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Study and modeling of separation methods H₂S from methane, selection of a method favoring H₂S valorization

Hamadi Cherif

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de l'Université de recherche Paris Sciences et Lettres
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Préparée à MINES ParisTech

Etude et modélisation de méthodes de séparation du méthane et de H₂S

Study and modeling of separation methods of H₂S from methane

Ecole doctorale n°432

SCIENCES DES METIERS DE L'INGENIEUR

Spécialité ENERGETIQUE ET GENIE DES PROCÉDES

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le 08 Décembre 2016

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Nomenclature

a	Specific geometric packing surface area	m^2/m^3
A	Constant used for the calculation of pressure drop	-
a_e	Effective interfacial area	m^2
b	Length of the corrugation base	m
B	Constant used for the calculation of pressure drop	-
C_{FI}	Specific packing constant for calculation of hydrodynamic parameters at flooding point	-
C_h	Specific packing constant for hydraulic area	-
C_L	Specific packing constant for mass transfer calculation in liquid phase	-
C_{lp}	Specific packing constant for calculation of hydrodynamic parameters at loading point	-
C_p	Specific packing constant for pressure drop calculation	-
C_V	Specific packing constant for mass transfer calculation in gas phase	-
D_V	Gas-phase diffusion coefficient	m^2/s
d	Packing diameter	m
d_h	Hydraulic diameter	m
d_{hV}	Diameter of the gas flow channel	m
d_p	Particle diameter	m
f	Approach to flood	-
F_c	Gas capacity factor	$(\text{m/s}) (\text{kg}/\text{m}^3)^{0.5}$
$F_{c,lp}$	Gas capacity factor at loading point	$(\text{m/s}) (\text{kg}/\text{m}^3)^{0.5}$
Fr_L	Liquid Froude number	-
F_I	Enhancement factor for pressure drop calculation	-
F_t	Correction factor	-
F_{SE}	Surface enhancement factor for effective interfacial area calculation	-
f_w	Wetting factor	-
g	Gravitational constant	m/s^2
g_{eff}	Effective gravitational constant	m/s^2
h	Corrugation height	m
h_L	Liquid holdup	m^3/m^3
$h_{L,FI}$	Liquid holdup at flooding point	m^3/m^3
$h_{L,lp}$	Liquid holdup at loading point	m^3/m^3
$h_{L,pl}$	Liquid holdup in preloading region	m^3/m^3
K	Wall factor	-
k_G	Gas-phase mass transfer coefficient	m/s
k_L	Liquid-phase mass transfer coefficient	m/s
L	Liquid mass flow	kg/h
M	Molar mass	kg/mol
n_{FI}	Exponent for calculation of liquid holdup at flooding point	-
n_{lp}	Exponent for calculation of liquid holdup at loading point	-
P	Pressure	Pa
P_c	Critical pressure	Pa
Pe	Peclet number	-
Re_L	Liquid Reynolds number	-
Re_V	Gas Reynolds number	-
$Re_{V,e}$	Effective Reynolds number for gas phase	-
$Re_{V,r}$	Relative Reynolds number for gas phase	-
s	Length of the corrugation side	m

Sc	Schmidt number	-
Sh	Sherwood number	-
T	Temperature	K
T _B	Boiling temperature	K
T _c	Critical temperature	K
u _L	Superficial liquid velocity	m/s
u _{L,e}	Effective liquid velocity	m/s
u _{L,lp}	Superficial liquid velocity at loading point	m/s
u _V	Superficial gas velocity	m/s
u _{V,e}	Effective gas velocity	m/s
u _{V,Fl}	Superficial gas velocity at flooding point	m/s
u _{V,lp}	Superficial gas velocity at loading point	m/s
V	Gas mass flow	kg/h
We _L	Liquid Weber number	-
z	Unit length	m
$\frac{dP}{dz}$	Pressure drop	Pa/m
$\left(\frac{dP}{dz}\right)_d$	Dry pressure drop	Pa/m
$\left(\frac{dP}{dz}\right)_{pl}$	Pressure drop in preloading region	Pa/m
α _L	Liquid flow angle	°
γ	Solid – liquid film contact angle	°
δ	Liquid film thickness	m
ε	Void fraction	-
ζ _{DC}	Coefficient for losses caused by direction change	-
ζ _{GG}	Coefficient for gas/gas friction losses	-
ζ _{GL}	Coefficient for gas/liquid friction losses	-
θ	Corrugation angle	°
μ _L	Dynamic viscosity of the liquid phase	kg/m.s
μ _V	Dynamic viscosity of the gas phase	kg/m.s
ν _L	Kinematic viscosity of the liquid phase	m ² /s
ξ _{bulk}	Direction change coefficient in the bulk zone	-
ξ _{GG}	Gas/Gas friction coefficient	-
ξ _{GL}	Gas/Liquid friction coefficient	-
ξ _{wall}	Direction change coefficient near the wall	-
ρ _L	Liquid density	kg/m ³
ρ _V	Gas density	kg/m ³
σ _L	Surface tension of the liquid phase	N/m
σ _w	Surface tension of water	N/m
φ	Fraction of the flow channel occupied by the liquid phase	-
ψ ₀	Resistance coefficient for dry pressure drop calculation	-
ψ _{Fl}	Resistance coefficient for pressure drop calculation at flooding point	-
ψ _L	Resistance coefficient for wet pressure drop calculation	-
ψ' _L	Resistance coefficient for wet pressure drop calculation	-
ψ _{lp}	Resistance coefficient for pressure drop calculation at loading point	-
Ω	Fraction of the packing surface occupied by holes	-

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General introduction

Recent decades were accompanied by economic growth and prosperity for humanity. This growth gained from oil and natural gas production has been accompanied by an environmental pollution that may trigger irreversible changes in the environment with catastrophic consequences for humans. Moreover, issues related to the reduction of fossil reserves are still relevant, and the global primary energy demand is increasing, pushing the international community to pursue the development of renewable energies.

The law on energy transition plans to reduce the share of fossil fuel consumption in France to 30 % in 2030, while the share of renewable energy should be increased to 23 % in 2020 and 32 % in 2030 against 16.1 % in 2012 as shown in the Fig. 1 [1].

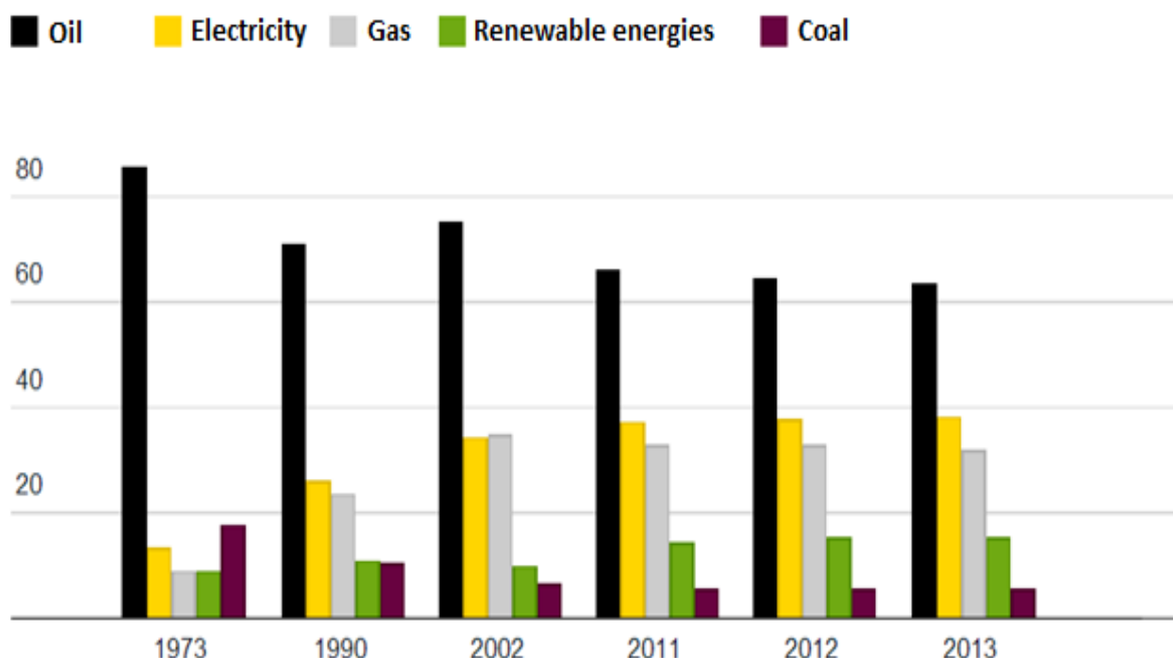


Fig. 1: Distribution of final energy consumption in France [1]

Renewable energy is derived directly from natural phenomena. It takes many forms: sunlight, wind, wood heat, water power ... The main renewable energies are: wind power, wave power, geothermal energy, solar energy, biomass, firewood and hydropower ...

According to the French Ministry of Energy [1], primary production of renewable energy rises to 22.4 Million tonnes of oil equivalent (Mtoe) in 2012. The distribution of renewable energies production by sector is depicted in Fig. 2.

Among renewable energies, biogas is a possibility. It is storable, transportable, not intermittent and substitutable for fossil energies. These strengths justify the consolidation of this emerging sector and the preparation of its future development by ambitious public policies. The European primary energy production from biogas reached 5901 kilotonnes of oil equivalent (ktoe) in 2007, 7500 ktoe in 2008, 10086 ktoe in 2011 and 13379 ktoe in 2013.

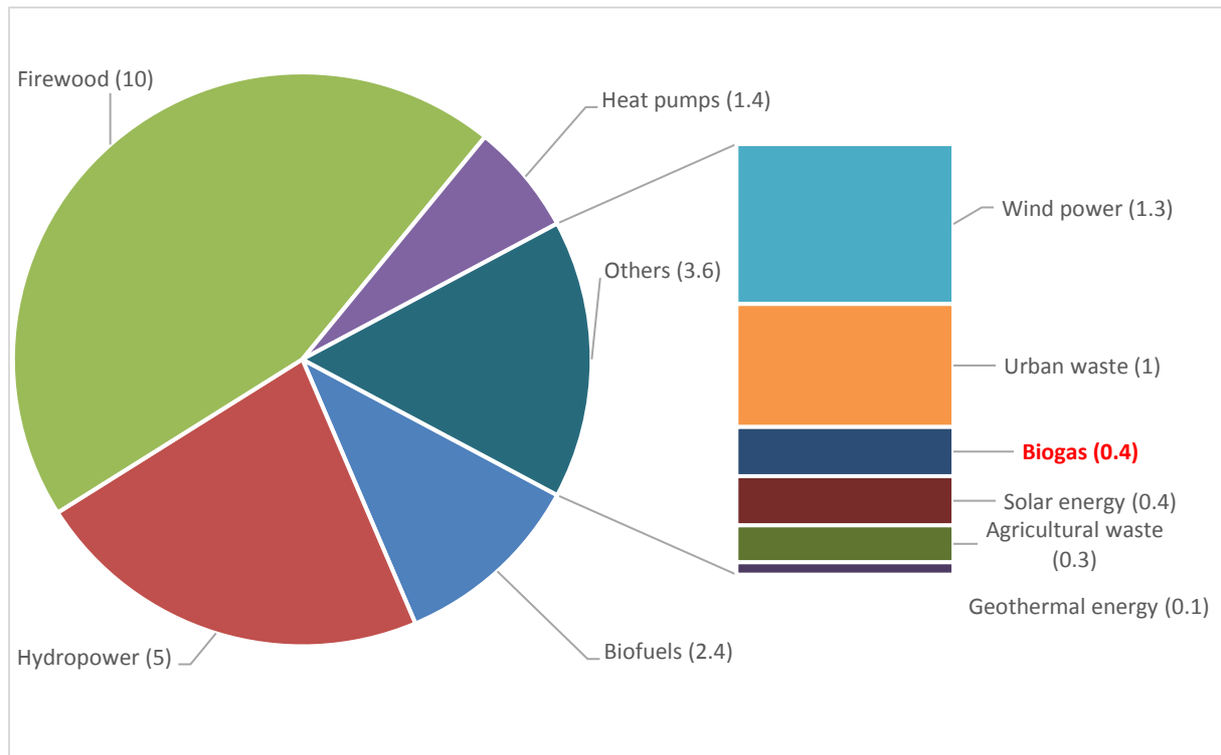


Fig. 2: Distribution of production of renewable energies by sector [1]

Biogas production is at the heart of the policy priorities for 2020, the European Union aims for a production that will cover 50 % of transport sector needs. Biogas is a growing sector. Fig. 3 provides a quick review of biogas production in Europe in 2013 [2].

Generally, this gas composed of methane and carbon dioxide, also contains other compounds as water, ammonia, volatile organic compounds and hydrogen sulfide.

Biogas can be valued in several applications such as production of heat and/or electricity, feed for fuel cells, injection into the natural gas grid and production of liquefied bio-methane. This last application presents an environmental and economic benefits. With hydrogen fuel, liquefied bio-methane is one of the best fuel for the reduction of carbon dioxide emissions with a reduction potential up to 97 % compared to Diesel. It can effectively reduce emissions of greenhouse gases and the pollution, responsible for 42000 premature deaths annually in France [3].

Primary production of biogas in the European Union in 2013** (in ktoe)

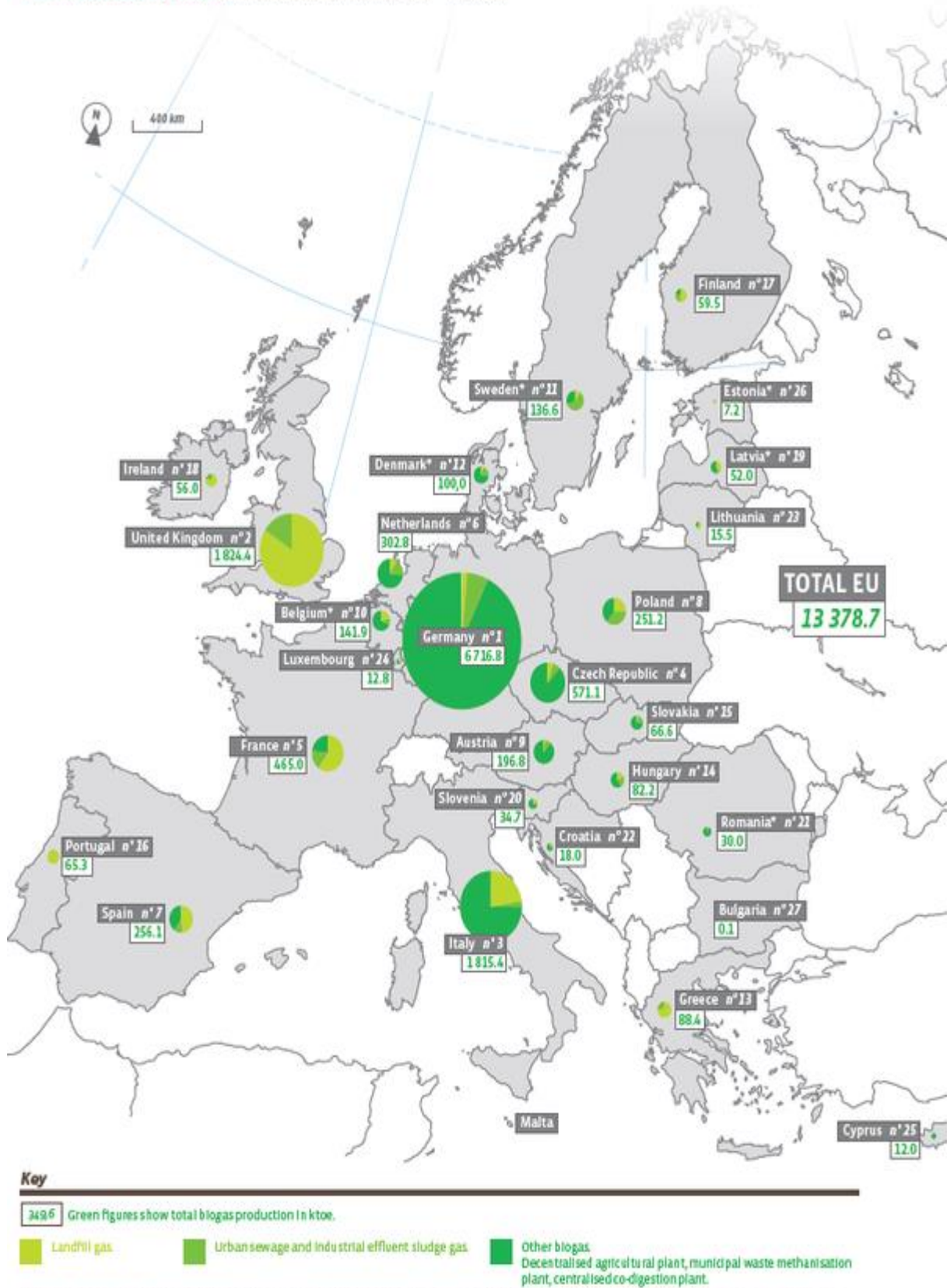


Fig. 3: Biogas primary production in 2013 [2]

Liquefied bio-methane requires very low temperatures which may lead to solidification of impurities and thus facilities malfunctions. These impurities must be separated from the biogas. This implies implementation of a purification process to remove from the raw biogas all unwanted substances in order to maximize its methane content. In particular, the complete desulfurization of biogas is peremptory in order to ensure an optimal operation and a high purity of other compounds to valorize, such as carbon dioxide. Moreover, the presence of hydrogen sulfide in the wet biogas, is a poison for installations. In case of leakage, the presence of hydrogen sulphide in the biogas characterized by a rotten egg smell can be dangerous for operators working on the site. Hence, the need to remove all traces of this compound. Several technologies are used for the removal of H₂S as adsorption on microporous solids, membrane technology, biological processes and absorption by means of liquid solvents.

The choice of the technique to implement is related to various parameters. The most important are the flow rate of biogas and hydrogen sulfide concentrations to treat. The solution must also respond to various economic, environmental and energetic imperatives as cost which must be reasonable, the threshold to reject which must be respected in an energy-efficient process. Therefore, there is no universal treatment method.

In this thesis, the choice fell on the use of a cryogenic method in order to combine biogas upgrading and biomethane liquefaction. The removal of H₂S will be performed upstream of the process either by reactive absorption using an aqueous solution of sodium hydroxide (NaOH) in a structured packed column or by adsorption using activated carbon. These two technologies will be tested and compared in order to choose the most effective and the most suitable for the process set up.

Experiments were performed on a industrial pilot plant developed by Cryo Pur[®] company called "BioGNVAL". This pilot plant treats 85 Nm³/h of biogas from waste water treatment plant, which contains around 20 – 100 ppm of H₂S.

For absorption technology, the hydrodynamic of flows in structured packing columns was studied in order to develop a model able to predict realistically the key hydrodynamic parameters as pressure drop, liquid holdup and transition points but also effective interfacial area and global mass transfer coefficients.

The remainder of the study is based on simulations using Aspen Plus[®] V8.0 to study realistically the effectiveness of a structured packed column which uses sodium hydroxide as a chemical solvent for the selective removal of hydrogen sulfide from biogas. Results were compared with data from BioGNVAL pilot plant.

Finally, the dynamic aspect of the adsorption phenomenon is modeled, by predicting the breakthrough curve in the case of an adsorption column used for the removal of H₂S.

Chapter 1: From biogas to biomethane

Résumé :

Après une introduction générale posant le contexte de la production de biogaz dans le paysage énergétique mondial, le premier chapitre est une introduction à la problématique de son obtention et de son usage sous une version raffinée en biométhane.

Le biogaz est produit par méthanisation lorsque des matières organiques sont décomposées dans des conditions bien définies et en absence d'oxygène. Il est constitué principalement de méthane et de dioxyde de carbone mais aussi de quantités variables de vapeur d'eau, de sulfure d'hydrogène et d'autres composés polluants. Afin de pouvoir utiliser le biogaz comme carburant pour véhicule, il doit être épuré en séparant le dioxyde de carbone et les autres composés contaminants du biogaz pour augmenter au maximum sa teneur en méthane.

Il existe différentes technologies utilisées dans le domaine du traitement des gaz. Les plus utilisées sont : l'absorption, l'adsorption, la technologie membranaire, la cryogénie, le traitement biologique et l'oxydation. Aujourd'hui, d'autres technologies sont développées telles que la biocatalyse, la photocatalyse et le plasma froid. Pour le moment, le manque d'informations à leur sujet limite leur application industrielle [18].

1.1. Introduction

Anaerobic digestion may be defined as the natural process of degradation by microorganisms under controlled conditions and in the absence of oxygen. This degradation results in the production of a gas mixture saturated with water outlet of the digester called biogas.

Anaerobic digestion is the result of four biochemical steps in which large carbon chains are converted into fatty acids and alcohols. These four steps are: Hydrolysis, acidogenesis, acetogenesis, methanogenesis [4].

- **Hydrolysis**

It takes place at the beginning of the fermentation and makes use of exo-enzymes in order to decompose the organic matter into simple substances.

- **Acidogenesis**

During this step, volatile fatty acids are formed. Carbon dioxide and hydrogen are also formed, they are used by microorganisms during the production of methane according to the chemical Reaction (R.1) shown below. The reaction enthalpy is about -567 kJ.mol^{-1} [5].

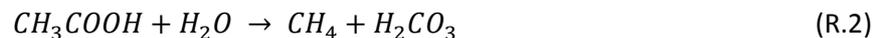


- **Acetogenesis**

This stage involves the production of acetate, an indispensable substrate for the synthesis of methane.

- **Methanogenesis**

This last step results in the production of methane. It is ensured by the methanogenic bacteria, which can only use a limited number of carbon compounds, including acetate responsible for 70 % of methane production according to the chemical Reaction (R.2) shown below. The reaction enthalpy is about -130 kJ.mol^{-1} [5].



The biogas needs to be purified and upgraded, which means that impurities are removed or valorized in order to produce a gas rich in methane called biomethane. There is a number of technologies available for this purpose as water scrubbing, membranes and pressure swing adsorption.

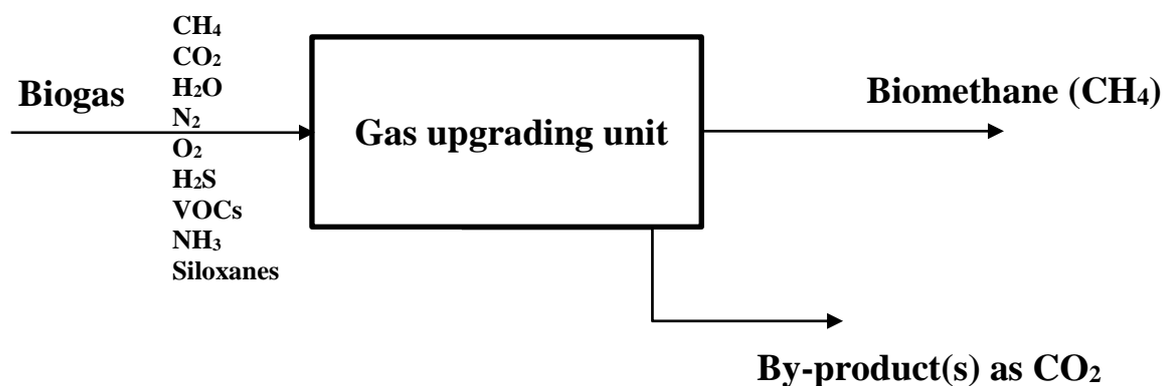


Fig. 4: Simplified diagram of production of biomethane

1.2. Biogas utilization

Biogas can be used in several applications. Sometimes it could be used raw, but almost always it has to be upgraded or as a minimum cleaned from its H₂S content because the presence of this compound in the biogas even at very low concentrations could damage the installations.

Biogas can be used in all applications designed for natural gas such as production of heat and electricity known as combined heat and power (CHP), production of chemicals and/or proteins, fueling internal combustion engines and fuel cells, it may also be used as vehicle fuel or injected in the natural gas grids.

1.2.1. Direct combustion

The simplest method of biogas utilization is direct combustion. The biogas burner can be installed in heaters for production of hot water and hot air, in dryers for various materials, and in boilers for production of steam for process heat or power generation [6]. Another limited application involves absorption heating and cooling to provide chilled water for refrigeration and hot water for industrial processes.

These applications based on direct combustion do not require a high gas quality.

Biogas can also be used to fuel internal combustion engines to supply electric power for pumps, blowers, elevator and conveyors, heat pumps and air conditioners [7].

1.2.2. Combined heat and power

Another application for biogas is CHP which involves the production of two or more forms of energy, generally electricity and thermal energy. This latter is generally the most used today but it could be a problem because the need of heat varies with season, and during summer for example, unused biogas is flared.

1.2.3. Injection into the natural gas grid

Upgrading of biogas to biomethane with a gas quality similar to natural gas and injecting it into the natural gas grid is an efficient way to integrate biogas into the energy sector. It allows the transport of large volume of biomethane and its utilization in wide areas where population is concentrated.

1.2.4. Vehicle fuel

Today, there is a big interest in using biogas as a vehicle fuel. But for this utilization, the raw biogas must be purified which means that contaminants are removed from biogas, and upgraded which means that CO₂ is eliminated leading to a raise in the energy content of biogas. Finally, the biomethane needs to be liquefied by chilling it to very low temperatures (≈ -161 °C at atmospheric pressure). In order to prevent corrosion and solid formation, some requirements concerning component concentrations are needed before liquefaction. They are presented in Table 1. The H₂S content must be lower than 4 ppm. This quantity is not in direct relation with corrosion or solidification aspects. This requirement ensures a high quality of biomethane and avoids odor problems caused by the presence of hydrogen sulfide.

Table 1. Concentration requirements before biogas liquefaction [8]

Compounds	Maximum concentration
CO ₂	25 ppm
H ₂ S	4 ppm
H ₂ O	1 ppm

Fig. 5 shows an example of use of biogas after its purification and upgrading using the Cryo Pur[®] system developed by Cryo Pur[®] Company.

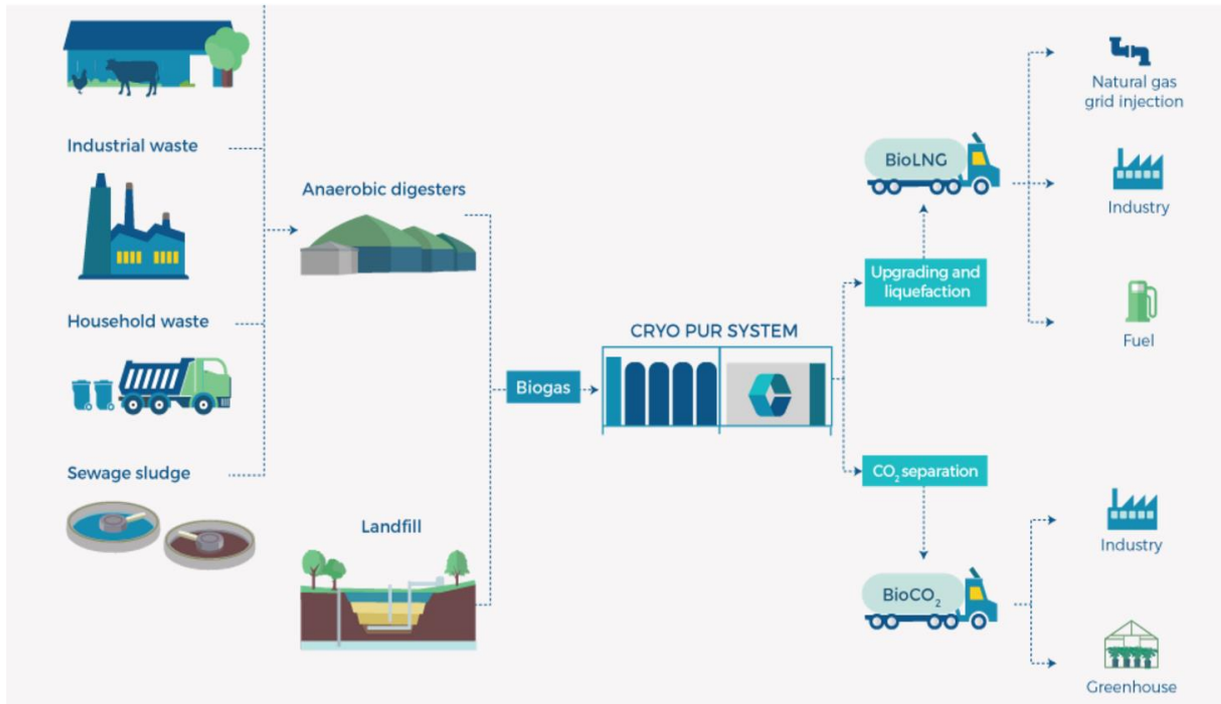


Fig. 5: Example of biogas utilization by Cryo Pur[®] Company [9]

1.3. Biogas composition

The composition of biogas depends mainly on the type of substrates which are segmented according to their origin. The following classification is commonly used:

- Household and industrial waste.
- Sludge from sewage water treatment plants.
- Agricultural and agro-industrial waste.

1.3.1. Household and industrial waste

To treat this type of waste, they are buried in landfill sites. The anaerobic conditions created by the landfill are sufficient to induce methanogenesis.

Biogas production from this type of waste is characterized by variations of flow rates and composition due to the variation of the feedstock. This biogas production is also dependent on the progression of degradation of waste, moisture and temperature. These parameters are not the same on the entire degradation zone leading to variations in the composition of biogas produced from the same landfill. This variability phenomenon is also noticeable on the concentration of methane in biogas which varies around 15 % during a year [10]. Two factors explain this: the high biological activity in summer and the rise of temperature. These conditions complicate the valorization of this type of biogas.

