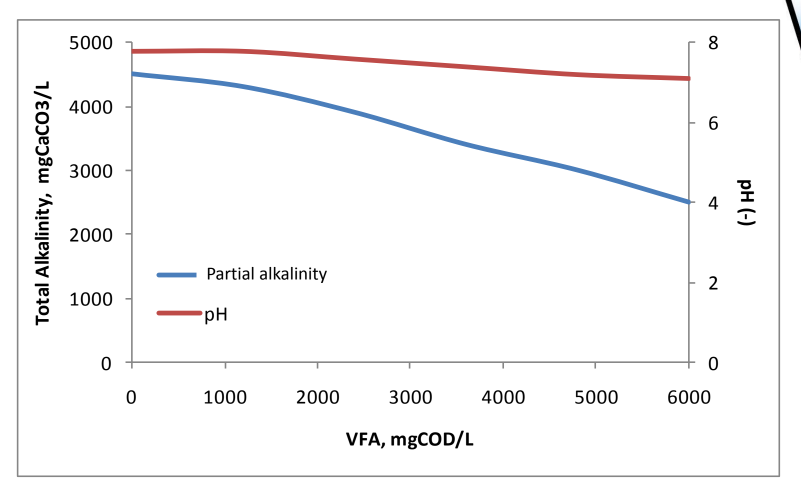
In particular, they suggested that values < 0.3 indicate a good state of the anaerobic process:

$$\alpha = \frac{IA}{PA} < 0.3$$

(this is also known as FOS/TAC in German, but obtained at different pH values)



Alkalinity determines the buffer capacity of the system and, differently from pH, is sensitive to process changes!



pH versus partial alkalinity for increasing VFA concentrations in the AD reactor



Clearly, different species in the bulk will contribute with a different "intensity" to the buffering capability of the system



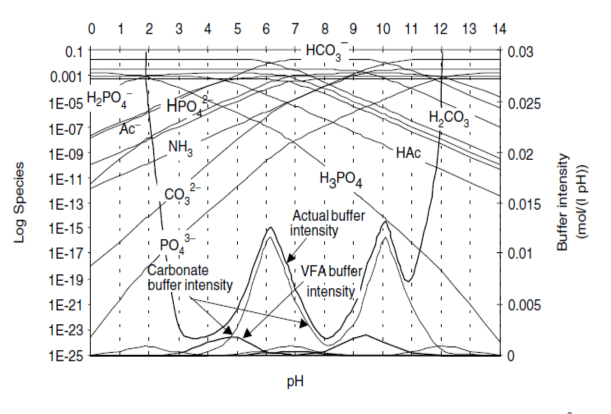


Figure 1. pH-log species and buffer intensity index diagrams in a typical anaerobic digestion sample ( $C_T = 1000 \, \text{mg} \, \text{dm}^{-3}$  as CaCO<sub>3</sub>, VFA =  $100 \, \text{mg} \, \text{dm}^{-3}$  as HAc, total phosphate concentration ( $P_T$ ) =  $50 \, \text{mg} \, \text{dm}^{-3}$  as P, total sulfide concentration ( $S_T$ ) =  $20 \, \text{mg} \, \text{dm}^{-3}$  as S, and total aqueous ammonium concentration ( $N_T$ ) =  $50 \, \text{mg} \, \text{dm}^{-3}$  as N, temperature =  $22 \, ^{\circ}$ C, TDS =  $3000 \, \text{mg} \, \text{dm}^{-3}$ ). Actual buffer intensity is the sum of buffer intensity curves of all subsystems.