Household waste sorting improves the yield of biogas production. This process by which waste is separated into different elements achieves higher yields compared to unsorted waste. The difference may reach 100 m³/t [11].

Household and industrial wastes are not all fermentable. The share of these fermentable wastes represents 45 % maximum [12].

Table 2 shows the principal compounds present in household waste.

Table 2. Composition and characterization of household waste [13]

Compounds	Composition [wt. %]	Dry matter [wt. %]	Dry organic matter [wt. %]
Putrescible	33.0	44	77
Papers	11.7	68	80
Cardboards	12.0	70	80
Various incombustible	8.5	90	1
Tetra brik	8.5	70	60
Glasses	5.4	98	2
Plastic	4.9	85	90
Green waste	4.5	50	79
Textiles	4.2	74	92
Metals	3.7	90	1
Special waste	2.0	90	1
Various fuels	1.6	85	75

1.3.2. Sludge from sewage water treatment plants

Biological treatment of urban wastewater is a widespread process which generates significant amounts of activated sludge. In order to stabilize these latter, an anaerobic degradation is used. It provides a solution to the storage and treatment of sludge.

A small part of the produced biogas ensures 100 % of the energy needs of the wastewater treatment plant. The by-products of anaerobic digestion, as the digestate can be valorized via fertilization and amendment of farmland.

Today, the injection into the natural gas grid of biomethane issued from sludge of wastewater treatment plant is subject to strong demand from local authorities. According to the French Ministry of Ecology, Sustainable Development and Energy, by 2020, more than 60 wastewater treatment plants could be provided with the necessary facilities for energy recovery of waste to allow the injection of 500 GWh/year of biomethane into the national gas grid, which is equivalent to the annual consumption of more than 40000 households [1].

Unlike the treatment of household and industrial waste, anaerobic digestion of sludge from wastewater treatment plants is a controlled process conducted under optimal conditions for biogas generation.

1.3.3. Agricultural and agro-industrial waste

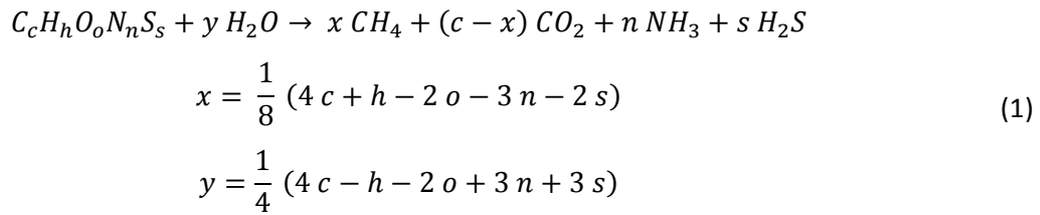
According to the European renewable energies Observatory [2], France has about 60 million tons of organic material that can be valued in biogas. These agricultural wastes are divided into two groups: liquid effluents and solid waste. For liquid effluents, dry matter and dry organic matter rates influence the production of biogas, as shown in Table 3. Digesters used for biogas production from agricultural and agro-industrial wastes are optimized systems, regulated and stable. According to Boulinguez [14], the composition of biogas from digesters varies with an amplitude of ± 5 % during the year.

Table 3. Production yields for agricultural and agro-industrial substrates [14]

Substrates	Dry matter [wt. %]	Dry organic matter [wt. %]	Production yield [m ³ /t of dry organic matter]	Production yield [m ³ /t of substrate]
Pig manure	6 to 25	75 to 80	300 to 450	22 to 60
Cow manure	8 to 30	80 to 82	350 to 700	21 to 168
Chicken manure	10 to 60	67 to 77	300 to 800	20 to 180
Horse manure	28	25	500	35
sheep manure	15 to 25	80 to 85	350	50 to 74
Grass silage	26 to 80	67 to 98	500 to 600	87 to 440
Hay	86 to 93	83 to 93	500	356 to 432
corn straw	86	72	500	310
Foliage	85	82	400	279
Sorghum	25	93	700	162
Helianthus annuus	35	88	750	231
Distillation residues	12	90	430	77
Brewery waste	15 to 21	66 to 95	500	50 to 100
Vegetable waste	5 to 20	76 to 90	600	23 to 108
oleaginous residues	92	97	600	536
Fruit residues	40 to 50	30 to 93	450 to 500	60 to 232
press cake	88	93	5550	450
slaughterhouse waste	15	80 to 90	450	58
bakery waste	50	80 to 95	450	665
Shortening waste	99	99	1200	1117

Table 3 shows that shortening (Alimentary fat) is the substrate with the highest methanogenic potential.

Assuming complete reaction without formation of by-products, Buswell Equation (1) shows that the theoretical yield of methane production may be estimated from the base elemental composition of a substrate [15].



The composition of biogas shown in Table 4 is defined according to the main types of substrates previously presented.

Table 4. Biogas composition depending on the type of substrate [14]

Compounds	Unit	Biogas from household and industrial waste	Biogas from wastewater treatment plants	Agricultural biogas
CH ₄	% mol.	40 - 55	65 - 75	45 - 75
CO ₂	% mol.	25 - 30	20 - 35	25 - 55
N ₂	% mol.	10	0 - 5	0 - 5
O ₂	% mol.	1 - 5	0.5	0 - 2
NH ₃	% mol.	Traces	Traces	0 - 3
COV	mg.Nm ⁻³	< 2500	< 3000	< 1500
H ₂ S	mg.Nm ⁻³	< 3000	< 4000	< 10000

1.4. Environmental and economic issues

Biogas is a very interesting energy from an ecological and economic point of view.

The environmental interest of reducing CO₂ emissions coincides with the approaches adopted to the production and consumption of renewable energies. The environmental impact of production and valorization of biogas is easily demonstrable.

From an economic point of view, digesters allow to value all the types of substrates discussed in the previous section at attractive cost. In addition, the digest that was recovered at the outlet of the digester could be valorized as fertilizer.

- **Environmental issues**

Anaerobic digestion is a natural phenomenon that rejects methane which is 23 times more potent as a greenhouse gas than carbon dioxide [16]. The simple conversion by combustion of CH₄ into CO₂ reduced to 8 % the initial potential of greenhouse gas from biogas, emitted directly into the atmosphere. Carbon dioxide footprint could be further improved when the biogas is purified and upgraded.

Moreover, the fossil energy used massively today releases large amounts of CO₂ and it is not an inexhaustible resource. The valorization of biogas is therefore interesting for environmental protection by saving fossil fuels and avoiding methane emissions into the atmosphere.

A medium agricultural digester allows the reduction of 1000 tons of CO₂ each year [1].

- **Economic issues**

The economic feasibility of a biogas sector depends mainly on the composition of the biogas.

The methane concentration in a biogas produced from household and industrial wastes is low as seen in Table 4. The CH₄ concentrations do not exceed 55 %. Moreover, the biogas produced can vary over time and it contains large amount of minor compounds, complicating biogas valorization. Biogas production from this type of waste is expected to be limited in future years in Europe. Restrictions and European standards are becoming more stringent, which makes the use of this source, unprofitable [12]. The most advanced biogas applications as vehicle fuel production or injection into the natural gas grid are hardly possible from this source, which requires advanced purification treatments.

Considering an anaerobic digestion unit treating 50000 t/year of household waste, the total costs vary between 50 and 95 €/t while the incomes do not exceed 30 €/t [14].

The economic balance of biogas production within wastewater treatment plants depends on the size and savings made on the sludge. This stable and controlled anaerobic digestion process is financially viable.

The wastewater treatment plant “Aquapol” situated in Grenoble, France treats 88 000 000 m³ of wastewater each year which is equivalent to 8000 dry tons of sludge processed per year. Part of the produced biogas is used for internal energy needs of the plant (8 GWh/year). The other part (14 GWh/year) will be injected in the natural gas grid after purification and compression [17].

The establishment of biogas purification process from agricultural waste is particularly attractive due to the reliability of the resource. The introduction of energy crops among the substrates improves the performance of anaerobic digestion giving a new economic aspect for the production of biogas.

1.5. From biogas to liquid biomethane

An advanced purification of biogas allows achieving an adequate quality threshold for use as vehicle fuel. The major step of this treatment is the separation of CO₂, in addition to dehumidification, desulfurization, reduction of oxygen and the removal of trace compounds in biogas. Qualities and tolerances in impurities, for the production of liquid biomethane to be used as vehicle fuel are listed in Table 5.

Table 5. Tolerances in impurities for the use of liquid biomethane as vehicle fuel [14]

Compounds	Unit	Liquid biomethane used as vehicle fuel
Methane – CH ₄	wt. %	> 96
Carbone dioxide – CO ₂	wt. %	< 3
Oxygen – O ₂	wt. %	< 3
Water – H ₂ O	mg.m ⁻³	< 30
Hydrogen sulfide – H ₂ S	mg.m ⁻³	< 5
Total sulfur	mg.m ⁻³	< 120
Organosulfur	mg.m ⁻³	< 15
Hydrocarbons	mg.m ⁻³	< 200
Critical size of particles	µm	< 1

Liquid biogas can be produced using a cryogenic upgrading technology, based on differences in condensation temperature for different compounds. It can also be produced by mean of a conventional technology connected with a small-scale liquefaction plant. When using the first method,

the carbon dioxide comes as a by-product which could be used in external applications bringing in extra income to the biogas upgrading unit.

The environmental benefits provided by the passage from the diesel to biomethane are impressive. Used as a fuel, Bio-LNG enables a considerable reduction of polluting emissions and represents a genuine alternative to diesel:

- Zero emission of fine particles responsible for 42000 premature deaths annually in France.
- -70 % NO_x emissions.
- -90 % CO₂ emissions.
- -99 % hydrocarbons emissions.
- -50 % noise pollution.

Bio-LNG is a renewable energy produced from waste which makes it neutral in terms of greenhouse gas emissions. Its use allows the decarbonization of the energy mix. Bio-LNG is the only sustainable solution for long-distance haulage operated by heavy goods vehicles.

1.6. Conclusion

Biogas is produced when organic material is decomposed under anaerobic conditions. The main constituents are methane and carbon dioxide. To be able to use the raw biogas as a vehicle fuel it must be purified and upgraded, which means that impurities and CO₂ respectively, are separated. There are a number of available upgrading technologies and the most commonly used are:

- Absorption
- Adsorption
- Membranes
- Cryogenic technology

Other technologies exist such as oxidation and biological treatment. These techniques are known in the field of gas treatment. Today other technologies are being developed such as biocatalysis, photocatalysis and cold plasma. For the moment, the lack of information about them limits their industrial application [18].

The choice of the technology to be used to purify, upgrade and liquefy the biogas requires the knowledge of the thermodynamic properties of biogas and the representation of phase equilibrium. These thermodynamics aspects of biogas will be discussed in the next chapter.

Chapter 2: Thermodynamic aspects of biogas

Résumé :

Le choix de la technologie à utiliser pour purifier et liquéfier le biogaz exige la connaissance des propriétés thermodynamiques du biogaz.

Ce chapitre recense les propriétés thermodynamiques des fluides qui sont essentielles pour concevoir et optimiser les technologies de purification de biogaz. Il montre aussi que les propriétés thermophysiques du biogaz dépendent fortement de sa composition, en particulier des concentrations de méthane et de dioxyde de carbone. La présence de composés à faibles concentrations comme l'azote ou l'hydrogène sulfuré pourrait modifier les propriétés physiques du biogaz. Par exemple, les gaz d'enfouissement comprennent de petites quantités d'azote et d'oxygène qui affectent le comportement de phase du système $\text{CH}_4 - \text{CO}_2$.

2.1. Introduction

Thermodynamics can be used as a powerful tool for setting and evaluating processes used for purification and upgrading of biogas. Therefore, this study will focus on the thermodynamic investigation of biogas. It will present the thermodynamic properties of pure compounds and of the gas mixture (biogas) at a pressure of 1.103 bar. This pressure is considered because the biogas treated in the experimental part (Chapter 4) comes from the wastewater treatment plant at a pressure slightly above atmospheric pressure in order to avoid air infiltration into the biogas pipe.

2.2. Thermodynamic properties of pure component present in biogas

Biogas refers to a mixture of different molecules of gases as seen in Table 4. The thermodynamic aspects of the main components of biogas will be presented in this section.

2.2.1. Hydrogen sulfide

Hydrogen sulfide is a colorless gas with the characteristic foul odor of rotten eggs. It is very poisonous, corrosive, flammable and explosive. Its olfactory threshold varies between 0.7 and 200 g.m⁻³, depending on the sensitivity of each individual. The olfactory sensation is not proportional to the concentration of H₂S in the air because it is possible that the smell felt at very low concentrations is attenuated or disappeared at high concentrations.

Hydrogen sulfide is created following the bacterial decomposition of organic matter in the absence of oxygen, such as in swamps and sewers. It also appears in volcanic gases and hot springs. Other sources of hydrogen sulfide are the industrial processes used in the oil and natural gas sectors, sewage treatment plants and factories producing pulp and paper ...
The thermos-physical properties of hydrogen sulfide are listed in Table 6.

Table 6. Thermo-physical properties of hydrogen sulfide [19]

Properties	Unit	Value
Molar mass	g.mol^{-1}	34.08
Auto-ignition temperature	$^{\circ}\text{C}$	270
Solubility in water (1.013 bar and 0°C)	vol/vol	4.67
Solid phase		
Melting point	$^{\circ}\text{C}$	-85.7
Latent heat of fusion (1.013 bar at melting point)	kJ.kg^{-1}	69.73
Liquid phase		
Boiling point at 1.013 bar	$^{\circ}\text{C}$	-60.3
Vapor pressure at 20°C	bar	17.81
Liquid phase density (1.013 bar at boiling point)	kg.m^{-3}	949.2
Latent heat of vaporization (1.013 bar at boiling point)	kJ.kg^{-1}	546.41
Gas phase		
Gas phase density (1.013 bar and 15°C)	kg.m^{-3}	1.45
Viscosity (1.013 bar and 0°C)	Pa.s	1.13×10^{-5}
Thermal conductivity (1.013 bar and 0°C)	$\text{mW.m}^{-1}.\text{K}^{-1}$	15.61
Specific volume (1.013 bar and 25°C)	$\text{m}^3.\text{kg}^{-1}$	0.7126
Heat capacity at constant pressure (1.013 bar and 25°C)	$\text{kJ.mol}^{-1}.\text{K}^{-1}$	0.0346
Heat capacity at constant volume (1.013 bar and 25°C)	$\text{kJ.mol}^{-1}.\text{K}^{-1}$	0.026
Critical point		
Critical temperature	$^{\circ}\text{C}$	99.95
Critical pressure	bar	90
Critical density	kg.m^{-3}	347.28
Triple point		
Triple point temperature	$^{\circ}\text{C}$	-85.45
Triple point pressure	bar	0.232

2.2.2. Carbon dioxide

Carbon dioxide is a colorless and odorless gas which is naturally present in the Earth's atmosphere. The concentration of carbon dioxide in the atmosphere reached 405 ppm at the end of 2016, against only 283 ppm in 1839.

Carbon dioxide is produced by all aerobic organisms when they metabolize carbohydrate and lipids to produce energy by respiration [20]. It is also produced by burning fossil fuels such as coal, natural gas and oil. Significant amounts of CO_2 are also released by volcanoes. The thermo-physical properties of carbon dioxide are listed in Table 7.

Table 7. Thermo-physical properties of carbon dioxide [19]

Properties	Unit	Value
Molar mass	g.mol^{-1}	44.01
Concentration in air	Vol %	0.0405
Solubility in water (1.013 bar and 0 °C)	vol/vol	1.7163
Solid phase		
Melting point	°C	-56.57
Latent heat of fusion (1.013 bar at melting point)	kJ.kg^{-1}	204.93
Solid phase density	kg.m^{-3}	1562
Liquid phase		
Boiling point	°C	-78.45
Vapor pressure at 20 °C	bar	57.291
Liquid phase density (19.7 bar at -20 °C)	kg.m^{-3}	1256.74
Gas phase		
Gas phase density (1.013 bar and 15 °C)	kg.m^{-3}	1.87
Viscosity (1.013 bar and 0 °C)	Pa.s	1.37×10^{-5}
Thermal conductivity (1.013 bar and 0 °C)	$\text{mW.m}^{-1}.\text{K}^{-1}$	14.67
Specific volume (1.013 bar and 25 °C)	$\text{m}^3.\text{kg}^{-1}$	0.5532
Heat capacity at constant pressure (1.013 bar and 25 °C)	$\text{kJ.mol}^{-1}.\text{K}^{-1}$	0.0374
Heat capacity at constant volume (1.013 bar and 25 °C)	$\text{kJ.mol}^{-1}.\text{K}^{-1}$	0.0289
Critical point		
Critical temperature	°C	30.98
Critical pressure	bar	73.77
Critical density	kg.m^{-3}	467.6
Triple point		
Triple point temperature	°C	-56.56
Triple point pressure	bar	5.187

2.2.3. Methane

Methane is a hydrocarbon which is naturally present in the Earth's atmosphere at very low concentrations (1.82 ppm in 2012) [21].

Huge amounts of methane are buried in the earth's crust in the form of natural gas and on the ocean floor in the form of methane hydrates. Moreover, mud volcanoes, landfills, and animal digestion release methane.

The thermo-physical properties of methane are listed in Table 8.

Table 8. Thermo-physical properties of methane [19]

Properties	Unit	Value
Molar mass	g.mol ⁻¹	16.043
Auto-ignition temperature	°C	595
Solubility in water (1.013 bar and 2 °C)	vol/vol	0.054
Solid phase		
Melting point	°C	-182.46
Latent heat of fusion (1.013 bar at melting point)	kJ.kg ⁻¹	58.682
Liquid phase		
Boiling point at 1.013 bar	°C	-161.48
Liquid phase density (1.013 bar at boiling point)	kg.m ⁻³	422.36
Gas phase		
Gas phase density (1.013 bar at boiling point)	kg.m ⁻³	1.816
Viscosity (1.013 bar and 0 °C)	Pa.s	1.0245 x 10 ⁻⁵
Thermal conductivity (1.013 bar and 0 °C)	mW.m ⁻¹ .K ⁻¹	30.57
Specific volume (1.013 bar and 25 °C)	m ³ .kg ⁻¹	1.5227
Heat capacity at constant pressure (1.013 bar and 25 °C)	kJ.mol ⁻¹ .K ⁻¹	0.0358
Heat capacity at constant volume (1.013 bar and 25 °C)	kJ.mol ⁻¹ .K ⁻¹	0.0274
Critical point		
Critical temperature	°C	-82.59
Critical pressure	bar	45.99
Critical density	kg.m ⁻³	162.7
Triple point		
Triple point temperature	°C	-182.46
Triple point pressure	bar	0.117

The vapor pressure is the basis of all equilibrium calculation. The vapor pressure curves for the three molecules of interest are shown in Fig. 6.

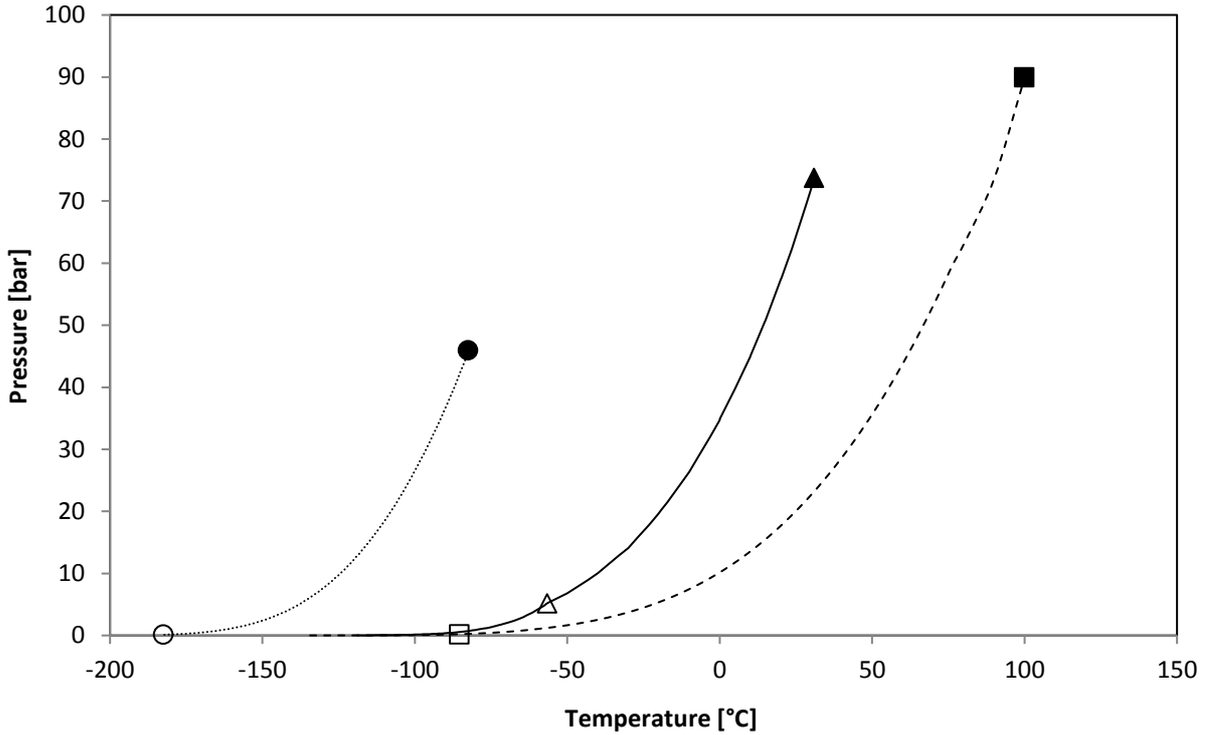


Fig. 6: Vapor pressure of the main components present in biogas

(■) H₂S critical point ; (▲) CO₂ critical point ; (●) CH₄ critical point ; (□) H₂S triple point ; (Δ) CO₂ triple point ; (○) CH₄ triple point ; (- - -) H₂S ; (—) CO₂ ; (.....) CH₄

The vapor pressures are calculated using Antoine Equation (2). The carbon dioxide vapor pressures above -76.36 °C were retrieved from the works of Kidnay [22], Yarym-Agaev [23], Miller [24] and Del Rio [25]. As shown in Fig. 6, the vapor pressure curve of CO₂ continues at temperatures lower than the triple point temperature (-56.56 °C) where it passes from Vapor – Liquid Equilibrium to Vapor – Solid Equilibrium.

$$\log P = A - \left(\frac{B}{T + C} \right) \quad (2)$$

Where:

P [bar]: Vapor pressure.

T [K]: Temperature.

A, B and C [-]: Component specific constants.

The constants used by Antoine Equation (2) are listed in Table 9 for each component.

Table 9. Constants used by Antoine Equation for the calculation of H₂S, CO₂ and CH₄ vapor pressures

Components	A	B	C	Temperature range [K]	References
H ₂ S	4.43681	829.439	-25.412	138.8 – 212.8	[26]
H ₂ S	4.52887	958.587	-0.539	212.8 – 349.5	[26]
CO ₂	6.81228	1301.679	-3.494	154.26 – 195.89	[27]
CH ₄	3.9895	443.028	-0.49	90.99 – 189.99	[28]

2.3. Thermodynamic properties of the gas mixture (biogas)

2.3.1. Phase equilibrium behavior of biogas

In this section, the gas mixture is assumed to consist of methane (1) and carbon dioxide (2) because all the other impurities are present at very low concentrations depending on the type of substrate as shown in Table 4. Further, the upgrading process is generally made after purification that consists in the elimination of all these pollutants such as H_2S , H_2O and siloxanes.

The liquefaction of biomethane requires very low temperatures, therefore a cryogenic technology for biogas upgrading could be envisaged. This technology requires correct design of heat exchangers to achieve maximal purity of biomethane, that's why the knowledge of the pressure – temperature behavior of the binary mixture methane – carbon dioxide is essential [29].

Fig. 7 presents the phase equilibrium behavior for the methane – carbon dioxide system.

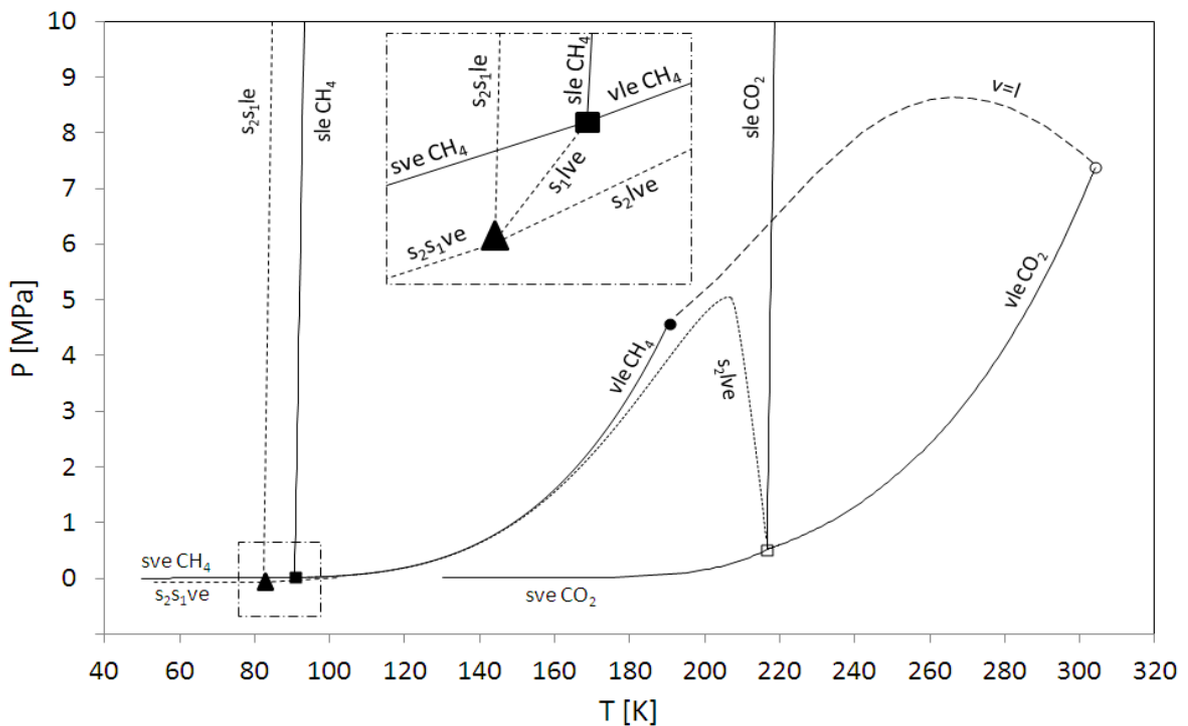


Fig. 7: Pressure – Temperature equilibrium behavior for the $CH_4 - CO_2$ system [29]

(■) CH_4 triple point ; (●) CH_4 critical point ; (□) CO_2 triple point ; (○) CO_2 critical point ; (▲) mixture quadruple point ; (---) vapor-liquid critical locus ; (-.-) three-phase loci ; (—) pure compound phase equilibria

In the context of CO_2 separation by solidification of CO_2 , the authors [29] have focused on the solid phase because thanks to the high triple point of carbon dioxide as seen in Table 7, CO_2 could be separated from methane directly from its gaseous phase into a solid phase for pressures close to the atmospheric pressure.

In order to understand the phase equilibrium behavior of the system CH_4 (1) – H_2S (2), Langè et al. [30] have studied this aspect for temperatures from 70 K up to the critical temperature of H_2S and pressures up to 250 MPa as seen in Fig. 8.

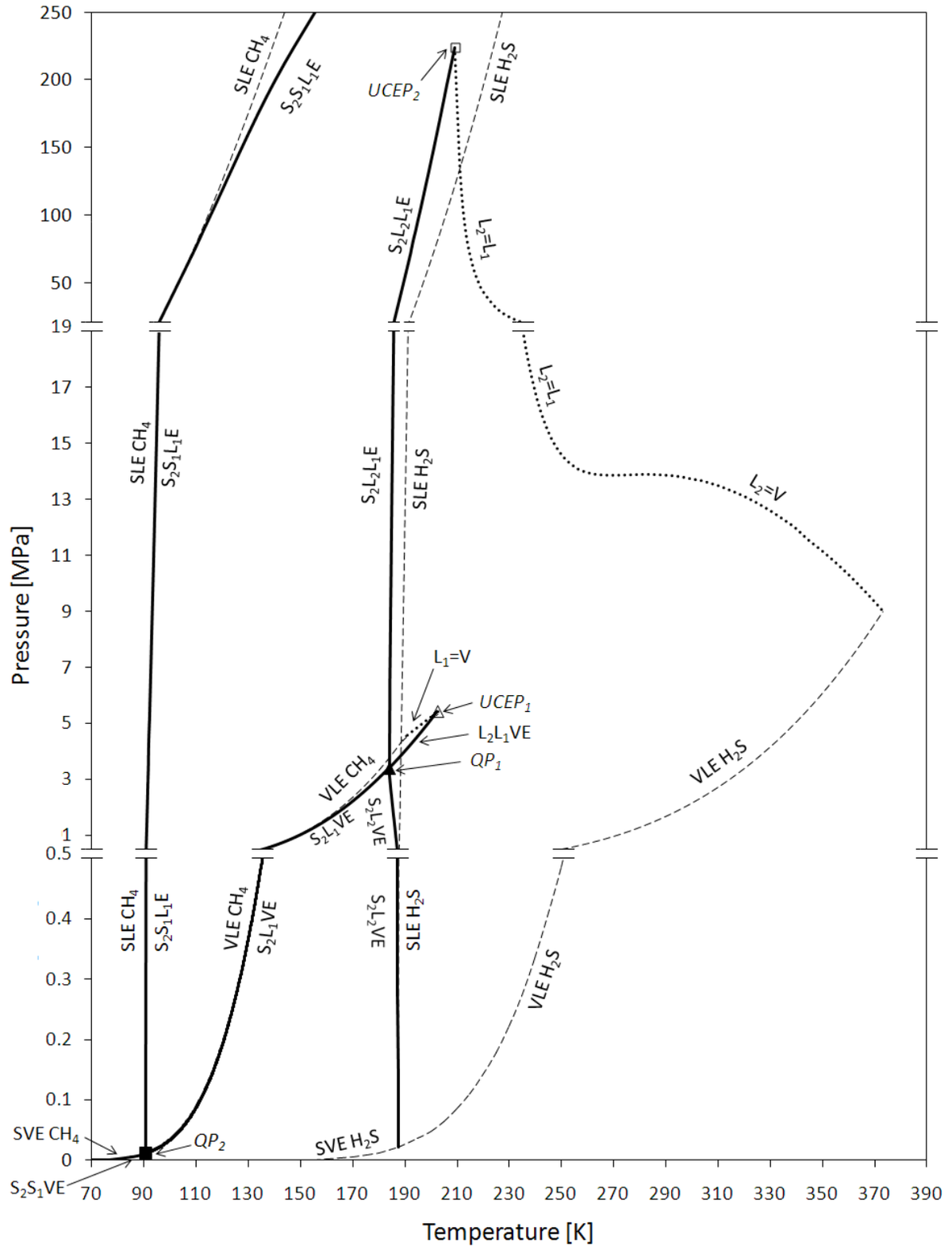


Fig. 8: Pressure – Temperature equilibrium behavior for the CH₄ – H₂S system [30]

(-) three-phase equilibrium boundaries ; (...) critical curves ; (- -) SVE, VLE and SLE of CH₄ and H₂S ; (■) quadruple point QP₂ ; (▲) quadruple point QP₁ ; (Δ) Upper Critical EndPoint UCEP₁ ; (□) Upper Critical EndPoint UCEP₂

2.3.2. Density and dynamic viscosity of biogas

All the thermophysical properties discussed in this section are calculated using the software REFPROP V9.0 which calculates the thermodynamic and transport properties of industrially important fluids and their mixtures. The equation of state for calculating these properties is the GERG (European Gas Research Group) – 2008 equation [31].

Depending on its composition, biogas has characteristics that it is interesting to investigate such as density and viscosity.

At a pressure of 1,103 bar, slightly higher than the atmospheric pressure, the variation of density as a function of temperature is shown in Fig. 9.

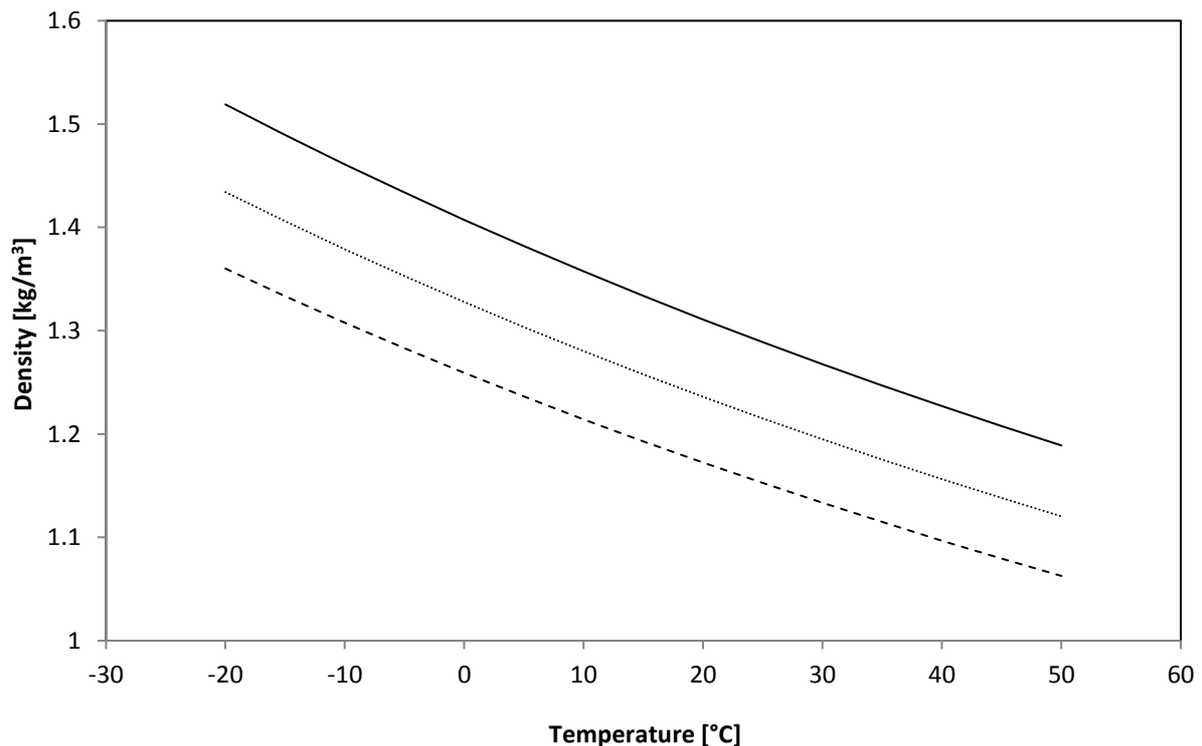


Fig. 9: Biogas density as a function of temperature

(—) Air ; (.....) Biogas with 40 mol% of CO₂ ; (- - -) Biogas with 35 mol% of CO₂

As seen in Fig. 9, biogas is lighter than air. Moreover, its density depends on the carbon dioxide content. The density of a biogas rich on carbon dioxide will be greater than that of a biogas containing an inferior CO₂ concentration. This is due to the molecular weight of CO₂ which is greater than that of methane as seen in Tables 7 and 8.

At a pressure of 1,103 bar, the evolution of the viscosity of the biogas as a function of temperature is depicted in Fig. 10.

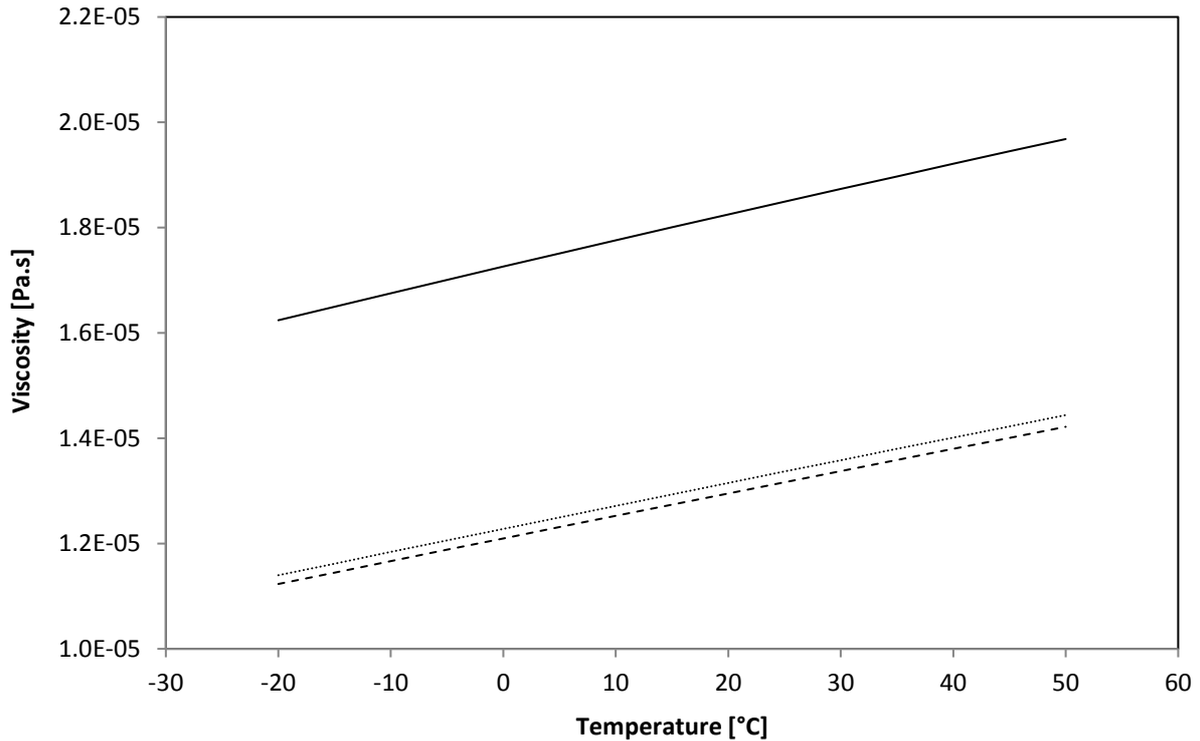


Fig. 10: Biogas viscosity as a function of temperature

(—) Air ; (.....) Biogas with 40 mol% of CO₂ ; (- - -) Biogas with 35 mol% of CO₂

As density, the biogas viscosity increases with the CO₂ concentration. This is due to the higher viscosity of carbon dioxide compared to methane, at equal pressure and temperature as shown in Tables 7 and 8.

2.3.3. Thermal conductivity of biogas

The thermal conductivity is a physical property which describes the ability of gases to conduct heat. The phenomenon of heat conduction in gases is explained by the kinetic gas theory which treats the collisions between the molecules.

The thermal conductivity depends on thermal capacity at constant volume and viscosity [32].

At a pressure of 1,103 bar, the variation of thermal conductivity of biogas as a function of temperature is shown in Fig. 11.

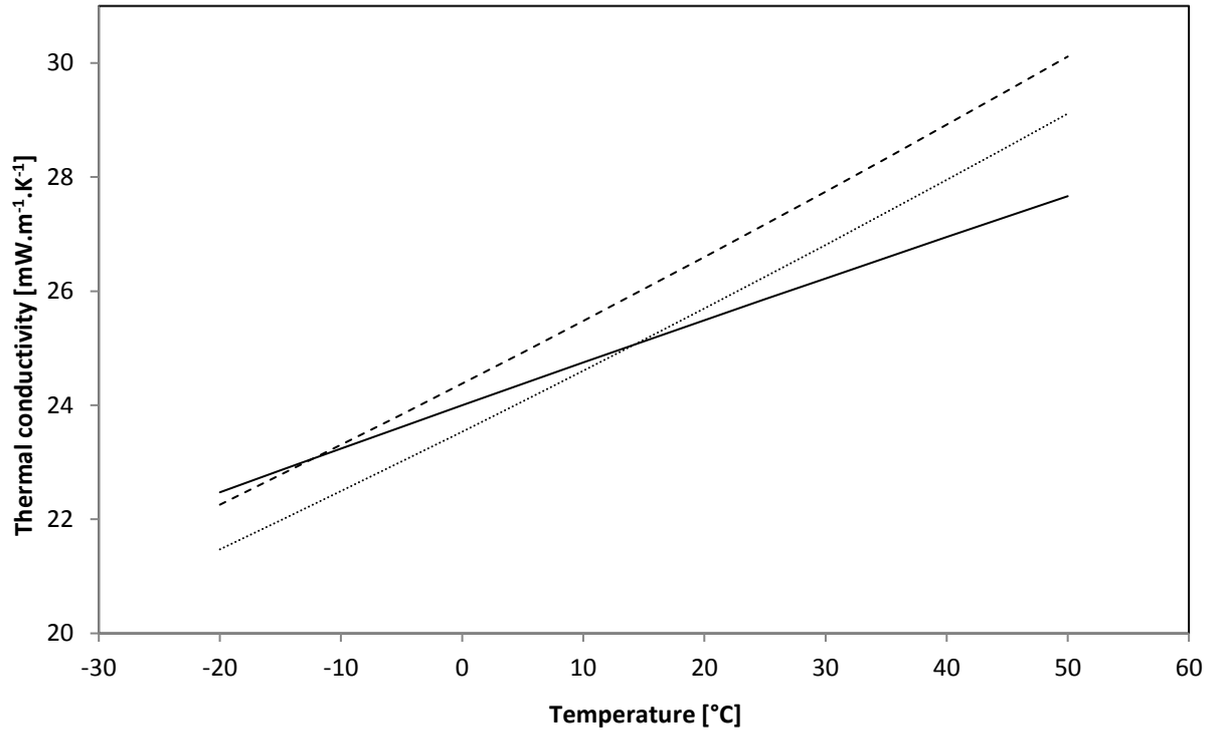


Fig. 11: Thermal conductivity of biogas as a function of temperature

(—) Air ; (.....) Biogas with 40 mol% of CO₂ ; (- - -) Biogas with 35 mol% of CO₂

As shown in Fig. 11, the thermal conductivity increases with temperature. This is due to collisions between the molecules of biogas that increase with temperature, resulting a thermal energy transfer increase.

2.3.4. Thermal capacities

The constant pressure heat capacity and the constant volume heat capacity are respectively related by the following thermodynamic Equations (3) and (4).

$$C_P = \left(\frac{\partial H}{\partial T} \right)_P \quad (3)$$

$$C_V = \left(\frac{\partial U}{\partial T} \right)_V \quad (4)$$

At a pressure of 1,103 bar, the variations of heat capacities of biogas as a function of temperature are shown in Fig. 12 and 13.

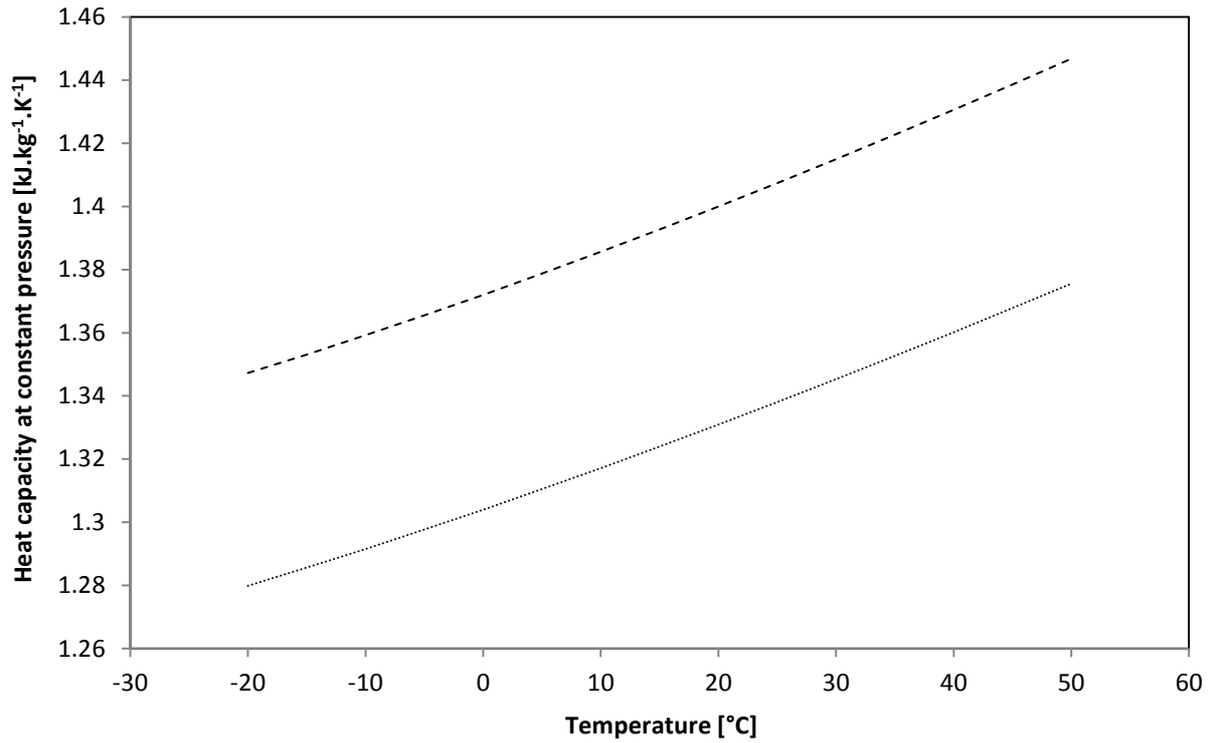


Fig. 12: Heat capacity at constant pressure of biogas as a function of temperature

(.....) Biogas with 60 mol% of CH₄; (- - -) Biogas with 65 mol% of CH₄

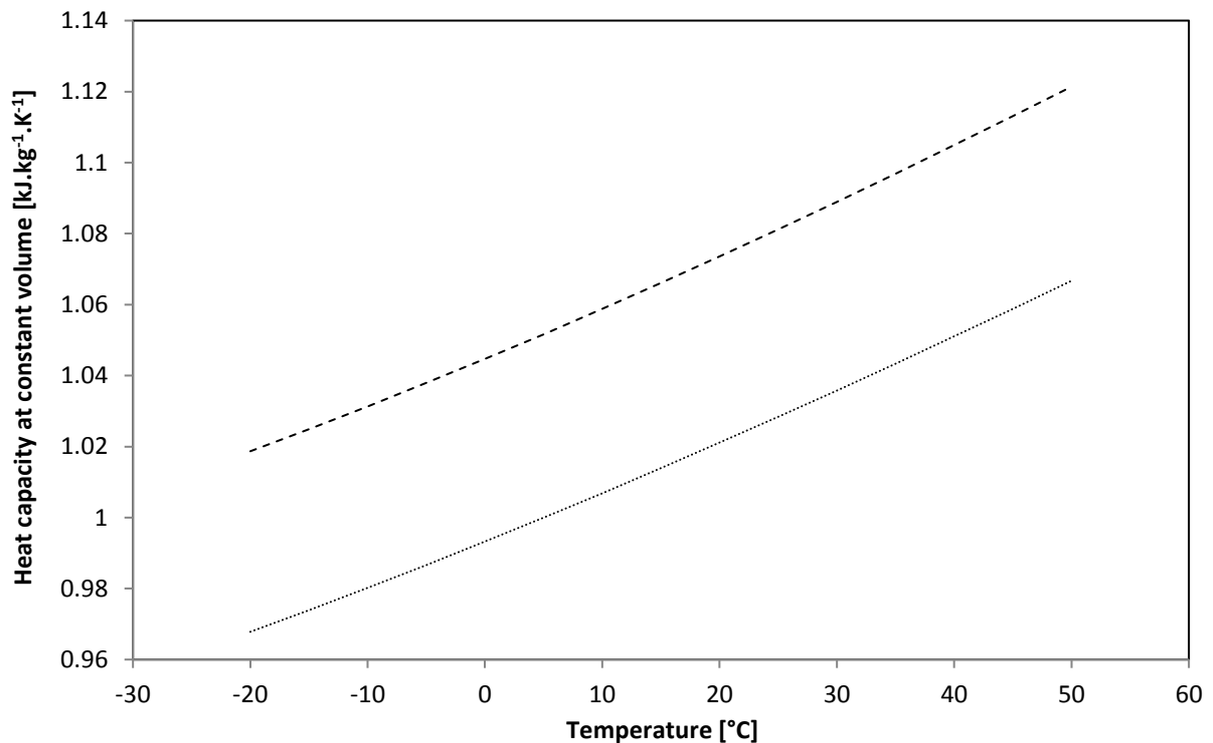


Fig. 13: Heat capacity at constant volume of biogas as a function of temperature

(.....) Biogas with 60 mol% of CH₄; (- - -) Biogas with 65 mol% of CH₄

Fig. 12 and 13 show that biogas specific heat capacities strongly depend on the methane concentration unlike the density, which is inversely proportional to the concentration of methane in the biogas.

According to Vardanjan et al. [33], an increase of methane concentration from 50 % to 75 % will result in the increase of heat capacity by 17 %. In Fig. 12 and 13, heat capacities increased by 5 % after the passage of the methane concentration from 60 to 65 %.

2.4. Energy content in biogas

An important property of biogas is the lower heating value (LHV) which measures its energy value. The LHV of biogas is proportional to its methane content. For example, a biogas containing 70 mol% of methane at 15 °C and at atmospheric pressure has a LHV equal to 6.6 kWh/m³ [34].

The energy value of biogas can also be evaluated by the Wobbe Index. Compared to the calorific value of biogas which has been treated by a lot of authors, only few studies have been conducted on the Wobbe index of biogas.

According to the Danish Technological Institute (DTI) [35], a biogas containing 60 mol% of methane, 38 mol% of carbon dioxide and 2 mol% of others has a Wobbe index of 19.5 MJ/m³.

2.5. Conclusion

This chapter has allowed identifying the thermophysical properties of biogas. These properties are essential to design and optimize technologies for biogas purification and upgrading.

The thermophysical properties of biogas discussed in this chapter strongly depend on the composition of biogas, especially the concentrations of methane and carbon dioxide.

The presence of minor compounds such as H₂O, N₂, O₂ and H₂S could change the physical properties of biogas. For example, landfill gas includes small amounts of nitrogen and oxygen which affect the phase behavior of the CH₄ – CO₂ system.

The technologies used for purification and upgrading of biogas will be discussed in the following section and a particular attention will be paid to the elimination of hydrogen sulfide.

Chapter 3: From molecules to the process

Résumé :

La valorisation du biogaz nécessite la mise en œuvre d'un procédé de purification qui consiste à éliminer du biogaz brut toutes les substances indésirables afin d'augmenter au maximum sa teneur en méthane.

Ce chapitre présente une étude comparative des méthodes connues de séparation de H₂S du biogaz. La méthode de séparation devra permettre de réduire la teneur résiduelle en H₂S à moins de 1 ppm.

La première technique de séparation étudiée est l'absorption dans des contacteurs gaz – liquide. L'accent a été porté sur les principes fondamentaux basés sur l'absorption avec des solvants de type physique et chimique ainsi que leur efficacité dans l'élimination de l'H₂S.

La deuxième technique de traitement de l'H₂S étudiée est l'adsorption sur des solides microporeux comme le charbon actif et les zéolithes.

Le troisième procédé discuté est la séparation membranaire dont la performance dépend de deux paramètres qui sont : la perméabilité et la sélectivité membranaires. En général, les membranes de perméabilité élevée présentent une faible sélectivité et inversement. Cette méthode présente quelques inconvénients comme le risque de rupture dû au gradient de pression qui constitue la force motrice, l'exposition à certains solvants qui peut endommager ou boucher la membrane, le prix élevé des membranes ou encore les pertes en méthane qui peuvent être considérables.

La dernière technologie séparative étudiée lors de ce chapitre est la condensation cryogénique basée sur la thermodynamique des équilibres de phases.

Cette synthèse bibliographique a permis de comparer les différents procédés selon plusieurs critères comme l'efficacité de séparation, l'impact environnemental et les coûts d'investissement et d'exploitation.

Finalement, le choix s'est porté sur les deux technologies suivantes utilisées pour l'élimination du sulfure d'hydrogène:

- Absorption chimique dans une colonne à garnissage structuré utilisant l'hydroxyde de sodium (NaOH) comme solvant.
- Adsorption dans un lit fixe de charbon actif.

3.1. Introduction

Biogas purification requires the removal of minor compounds despite their low amount relative to methane. Among these compounds, the removal of water vapor and ammonia are sufficiently documented in the literature and do not constitute a technical difficulty. However, the treatment of hydrogen sulfide present at low concentration is a challenge. This gas is formed when organic material containing sulfur is decomposed under anaerobic conditions. It is very corrosive on most metals which negatively affects the operation and viability of equipment especially pumps, heat exchangers and pipes. In the following section, conventional purification and upgrading technologies will be described.

3.2. Absorption technology

In chemistry, absorption is an operation by which a substance combined in one state is transferred into another substance of a different state. The most frequent use of absorption is the separation of a gas mixture by the absorption of part of the mixture in a solvent. The two phases are brought into contact in an absorption column and are allowed to exchange mass and energy across their common interface, where the flux of H₂S transferred is calculated using Equation (5) by applying the two-film theory presented in Fig. 14. This theory assumes that the mass transfer resistance is located on the boundary layer on the gas side and the liquid side respectively.

The mass transfer between the liquid and the vapor phases heavily depends on the effective interfacial area.

$$N_{H_2S, z} = \frac{k_G}{RT} (p_{H_2S} - p_{H_2S}^*) = k_L (C_{H_2S}^* - C_{H_2S}) \quad (5)$$

Where the gas-phase and liquid-phase mass transfer coefficients are calculated using Equations (6) and (7) respectively.

$$k_G = \frac{D_{H_2S,G}}{\delta_G} \quad (6)$$

$$k_L = \frac{D_{H_2S,L}}{\delta_L} \quad (7)$$

Where:

$N_{H_2S, z}$ [mol.s⁻¹.m⁻²]: flux of H₂S transferred by unit area.

p_{H_2S} [Pa]: partial pressure of H₂S in the gas phase.

$p_{H_2S}^*$ [Pa]: partial pressure of H₂S at the interface.

C_{H_2S} [mol.m⁻³]: concentration of H₂S in the liquid phase.

$C_{H_2S}^*$ [mol.m⁻³]: concentration of H₂S at the interface.

δ_G and δ_L [m]: thickness of the stagnant film on the gas side and the liquid side respectively.

$D_{H_2S,G}$ and $D_{H_2S,L}$ [m².s⁻¹]: H₂S diffusion coefficients in gas phase and liquid phase respectively.

This theory assumes also that the thermodynamic equilibrium is reached in the interface. Knowing that the H₂S concentration is low, therefore the Henry's law presented in Equation (8) is applicable.

$$p_{H_2S}^* = H_{H_2S} x_{H_2S} \quad (8)$$

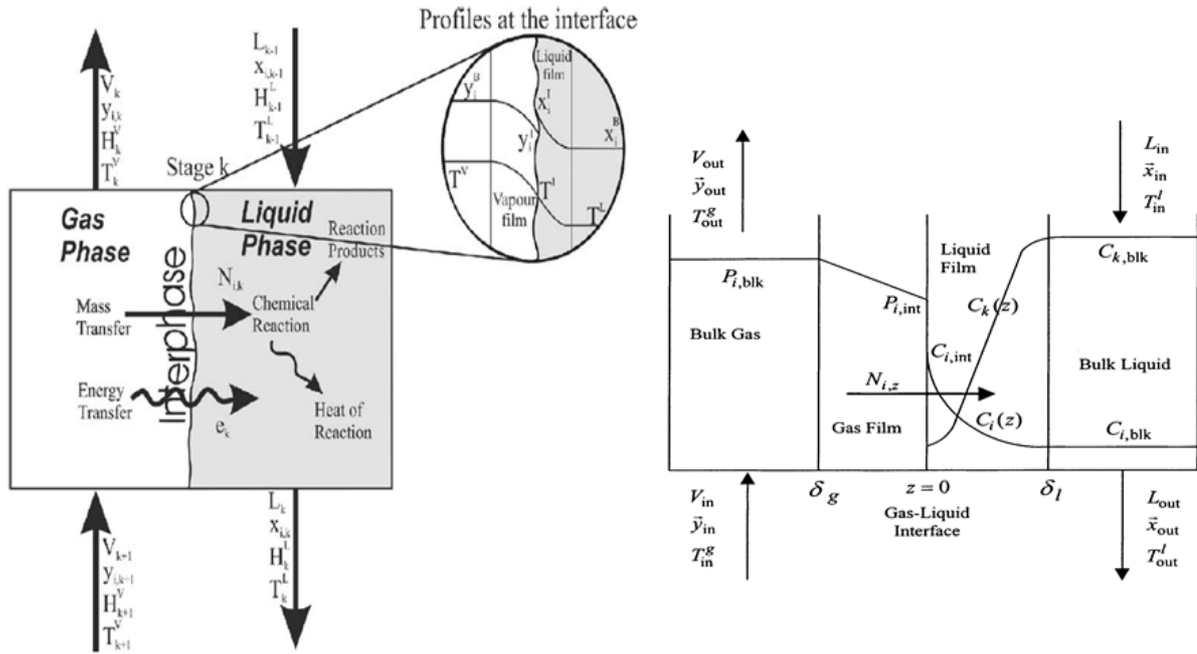


Fig. 14: Two-film theory [36]

There are two types of absorption processes: physical absorption and chemical absorption, depending on whether there is any chemical reaction between the pollutant and the absorbent.

For example, when water absorbs oxygen from the air, a mass of the gas moves into the liquid, and no significant chemical reactions occur between the solvent and the solute. In this case, the process is commonly referred to as physical absorption.

Chemical absorption occurs, when a chemical reaction is carried out in the liquid phase, to dissolve the compound to be removed, and thus enhance the efficiency of the process. An example of chemical absorption is the process for absorbing CO₂ and/or H₂S with aqueous solution of sodium hydroxide. Chemical solvents are favored over physical solvents when the concentration of pollutants at low partial pressure has to be reduced to a very low level [37]. However, if the impurity is available in the feed gas at high partial pressure, physical solvents might be preferred to chemical solvents [38]

In the presence of a chemical reaction, the rate of absorption increases. The flux of H₂S transferred depends on an acceleration factor E, and is expressed in Equation (9).

$$N_{H_2S} = E k_L C_{H_2S}^* \quad (9)$$

The acceleration factor E is equal to the ratio between the flux of H₂S transferred in the presence of a chemical reaction and the flux transferred in the absence of a chemical reaction. This factor characterizes the importance of the chemical reaction on the transfer compared to the diffusion process.