Lahav & Morgan, JCTB, 79(2004), 1331-1341



#### Monitoring of volatile fatty acids (VFA)

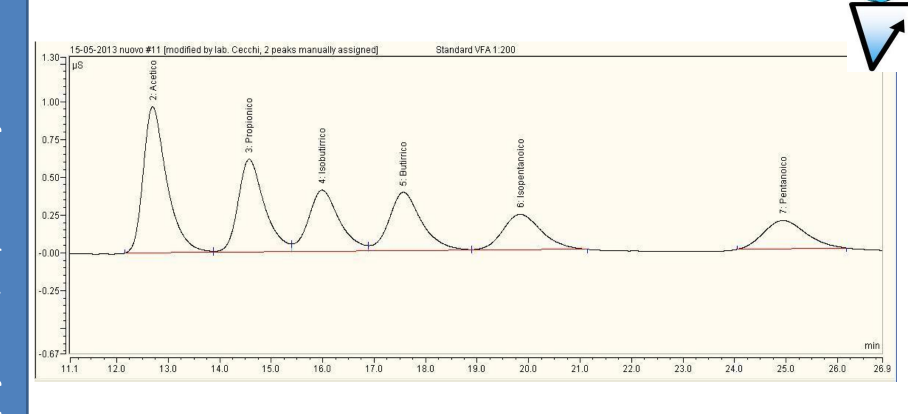
The fatty acids used for monitoring purposes are the so called short-chain volatile fatty acids (SC-VFA) with less than 6 atoms of carbon, from acetate (C2) to valerate (C5).

In a stable reactor, where the hydrolytic and methanogenic rates are balanced, with  $k_{methane}$   $k_{hydrolysis}$  their presence will be constant, and normally at a very low level of concentration (however, AD reactors treating very biodegradable substrates can be an exception).



Substrate (particulate organic matter)





#### VFA presence in an AD reactor (standard)



Concentration levels in the system can be very different, from 100 to X,000 mg/L ....
Without upsetting the process stability .....

..... but .....

The higher the VFA presence in the digestate, the lower the biogas production ....

(1 g COD as acetate produce 0.35 L CH<sub>4</sub>)



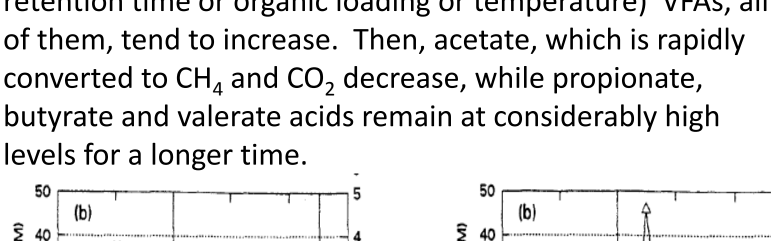


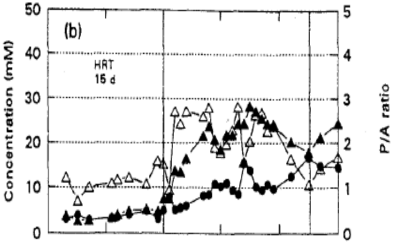
Current methods for VFA measurement include distillation, colorimetry, gas or liquid chromatography and various titration techniques.

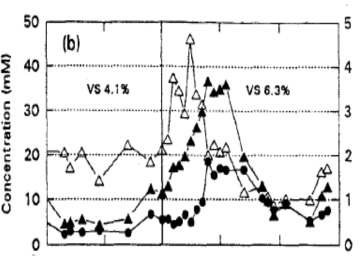
Despite titration techniques are simple and low in time and money consuming, only GC and LC analysis allow for the definition of the different VFA species present in the reactor, and this info is particularly valuable ....



After a perturbation of a stable system (change in hydraulic retention time or organic loading or temperature) VFAs, all of them, tend to increase. Then, acetate, which is rapidly converted to CH<sub>4</sub> and CO<sub>2</sub> decrease, while propionate, butyrate and valerate acids remain at considerably high



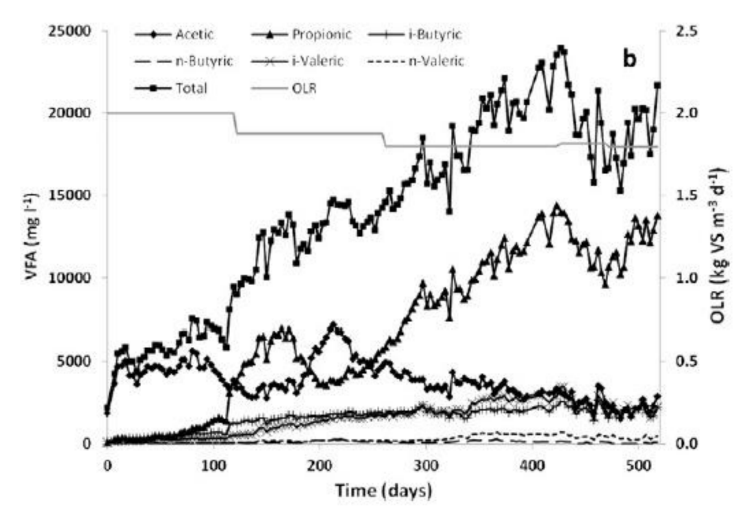




Changes in acetate ( $\Delta$ ) and propionate ( $\pm$ ) concentrations and P/A ratio ( $\bullet$ ) after a change in the HRT (left) and organic loading (right) (from Ahring et al., Appl Microbiol Biotechnol 43(1995), 559)

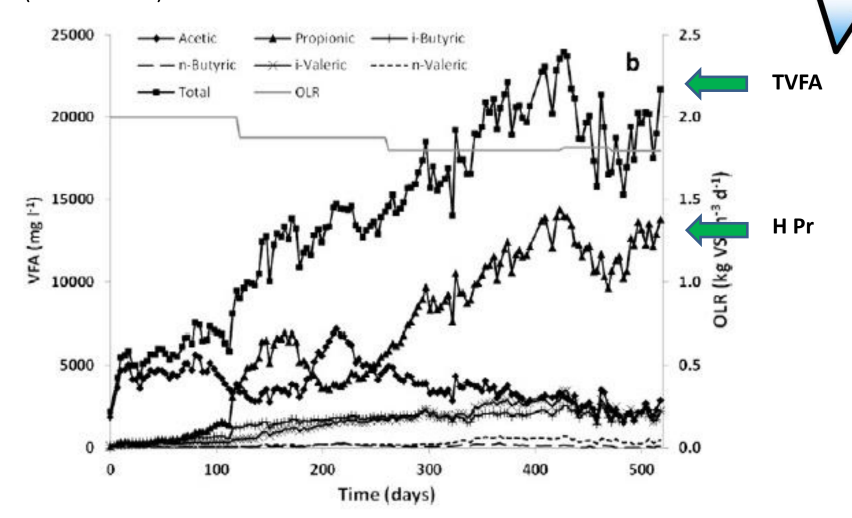


Propionic acid is typically the major compound found in unbalanced situations ... (+ formic acid).

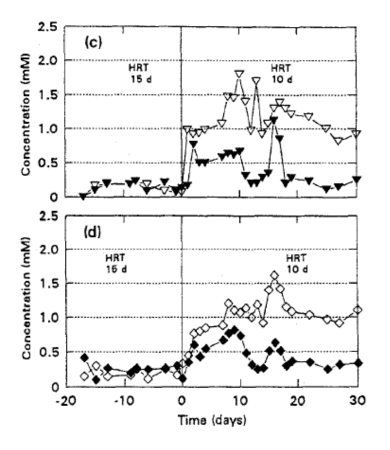


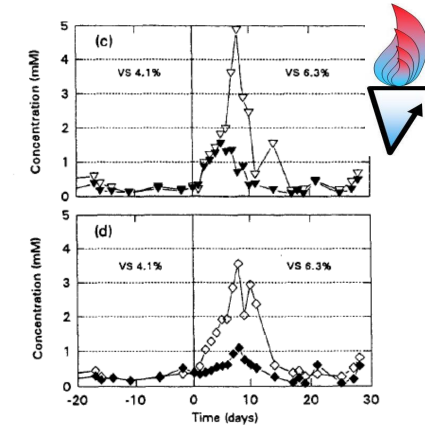


Propionic acid is typically the major compound found in unbalanced situations .... (+ formic acid).