The addition of a chemical solvent such as sodium hydroxide for the removal of hydrogen sulfide allows a significant increase of the acceleration factor which becomes controlled by the reaction rate. It will also cause an increase in temperature because of the exothermicity of the reaction. For low concentrations, the increase in temperature will be small.

3.2.1. Physical solvents

The first solvent studied and used in absorption process is water. Hydrophilic compounds present in the biogas such as CO₂ and H₂S are absorbed better in water than the hydrophobic and non-polar compounds as methane.

Table 10 shows the solubility of some compounds of biogas in water.

Table 10. Solubility of the main compounds of biogas in water [39]

Compound	Solubility in water at 1 bar of partial pressure	
	[mg.l ⁻¹] at 0 °C	[mg.l ⁻¹] at 25 °C
NH ₃	53 000	28 000
H ₂ S	205	102
CO ₂	75	34
CH ₄	2.45	1.32
Siloxanes	< 1	< 1

In order to improve the effectiveness of absorption processes, other solvents were tested for purification and upgrading of biogas. They allow the reduction of the size of columns, the energy used for the treatment and the volumes of solvents involved.

Selexol® is a physical absorption process developed by Allied Chemical Corporation, then improved by Norton. Today, it is owned by Universal Oil Products (UOP). It is made up of Dimethyl Ethers of Polyethylene Glycol (DMEPG) whose molecular weight is about 272 g.mol⁻¹.

Rectisol® is one of the older physical absorption processes. It was developed by Linde and Lurgi in order to separate the acid gases present in the syngas from gasification of coal. It has also been used for separation of CO₂ in the syngas of units producing hydrogen and ammonia. This process uses a methanol-based solvent (MeOH) whose chemical formula is CH₃OH.

These solvents exhibit a high selectivity for H₂S removal compared to the other compounds, in particular CO₂.

Other processes exist such as Purisol® based on the use of N-Methyl-2-Pyrrolidone (NMP), and Morphysorb® destined for the separation of acid gases at high concentrations, using N-Formyl-Morpholine (NFM).

The main physical solvents are summarized in Table 11 where selected physical properties are compared.

Table 11. Properties of physical solvents [40]

	Selexol®	Purisol®	Rectisol®	Morphysorb®	Fluor solvent®
Solvent	Dimethyl Ethers of Polyethylene Glycol	N-Methyl-2-Pyrrolidone	Methanol	N-Formyl-Morpholine	Propylene Carbonate
Chemical formula	$(\text{CH}_3\text{O}(\text{CH}_2\text{CH}_2\text{O})_x\text{CH}_3)$ $3 \leq x \leq 9$	$\text{C}_5\text{H}_9\text{NO}$	CH_3OH	$\text{C}_5\text{H}_9\text{NO}_2$	$\text{C}_4\text{H}_6\text{O}_3$
Maximum temperature [°C]	175				65
Vapor pressure at 25 °C [kPa]	0.097	32	1.3 (at 20 °C)	9	3
Viscosity at 25 °C [Pa.s]	5.8	1.65	0.6	9.5	3
Boiling point [°C]	240	202	110	242	240
Melting point [°C]	-23 / -29	-24	-98	21	-48
Molar mass [g.mol⁻¹]	280	99	32	115.3	102

The solubility of a compound in a physical solvent is often expressed as the volume of gas absorbed by the solvent volume. The absorption capacity of H₂S in physical solvents is frequently higher than that of CO₂. As shown in Table 12, the solubility of H₂S in NMP is very important. Moreover, this physical solvent has a high selectivity for H₂S.

Table 12 also shows a very high solubility of water in some physical solvents. This can be harmful to these physical absorbents because the presence of water in the gas phase will cause its accumulation in the solvent and therefore the reduction of the solvent absorption capacity.

Table 12. Solubility of gases in physical solvents at 25 °C and 0.1 MPa [41]

vol.vol ⁻¹	DMEPG	PC	NMP	MeOH (at -25° C)
H ₂	0.013	0.0078	0.0064	0.0054
N ₂	0.02	0.0084		0.012
O ₂		0.026	0.035	0.02
CO	0.028	0.021	0.021	0.02
CH ₄	0.066	0.038	0.072	0.051
CO ₂	1	1	1	1
NH ₃	4.8			23.2
H ₂ S	8.82	3.2	10.2	7.06
SO ₂	92.1	68.6		
H ₂ O	730	300	4000	

Operating at ambient temperature or higher should be avoided during use of some physical solvents in absorption operations because it leads to solvent losses by volatility. A chiller is necessary because the solubility of acid gases in physical solvents is favored by low temperatures.

The two most volatile physical solvents are methanol and NMP. They use absorption columns with respective temperatures of -30 °C and -5 °C to prevent evaporation of the product and improve the solubilization of the acid gases.

However, the DMEPG shows minimal evaporation losses at ambient temperature due to its very low vapor pressure. Furthermore, it becomes viscous at low temperatures. The physical solvents such as DMEPG and NFM must be used at ambient or higher temperatures.

3.2.2. Chemical solvents

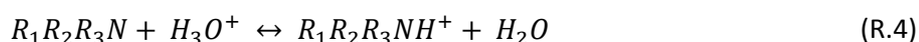
Alkanolamines are most commonly used in acid gas absorption processes. Their molecular structure contains at least both amino (-N) and hydroxyl (-OH) functional groups. The hydroxyl functional group increases the solubility of acid gases in water and reduces the solvent vapor pressure. The amino functional group provides the necessary alkalinity in aqueous solution to ensure the absorption of acid gases.

At equilibrium, in aqueous solution, the reactions between the alkanolamines (R₁R₂R₃N) and acid gases, particularly CO₂ and H₂S are described by the following chemical equilibria [42]:

– Self ionization of water:



– Protonation of alkanolamine:



– Hydrolysis of hydrogen sulfide:



– Bisulfide ion dissociation:



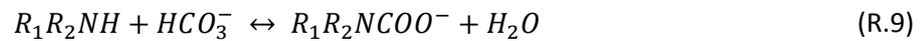
– Carbon dioxide hydrolysis:



– Bicarbonate ion dissociation:



– Carbamate hydrolysis for primary and secondary amines:



The most used amines in industry are:

- Primary amines: monoethanolamine (MEA) and diglycolamine (DGA).
- Secondary amines: diethanolamine (DEA) and diisopropanolamine (DIPA).
- Tertiary amines: méthyldiethanolamine (MDEA) and triethanolamine (TEA).

The triethanolamine was the first used in gas processing industry.

Table 13 lists the physical properties of some of these chemical solvents.

Table 13. Physical properties of some chemical solvents [43]

	MEA	DGA	DEA	DIPA	TEA
Chemical formula	$HOC_2H_4NH_2$	$H(OC_2H_4)_2NH_2$	$(HOC_2H_4)_2NH$	$(HOC_3H_6)_2NH$	$(HOC_2H_4)_3N$
Molecular weight	61.08	105.14	105.14	133.19	148.19
Boiling point [°C]	170.50	221.11	269.00	248.72	360.00
Melting point [°C]	10.5	-12.5	28.0	42.0	22.4
Viscosity [cP]	24 (20 °C)	40 (15.6 °C)	350 (20 °C)	870 (30 °C)	1013 (20 °C)

Fig. 15 shows the chemical structure of some alkanolamines. The primary, secondary or tertiary amines are distinguished according to the degree of substitution of the nitrogen atom.

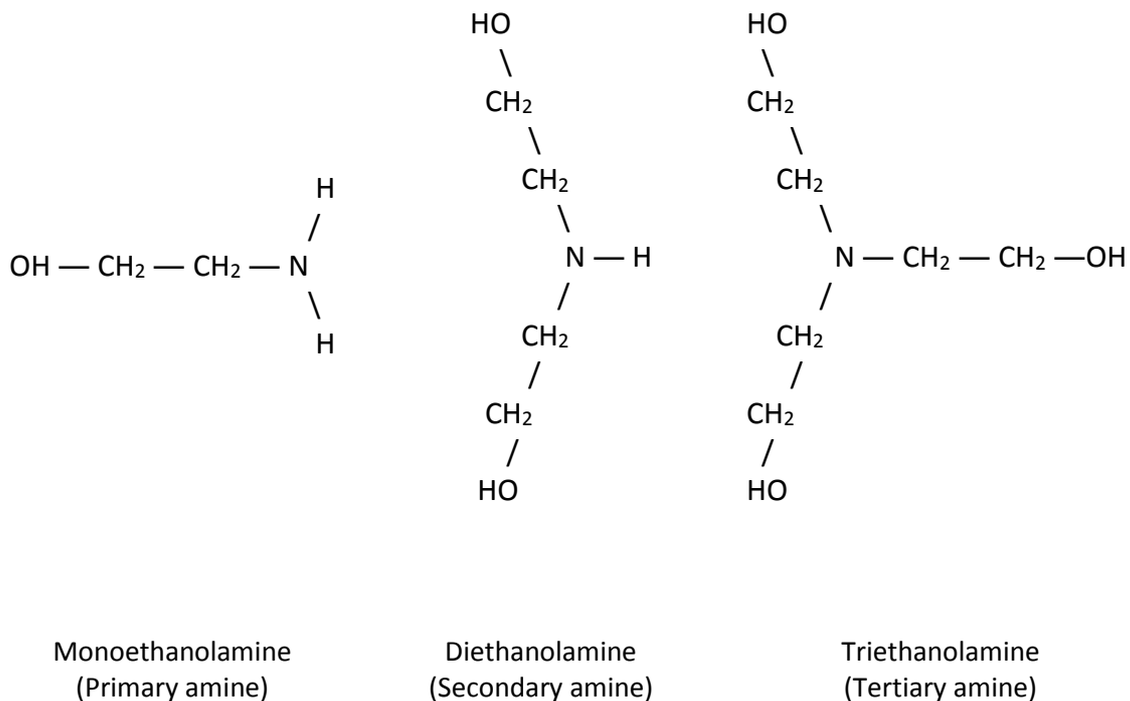


Fig. 15: Chemical structure of some alkanolamines [43]

Primary amines such as monoethanolamine (MEA) are very reactive with H_2S and CO_2 , but they react also with other impurities present in the biogas, which leads to a significant energy requirement for regeneration. They are also susceptible to degradation and corrosion.

Secondary and tertiary amines require less energy for regeneration and are less susceptible to degradation. However, they are less reactive than primary amines and are therefore used for less demanding objectives in terms of purity.

Bottoms [44] was the first in 1930 to study the amines in the gas processing industry. He filed the first patent for the process of absorption of acid gases by ethanolamines.

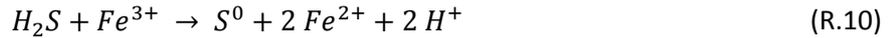
In 1997, Pani et al. [45] experimentally studied the absorption of hydrogen sulphide by a MDEA solution in a temperature range varying between 296 and 343 K. They developed a device to determine the kinetics of absorption of acid gases by alkanolamine solutions, The H_2S concentrations used vary between 0 and 0.44 moles of gas per mole of amine. A mass transfer model incorporating a reversible reaction was used to test the experimental flow absorbed and to determine the diffusion coefficient of the MDEA.

In 1984, Blauwhoff et al. [46] investigated the selective absorption of hydrogen sulphide and have shown that it significantly reduces the cost of gas treatment, by reducing the CO_2 flux transferred.

It is important to note that the solvents presented in this section are more suitable for the elimination of high H_2S concentrations present in natural gas.

Other chemical solvents more suitable for the treatment of biogas are available for the absorption of hydrogen sulfide:

- Aqueous solution of potassium carbonate (K_2CO_3), to which are added additives such as amines. This type of chemical solvent is used in many gas treatment processes such as Flexsorb HP and Catacarb processes, developed respectively by ExxonMobil® and Eickmeyer®.
- Aqueous solution of sodium hydroxide (NaOH), also known as caustic soda.
- Iron chelate solution Fe (III) according to the following reaction (R.10).



According to Neumann and Lynn [47], absorption of hydrogen sulfide by iron chelate solutions are advantageous for achieving high reaction rates (99.99 % H₂S removal).

3.2.3. Hybrid solvents

There are processes that combine both physical and chemical absorption, implementing aqueous mixtures that contain water, amine and an organic solvent. This type of processes has been developed to treat gases containing significant fractions of acid gases.

An example of such processes is the Hi-Pure process, where the gas to treat is put into contact, firstly with an aqueous solution of potassium carbonate, then with an aqueous solution of amine.

Patented by Shell, Sulfinol process uses a hybrid solvent containing a physical solvent called sulfolane, water and a chemical solvent. This latter determines the name of the mixture. It is called Sulfinol-D when DIPA is used as chemical solvent and Sulfinol-M when MDEA is used.

Sulfinol-M is used for the selective removal of H₂S in the presence of CO₂.

Amisol is a process also used for the selective removal of H₂S in the presence of CO₂. It was developed by Bratzler and Doerges in 1974. The mixture used is composed of methanol as a physical solvent, and DEA or MEA as a chemical solvent.

3.2.4. Gas-liquid contactors

Application of the principle of absorption is based on contacting the gas and liquid phases in a gas-liquid contactor. This latter, also called absorber, aims to achieve better mass exchange between the two phases in contact.

The efficiency of a gas-liquid contactor is dependent on phenomena involved in the absorption process:

- Transfer laws in the vicinities of interfaces, in particular the transfer coefficients and the interfacial area.
- Transport laws, in particular diffusion coefficients.
- The thermodynamic equilibrium at the interface, especially the solubility of acid gases in the solvent.
- The chemical reaction kinetics: the reaction schemes, the kinetic constants and orders of reactions.

Film thickness, residence time and flow regime also all have a very important impact on the efficiency of the contactor.

The most common concept to evaluate the separation efficiency of packed columns is expressed in terms of Height Equivalent to a Theoretical Plate (HETP).

There is a large number of gas-liquid contactors in the industry, for mass and heat transfer between the two phases, as seen in Fig. 16. Generally, the gas and the liquid flow counter-currently in order to obtain significant concentration gradients and better absorption rate.

The gas-liquid contactors can be classified according to the dispersion mode of phases. Despite a few exceptions, the liquid phase is naturally the dispersed phase in gas treatment application. The choice of the absorber is mainly related to the physicochemical properties of the gas to be treated and to the chemical reactions involved, as well as gas and liquid flow rates implemented.

Table 14 ranks the gas-liquid contactors according to the continuous phase, the fluid inclusion type and the main associated applications (See Fig. 16).

Table 14. Classification of main gas-liquid contactors [48]

Gas-liquid contactors	Continuous phase	Fluid inclusion type	Main applications
Bubble column	Liquid	Bubbles	Oxidation / Chlorination
Gas-liquid agitated vessel	Liquid	Bubbles	Oxidation / Fermentation
Spray tower	Gas	Drops and liquid films	Gas scrubbing dust-laden
Packed column	Gas	Drops and liquid films	Gas scrubbing
Venturi tube	Gas	Drops and liquid films	Gas scrubbing dust-laden
Plate column	Liquid	the flow is stratified, with entrained bubbles in the liquid and a mist or spray of liquid droplets in the vapour	Manufacture of nitric acid

To promote the mass transfer, the absorbers are usually equipped with internal devices to generate the largest interfacial area in order to achieve better mass exchange between the two phases in contact.

In prior years, plate columns were heavily favored over packed columns. But, nowadays, packing columns are the most used in gas absorption applications. Only few specific applications with special design requirement can lead to different choices as in the case of very large flow rates or very soluble compounds where it is preferable to use plate or spray columns.

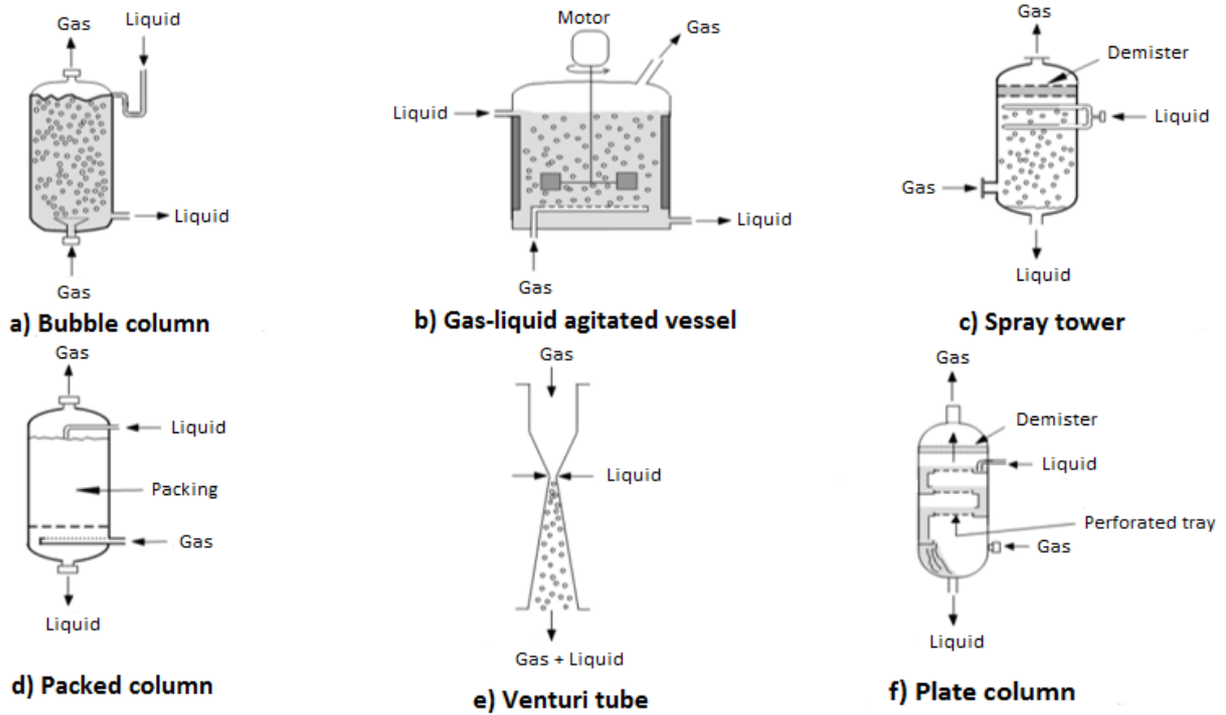


Fig. 16: Main gas-liquid contactors [48]

In a packed column, the gas and liquid normally flow counter currently as seen in Fig. 17. The liquid is sprayed from the top of the column to flow by gravity on the packing forming a large-area liquid film. The liquid enters in contact with the gas injected from the bottom of the column. Liquid flow must be sufficient to ensure uniform wetting of the packing and must not exceed a certain threshold in order to avoid flooding of the column.

The selection of the packing type and material is a very important issue in packed column design. The material should respect certain requirements as weight, pressure drop and especially corrosion resistance.

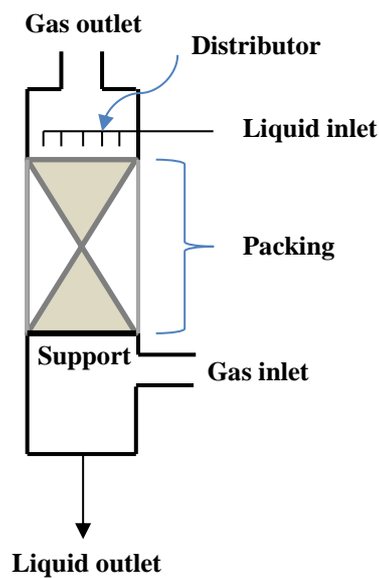


Fig. 17: Schematic representation of a packed column

It exists two types of packings: Those consisting of packing elements placed in a random disposition and those containing corrugated sheets arranged in an orderly manner. The first one is called random packing and the latter is called structured packing.

Fig. 18 shows the two types of packing produced by Sulzer®.

Today, structured packings are much more used than random packings. structured packings ensure a better transfer with a minimal pressure drop.



Fig. 18.a: Random packing Nutter Ring [Sulzer]



Fig. 18.b : Structured packing Mellapak [Sulzer]

3.3. Adsorption technology

Adsorption is a growing process, that is increasingly used in biogas purification. It removes water vapor, odors and other impurities as hydrogen sulfide, from biogas streams. Adsorption is a surface phenomenon that occurs between a vapor or liquid phase and a solid.

Molecules, ions or atoms forming a solid surface, are subjected to asymmetric forces that result in an attractive force field. This attractive force has a limited range, but enough to attract gas or liquid molecules located in the immediate vicinity of the interface. These are forces that cause fixing of molecules on the surface.

Adsorption is classified according to the nature of the interactions that allow the fixing of the adsorbate on the surface of the adsorbent.

There are two types of adsorption, according to the nature of the interactions: physical and chemical adsorption.

Physical adsorption is a reversible phenomenon characterized by weak interaction forces as Van der Waals' forces, while chemical adsorption is usually irreversible involving strong binding energies.

In contrast to physical adsorption, chemisorption grows at high temperatures and causes the formation of a chemical compound on the surface of the adsorbent.

Table 15 lists the criteria to differentiate between physical and chemical adsorption.

Table 15. Criteria to differentiate between physical and chemical adsorption [49]

Criteria	Physical adsorption	Chemical adsorption
Process temperature	Low	High
Type of bond	Van der Waals	Chemical
Interaction forces	30 to 40 kJ.mol ⁻¹	80 to 800 kJ.mol ⁻¹
Layers	Monolayers or multilayers	Monolayers only
Kinetic	Fast and reversible	Slow and irreversible
Desorption	Easy	Difficult

3.3.1. Mechanism of adsorption

During adsorption process, the fluid molecules bind to the surface of a solid following three steps, describing the mass transfer from the fluid phase to the solid surface.

During the first step called external diffusion, the molecules of the fluid phase migrate to the vicinity of the outer surface of the solid particles. To model the transfer of the fluid phase towards the outer surface of the solid phase, the Equation (10) is often used:

$$-\frac{dC_t}{dt} = k_f \left(\frac{a_{ads}}{V} \right) (C_t - C_e) \quad (10)$$

Where:

C_t [mol.m⁻³]: the concentration of the compound in the fluid phase.

C_e [mol.m⁻³]: the concentration of the compound at the surface of the adsorbent.

k_f [m.s⁻¹]: the external mass transfer coefficient.

a_{ads} [m²]: the useful surface area for external transfer.

V [m³]: Volume of the adsorption bed.

The second step is called internal diffusion. It results from the transfer of fluid phase particles from the outside of the solid surface inside the pores. To simplify the problem to one spatial dimension, the pore is assumed spherical. The flow transferred into the pore is expressed by Equation (11).

$$J = -D_p \frac{\varepsilon_p}{\tau} \frac{\partial C_p}{\partial r} \quad (11)$$

Where:

J [mol.m⁻².s⁻¹]: the transferred flux.

D_p [m².s⁻¹]: the pore diffusion coefficient.

ε_p [-]: the porosity.

τ [-]: the tortuosity.

C_p [mol.m⁻³]: the compound concentration in the pore.

r [m]: radius of the pore.

The last step of the adsorption mechanism is the surface diffusion. It corresponds to the attachment of the vapor phase particles on the surface of the adsorbent. This step is very quick and independent of the overall process. The flux is then defined by Equation (12).

$$J = -D_s \frac{\varepsilon_p}{\tau} \frac{\partial q}{\partial r} \quad (12)$$

Where D_s [$\text{m}^2 \cdot \text{s}^{-1}$] is the surface diffusivity and q [$\text{mol} \cdot \text{kg}^{-1}$] is the amount adsorbed.

Fig. 19 shows the different steps of the adsorption mechanism.

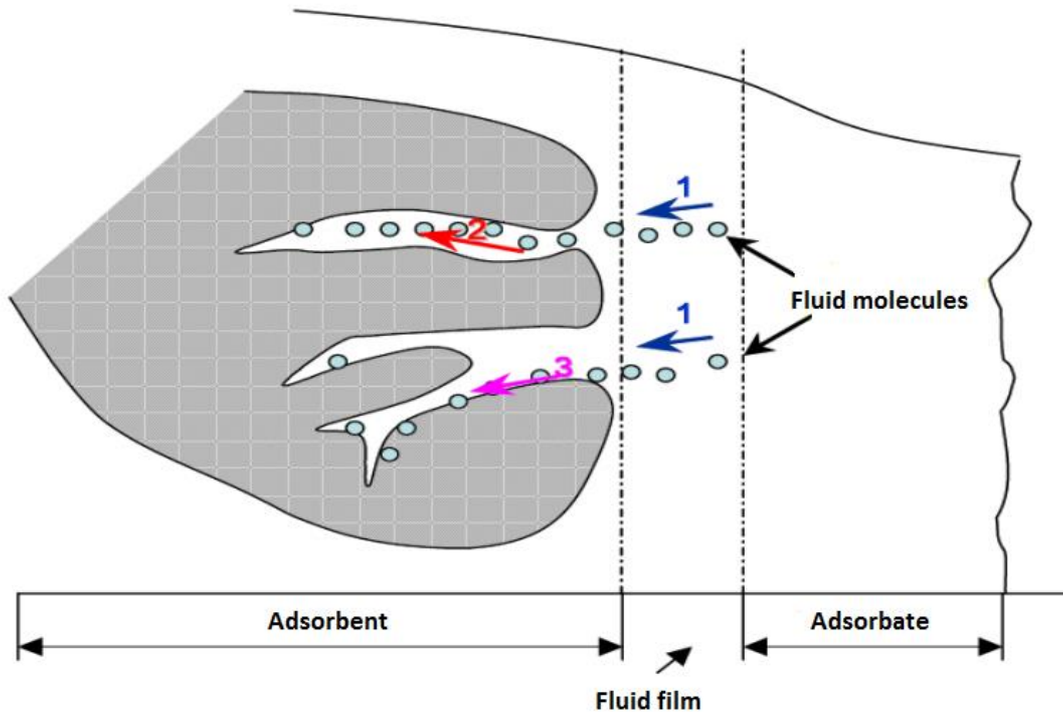


Fig. 19: Transport mechanism of the adsorbate molecules on the adsorbent surface

3.3.2. Materials used for H_2S adsorption

The adsorbents are microporous solids, characterized by high surface area per unit weight, from 100 to over $2000 \text{ m}^2 \cdot \text{g}^{-1}$. The classification of the International Union of Pure and Applied Chemistry (IUPAC) defines three kinds of pores by their size:

- Microporous materials have pore diameters of less than 2 nm.
- Mesoporous materials have pore diameters between 2 and 50 nm.
- Macroporous materials have pore diameters of greater than 50 nm.

The most widely used adsorbents in industry remain: activated carbon, zeolite, silica gel and activated alumina. Other materials rarely used today as red mud may have good adsorption performance in H_2S removal.

- **Activated carbon**

Activated carbon is characterized by a high degree of microporosity. This structure gives the activated carbon its very large surface area. This essential feature allows the activated carbon to be by far, the most widely used adsorbent in industry. Pores size between of 0.5 and 1 nm were found by Yan et al. [50] to have the best adsorption capacity.

Activated carbon is characterized by a non-polar surface allowing it to preferentially adsorb non-polar compounds.

Activated carbon can be impregnated with potassium hydroxide (KOH) or sodium hydroxide (NaOH) which acts as catalysts to remove H₂S, because non-impregnated activated carbon is a weak catalyst, and however, removes hydrogen sulfide at a much slower rate.

Bandosz [51] has shown that using low hydrogen sulfide concentrations, with a sufficient time in laboratory tests leads to comparable removal capacities of both impregnated and non-impregnated activated carbons. But in on-site applications, removal capacities vary greatly because of the presence of other constituents such as volatile organic compounds (VOCs) which may inhibit the removal capacity. According to Abatzoglou and Boivin [52], the typical H₂S adsorption capacities for respectively impregnated and non-impregnated activated carbons are 150 and 20 mg H₂S/g of activated carbon.

Activated carbon is produced in different shapes and sizes depending on the application for which it is used:

- Extruded activated carbon (EAC).
- Granular activated carbon (GAC).
- Powder activated carbon (PAC).

Fig. 20 shows the different shapes of commercial activated carbons.



Fig. 20: Shapes of commercial activated carbons [53]

- **Zeolites**

Zeolites are microporous adsorbent materials. The size of the pores can be adjusted by ion exchange to catalyze selective reactions. According to the International Zeolite Association (IZA), in 2007, there are 176 crystal structures identified by a three letter code. The vast majority of these structures are synthetic while the rest exists in nature. The zeolites are particularly effective for removing polar compounds such as water and H₂S, from non-polar gas streams, such as methane. Zeolites are low capacity adsorbents, with a surface area not exceeding 900 m².g⁻¹ but they have a good selectivity. Compared to activated carbon, they are less sensitive to heat.

The adsorption of H₂S present in biogas is one of the application envisaged with zeolites. The measured adsorption capacities remain well below the purification yields obtainable with other systems. Yasyerli et al. [54] determined an adsorption capacity of 30 mg.g⁻¹ with clinoptilolite, a natural zeolite during the treatment of a real biogas. Cosoli et al. [55] reported an adsorption capacity of 40 mg.g⁻¹ on synthetic zeolites, for H₂S concentration of 1000 mg.m⁻³.

3.3.3. Factors affecting the adsorption

Literature concerning the factors affecting adsorption dates back to 1914 [56]. The main factors affecting the adsorption rate are the temperature, the surface area and the porosity of the adsorbent, the competition between species, and the polarity of the adsorbent and the adsorbate.