Changes in isobutyrate (△) and butyrate (•) and isovalerate (◊) and valerate (•) concentrations after a change in the HRT (left) and organic loading (right) (from Ahring et al., Appl Microbiol Biotechnol 43(1995), 559)



However, more properly, the three parameters pH, alkalinity and VFA should be considered as a whole (then H<sub>2</sub>/CO<sub>2</sub>/CH<sub>4</sub> presence in biogas complete the set of info).

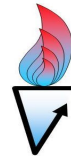


In fact, under stable operating conditions, H<sub>2</sub> (with CO<sub>2</sub>) and acetic acid formed by acidogenic and acetogenic bacterial activity are utilized immediately by the methanogens and converted to methane.

Consequently, the VFA concentration in properly running anaerobic digesters is typically fairly stable and (generally) low as *carbonate alkalinity* is not consumed in excess and the pH is stable.



#### Monitoring of biogas flowrate and composition



Biogas flowrate and its composition in terms of methane, dioxide carbon and hydrogen is strongly dependent on the substrate characteristics of course but also feeding conditions and the state of the reactor. These are therefore fundamental parameters to be known.

It is clear that an increase in terms of  $\rm H_2$  and  $\rm CO_2$  presence in biogas is related to an over-loading situation and, probably, SC-VFA accumulation.



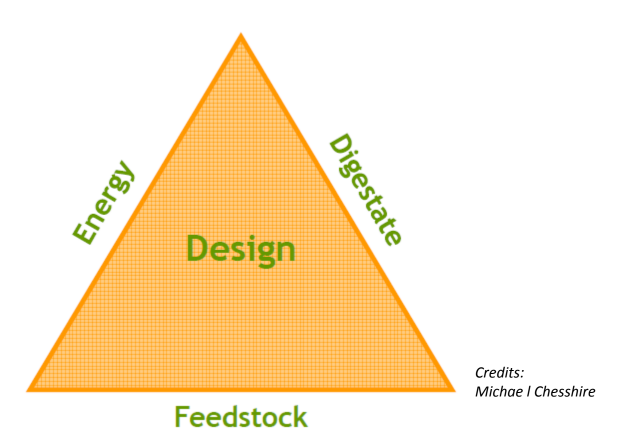


# Determination of the Biochemical Methane Potential (BMP)



Biomethane potential of substrates is important as energy is one of the 3 pillars of AD engineering (tariff+heat)







 AD is a proven technology which is gaining more and more appeal in recent years. Now, several "new" substrates are fed to reactors



- @ Because of the necessity of defining yields and economic balances, the assessment of the biomethane potential of a given substrate is very important
- Several standard methods have been proposed by different agencies during the last years. They generally focus on the biodegradability concept of single molecules rather than the biogas potential and some of them are really questionable
- Prom the scientific stand point, hundreds of papers have been publishing reporting different methods and using different units for the definition of the BMP.....



In 2007 the ABAI task group of the International Water Association proposed a protocol for the BMP determination (see Angelidaky et al. (2009), WST 59(5), 927)

The objective of the protocol is to provide important experimental guidelines to carry out an accurate assessment of the anaerobic biodegradability of any compound or material to methane and carbon dioxide and define the ultimate methane production of a given substrate in terms of m<sup>3</sup>CH<sub>4</sub>/kgVS





## Defining the biomethane potential (BMP) of solid organic wastes and energy crops: a proposed protocol for batch assays

I. Angelidaki, M. Alves, D. Bolzonella, L. Borzacconi, J. L. Campos, A. J. Guwy, S. Kalyuzhnyi, P. Jenicek and J. B. van Lier

#### **ABSTRACT**

The application of anaerobic digestion technology is growing worldwide because of its economic and environmental benefits. As a consequence, a number of studies and research activities dealing with the determination of the biogas potential of solid organic substrates have been carrying out in the recent years. Therefore, it is of particular importance to define a protocol for the determination of the ultimate methane potential for a given solid substrates. In fact, this parameter determines, to a certain extent, both design and economic details of a biogas plant. Furthermore, the definition of common units to be used in anaerobic assays is increasingly requested from the scientific and engineering community. This paper presents some guidelines for biomethane potential assays prepared by the Task Group for the Anaerobic Biodegradation, Activity and Inhibition Assays of the Anaerobic Digestion Specialist Group of the International Water Association. This is the first step for the definition of a standard protocol.