- **Temperature**

Fig. 21 shows that during adsorption processes, the adsorbed amount increases as the temperature decreases. Moreover, physisorption releases heat. So, as any exothermic reaction, it is favored by low temperatures.

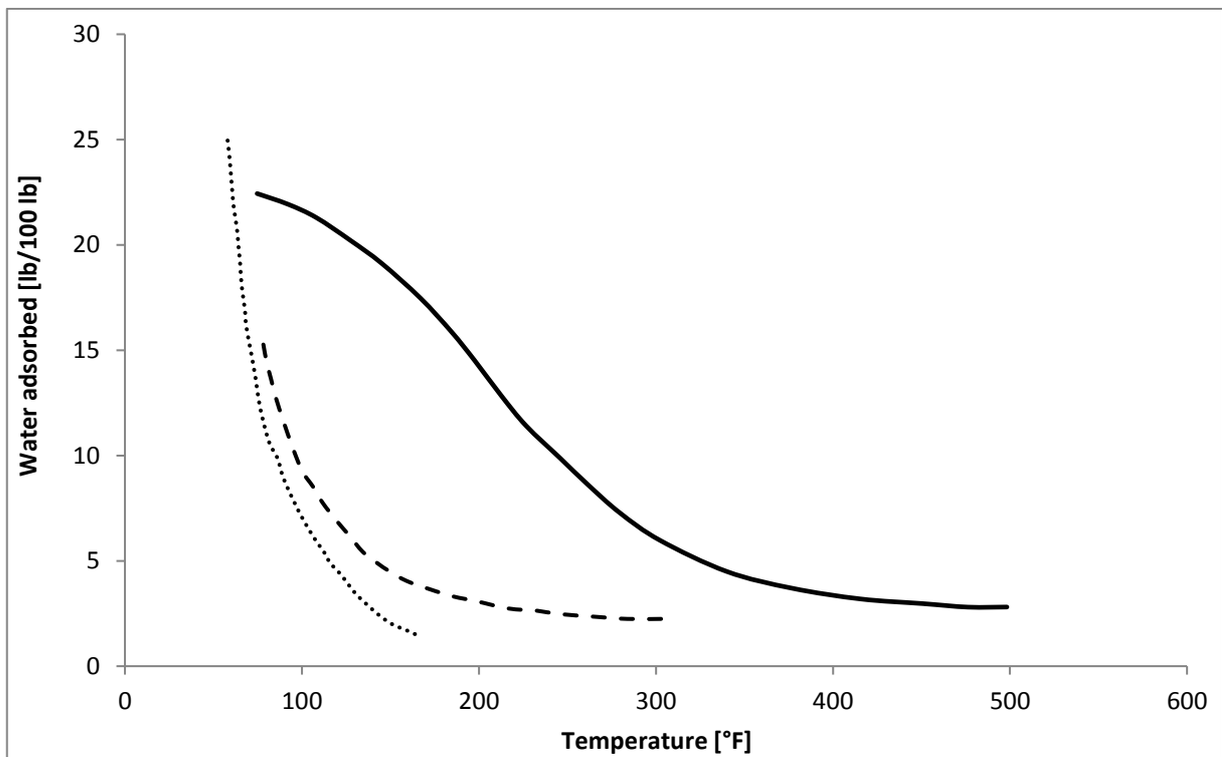


Fig. 21: Effect of temperature on some adsorbents [57]

Adsorbents: (—) Molecular sieves ; (- - -) Activated alumina ; (.....) Silica gel

Unlike physical adsorption, chemical adsorption requires higher temperatures, because it is an endothermic phenomenon.

- **Specific surface area**

Adsorption performance increases with the specific surface area of the adsorbent. This proportionality was demonstrated by Bouchemal and Achour [58] during adsorption study of tyrosine on activated carbon.

- **Selectivity**

The concept of selectivity is crucial in the design of adsorption processes. The presence of competitive species at the surface of the adsorbent decreases the capacity of each species to be adsorbed. However, The higher the selectivity, the easier would be the separation.

- **Pore size distribution**

Adsorption is a surface phenomenon, hence the interest of porous structures. The porosity of the adsorbent material is therefore an important physical property. For example, the microporous activated carbon has a better adsorption capacity than the mesoporous activated carbon in the case of macromolecules.

The thermal regeneration and impregnation could modify the pore volume of the adsorbent.

The resolution of Equation (13) allows access to the porous distribution [59].

$$N(p/p_0) = \int_{w_{min}}^{w_{max}} N(p/p_0, w) f(w) dw \quad (13)$$

Where:

$N(p/p_0)$ [-], is the experimental adsorption isotherm.

$N(p/p_0, w)$ [-], is the local isotherm in a pore size of w .

$f(w)$ [-], is the pore distribution function.

w [nm], is the opening dimension of a pore.

- **Molecular weight and structure**

If the molecular weight of particles is low, it means that they are light and move faster than those with high molecular weight. The probability of being adsorbed is therefore much greater.

If the molecular structure of particle is large, pores are filled rapidly with low yields to saturation, causing the decrease of free sites for other molecules.

- **Polarity**

For more affinity between the adsorbent and the adsorbate, they must have the same polarity [60]. For example, the structure of activated carbons is non-polar and therefore promotes the adsorption of nonpolar molecules. Hydrogen sulfide is a polar gas, it is adsorbed on the polar surfaces in the absence of water vapor. In the presence of water vapor in the gas, there is competitive adsorption to the advantage of water vapor which has a much higher partial pressure and which is much more polar than hydrogen sulfide.

3.3.4. Adsorption isotherms

In order to model the binding of a gas over a bed of adsorbent, it is necessary to choose a model to represent interactions between the gas and the solid.

An adsorption isotherm is the curve presenting the static adsorption capacity of an adsorbate/adsorbent system at a given temperature. The curve presents the specific amount adsorbed, N_a as a function of the relative pressure, P/P_0 as seen in Fig.22.

According to the classification of IUPAC based on the one established by Brunauer [61], there are six different isotherms profiles represented in Fig. 22.

The adsorption isotherm type I, is distinguished by the existence of a horizontal line, which results in the saturation of the adsorbent. This isotherm is characteristic of adsorbent having micropores, which are filled at low relative pressures. This is essentially a monolayer adsorption, often described

by the Langmuir isotherm, where there may be strong interactions involved. The equation that describes the Langmuir isotherm is presented in Table 16.

The adsorption isotherm type II are widespread for non-porous or macroporous solids. The absence of a clearly identifiable inflection point on the curve corresponding to the filling of a monolayer and the absence of a continuous increase of the amount adsorbed are indicative of energy heterogeneity of the surface regarding the interactions adsorbate / adsorbent.

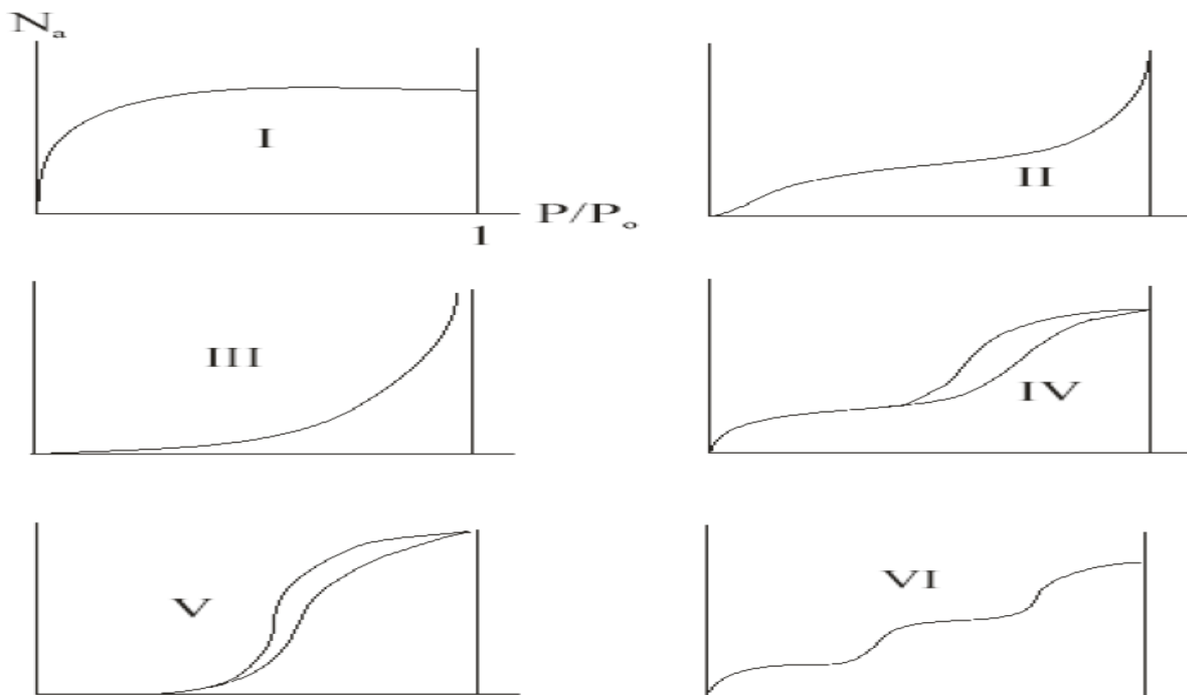


Fig. 22: Classification of adsorption isotherms [IUPAC]

The adsorption isotherm type III corresponds to non-porous or macroporous solids. This isotherm is characterized by weak interactions adsorbate / adsorbent.

The adsorption isotherms type IV and V are characterized by a filling of mesopores, and a capillary condensation in the pores. The interactions adsorbate / adsorbent for type V isotherm are weaker than those of type IV.

The adsorption isotherm type VI is very rare. It is encountered in the case of very homogeneous surfaces.

- **Modeling of adsorption isotherms**

Several models have been proposed to describe the experimental adsorption isotherms. Despite their common interest, the assumptions defined for each model are different such as those concerning interactions that hold the fluid molecules on the surface of the adsorbent.

Most of these models are described in the literature such as: Freundlich model, Elovich model, Temkin model, Toth model and Langmuir model.

The latter is the best known, and probably the most widely used to describe the adsorption isotherm. It was developed by Irving Langmuir in 1916 [62], sixteen years before obtaining the Nobel Prize in chemistry.

The Langmuir model assumes uniform energies of adsorption onto the surface and no transmigration of adsorbate in the plane of the surface [62].

The Freundlich model assumes that as the adsorbate concentration increases, the concentration of adsorbate on the adsorbent surface also increases [63]. This proportionality is explained by the Freundlich expression which is an exponential equation.

The Temkin model assumes that in adsorption, the binding energies are distributed uniformly, and that due to interactions between the adsorbent and the adsorbate, the heat of adsorption of all the molecules in the layer decreases linearly with coverage [64].

The Elovich model is derived from a kinetic principle assuming that the adsorption sites increase exponentially with adsorption, implying a multilayer adsorption [65].

The Toth model [66] was developed based on an improvement of the Langmuir model to reduce the error between experimental and predicted data. This model is applied in the case of multilayer adsorption.

The equations defining the main models are shown in Table 16.

Table 16. Equations governing adsorption isotherm models, and their linear forms [67]

Isotherm	Equation	Linear form
Langmuir	$q_e = \frac{q_m b C_e}{1 + b C_e}$	$\frac{1}{C_e} = b \left(\frac{q_m}{q_e} - 1 \right)$
Freundlich	$q_e = K_F C_e^{1/n}$	$\ln q_e = \ln K_F + \frac{1}{n} \ln C_e$
Temkin	$\theta = \frac{R T}{\Delta Q} \ln K_0 C_e$	$\theta = \frac{R T}{\Delta Q} (\ln K_0 + \ln C_e)$
Elovich	$\frac{q_e}{q_m} = K_E C_e e^{\left(-\frac{q_e}{q_m}\right)}$	$\ln \frac{q_e}{C_e} = \ln K_E q_m - \frac{q_e}{q_m}$
Kiselev	$k_1 C_e = \frac{\theta}{(1 - \theta) (1 + k_n \theta)}$	$\frac{1}{C_e(1 - \theta)} = k_1 \left(k_n + \frac{1}{\theta} \right)$
Fowler-Guggenheim	$K_{FG} C_e = \frac{\theta}{1 - \theta} e^{\left(\frac{2\theta W}{R T}\right)}$	$\ln \left[\frac{C_e(1 - \theta)}{\theta} \right] = \frac{2\theta W}{R T} - \ln K_{FG}$
Hill-de Boer	$K_1 C_e = \frac{\theta}{1 - \theta} e^{\left(\frac{\theta}{1 - \theta} - \frac{K_2 \theta}{R T}\right)}$	$\ln \left[\frac{C_e(1 - \theta)}{\theta} \right] - \frac{\theta}{1 - \theta} = - \left(\ln K_1 + \frac{K_2 \theta}{R T} \right)$
Toth	$\frac{q_e}{q_{mT}} = \frac{C_e}{\left(\frac{1}{K_T} + C_e^{mT} \right)^{\frac{1}{mT}}}$	$\ln \left(\frac{q_e^{mT}}{q_{mT}^{mT} - q_e^{mT}} \right) = mT (\ln C_e + \ln K_T)$

Where :

q_e [g.kg⁻¹] is the amount of solute adsorbed per unit weight of adsorbent at equilibrium.

q_m [g.kg⁻¹] is the maximum adsorption capacity.

q_{mT} [g.kg⁻¹] is the Toth maximal adsorption capacity.

b [m³.g⁻¹] is the Langmuir constant related to the free energy of adsorption.

C_e [g.m⁻³] is the equilibrium concentration of the solute in the bulk solution.

C_0 [g.m⁻³] is the initial concentration of the solute in the bulk solution.

$K_F [g^{1-(1/n)} \cdot m^{(3/n)} \cdot kg^{-1}]$ is the Freundlich constant indicative of the relative adsorption capacity of the adsorbent.

$K_T [-]$ is the Toth equilibrium constant.

$n [-]$ is the Freundlich constant indicative of the intensity of the adsorption.

$mT [-]$ is the Toth model exponent.

$\theta [-] = q_e / q_m$, is the surface coverage.

$\Delta Q [kJ \cdot mol^{-1}]$ is the variation of adsorption energy.

K_0 and $K_1 [m^3 \cdot g^{-1}]$ are the Temkin and Hill-de Boer equilibrium constants respectively.

$K_2 [kJ \cdot mol^{-1}]$ is the energetic constant of the interaction between adsorbed molecules.

$K_{FG} [m^3 \cdot mol^{-1}]$ is the Fowler-Guggenheim equilibrium constant.

$W [kJ \cdot mol^{-1}]$ is the interaction energy between adsorbed molecules.

3.3.5. Adsorption processes

There are two major classes of adsorption processes: the temperature swing adsorption processes (TSA) and the pressure swing adsorption processes (PSA).

- **Temperature swing adsorption processes**

The temperature swing adsorption is the oldest cyclic adsorption process. It consists of two main phases: the adsorption phase and the desorption phase, during which the adsorber is heated. A pre-cooling phase is therefore commonly added to bring the temperature to a level similar to that desired for adsorption.

The main advantage of temperature swing adsorption process compared to pressure swing adsorption is to desorb more easily species strongly adsorbed. For this reason, the temperature swing adsorption processes are used, for example, to capture volatile organic compounds present in many effluents.

Temperature swing adsorption is used whenever the energy required to regenerate a bed is sufficiently large that long high thermal cycles are needed given the strength of the bond between the adsorbate and adsorbent. For example, adsorption of H_2O on zeolites.

However, significant time required to heat and cool the adsorber, prevent the use of temperature swing adsorption process in fast cycle. Moreover, the adsorption columns used for temperature swing adsorption cycles are large in size, which has an impact on the cost of the installation. However, unlike pressure swing adsorption processes, that use mechanical energy, temperature swing adsorption processes can use residual heat, which then reduce their operating cost.

- **Pressure swing adsorption processes**

The pressure swing adsorption process was initially introduced as an alternative to temperature swing adsorption process. It is mainly used in separation of some gas species from a mixture of gases, this is another option that complements the traditional separation processes as absorption and cryogenic distillation.

In a pressure swing adsorption process, the feed pressure is generally greater than atmospheric pressure. The regeneration pressure may be less than the atmospheric pressure, in this case the process is called vacuum swing adsorption (VSA).

Since the adsorption step is performed at a higher pressure than the pressure of desorption step, intermediate steps are necessary: a compression is required to move from the low to the high pressure at the end of the regeneration step. A decompression step is also necessary to reduce the pressure at the end of the adsorption step.

These four steps are the elements of a basic cycle called the Skarstrom cycle. Steps which constitute this cycle, and the variation of the pressure as a function of the different phases, are shown in Fig. 23.

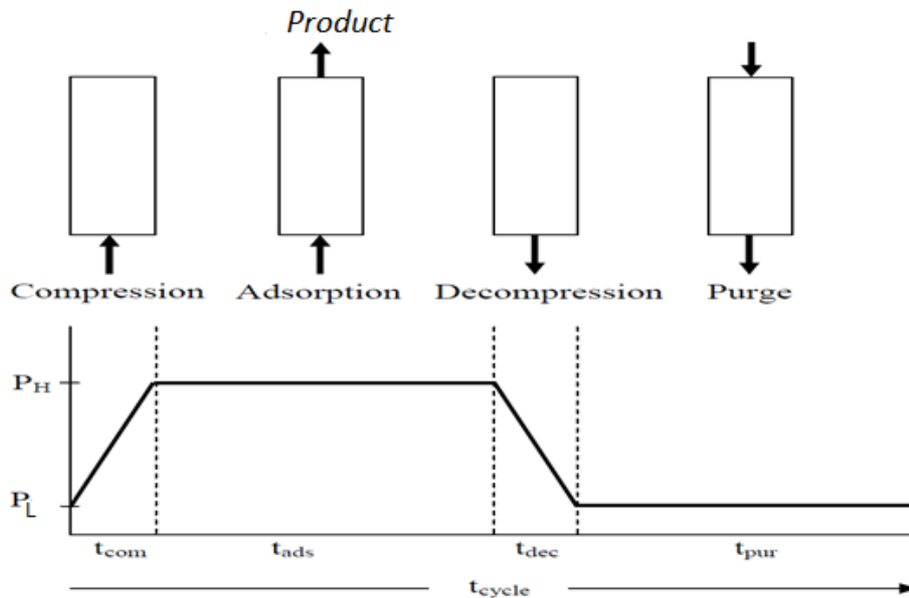


Fig. 23: Skarstrom cycle stages and pressure variations [68]

To ensure continuous production, the pressure swing adsorption process must have at least two separation columns. These columns suffer the four steps mentioned above, but with a temporal phase shift leading to a cyclic operation of the pressure swing adsorption process, in which one of the columns is regenerated, while the gas mixture is separated in the other one.

Currently, there are several hundred thousand pressure swing adsorption processes installed worldwide. Their size varies from $6 \text{ l}\cdot\text{hr}^{-1}$, often for the production of medical grade oxygen with a purity of 90 % to $2000 \text{ m}^3\cdot\text{h}^{-1}$ typically for the production of pure hydrogen at 99.999 % [69].

For purification, temperature swing adsorption is generally the process of choice. For bulk separation, pressure swing adsorption is more suitable.

3.4. Membranes technology

A membrane can be defined as a physical barrier for the selective transport of chemical species. As seen in Fig. 24, it allows the restricted passage of one or more constituents. The flux passing through the membrane is called permeate, while the retained is called retentate or concentrate. One or the other of these two flows may be advantageous according to the intended application, and can therefore be used as final product [70].

The driving force in a membrane can be a pressure gradient, a concentration gradient, a temperature gradient, or an electrochemical gradient. Thus, the membranes include a wide variety of materials and structures. Table 17 shows the main materials used by membranes manufacturers.

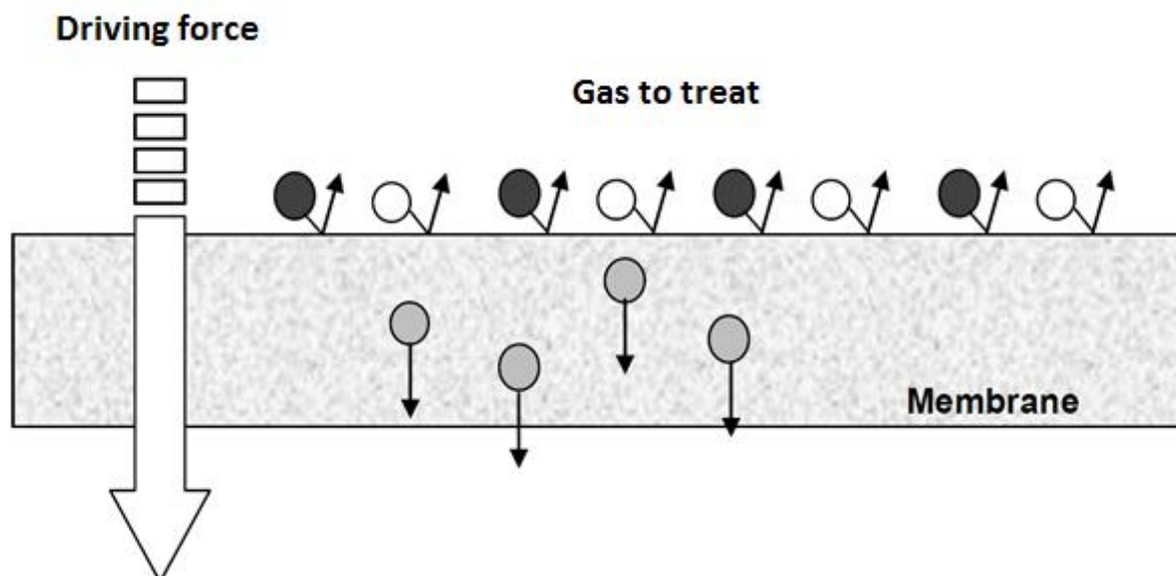


Fig. 24: Schematic representation of a membrane [71]

The membrane performance is evaluated for its ability to separate different species from a gas mixture, and transport a maximum quantity of gas at high speed. These two criteria are generally in competition, a membrane is more efficient when it presents the best compromise: flow / selectivity.

A membrane can be gaseous, liquid, solid or a combination thereof [72]. It covers a wide range of applications such as ultrafiltration, microfiltration, reverse osmosis, pervaporation, electrodialysis and gas separation. In addition, other medical applications such as blood oxygenators and artificial kidneys require the use of membrane technology. Of the applications listed in Table 18, reverse osmosis and ultrafiltration are the most widely used industrially.

Table 17. The membrane materials used by manufacturer [71]

The membrane materials	manufacturers
Cellulose acetate	Grace
Hydrin C	Zeon
Pebax	Atochem
Polyacrylate	Röhm
Polydimethylsiloxane	Wacker, GKSS
Polyhydantoin	Bayer
Polyetherimide	General Electric
polyethersulfone	Bayer, BASF, Monsanto

The two essential parameters in the operation of a membrane are: permeability and selectivity. These parameters are used to provide informations on the membrane, and to describe its performance on the transfer of material through the barrier and its ability to separate one or more chemical species in a gas mixture.

The selectivity expressed most often by separation factor is defined as the ratio of the compositions of components i and j in the permeate relative to the composition ratio of these components in the retentate.

$$S_{f i,j} = \frac{\left(\frac{x_i}{x_j}\right)_{\text{permeate}}}{\left(\frac{x_i}{x_j}\right)_{\text{retentate}}} \quad (14)$$

The permeability is used to indicate the ability of the membrane to feed the permeate. In order to ensure an attractive performance, the permeability Π_i [$\text{mol.m}^{-2}.\text{Pa}^{-1}.\text{s}^{-1}$] must be high to ensure a large transmembrane flux.

$$\Pi_i = \frac{J_i}{\Delta P_i} \quad (15)$$

Where J_i [$\text{mol.m}^{-2}.\text{s}^{-1}$] is the transferred flux and ΔP_i [Pa] is the partial pressure difference of the constituent i through the membrane.

Generally, the membranes with a high permeability have a low selectivity and vice versa.

Table 18. Classification of membrane separation processes [73]

Application	Driving force	Separation size range
Microfiltration	Pressure gradient	10 – 0.1 μm
Ultrafiltration	Pressure gradient	< 0.1 μm – 5 nm
Reverse osmosis	Pressure gradient	< 5 nm
Electrodialysis	Electric field gradient	< 5 nm
Dialysis	Concentration gradient	< 5 nm

- **Membranes for gas separation**

The membranes can be classified according to different viewpoints. They can be divided according to the nature: biological or synthetic, according to the morphology, or according to the structure.

Several scientific journals classify membranes into porous and non-porous membranes depending on the structure of the material.

In general, a membrane may be thick or thin, and its structure may be homogeneous or heterogeneous with a transfer mechanism which may be active or passive [74].

Fig. 25 shows the main types of membrane. The figure shows a thin interface that forms the membrane. This interface can be homogeneous at the molecular level, that is to say, completely uniform in composition and structure. It can also be heterogeneous, comprising for example pores.

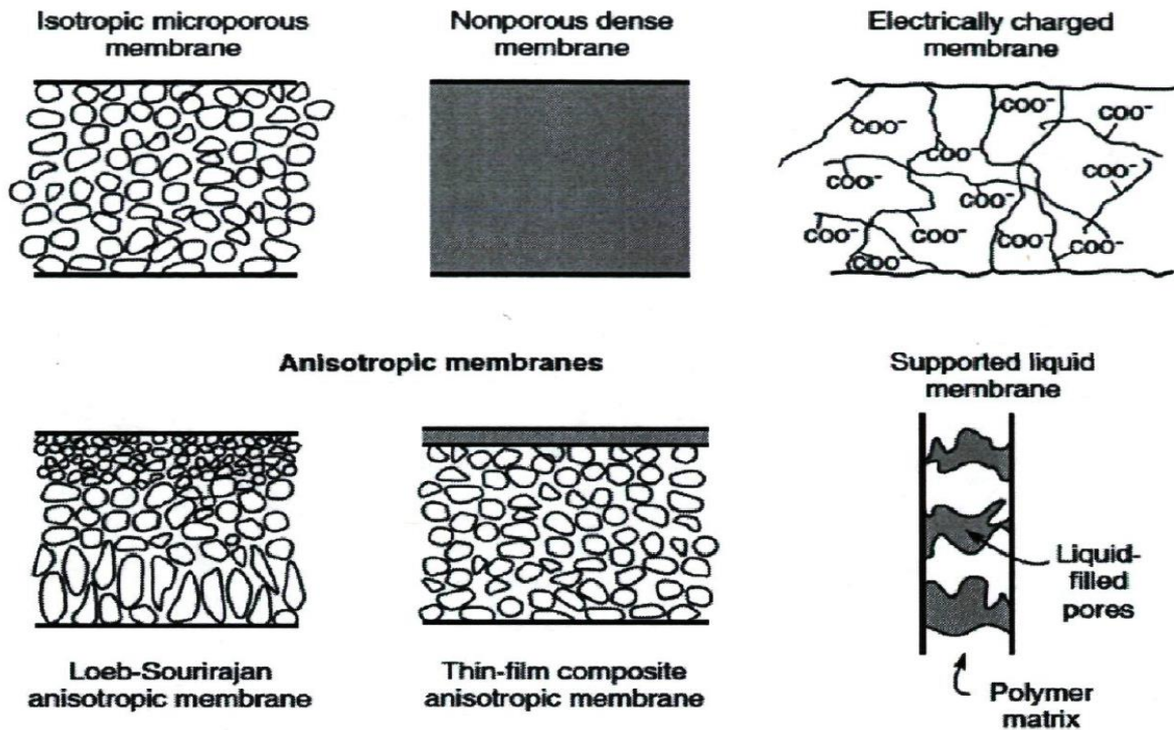


Fig. 25: Diagram of the main types of membranes [75]

The main types of membranes used today for gas separation are membranes with dense polymeric materials, where transfers follow a solute solubility and diffusion mechanism based on Fick's law.

$$J_i = D_{ij} \frac{\partial c_i}{\partial x} \quad (16)$$

The structure of a porous membrane is like a sponge, it is also very similar to a conventional filter [74]. Most of the materials used are characterized by tortuous and interconnected pores, whose precise geometry is inaccessible.

Table 19 shows the different categories of membranes according to the size of their pores provided by the International Union of Pure and Applied Chemistry.

Table 19. Distribution of membranes according to pore size [IUPAC]

Membrane type	Pores size (Å)	Physical mechanism	Application
Dense		Diffusion	Reaction, gas separation
Microporous	≤ 20	Micropore diffusion	Gas separation
Mesoporous	20 – 500	Knudsen diffusion	Ultrafiltration, Nanofiltration, Gas separation
Macroporous	≥ 500	Molecular sieve	Ultrafiltration

- **Geometric configuration of membranes**

The geometric design is an essential step in any membrane process as this factor defines the active area of the module. The first membranes used are: flat sheet and tubular membranes. Today, these

systems are still available, but their use has declined because of their low efficiency and high cost. They are mostly replaced by spiral wound, and hollow fibers membranes.

Fig. 26 shows the four membrane contactors mentioned above.

The flat sheet membranes are the oldest and simplest to use. They have planar configuration and are mainly rectangular, though other geometries exist for membrane modules designed to rotate. Modules may be stacked to provide a double deck.

Tubular membranes consist of tubes having an inner diameter between 4 and 25 mm [76]. They are based on a simple technology, easy to use and clean, but they are large energy consumers for a low exchange surface area per unit volume.