Key words | anaerobic digestion, batch assays, biomethane potential (BMP), energy crops, organic solid waste

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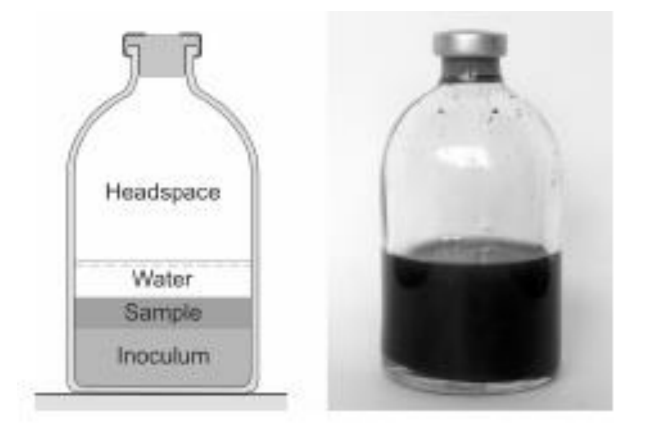
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<sup>\*</sup>TG-ABAI - Task Group for the Anaerobic Biodegradation, Activity and Inhibition of the Anaerobic Digestion Specialist Group of the International Water Association. doi: 10.2166/wst.2009.040

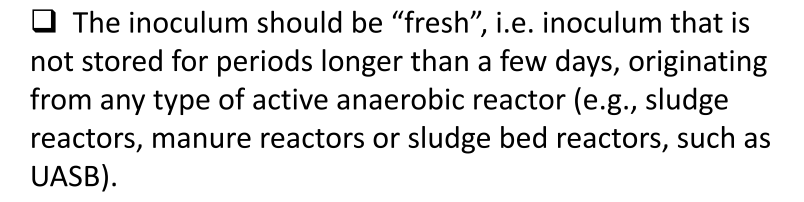
## What you need for a BMP trial at a given temperature is :

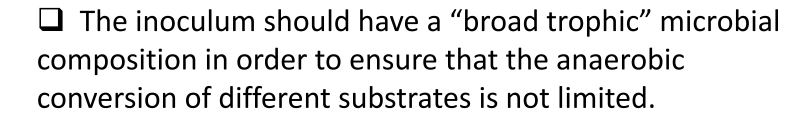
- . The substrate
- II. An active and sufficient inoculum
- III. A device for biogas measurement





## Inoculum





☐ Characteristics in terms of TS, TVS, COD and activity (on acetate) should be always stated



Inoculum pre-treatment (some suggestion ...)

The inoculum should be "degassed" (i.e., pre-incubated) in order to deplete the residual biodegradable organic material (endogenous)

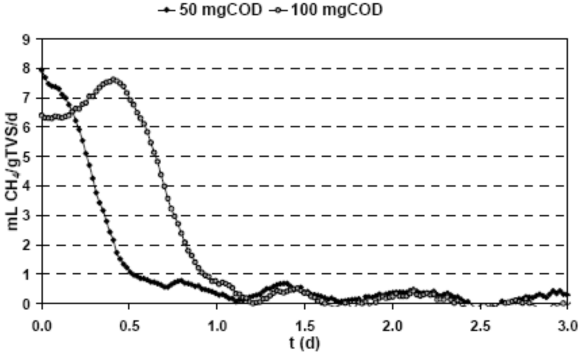
The inoculum should be a close representation of the one sampled from the reactor, and SHOULD NOT (as described on previous ISO 11734, ASTM E 2170 (2001)) be washed to remove residual substrate material and inorganic carbon compounds.



## Activity of the inoculum

The quality of inoculum could be examined by performing activity tests on acetate and cellulose (or gelatine or both)





Specific Methanogenic Activity on acetate



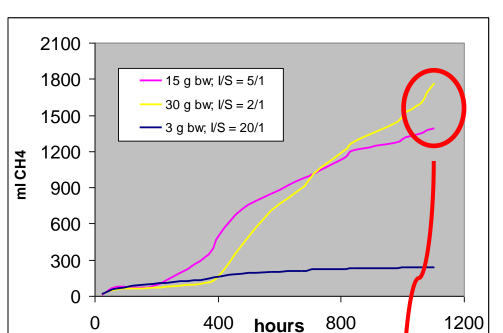


- □ Typical SMA determined on acetate are in the range 30 40 mlCH<sub>4</sub>/gVSSd but can be as low as 10 mlCH<sub>4</sub>/gVSSd; in last case the inoculum concentration in the test vessel should be very high (typically being 70% to 80% of the total liquid used in the test).
- High inoculum concentrations should always be preferred. This offers an excess of active biomass and buffer capacity, conditions similar to those of real anaerobic reactors (generally CSTR)



# Inoculum to substrate ratio

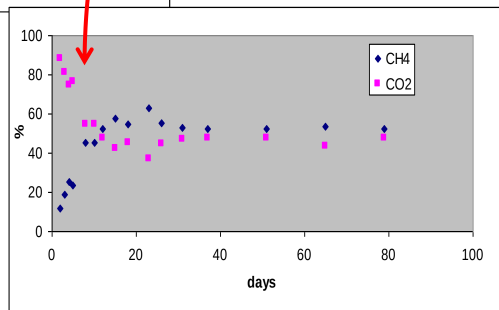




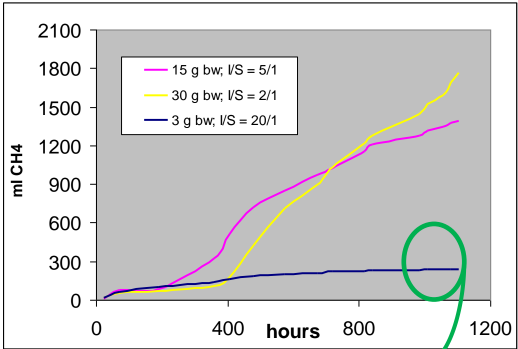
Too much substrate ...

Initial flat curve and CO<sub>2</sub> evolution ...

Acidogenic conditions!



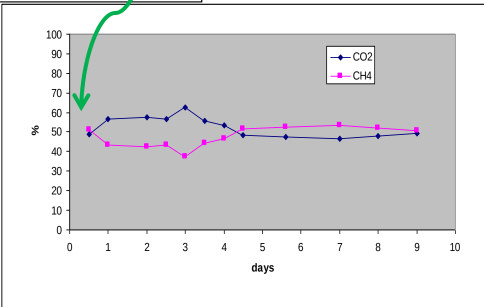






Right proportion ...

Good curve, useful also for kinetic tests and  $CH_4$  evolution ...





If the specific activity and hydrolysis constant are known (or at least estimated) the correct ratio can eventually be determined according the mass balance for VS, which gives:



$$Vinoculum = \frac{X_{SS}V_{ww}k_h}{VSS_{inoculum}SMA_{inoculum}}$$

 $X_{SS}$  concentration of hydrolysable substrate (g/L),  $V_{ww}$  volume of waste(water) in the assay vessel (L),  $k_h$  first order hydrolysis constant (day<sup>-1</sup>),  $VSS_{inoculum}$  content of the inoculum (gVSS/L) and  $SMA_{inoculum}$  specific methanogenic activity of the inoculum (g COD-CH<sub>4</sub>/(gVSSday)).