Spiral wound membranes consist of a flat sheet membrane coiled on itself around a perforated tube, which collects the residue. As seen in Fig. 26.c, the feed flows axially in the channels, while the permeate flows along a spiral path towards the porous tube [76].

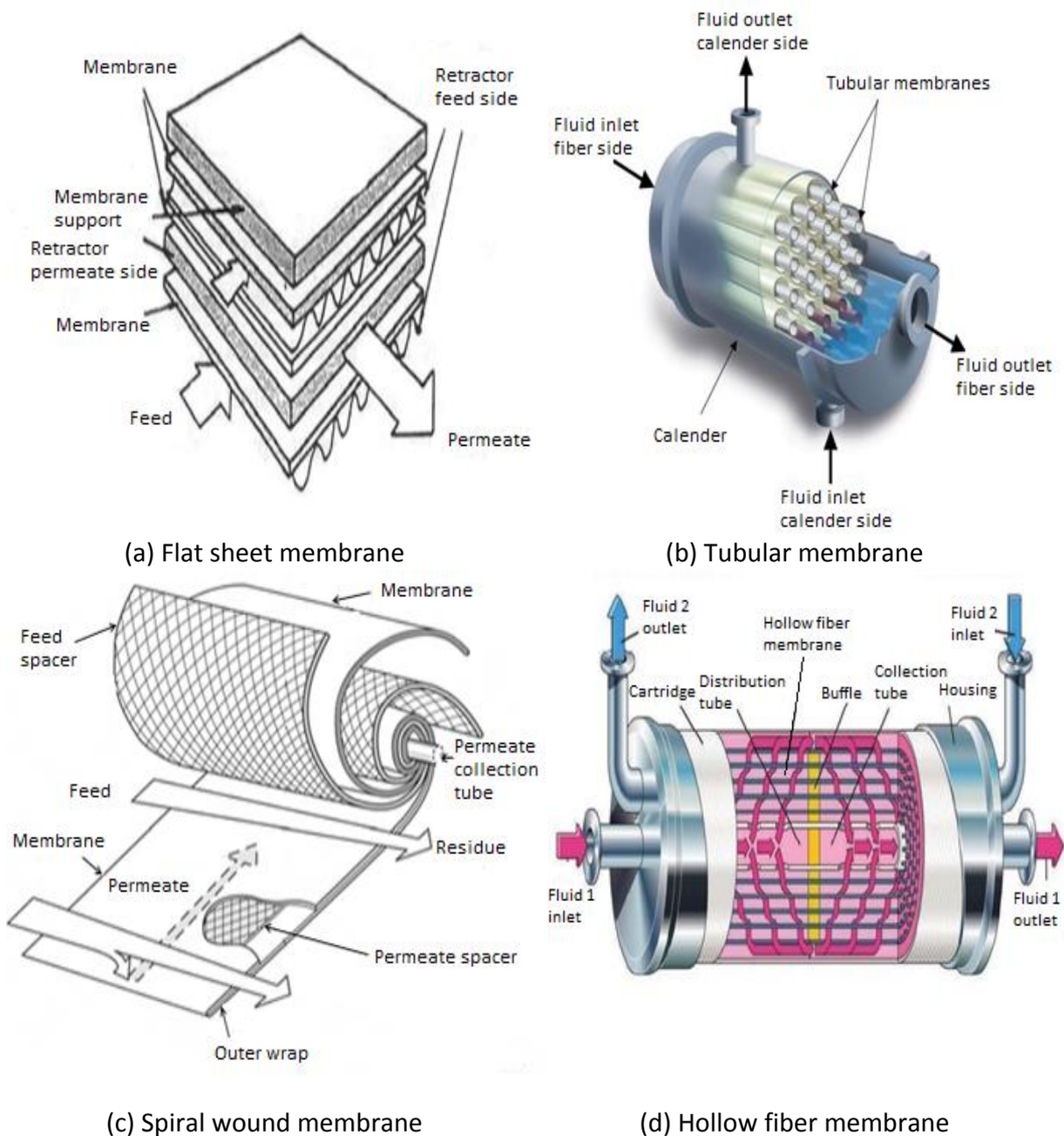


Fig. 26: The geometric configurations of membrane contactors [76]

The hollow fiber modules consist of a bundle of hollow fibers of outer diameter less than 1 mm. The main advantage of hollow fiber membrane is their compactness, due to the membrane's high packing density. Also, They can be operated at very high pressures, due to the absence of membrane support. One disadvantage of hollow fiber membrane is the pressure drop. Hence the importance of fiber length criterion in the design of a separation unit.

Overall, the selection of a given configuration should be addressed individually based on membrane properties and the throughput rates desired [77]. Table 20 shows the characteristics of different geometries of membranes.

Table 20. Characteristics of the different geometries of membranes [76]

Property	Flat sheet membrane	Tubular membrane	Spiral wound membrane	Hollow fiber membrane
Interfacial area [m ² /m ³]	≈ 100	≈ 1000	≈ 500	5 - 10000
Filling density	Low	High	Moderate	Very high
Resistance to soiling	Good	Low	Moderate	Low
Use at high pressures	Difficult	Easy	Easy	Easy
manufacturing cost	High	Moderate	High	Moderate
Application limited to membranes	No	Yes	No	Yes

- **Biogas purification by membrane processes**

In the case of biogas, separation of compounds is often limited to three species: methane, carbon dioxide and hydrogen sulfide. Polyurethane membranes show significant selectivity between methane and hydrogen sulfide compared to that of methane and carbon dioxide. The other types of membranes such as polyimide, polyamide, polysulfone and cellulose acetate membranes show a significant selectivity between methane and carbon dioxide, at the expense of selectivity between methane and hydrogen sulfide.

Today, despite the limited studies and few results, it is possible to obtain methane concentrations above 95 % in the retentate. However, this residue may contain significant hydrogen sulfide concentrations. But the main disadvantage of the membrane process remains the methane loss, with methane weight percentages which can reach 15 % in the permeate. In addition, the membrane processes require treatment upstream, to separate the volatile organic compounds and water vapor.

The membrane's resistance to breaking due to the pressure gradient is one important technical limitation. Exposure to certain solvents and materials causes the membrane to get either damaged or blocked up. These limitations are of great importance since membranes usually are expensive.

3.5. Cryogenic technology

The term cryogenic refers to the science of very low temperatures. The cryogenic separation process consists on passing the pollutant from the gas phase to the liquid or solid phase by lowering the

temperature, in order to separate it from the carrier gas which is the methane. The pollutant is recovered, and will then be destroyed or valorized for possible use.

This technology is based on the thermodynamics of phase equilibria. The thermodynamic equilibrium between the different phases results in a graph called phase diagram, which generally uses pressure and temperature as variables.

To purify and upgrade biogas with the cryogenic technology, the gas is chilled and the differences in condensation or solidification temperatures for different compounds are used to separate impurities and carbon dioxide from biogas, which can be seen in Table 21.

Table 21. Condensation or solidification temperatures, at atmospheric pressure, for the different compounds present in biogas

Compound	Condensation temperature [°C]
H ₂ S	-60.3
CO ₂	-78.5
CH ₄	-161.5
N ₂	-195.8

The technology can be used to upgrade biogas by cooling it at atmospheric pressure in order to separate carbon dioxide at temperatures related to the CO₂ partial pressures upstream and downstream the refrigeration unit, typically from -90 °C to -120 °C. Then, the biogas is chilled to produce liquid biogas (LBG) at temperatures which depend on the pressure: -120 °C at 1.5 MPa to -162 °C at atmospheric pressure.

To ensure a high purity of the products and an optimal operation, all traces of hydrogen sulfide should be removed upstream of the process using one of the conventional technologies presented above.

Depending on the temperature of the process, different purity grades can be reached. A lower temperature results in a higher removal efficiency.

A study was performed to condense the volatile organic silicon compounds (VOSiC) contained in biogas. This process involves cooling the biogas at different temperatures, in order to evaluate the effect of cold on volatile organic silicon compounds removal. Table 22 shows the results obtained by different authors using this process at different temperatures.

Table 22. Effect of temperature on the abatement of volatile organic silicon compounds

Condensation temperature [°C]	Abatement rate of VOSiC [%]	References
5	12.0	[78]
-25	25.9	[79]
-70	99.3	[79]

Production of liquid biogas is a suitable upgrading technology for landfill gas, which usually consists of significant amount of nitrogen, hard to separate from methane with conventional technologies. However, when the methane is liquefied, nitrogen can be separated due to its lower condensation temperature [80].

Cooling biogas to very low temperatures is energy intensive but in some occasions the product is more valuable. If the biogas production plant is situated on the countryside, far from the end users, it is more space efficient to transport biogas in its liquid state. Today pressurised (200 bar) gas is delivered in gas vessels stored on a mobile compressed biogas (CBG) storage, leading to transportation of a huge share of steel, compared to gas [81].

Producing liquid biogas also leads to a renewable fuel available for heavy duty vehicles. The fuel can be stored as liquid biogas on the vehicle, which increase the driving distance per tank. The requirement is that the vehicle is running frequently, otherwise liquid biogas will vapourize and the methane will be vented to the atmosphere [82].

An advantage of the cryogenic technology is that it does not need any water or solvent to function, although it requires external cooling equipment such as a refrigeration system.

3.6. Choice of the separation process

Various technologies are available in order to purify and upgrade biogas. Water scrubbing and pressure swing adsorption dominated the market until 2008. But lately, membrane separation units and chemical scrubbers have increased their market share as seen in Fig. 27.

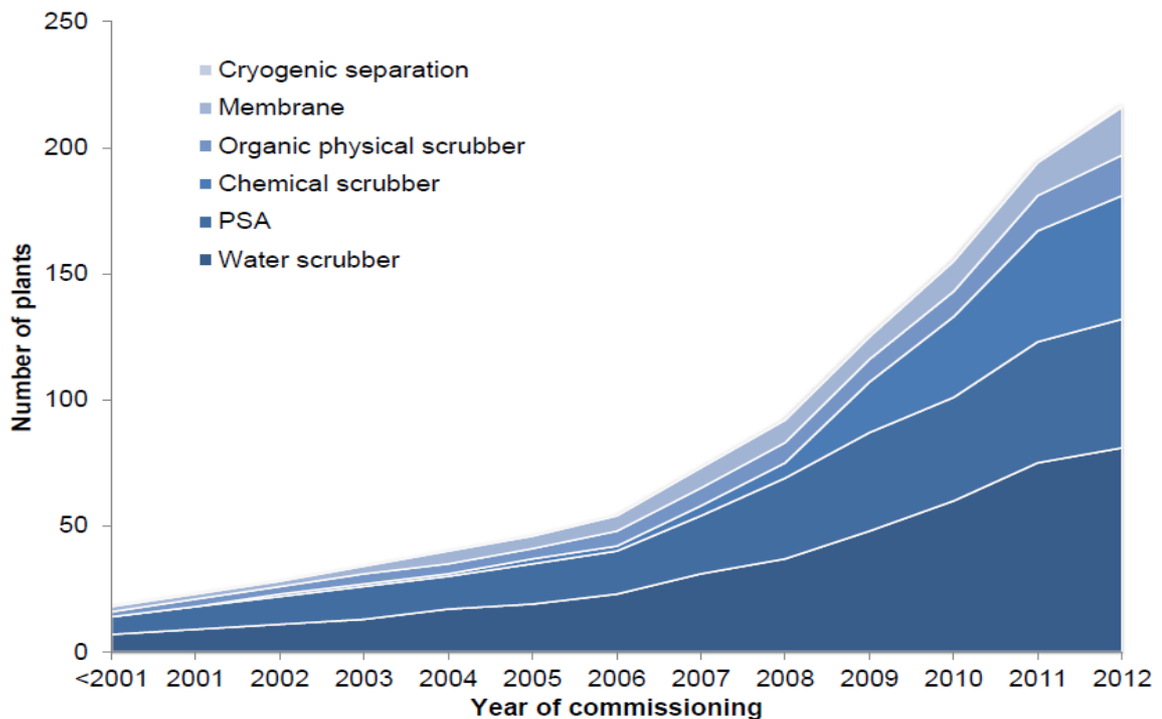


Fig. 27: Most technologies used for the purification and upgrading of biogas [International Energy Agency]

The choice of the technology to be used depends on multiple parameters, such as the final use, the incentives, the flow rate, the nature and diversity of species present, and the concentrations to treat. The solution chosen has to meet different requirements, both technical and economic. Other criteria can sometimes be decisive: environment, maintenance, temperature and pressure ... There is therefore, no universal treatment technology.

Table 23 provides an overview of the different biogas purification and upgrading techniques. It indicates the methane concentration in the purified gas, loss of methane and the substances used in the process such as water and chemicals solvents.

The most important criterion for environmental impact of technology is methane losses. The portion of methane which slips away from raw biogas because of the separation technology itself, contributes to global warming, so regulations of most European countries require that the methane slip has to be burnt.

Table 23. Comparison of the different biogas purification and upgrading technologies

Separation technology	Methane concentration obtained [%]	Methane losses [%]	Process needs
Chemical absorption	95 – 98 [83] > 99.5 [84]	0.1 – 0.2	Amines or chemical solvent recharges
Water wash at high pressure	96 – 98 [83] > 98 [84]	10 – 20 (The high pressure increases the methane solubility in water)	Large water requirement
Adsorption	95 – 98 [83] 98 [84]	2	
Membrane separation	76 – 95 [83] 90 – 93.5 [85] 98 [86]	6.5 – 10 [85] 2 [86]	Change membranes
Cryogenic technology	> 97 [83]		Refrigerants

The two main selling points of a biogas treatment unit, are its efficiency and cost. Table 24 compares the main purification and upgrading technologies according to these two criteria. Operating and investment costs are very variable depending on the technology, for abatement performance, often above 90 %.

Generally, the more expensive technologies, both in investment and in operation are those of oxidation, but during the work of this thesis, only anaerobic processes are studied.

Table 24. Performances, costs, advantages and disadvantages of separation processes [87]

Separation technology	Efficiency [%]	Investment cost [€/m ³ .h ⁻¹]	Operating cost [€/1000 m ³]	Advantages	Disadvantages
Absorption	95 – 98	7 – 32	1.7 – 8.2	<ul style="list-style-type: none"> - Simple operation for a wide range of flow rates, concentrations and compounds. - Possible pollutant recovery. 	<ul style="list-style-type: none"> - High operating costs, related to the liquid phase in general. - Generating a polluted aqueous effluent. - Can generate additional separation operations.
Adsorption on activated carbon	80 – 90	7 – 55	0.7 – 2.4	<ul style="list-style-type: none"> - Very easy to use. - Tolerates flow variations. - Possible pollutant recovery. 	<ul style="list-style-type: none"> - Add an operating cost, associated with the regeneration of the adsorbent.
Condensation	50 – 90	5 – 37	1.4 – 8.2	<ul style="list-style-type: none"> - Possible pollutant recovery. 	<ul style="list-style-type: none"> - Icing Possibility

The investment costs of the different technologies do not differ greatly, especially at high flow rates. They are presented in Fig. 28.

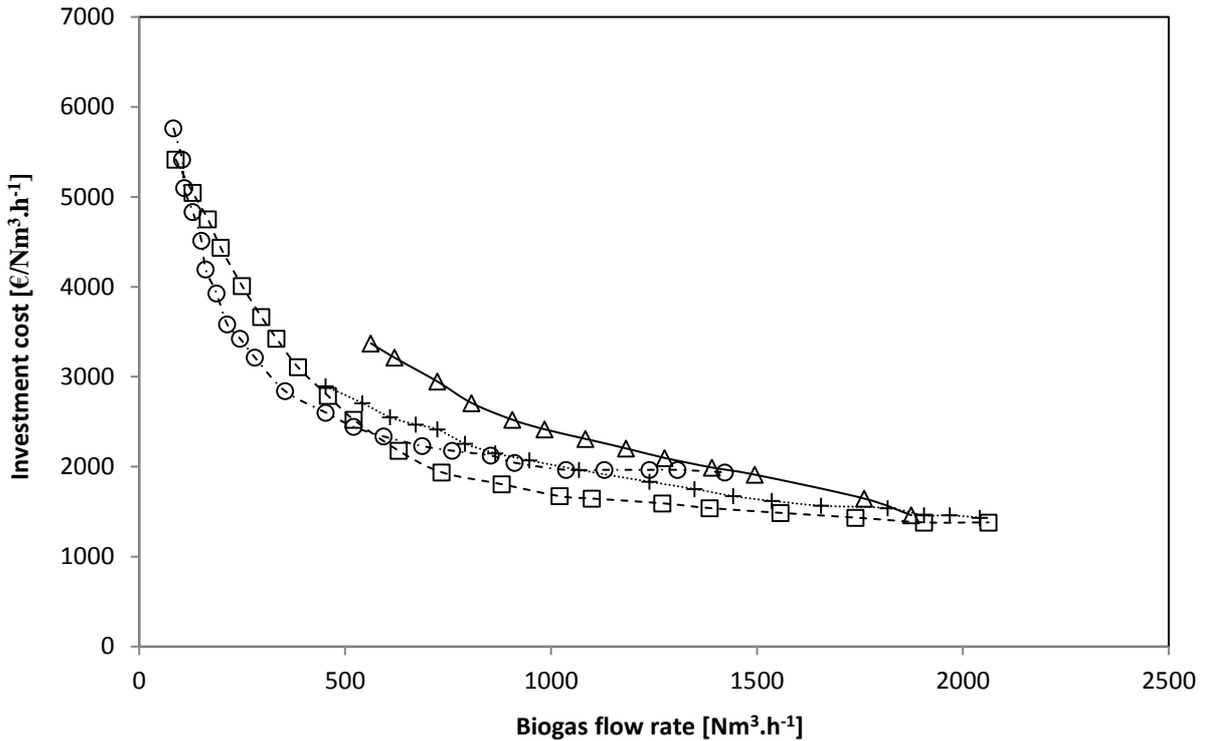


Fig. 28: Comparison of investment costs of different biogas purification and upgrading technologies [88]

Separation technologies: (—Δ—) Chemical absorption (Amines) ; (- -□ - -) Water wash ; (- ·○· ·) Membranes ; (··+··) Adsorption (PSA)

Chemical absorption using amines as chemical solvent is slightly more expensive in terms of investment, and membrane separation process is less costly for low flow rates. This investment cost criterion begins to converge for all technologies at higher flow rates.

3.7. Conclusion

This chapter has introduced the various technologies used for the purification and upgrading of biogas. These separation methods were compared according to several criteria such as separation efficiency, the environmental impact and investment and operating costs.

The work of this thesis is part of the project led by the company Cryo Pur[®], which aims to create an innovative biogas cryogenic purification and upgrading process for the production of a renewable fuel and liquid carbon dioxide.

For carbon dioxide capture, Cryo Pur[®] company has developed a new technology which consists of anti-sublimating the carbon dioxide on a low temperature surface (from -90 °C to -120 °C), thus transforming CO₂ directly from its gaseous phase into a solid phase frosted on the cold surface [89]. Having regard to the need to liquefy the biogas at low temperatures to be used as vehicle fuel, the best alternative to the conventional technologies is therefore to upgrade biogas with cryogenic technology.

Biogas upgrading depends on the concentrations of hydrogen sulfide present. Indeed, this compound must be completely eliminated upstream of the process to ensure high quality products and to prevent corrosion of equipment such as heat exchangers used in the cryogenic process. Hence the need to use a process with a very high efficiency, to be able to eliminate all the hydrogen sulfide present in the biogas. On the whole, the lowest methane losses are indicated for chemical absorption and adsorption processes, and the highest one relates to membranes and water wash.

Finally, the choice was focused on the following two technologies used for the removal of hydrogen sulfide:

- Chemical absorption in a structured packed column using sodium hydroxide (NaOH) as solvent.
- Adsorption in a fixed bed using activated carbon.

The next chapter will present the technology developed by Cryo Pur[®] Company for purification and upgrading of biogas.

Chapter 4: Industrial demonstrator description

Résumé :

Ce chapitre présente le procédé industriel dans lequel se situe l'étape de désulfuration étudiée. Il s'agit d'un démonstrateur industriel appelé « BioGNVAL » développé par la société Cryo Pur® en partenariat avec SUEZ et l'Agence De l'Environnement et de la Maitrise de l'Energie (ADEME). GNVert et IVECO sont également partenaires du projet BioGNVAL.

Ce pilote traite 85 Nm³/h de biogaz issu de la station d'épuration du Syndicat Interdépartemental pour l'Assainissement de l'Agglomération Parisienne (SIAAP) à Valenton, la deuxième plus grande en France.

Une des voies de valorisation du biogaz est la production de bio-GNL (GNL : Gaz Naturel liquéfié) qui se présente comme un carburant neutre en émissions de gaz à effet de serre avec plusieurs avantages économiques et environnementaux. La production de ce type de carburant requiert de très basses températures afin de liquéfier le bio-méthane, ce qui peut induire la solidification des impuretés et donc des problèmes de fonctionnement des installations. Ces impuretés doivent donc être séparées du biogaz, en particulier l'élimination de l'hydrogène sulfuré est impérative afin de garantir un fonctionnement optimal et une grande pureté des autres composés à valoriser comme le dioxyde de carbone. De surcroît, la présence de H₂S dans le biogaz est une source de corrosion pour les équipements comme les pompes et les échangeurs de chaleur.

4.1. Introduction

Experiments were performed on the industrial demonstrator “BioGNVAL” treating 85 Nm³/h of biogas from the Valenton water treatment plant, the second biggest in France run by the SIAAP (Public society serving the Paris region). The demonstrator shown in Fig. 29 was developed by the Cryo Pur[®] Company. It was built in partnership with SUEZ as part of the BioGNVAL project, and partially funded by the ‘Invest in the Future’ program run by the ADEME (French Environment and Energy Management Agency). GNVert (Engie) and IVECO are also partners in the BioGNVAL project, providing the Bio-LNG distribution station and the heavy goods vehicle Flex Fuel gas / Bio-LNG respectively.



Fig. 29: BioGNVAL demonstrator located at Valenton water treatment plant [9]

The BioGNVAL pilot plant uses a cryogenic method to purify and liquefy biogas efficiently without loss of methane and without emitting greenhouse gases. The system generates two products from biogas: liquid bio-methane and bioCO₂ at purity level greater than 99.995 % respecting EIGA (European Industrial Gases Association) specifications [90].

The general principle of operation of the BioGNVAL pilot plant is depicted in Fig. 30. It takes place in three main stages:

- Pretreatment or purification which is to remove trace compounds present in the biogas as hydrogen sulfide, water vapor and siloxanes.

- CO₂ capture or biogas upgrading which consists in separating carbon dioxide from biogas. The content of carbon dioxide in biogas is typically greater than 30 %.
- Liquefaction of the biomethane after purification and upgrading biogas.

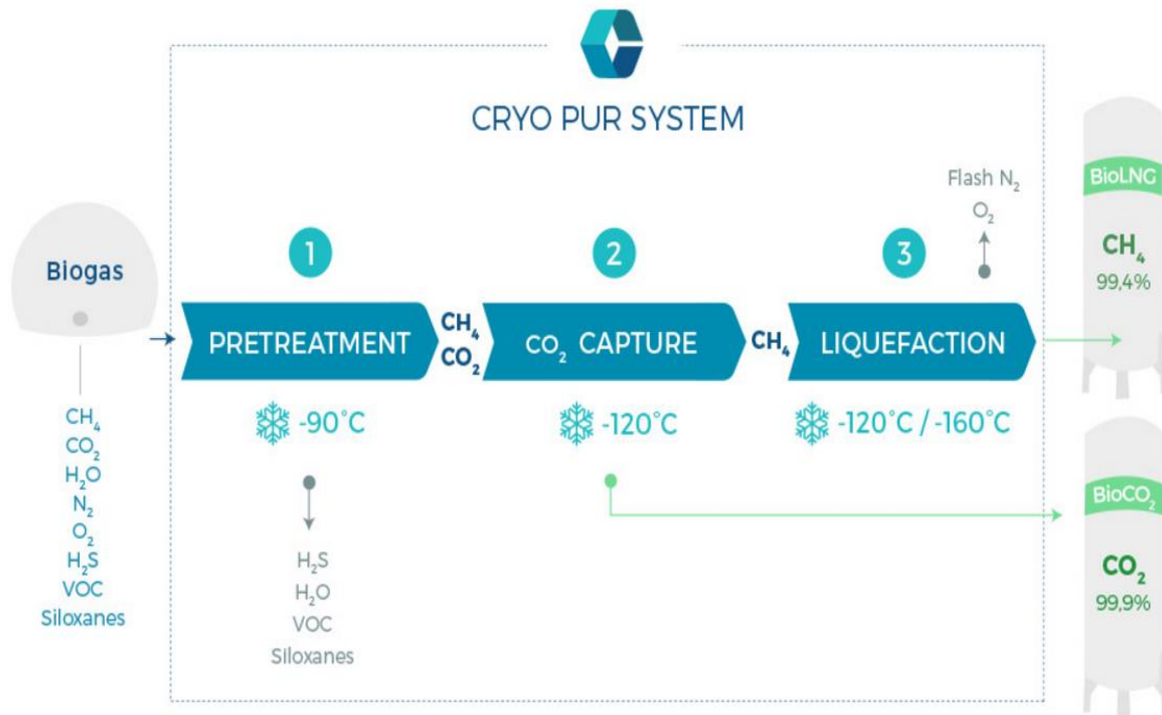


Fig. 30: Schematic representation of Cryo Pur[®] system [9]

This chapter is divided into two sections: The first part consists of a general presentation of the pilot plant. The second part is dedicated to the operating principle of the subsystems.

4.2. General presentation of the BioGNVAL pilot plant

Fig. 31 shows the general operating principle of the BioGNVAL demonstrator. It consists of 7 subsystems:

- Chiller.
- Desulfurization subsystem, either by chemical absorption in a structured packing column using sodium hydroxide, or by adsorption onto activated carbon in a fixed bed respectively.
- Biogas dehumidification and siloxanes icing subsystem.
- Subsystem for the capture of carbon dioxide.
- Biogas liquefaction subsystem.
- Biogas treatment line with flaring output.

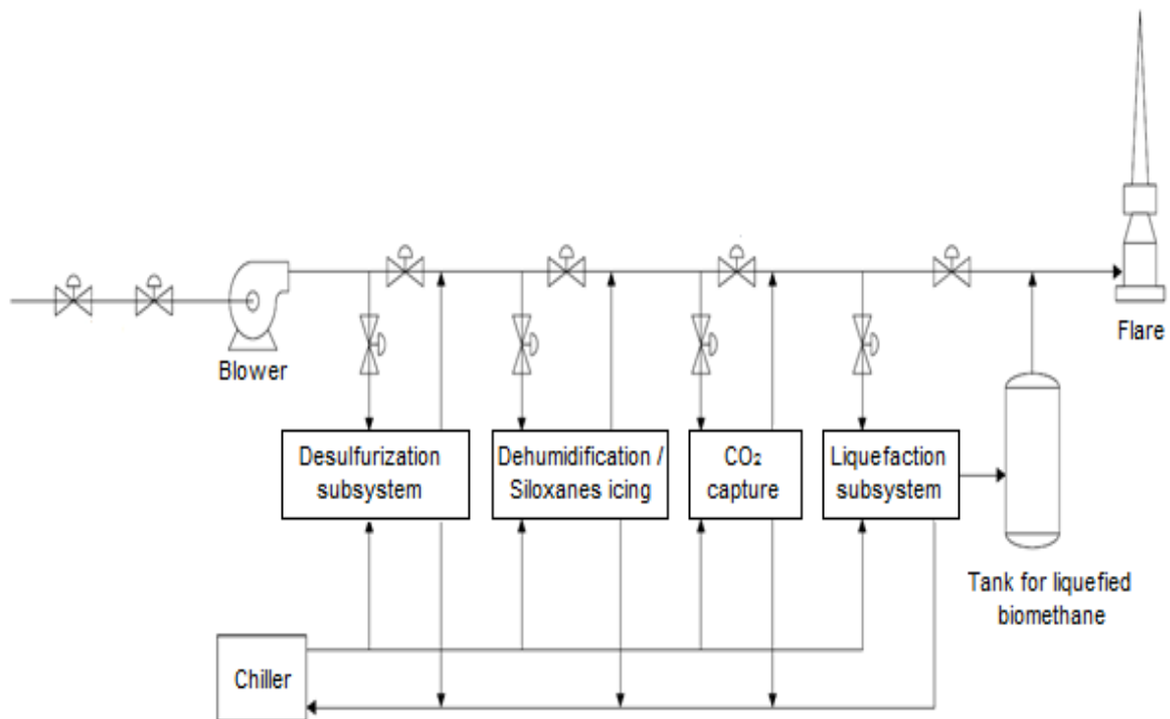


Fig. 31: Simplified flowsheet of the BioGNVAL pilot plant [9]

The raw biogas, whose conditions and composition are presented in Table 25, comes from the anaerobic digester through the line of biogas. The latter distributes the biogas in all sub-systems, starting with the desulfurization subsystem, until the liquefaction subsystem.