Angelidaki and Sanders, 2004, Rev in Env Sci & BioTechnol, 3(2), 117



## Medium

Necessary nutrients/micronutrient/vitamins/buffers are generally required for optimal performance of anaerobic microorganisms

Table 4.2. Basic Anaerobic Medium (Angelidaki and Sanders 2004).

## Description of Anaerobic Basic Medium

The basic medium is prepared from the following stock solutions, (chemicals given below are concentrations in g/l, in distilled water).

- (A) NH<sub>4</sub>Cl, 100; NaCl, 10; MgCl<sub>2</sub> 6H<sub>2</sub>O, 10; CaCl<sub>2</sub> 2H<sub>2</sub>O, 5
- (B) K<sub>2</sub>HPO<sub>4</sub> 3H<sub>2</sub>O, 200
- (C) resazurin 0.5
- (D) trace-metal and selenite solution: FeCl<sub>2</sub> 4H<sub>2</sub>O, 2; H<sub>3</sub>BO<sub>3</sub>, 0.05; ZnCl<sub>2</sub>, 0.05; CuCl<sub>2</sub> 2H<sub>2</sub>O, 0.038; MnCl<sub>2</sub> 4H<sub>2</sub>O, 0.05; (NH<sub>4</sub>)<sub>6</sub>Mo<sub>7</sub>O<sub>24</sub> 4H<sub>2</sub>O, 0.05; AlCl<sub>3</sub>, 0.05; CoCl<sub>2</sub> 6H<sub>2</sub>O, 0.05; NiCl<sub>2</sub> 6H<sub>2</sub>O, 0.092; ethylenediaminetetraacetate, 0.5; concentrated HCl, 1 ml; Na<sub>2</sub>SeO<sub>3</sub> 5H<sub>2</sub>O, 0.1
- (E) vitamin mixture (componets are given in mg/l): Biotin, 2; folic acid, 2; pyridoxine acid, 10; ridoflavin, 5; thiamine hydrochloride, 5; cyanocobalamine, 0.1; nicotinic acid, 5; P-aminobenzoic acid, 5; lipoic acid, 5; DL-pantothenic acid.

To 974 ml of distilled water, the following stock solutions should be added (A), 10 ml; (B), 2 ml; (C), 1 ml; (D), 1 ml and (E), 1 ml. The mixture is gassed with 80% N<sub>2</sub> - 20% CO<sub>2</sub>. Cysteine hydrochloride, 0.5 g and NaHCO<sub>3</sub>, 2.6 g, are added and the medium is dispensed to serum vials and autoclaved if necessary. Before inoculation the vials are reduced with Na<sub>2</sub>S 9H<sub>2</sub>O to a final concentration of 0.025%.

## Data collection of produced biogas

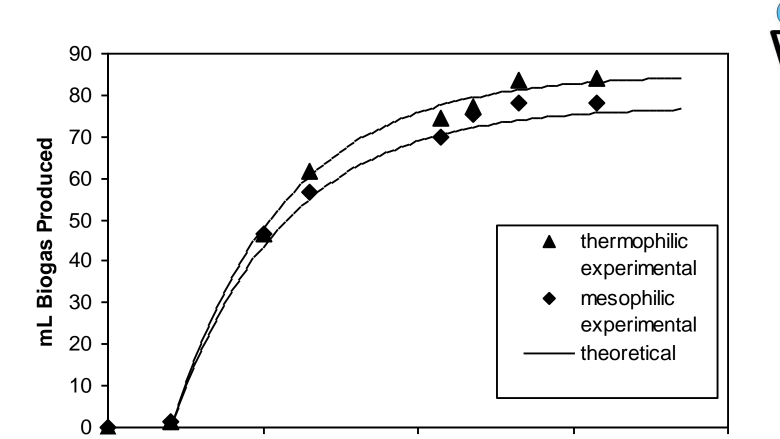
Method	Comments
Volumetric	Inaccuracy due to variations of atmospheric pressure
	Inaccuracy due to inorganic carbon in liquid phase
	Evaporation of water in displacement systems
	<ul><li>Simple and cheap</li></ul>
Manometric	<ul> <li>Manometric transducers limited range of accuracy</li> </ul>
	Inaccuracy due to inorganic carbon in liquid phase
GC- TDC	<ul> <li>Special equipment</li> </ul>
	Time and labour requiring
	<ul><li>Many simultaneous</li></ul>
	Direct measurement, precise
GC- FID	Special equipment
	◆ Fast
	<ul><li>Many simultaneous</li></ul>
	Direct measurement, precise



The methane accumulated in the headspace of the closed vessel should be measured by gas chromatography (GC). For that, a sample volume of e.g. 100 μL should be collected with a gas-tight syringe and injected into the GC. Either a Thermal Conductivity Detector (TCD) or a Flame Ionization Detector (FID) can be used. The obtained peak area should be compared to that obtained by injecting the same volume of a standard gas mixture of the known composition.

The volume of methane produced is obtained by multiplying the headspace volume by the % of  $CH_4$  in the headspace as determined by GC analysis. For publication and comparison with other studies, the values are often calculated to STP conditions, i.e. converted to 0°C and 1 atm.

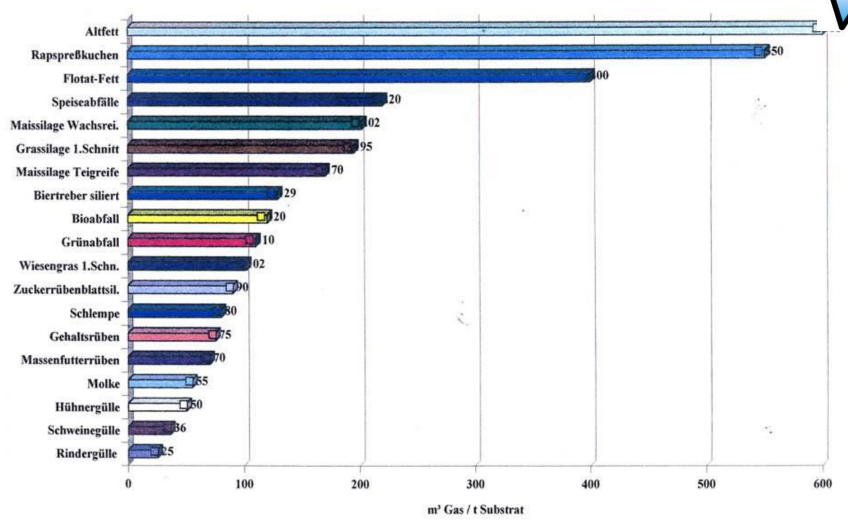




hours



## Biogas potential for different substrates





## Determination of the hydrolysis constant (d<sup>-1</sup>) (first order kinetic)

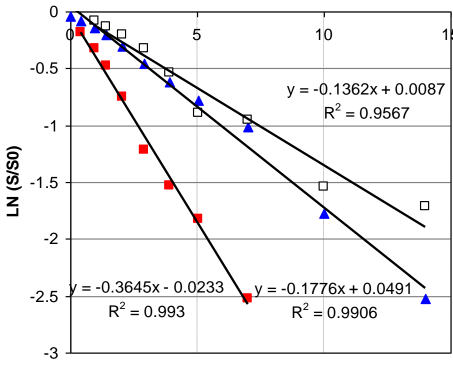


$$\frac{\mathrm{d}S}{\mathrm{d}t} = -k_h S$$



$$n\frac{B_{\infty}-B}{B_{\infty}}=-k_h t$$

## Fit on Methane







As a first indication, the following range of values can be considered:

 $k_h > 0.5 d^{-1}$ , very biodegradable substrates (e.g., food waste, material rich in carbohydrates ....)

 $k_h = 0.2-0.3 d^{-1}$ , intermediate biodegradable substrates (e.g., food waste, material rich in carbohydrates ....)

 $k_h = 0.1 d^{-1}$ , slowly biodegradable substrates (e.g., waste activated sludge, or biological sludge)



## Round-test in 16 Italian labs



Box plot: median + 25° and 75° percentile