Table 25. Conditions and composition of the raw biogas treated by BioGNVAL pilot plant

Conditions	Unit	Value
Pressure	[mbara]	1000 – 1020
Temperature	[°C]	< 37
Volume flow rate	[Nm ³ /h]	85
Mass flow rate	[kg/h]	94
Composition	Unit	Value
CH ₄	[%]	64 ± 2
CO ₂	[%]	34 ± 2
H ₂ O	[%]	< 5
N ₂	[%]	< 2
O ₂	[%]	< 1
H₂S	ppm	< 100 (Average ≈ 30)
Siloxanes	mg/Nm ³	< 15

Conditions at the outputs of each sub-system in terms of composition and temperature are defined. If one of the conditions is not met, the biogas is routed through the biogas treatment line to the flare. These conditions are shown in Table 26. Once the full treatment is performed, the liquefied biogas is stored in a mobile container presented in Fig. 31.

Table 26. conditions of passage from one subsystem to another [9]

Passage	Conditions	
	Temperature [°C]	Composition
Dehumidification and siloxanes icing subsystem → Carbon dioxide capture subsystem	< -85	H ₂ S < 1 ppm
Carbon dioxide capture subsystem → Liquefaction subsystem	< -115	CO ₂ < 2.5 %

4.3. The operating principle of the BioGNVAL subsystems

4.3.1. Desulfurization subsystems

The operating principle of reactive absorption is simple. The system mainly comprises a structured packed column, a water circuit and two heat exchangers, used to cool biogas and the liquid phase. Fig. 32 shows the apparatus setup for the desulfurization process.

The biogas is saturated with water vapor at the input of the pilot plant. It firstly passes through a heat exchanger (Green line) to cool the gas phase and condense a portion of the water vapor contained therein.

A phase separator recovers the condensed water vapor and sends it to a drainage tank.

Thereafter, the biogas enters the bottom of the absorption column where it is contacted in a counter-current with the washing water sprayed from the top (Violet line). This water is neutralized by an aqueous solution of sodium hydroxide. The injected quantity is controlled by a pH meter placed on the tank TK-230-04. This tank showed in Fig. 32 provides the column with the liquid phase.

The liquid phase is recirculated by a pump and is cooled by an exchanger to a temperature slightly higher than 2 °C to prevent freezing.

Biogas finally exits from the top of the column at a temperature of about 5 °C. The cooling duty is provided by the liquid phase on the surface of the packing in direct contact within the column. High efficient mass and heat transfer between the liquid and the biogas are achieved thanks to the packing surface.

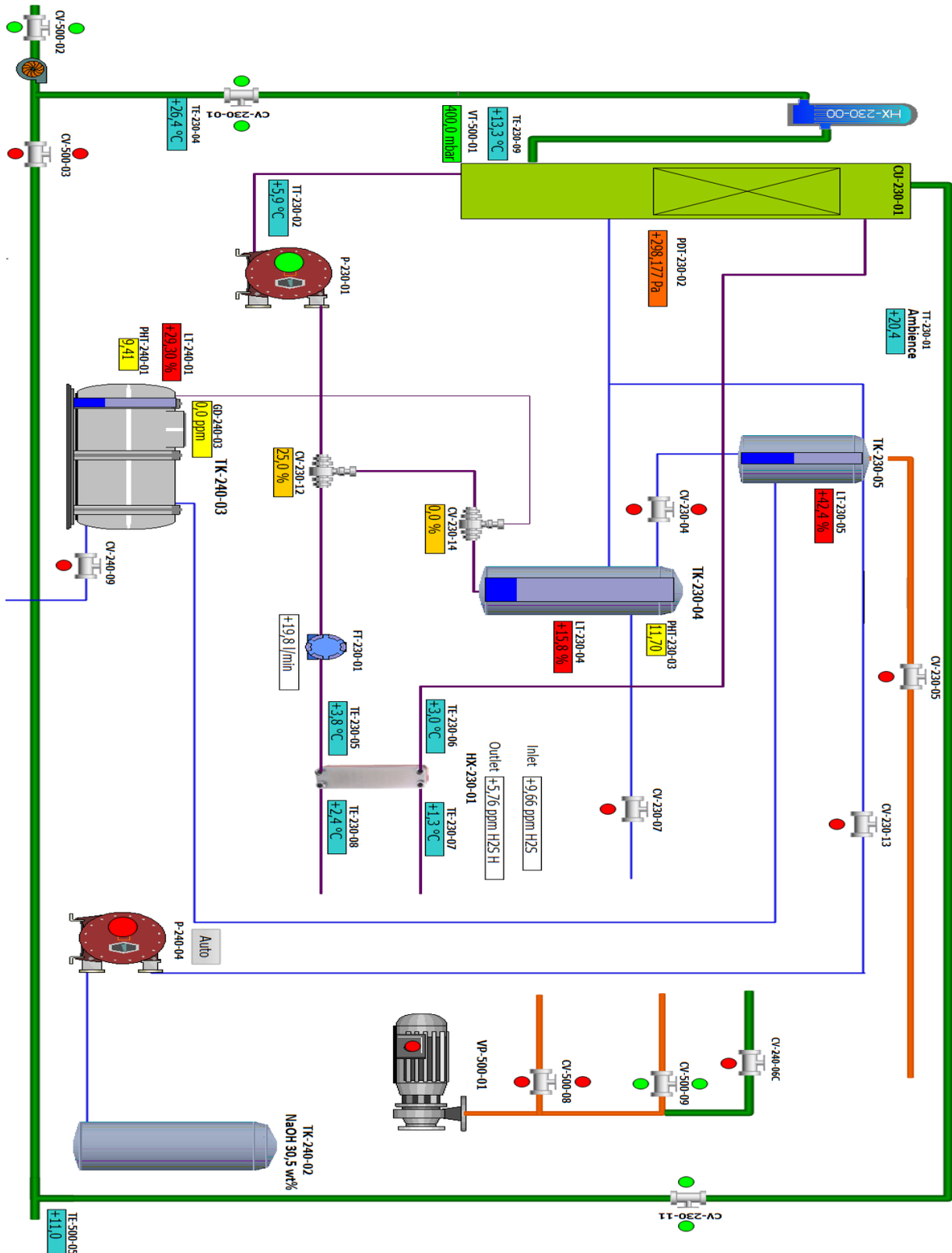


Fig. 32: Schematic diagram of the absorption subsystem for elimination of hydrogen sulfide [9]

Equipment: CV (Control Valve) ; HX (Heat Exchanger) ; CU (Condensing Unit) ; TK (Tank) ; P (Pump) ; VP (Vaccum Pump) ;

Instrumentation: FT (Flow Transmitter) ; TT (Temperature Transmitter) ; TE (Temperature Element) ; PT (Pressure Transmitter) ; PDT (Pressure Difference Transmitter) ; PHT (pH Analyzer Transmitter) ; LT (Level Transmitter) ; GD (Gas Detector)

In order to test the adsorption technology, the absorption process could be bypassed. Therefore, the biogas pass through two fixed bed adsorption columns placed in series, as seen in Fig. 33.

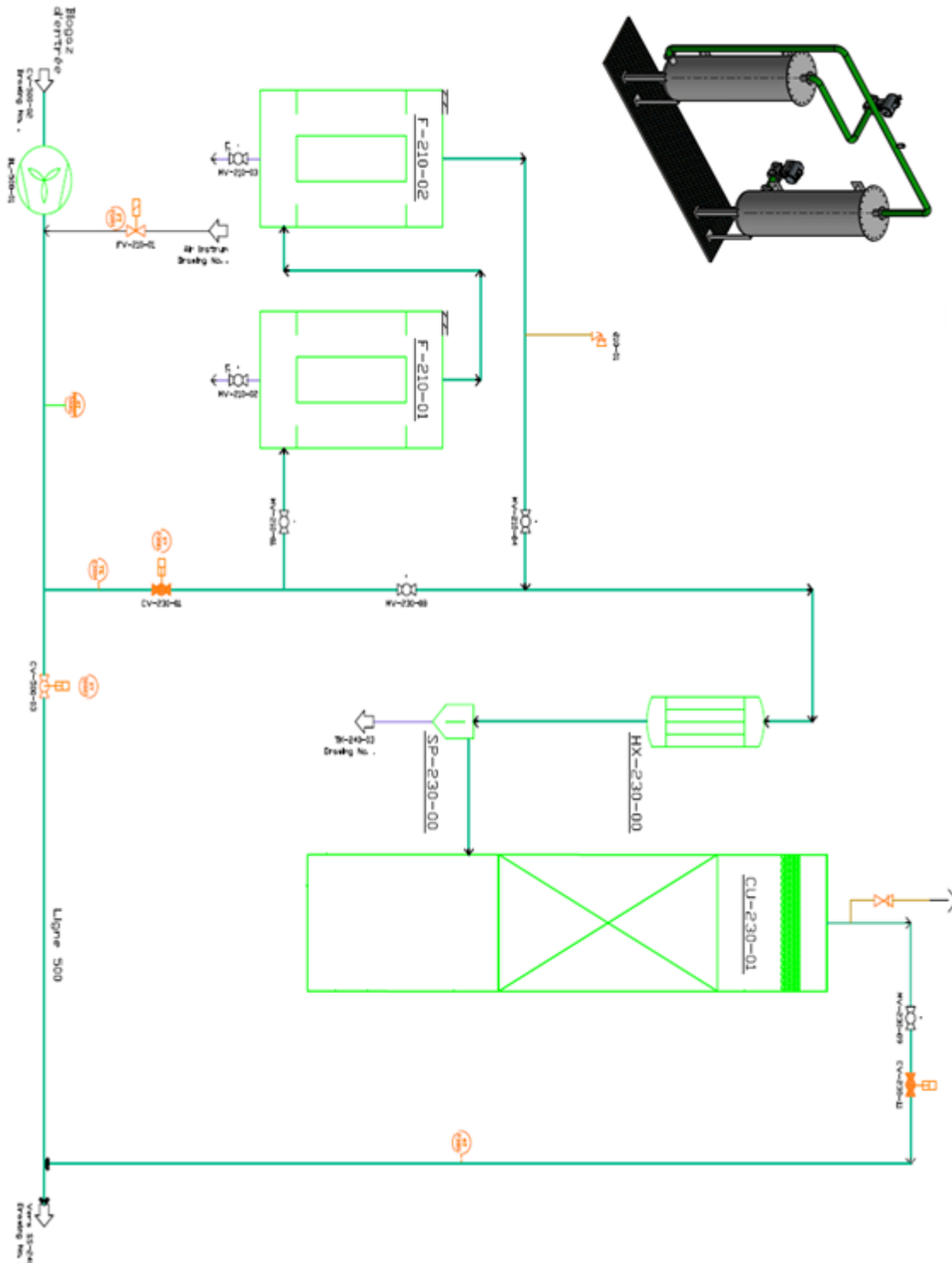


Fig. 33: Piping and instrumentation diagram of adsorption subsystem for the removal of hydrogen sulfide [9]

Equipment: CV (Control Valve) ; FV (Flow Valve) ; MV (Manual Valve) ; PSV (Pressure Safety Valve) ; BL (Blower) ; F (Filter) ; HX (Heat Exchanger) ; SP (Separator) ; CU (Condensing Unit) ; TK (Tank).
Instrumentation: FT (Flow Transmitter) ; TE (Temperature Element) ; AT (Analyzer Transmitter) ; XY (Limit Switch)

These two columns packed with impregnated activated carbon, provide a continuous treatment of hydrogen sulphide. To further improve the functioning of activated carbon, a small amount of oxygen is added to oxidize the hydrogen sulfide, which will make larger molecules and allow to block them into the pores.

4.3.2. Dehumidification and siloxanes icing subsystem

The biogas leaves the desulfurization subsystem at 5 °C and with a H₂S content lower than 1 ppm. Dehumidification and cooling the biogas continue until -40 °C. At this temperature, only 125 ppm of water vapor remains in the biogas and the heaviest Siloxanes are removed.

Biogas cooling continues until a temperature of -87 ° C. This eliminates the siloxanes, as well as water vapor, whose content is reduced to less than 1 ppm at the outlet of the subsystem.

To ensure a continuous operation and to prevent the accumulation of ice that could block the passage of biogas, the chillers used contain two evaporators placed in parallel to alternate operation in frosting and defrosting mode.

4.3.3. Carbon dioxide capture subsystem

After having been purified of hydrogen sulphide, water vapor and siloxane, the biogas is fed into the CO₂ capture subsystem to be upgraded. The biogas is now composed of methane and carbon dioxide at atmospheric pressure, that's to say a carbon dioxide partial pressure below its triple pressure. After cooling the biogas to -120 ° C in this subsystem, the carbon dioxide thus undergoes the phenomenon of antisublimation, which means that it is transformed directly from its gaseous phase into a solid phase frosted on the cold surface of the heat exchanger as seen in Fig. 34.

To allow continuous operation of the upgrading system, two heat exchangers are used. When the first is frosting the carbon dioxide, the second operates in defrost mode. Thus carbon dioxide is recovered in the liquid phase and then stored in a cryogenic vessel. This strategy allows recovery of CO₂ at a very high purity level (99.995 %) which could be used for industrial and food applications.

4.3.4. Biogas liquefaction subsystem

After purification and upgrading of biogas, the biomethane is sent to the liquefaction subsystem which is composed of a compression unit, a cooling and liquefaction unit and a storage unit for the liquefied biomethane.

At the outlet of the upgrading unit, a gas analyser ensures that the biomethane produced is composed of less than 2.5 % of residual carbon dioxide. Once the required biomethane quality reached, it is supplied to the liquefaction subsystem to be compressed and liquefied. Compression of biomethane increases its liquid-vapor saturation temperature, which means its liquefaction at a high enough temperature level reducing electrical consumption of the refrigeration machine.

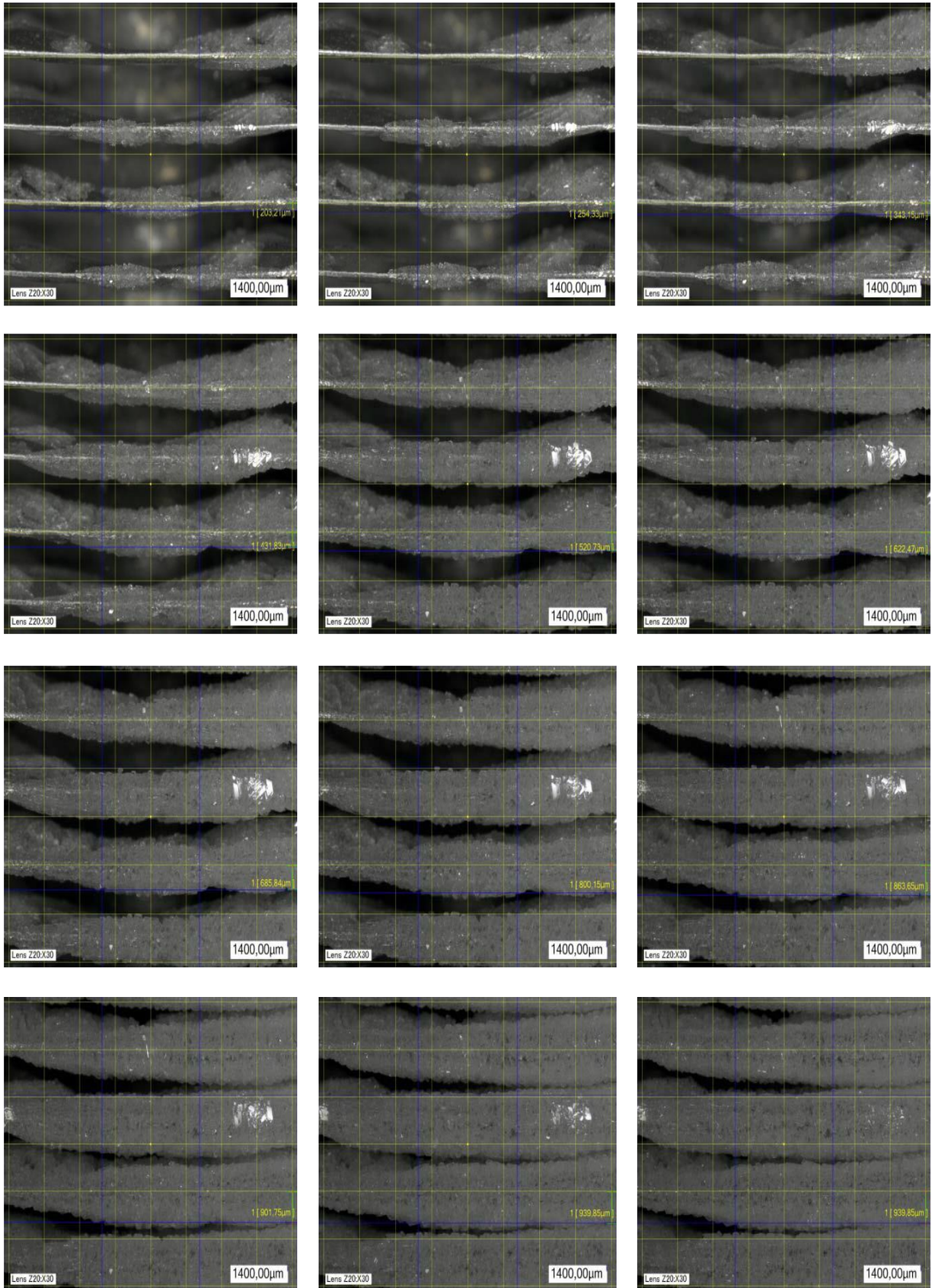


Fig. 34: Carbon dioxide antisublimation [9]

4.4. Experimental results concerning the removal of hydrogen sulfide by chemical absorption using sodium hydroxide

The main objectives of the experiments on the absorption column is to maintain the concentration of hydrogen sulfide between 0 and 1 ppm at the outlet of the column throughout the testing period, for a content at the entrance equal to 20 ppm. The biogas temperature at the outlet of the column should be slightly greater than 2 °C whatever its temperature at the inlet.

These objectives have been reached on the demonstrator. The hydrogen sulfide content at the outlet of the absorption column was maintained between 0 and 1 ppm as seen in Fig. 35. The biogas temperature at the outlet of the packing column was maintained between 5 and 6 °C even when the biogas temperature at the inlet reaches 35 °C as shown in Fig. 36.

The concentrations measurement are made using a biogas analyser, model “Gas 3200 R Biogas” bought from “Gas Engineering and Instrumentation Technologies Europe” GEIT® company. The measurements of H₂S concentration are made with 3-electrodes electrochemical cell designed for biogas applications with several measuring range: from 0 – 50 ppm to 0 – 9999 ppm. The sensitivities of measurement tools used in the experiments are shown in Table 27.

Table 27. Sensitivities of measurement tools

Properties to measure	Measurement tools	Accuracy
Pressure drop in the packing column	Differential Pressure Transmitter	± 0.065 % Full Scale (FS)
Pressure	Pressure sensor	< 0.5 % FS
H ₂ S concentration	Biogas analyser	± 3 % FS
Flow rates	Flowmeter	< 1 % FS

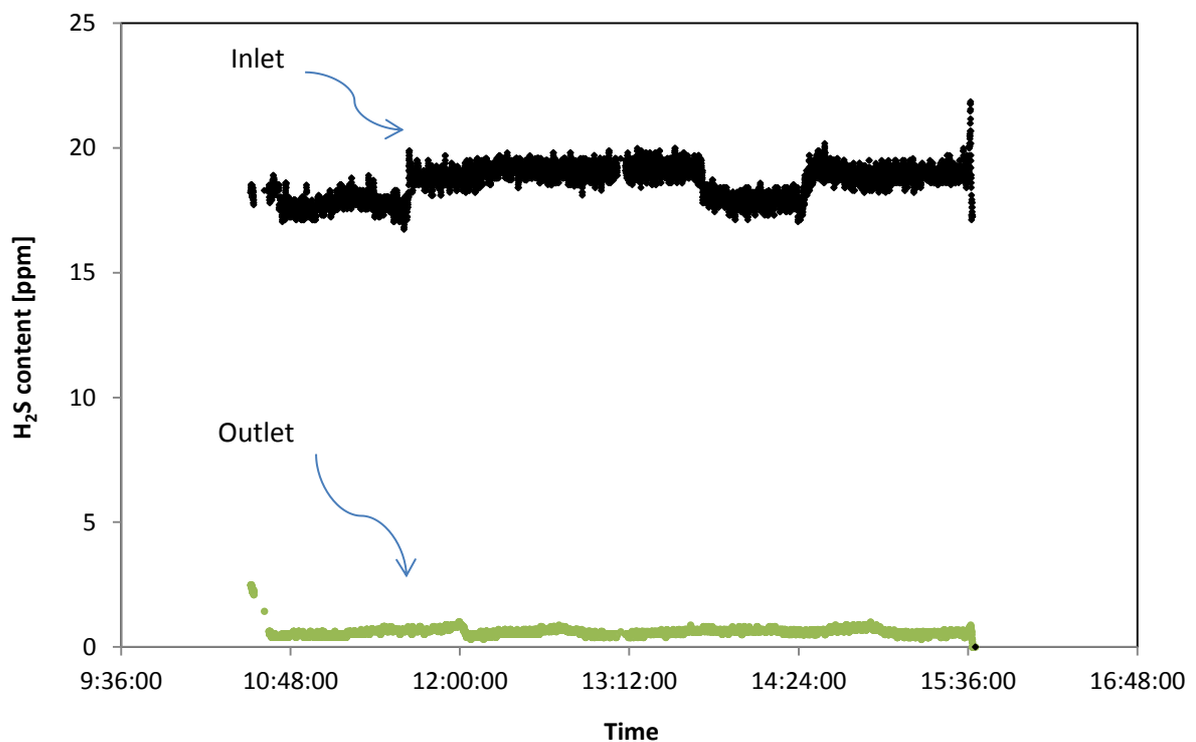


Fig. 35: Variation of the H₂S content at the inlet and at the outlet of the absorption column

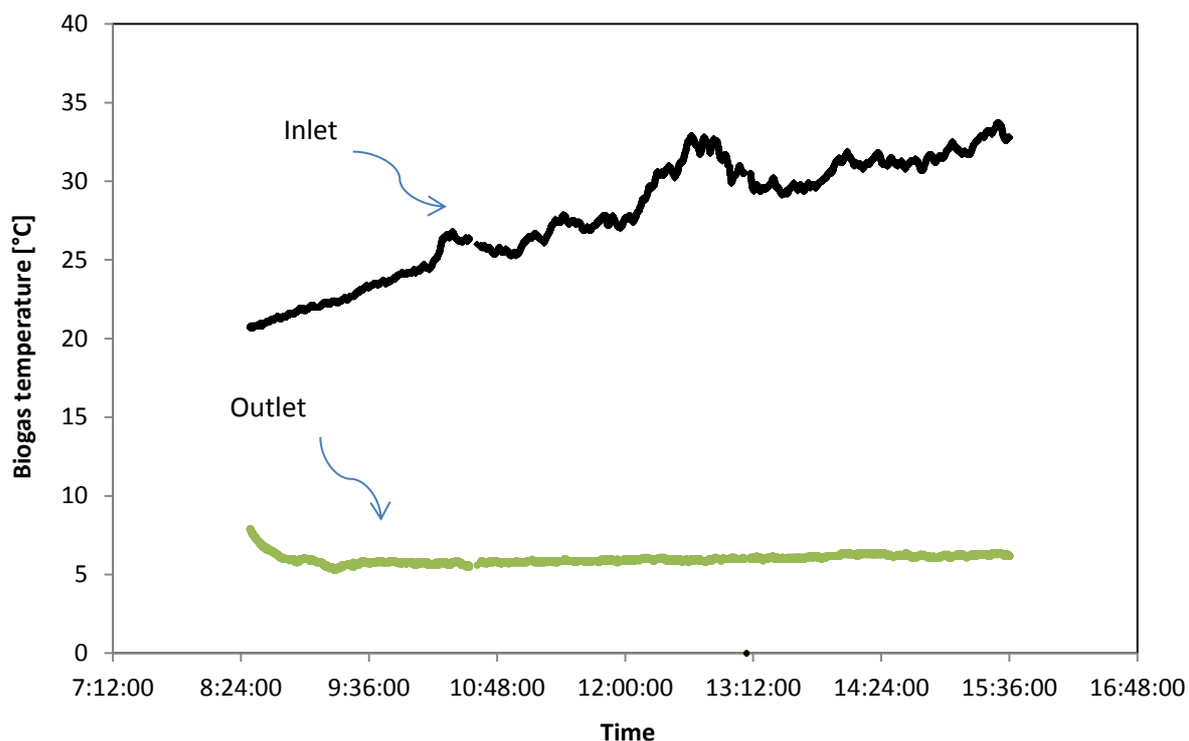


Fig. 36: Variation of the biogas temperature at the inlet of the demonstrator and at the outlet of the absorption column

Sodium hydroxide was used as chemical solvent for the removal of hydrogen sulfide. The weight percentage of NaOH in water is equal to 30.5 wt%.

Averaged over all experiments, sodium hydroxide consumption was assessed at 6 l/h (solution containing 30.5 wt% of NaOH and 69.5 wt% of H₂O). This consumption was calculated with an indicator allowing the NaOH level measurement in the tank TK-240-02 (See Fig. 32).

This consumption is very excessive relative to the theoretical consumption of 1.35 l/hr of commercial caustic soda.

This overconsumption is explained by desorption of hydrogen sulfide, because the reaction between hydrogen sulphide and sodium hydroxide is reversible (R.11). It leads to the formation of sodium sulfide (Na₂S) that is unstable in water.

Fig. 37 shows the desorption phenomenon observed during experiments. Indeed, stopping the injection of sodium hydroxide at 13:45:00 (See Fig. 37) shows that the hydrogen sulfide content at the outlet of the column is greater than its content at the inlet. This explains the overconsumption of the solvent.

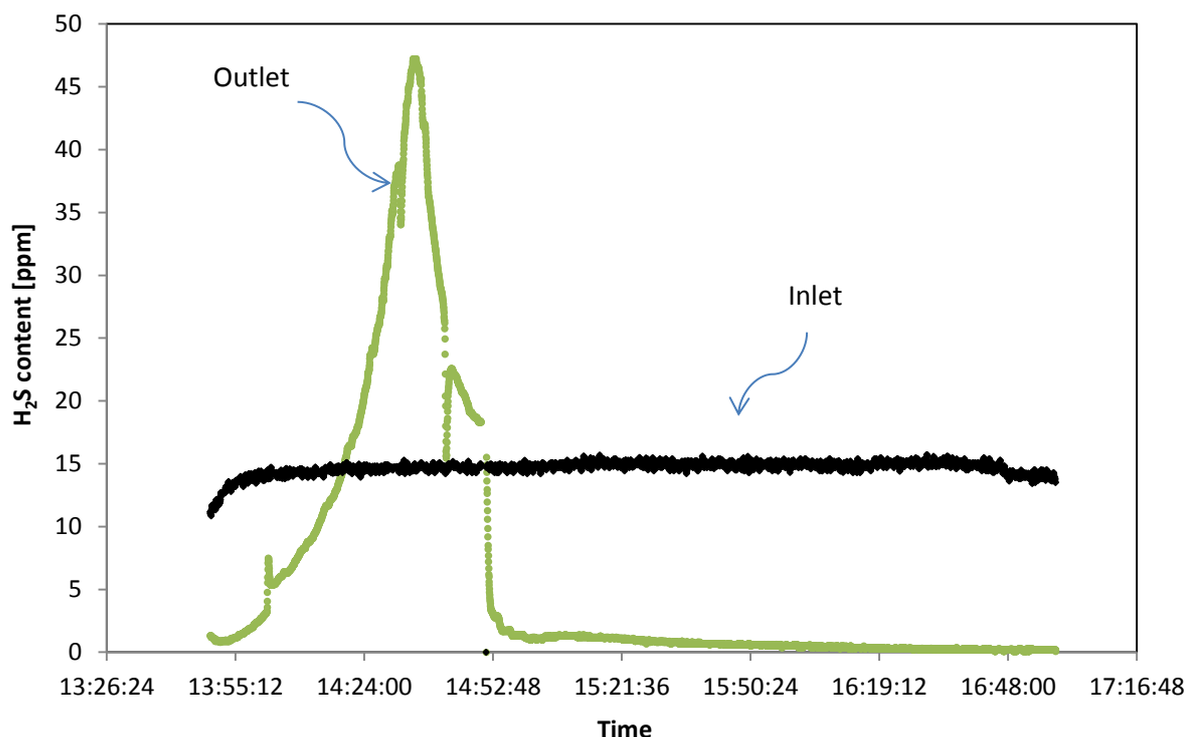


Fig. 37: Observation of the desorption phenomenon of H₂S

To overcome this problem, the injection of sodium hypochlorite (NaOCl) has been proposed to prevent the regeneration of hydrogen sulfide, and to obtain soluble by-products in water (Na₂SO₄ and NaCl) as seen in Reaction 2.



The injections of sodium hydroxide and sodium hypochlorite will respectively be controlled by the pH and conductivity measurements. This solution will reduce the consumption of the solvents.

4.5. Conclusion

This section has presented two different processes (absorption and adsorption) used for the removal of H₂S in order to reduce its content to less than 1 ppm throughout the operation.

The other undesirable components present in the biogas such as H₂O, siloxanes and CO₂ are captured through cryo-condensation, freezing each component.

The CO₂ is retrieved in liquid form at a high level of purity enabling revalorization. Once purified, the biomethane fulfills the characteristics necessary to be used as fuel for heavy goods vehicles (HGV).

To allow the absorption column to operate at full capacity, an hydrodynamic study is necessary. This study will optimise the flow rates involved and ensure a better mass transfer in the packing column, with a lower consumption of the liquid phase and a lower pressure drop. This study which will be presented in the next chapter, is also of great interest for the design of packed columns.

Chapter 5: Hydrodynamic study of gas-liquid countercurrent flows in structured packing columns

Résumé :

Ce chapitre compare trois modèles existants utilisés pour la prédiction des paramètres hydrodynamiques dans des colonnes à garnissage structuré.

Ces modèles sont utilisés pour évaluer la perte de pression, la rétention liquide, l'aire interfaciale effective, les coefficients de transfert de masse et les points de transition. Les résultats de ces modèles sont comparés à des données expérimentales afin de choisir celui avec le meilleur ajustement.

Les comparaisons ont été effectuées en utilisant deux systèmes : Air - Eau et Air - Kerosol 200 et un garnissage structuré de type Flexipac 350Y.

Le modèle choisi est basé sur des corrélations semi-empiriques contenant des constantes et des exposants définis selon des mesures expérimentales. Pour rendre le modèle plus représentatif du système d'intérêt (biogaz contenant du H₂S / solution aqueuse d'hydroxyde de sodium), ces constantes ont été modifiées et certains exposants ont été ajustés en fonction de la vitesse superficielle du liquide et la densité.

Une fois le modèle modifié, les résultats de la perte de pression ont été comparés aux données expérimentales obtenues sur le démonstrateur BioGNVAL. Les résultats obtenus sont en bon accord mais il est judicieux de noter que ce modèle peut perdre de sa précision en variant les applications.

Par conséquent, ce modèle est idéal pour prédire avec précision les trois régions opérationnelles d'une colonne à garnissage structuré à petite échelle utilisée pour des applications de biogaz ou de gaz naturel.

5.1. Introduction

Today, in modern absorption columns, structured packings are widely used, thanks to their higher capacity and lower pressure drop compared to random packings.

Structured packings were used for the first time in 1950 [91]. They are in continuous development to expand their use and improve their efficiency. They provide a large surface area for the liquid and gas phases to be in direct contact within the column. High efficient mass transfer between the two phases is achieved thanks to the packing surface.

This work compares three existing models used for the prediction of hydrodynamic parameters in structured packing columns. These models are used to evaluate pressure drop, liquid holdup, effective interfacial area, mass transfer coefficients and transition points. The results obtained with these models are compared to experimental data in order to choose the one with the best fit. Comparisons were made using Flexipac 350Y structured packing and two systems: Air – Water and Air – Kerosol 200. The model chosen is based on semi-empirical correlations using constants and exponents defined according to experimental measurements. To adapt the model to biogas application and to make it more representative of the system of interest, these constants were optimized and some exponents have been adjusted. Once the model modified, the results of pressure drop were compared to data from BioGNVAL pilot plant.

5.2. Theoretical principles

In a packed column, hydrodynamics and mass transfer processes occur simultaneously. They are correlated and the link parameter is liquid holdup h_L defined as the volume of the liquid per unit volume of the column. Equation (17) defined by Chan and Fair [92] for sieve trays illustrated the relation between the two processes.

$$k_V a_e = \frac{316 D_V^{0,5} (1030 f + 867 f^2)}{h_L^{0,5}} \quad (17)$$

Regarding the hydrodynamic analysis, increasing the velocity of liquid and gas, results in an increase of the liquid holdup and the thickness of the liquid film which leads to an increase in pressure drop.

About mass transfer analysis, increasing liquid holdup causes the enlargement of the interfacial area leading to higher mass transfer rates.

The curve which represents the evolution of the pressure drop or the liquid holdup as a function of the gas capacity factor F_c is divided by two points (loading and flooding points) into three operating regions as seen in Fig. 38.

The liquid flow rate is not influenced by the counter-current flow of the gas in the preloading region, as can be seen in Fig. 38.b which presents the evolution of the liquid holdup as a function of the gas capacity factor defined by Equation (18).

$$F_c = u_V \times \sqrt{\rho_V} \quad (18)$$

The loading point represented by the line AA in Fig. 38 is reached when the slope of the liquid holdup curve starts to increase, or when the wet pressure drop curve starts to deviate from the pressure drop in a dry column.

The flooding point is represented by the line BB in Fig. 38. It is the point where the slope of pressure drop and liquid holdup curves tends toward infinity.

Therefore, it is necessary to predict accurately the transition points because they characterize the capacity of a packing column. According to Paquet [93], under-predicting the flooding point will prevent the column to operate at its optimal conditions, and its capacity could be very low. However, over-predicting the flooding point may lead to higher pressure drop which could be problematic.

Because of the lack of predictive models, and because of the imprecision of existing ones to accurately predict the hydrodynamic parameters for some specific applications such as biogas purification, most distillation and packing columns are still being designed based on experimental data from a pilot plant [94].

The objective of this work is to find a model adapted for the representation of the experimental results obtained on the BioGNVAL pilot plant. To this aim, three literature models for the hydrodynamics in structured packing columns have been compared: Billet and Schultes model [95], SRP model [96] and Delft model [97]. These models have been developed on dimensionless analysis and experimental data obtained using a distillation column. The two first models are implemented in the process simulator Aspen Plus®. The three models are described in detail in the following section.

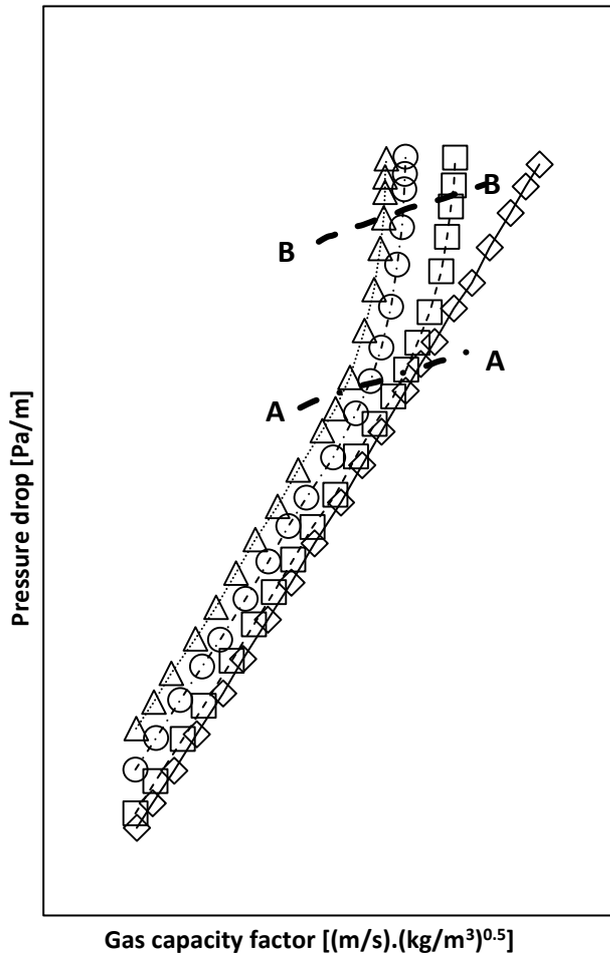


Fig. 38.a: Pressure drop evolution in a packing column

(—Δ—) Dry bed ; (- -□- -) 1 ; (- ·○· ·) 2 ; (··Δ··) 3
 (AA) Loading point ; (BB) Flooding point
 Liquid load: 1 < 2 < 3

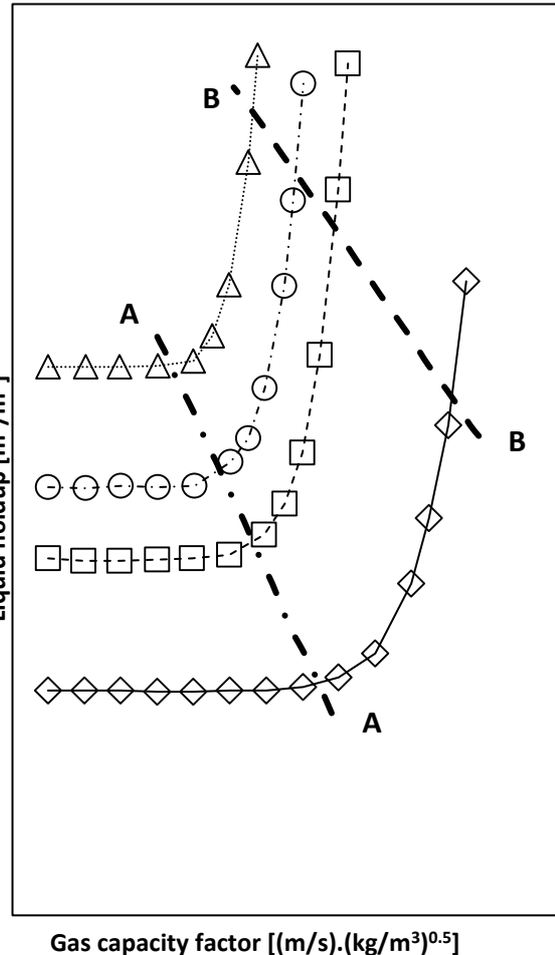


Fig. 38.b: Liquid holdup evolution in a packing column

(—Δ—) 1 ; (- -□- -) 2 ; (- ·○· ·) 3 ; (··Δ··) 4
 (AA) Loading point ; (BB) Flooding point
 Liquid load: 1 < 2 < 3 < 4

5.2.1. Billet and Schultes model

The Billet and Schultes model [95] was at the base founded for random packings. Then, it was extended to cover structured packings. Based on semi-empirical correlations, this model assumes that the packing void fraction is represented by vertical tubes where the liquid is sprayed from the top as a film that meets the gas flow in a counter-current configuration. The angle between the corrugations of the packing is not taken into account by the Billet and Schultes model.

As reported by Paquet [93], the main disadvantage of this model is that it requires six specific constants for each type of packing. The ones needed for Flexipac 350Y and for some other types of packing, are presented in Table 28.

Table 28. Constants for the Billet and Schultes model [98]

Manufacture	Material	Description	a [m ² .m ⁻³]	ε	C _{ip}	C _{Fl}	C _h	C _p	C _L	C _V
Flexipac	Metal	350Y	350	0.985	3.157					
Mellapak	Metal	250Y	250	0.970	3.157	2.464	0.554	0.292	-	-
Ralu pak	Metal	YC-250	250	0.945	3.178	2.558	-	0.191	1.334	0.385
Gempack	Metal	A2T-304	202	0.977	2.986	2.099	0.678	0.344	-	-
Euroform	Plastic	PN-110	110	0.936	3.075	1.975	0.511	0.250	0.973	0.167
Impulse packing	Metal	250	250	0.975	2.610	1.996	0.431	0.262	0.983	0.270
	Ceramic	100	91.4	0.838	2.664	1.655	1.900	0.417	1.317	0.327
Montz packing	Metal	B1-200	200	0.979	3.116	2.339	0.547	0.355	0.971	0.390
		B2-300	300	0.930	3.098	2.464	0.482	0.295	1.165	0.422
	Plastic	C1-200	200	0.954	-	-	-	0.453	1.006	0.412
		C2-200	200	0.900	2.653	1.973	-	0.481	0.739	-

Correlations (19), (20) and (21) illustrated in Table 29 are used by Billet and Schultes to calculate the effective interfacial area respectively at loading point, in the loading region and at the flooding point.

Table 29. Effective interfacial area in packing columns using Billet and Schultes model [98]

Parameter	Correlation	
Effective interfacial area at loading point	$\left(\frac{a_e}{a}\right)_{lp} = 1.5 (a d_h)^{-0.5} \left(\frac{u_L d_h}{\nu_L}\right)^{-0.2} \left(\frac{u_L^2 \rho_L d_h}{\sigma_L}\right)^{0.75} \left(\frac{u_L^2}{g d_h}\right)^{-0.45}$ $= 1.5 (a d_h)^{-0.5} Re_L^{-0.2} We_L^{0.75} Fr_L^{-0.45}$	(19)
Effective interfacial area in loading region	$\left(\frac{a_e}{a}\right)_{u_V > u_{V,lp}} = \left(\frac{a_e}{a}\right)_{lp} + \left[\left(\frac{a_e}{a}\right)_{Fl} - \left(\frac{a_e}{a}\right)_{lp} \right] \left(\frac{u_{V,lp}}{u_{V,Fl}}\right)^{13}$	(20)
Effective interfacial area at flooding point	$\left(\frac{a_e}{a}\right)_{Fl} = 7 \left(\frac{\sigma_L}{\sigma_W}\right)^{0.56} \left(\frac{a_e}{a}\right)_{lp}$	(21)

The Billet and Schultes model is composed of several correlations that describe liquid holdup and pressure drop in the preloading, loading and flooding regions. Velocities and liquid holdup at loading and flooding points are calculated using the equations listed in Table 30.

Table 30. Liquid holdup and velocities at loading and flooding point [99]

Parameter	Correlation	
Gas velocity at loading point	$u_{V,lp} = \sqrt{\frac{g}{\psi_{lp}}} \left(\frac{\varepsilon}{a^6} - a^{0.5} \left(\frac{12 \mu_L}{g \rho_L} u_{L,lp} \right)^{\frac{1}{3}} \right) \left(\frac{12 \mu_L}{g \rho_L} u_{L,lp} \right)^{\frac{1}{6}} \sqrt{\frac{\rho_L}{\rho_V}}$	(22)
Liquid velocity at loading point	$u_{L,lp} = \frac{\rho_V}{\rho_L} \frac{L}{V} u_{V,lp}$	(23)
Gas velocity at flooding point	$u_{V,Fl} = \sqrt{\frac{2g}{\psi_{Fl}}} \frac{(\varepsilon - h_{L,Fl})^{\frac{3}{2}}}{\varepsilon^{0.5}} - \sqrt{\frac{h_{L,Fl}}{a}} \sqrt{\frac{\rho_L}{\rho_V}}$	(24)
Resistance coefficient at loading point	$\psi_{lp} = \frac{g}{C_{lp}^2 \left(\frac{L}{V} \sqrt{\frac{\rho_V}{\rho_L}} \left(\frac{\mu_L}{\mu_V} \right)^{0.4} \right)^{2n_{lp}}}$	(25)
Resistance coefficient at flooding point	$\psi_{Fl} = \frac{g}{C_{Fl}^2 \left(\frac{L}{V} \sqrt{\frac{\rho_V}{\rho_L}} \left(\frac{\mu_L}{\mu_V} \right)^{0.2} \right)^{2n_{Fl}}}$	(26)
Packing specific constant at loading point	$\frac{L}{V} \sqrt{\frac{\rho_V}{\rho_L}} \leq 0.4 \rightarrow n_{lp} = -0.326 \rightarrow C_{lp} = C_{lp}$	(27)
	$\frac{L}{V} \sqrt{\frac{\rho_V}{\rho_L}} > 0.4 \rightarrow n_{lp} = -0.723 \rightarrow C_{lp} = 0.695 C_{lp} \left(\frac{\mu_L}{\mu_V} \right)^{0.1588}$	
Packing specific constant at flooding point	$\frac{L}{V} \sqrt{\frac{\rho_V}{\rho_L}} \leq 0.4 \rightarrow n_{Fl} = -0.194 \rightarrow C_{Fl} = C_{Fl}$	(28)
	$\frac{L}{V} \sqrt{\frac{\rho_V}{\rho_L}} > 0.4 \rightarrow n_{Fl} = -0.708 \rightarrow C_{Fl} = 0.6244 C_{Fl} \left(\frac{\mu_L}{\mu_V} \right)^{0.1028}$	
Liquid holdup at the loading point	$h_{L,lp} = \left(\frac{12 a^2 \mu_L u_L}{\rho_L g} \right)^{\frac{1}{3}} \left(\frac{a_h}{a} \right)^{\frac{2}{3}}$	(29)
Liquid holdup at the flooding point	$h_{L,Fl}^3 (3h_{L,Fl} - \varepsilon) = \frac{6}{g} a^2 \varepsilon \frac{\mu_L}{\rho_L} \frac{L}{V} \frac{\rho_V}{\rho_L} u_{V,Fl}$	(30)

Table 31 presents the correlations used by Billet and Schultes to calculate the liquid holdup in the loading region. This property depends on the liquid holdup in the preloading region and at the flooding point. The first one is theoretically derived from a force balance, while the second is purely empirical. The liquid holdup in the preloading region does not depend on the gas properties. It is only a function of the liquid properties and its velocity, as seen in Equation (31).

As stated in the thesis of Paquet [93], the hydraulic area of the packing accounts for the surfaces that were not completely wetted by the liquid flow.

Table 31. Liquid holdup in preloading and loading regions [98]

Parameter	Correlation	
Liquid holdup in preloading region	$h_{L,pl} = \left(12 \frac{\mu_L a^2 u_L}{\rho_L g} \right)^{\frac{1}{3}} \left(\frac{a_h}{a} \right)^{\frac{2}{3}}$	(31)
Hydraulic area of the packing	$\frac{a_h}{a} = C_h Re_L^{0.15} Fr_L^{0.1} \quad \text{For } Re_L < 5$	(32)
	$\frac{a_h}{a} = 0.85 C_h Re_L^{0.25} Fr_L^{0.1} \quad \text{For } Re_L \geq 5$	
Liquid Reynolds number	$Re_L = \frac{u_L \rho_L}{a \mu_L}$	(33)
Liquid Froude number	$Fr_L = \frac{u_L^2 a}{g}$	(34)
Liquid holdup at flooding point	$h_{L,Fl} = 2.2 h_{L,pl}$	(35)
Liquid holdup in loading region	$h_L = h_{L,pl} + (h_{L,Fl} - h_{L,pl}) \left(\frac{u_V}{u_{V,Fl}} \right)^{13}$	(36)

The equations used to calculate pressure drop are listed in Table 32.

Table 32. Pressure drop in packing columns using Billet and Schultes model [98]

Parameter	Correlation	
Dry pressure drop	$\left(\frac{dP}{dz} \right)_d = \psi_0 \frac{a}{\varepsilon^3} \frac{F_c^2}{2} \frac{1}{K}$	(37)
Resistance coefficient	$\psi_0 = C_p \left(\frac{64}{Re_V} + \frac{1.8}{Re_V^{0.08}} \right)$	(38)
Gas capacity factor	$F_c = u_V \sqrt{\rho_V}$	(39)
Wall factor	$\frac{1}{K} = 1 + \frac{2}{3} \frac{1}{(1 - \varepsilon)} \frac{d_p}{d}$	(40)
Particle diameter	$d_p = 6 \frac{1 - \varepsilon}{a}$	(41)
Gas Reynolds number	$Re_V = \frac{u_V d_p \rho_V}{(1 - \varepsilon) \mu_V} K$	(42)
Wet pressure drop	$\frac{dP}{dz} = \psi_L \frac{f_w a}{(\varepsilon - h_L)^3} \frac{F_c^2}{2} \frac{1}{K}$	(43)
Resistance factor	$\psi'_L = \psi_L f_w = C_p f_s \left(\frac{64}{Re_V} + \frac{1.8}{Re_V^{0.08}} \right) \left(\frac{\varepsilon - h_L}{\varepsilon} \right)^{(1.5)}$	(44)
f_s	$f(s) = \left(\frac{h_L}{h_{L,lp}} \right)^{0.3} \exp \left(\frac{Re_L}{200} \right)$	(45)

The expression of dry pressure drop is obtained by applying a force balance. The wall factor K is used to take into account the free spaces more available at the wall. The constant C_p used to calculate the resistance coefficient ψ_0 characterizes the geometry of the packing.

For the wetted packing column, Equation (43) used to calculate pressure drop replaces the void fraction (ϵ) by an effective void fraction ($\epsilon - h_L$) which depends on liquid holdup, reducing the volume available for the gas flow. This equation introduces a wetting factor f_w to account for any change in the surface of the packing caused by the wetting action [93].

5.2.2. SRP model

The SRP (Separations Research Program) model [96] was developed at the University of Texas [100]. The latest version of this model was published in the work of Fair et al. in 2000 [96]. According to Paquet [93], the SRP model considers the void fraction as a series of wet columns where the gas flow passes through. Unlike the Billet and Schultes model, the geometry depends on the angle and dimensions of corrugations.

To calculate liquid holdup and effective interfacial area, the SRP model uses a correction factor that takes into account the packing surface that is not completely wetted by the liquid flow.

The prediction of the effective interfacial area is based on a simple equation that depends on the liquid holdup correction factor and a surface enhancement factor as seen in Table 33. The surface enhancement factor is equal to 0.35 for stainless steel sheet metal packing [100].

Table 33. Effective interfacial area in packing columns using SRP model [96]

Parameter	Correlation	
Reynolds number	$Re_L = \frac{s u_L \rho_L}{\mu_L}$	(46)
Froude number	$Fr_L = \frac{u_L^2}{s g}$	(47)
Weber number	$We_L = \frac{s \rho_L u_L^2}{g \sigma_L}$	(48)
Solid – liquid film contact angle	For $\sigma_L \leq 0,055 N \cdot m^{-1}$ $\cos\gamma = 0,9$ For $\sigma_L > 0,055 N \cdot m^{-1}$ $\cos\gamma = 5,211 \times 10^{-16,835 \sigma_L}$	(49)
Correction factor	$F_t = \frac{29,12 s^{0,359} (We_L Fr_L)^{0,15}}{\epsilon^{0,6} Re_L^{0,2} (\sin\theta)^{0,3} (1 - 0,93 \cos\gamma)}$	(50)
Effective interfacial area	$\frac{a_e}{a} = F_t F_{SE}$	(51)

The SRP model uses the effective gravity which takes into account forces that oppose the flow of the liquid film over the packing. These forces are caused by the pressure gradient, buoyancy and shear stress in the gas phase [93].

An iterative approach exploiting this effective gravity is used to calculate liquid holdup. The calculation steps followed for predicting liquid holdup in a packing column are shown in Table 34.

Table 34. Liquid holdup in packing columns using SRP model [101]

Parameter	Correlation	
Dry pressure drop	$\left(\frac{\Delta P}{\Delta z}\right)_d = \frac{A \rho_V}{s \varepsilon^2 (\sin\theta)^2} u_V^2 + \frac{B \mu_V}{s^2 \varepsilon \sin\theta} u_V$	(52)
Initial condition for the iterative approach	$\left(\frac{\Delta P}{\Delta z}\right)_{iter} = \left(\frac{\Delta P}{\Delta z}\right)_d$	(53)
Iterative approach	$h_L = \left(\frac{4 F_t}{s}\right)^{\frac{2}{3}} \left[\frac{3 \mu_L u_L}{\rho_L \varepsilon g_{eff} \sin\theta}\right]^{\frac{1}{3}}$ $\frac{\Delta P}{\Delta z} = \frac{\left(\frac{\Delta P}{\Delta z}\right)_d}{[1 - h_L(71,35 s + 0,614)]^5}$	(54)
Convergence	$\text{If } \frac{\Delta P}{\Delta z} \neq \left(\frac{\Delta P}{\Delta z}\right)_{iter} \rightarrow \left(\frac{\Delta P}{\Delta z}\right)_{iter} = \frac{\Delta P}{\Delta z} \text{ and restart from (52)}$ $\text{If } \frac{\Delta P}{\Delta z} \approx \left(\frac{\Delta P}{\Delta z}\right)_{iter} \rightarrow \text{Convergence}$	(55)

The constants A and B used to calculate the pressure drop in a dry column depend on the type of the packing. For metal structured packings, A and B are equal to 0.177 and 88.77 respectively [101]. Table 35 presents the equations used for the prediction of pressure drop in preloading and loading regions.

Table 35. Pressure drop in packing columns using SRP model [96]

Parameter	Correlation	
Liquid film thickness	$\delta = \left(\frac{3 \mu_L u_L}{a g \rho_L \sin\theta}\right)^{\frac{1}{3}}$	(56)
Gas flow channel diameter	$d_{hV} = \frac{(b h - 2 s \delta)^2}{\frac{b h}{\left[\left(\frac{b h - 2 s \delta}{2 h}\right)^2 + \left(\frac{b h - 2 s \delta}{b}\right)^2\right]^{0,5}} + \frac{b h - 2 s \delta}{2 h}}$	(57)
Gas capacity factor at loading point	$F_{c,lp} = \left[0,053 g d_{hV} \varepsilon^2 (\sin\theta)^{1,15} (\rho_L - \rho_V) \left(\frac{u_L}{u_V} \sqrt{\frac{\rho_L}{\rho_V}}\right)^{-0,25}\right]^{0,5}$	(58)
Pressure drop enhancement factor	$F_l = 3,8 \left(\frac{F_c}{F_{c,lp}}\right)^{\frac{2}{\sin\theta}} \left(\frac{u_L^2}{g d_{hV} \varepsilon^2}\right)^{0,13}$	(59)
Pressure drop in preloading region	$\left(\frac{\Delta P}{\Delta z}\right)_{pl} = \left(\frac{\Delta P}{\Delta z}\right)_d \left(\frac{1}{[1 - h_L(71,35 s + 0,614)]}\right)$	(60)
Pressure drop in loading region	$\left(\frac{\Delta P}{\Delta z}\right) = F_l \left(\frac{\Delta P}{\Delta z}\right)_{pl}$	(61)

5.2.3. Delft model

The Delft model [97] was developed in a joint academic project between Montz Company and Delft University of Technology. The Delft model considers that all the packing surface area is wetted by the liquid film [93].

The prediction of the effective interfacial area with the Delft model is based on an empirical correlation presented in Equation (62).

$$a_e = a \frac{(1 - \Omega)}{\left(1 + \frac{A}{u_L^B}\right)} \quad (62)$$

According to Paquet [93], Ω is equal to 0.1 for Montz Packing and for most packing with holes as Flexipac and Mellapak. A and B are constants specific to the type and size of the packing. For example, these two constants are respectively equal to 2.143×10^{-6} and 1.5 for Montz® Packing B1-250 [100].

The Delft model introduces a new expression to define the effective liquid flow angle as seen in Equation (63).

$$\alpha_L = \arctan \left[\frac{\cos(90 - \theta)}{\sin(90 - \theta) \cos \left[\arctan \left(\frac{b}{2h} \right) \right]} \right] \quad (63)$$

This model uses a simple function for predicting liquid holdup consisting on the product of the specific surface of the packing and the thickness of the liquid film.

$$h_L = \delta a \quad (64)$$

The expression of the liquid film thickness is the same adapted by the SRP model except that it uses the effective liquid flow angle.

For the prediction of the pressure drop, the Delft model uses the same equations as the SRP model. The only difference is situated in the preloading region. As reported by Paquet [93], the Delft model assumes that the gas flows in a regular zigzag pattern through the packed column. It uses three parameters which contribute to the calculation of the pressure drop in the preloading region. The details of calculation of pressure drop in the preloading region are summarized in Table 36.

Table 36. Pressure drop in preloading region using Delft model [97]

Parameter	Correlation	
Effective gas velocity	$u_{V,e} = \frac{u_V}{\varepsilon (1 - h_L) \sin\theta}$	(65)
Effective liquid velocity	$u_{L,e} = \frac{u_L}{h_L \varepsilon \sin\alpha_L}$	(66)
Relative Reynolds number for gas phase	$Re_{Vr} = \frac{\rho_V d_{hV} (u_{V,e} + u_{L,e})}{\mu_V}$	(67)
Effective Reynolds number for gas phase	$Re_{Ve} = \frac{\rho_V d_{hV} u_{V,e}}{\mu_V}$	(68)
Fraction of the flow channel occupied by the liquid phase	$\varphi = \frac{2s}{2s + b}$	(69)
Fraction of the channels ending at the column wall	$\psi = \frac{2}{\Pi} \arcsin\left(\frac{h_{pe}}{d_c \tan\theta}\right) + \frac{2 h_{pe}}{\Pi d_c^2 \tan\theta} \left(d_c^2 - \frac{h_{pe}^2}{\tan^2\theta}\right)^{0,5}$	(70)
Gas/Liquid friction coefficient	$\xi_{GL} = \left[-2 \log_{10} \left[\frac{\delta}{3,7 d_{hV}} - \frac{5,02}{Re_{Vr}} \log_{10} \left(\frac{\delta}{3,7 d_{hV}} + \frac{14,5}{Re_{Vr}} \right) \right] \right]^{-2}$	(71)
Gas/Gas friction coefficient	$\xi_{GG} = 0,722 (\cos\theta)^{3,14}$	(72)
Direction change factor for bulk zone	$\xi_{bulk} = 1,76 (\cos\theta)^{1,63}$	(73)
Direction change coefficient for wall zone	$\xi_{wall} = 34,19 u_L^{0,44} (\cos\theta)^{0,779} + \frac{4092 u_L^{0,31} + 4715 (\cos\theta)^{0,445}}{Re_{Ve}}$	(74)
Coefficient for gas/liquid friction losses	$\zeta_{GL} = \xi_{GL} \varphi \frac{h_{pb}}{d_{hV} \sin\theta}$	(75)
Coefficient for gas/gas friction losses	$\zeta_{GG} = \xi_{GG} \frac{h_{pb}}{d_{hV} \sin\theta} (1 - \varphi)$	(76)
Coefficient for losses caused by direction change	$\zeta_{DC} = \frac{h_{pb}}{h_{pe}} (\xi_{bulk} + \psi \xi_{wall})$	(77)
Pressure drop in preloading region	$\Delta P_{pl} = \Delta P_{GG} + \Delta P_{GL} + \Delta P_{DC} = \frac{1}{2} \rho_V u_{V,e}^2 (\zeta_{GG} + \zeta_{GL} + \zeta_{DC})$	(78)

5.3. Models evaluation

The three models introduced in the previous section are evaluated and compared in order to choose the most effective in the prediction of hydrodynamic properties. To achieve this, the models are compared using two systems: Air / Water and Air / Kerosol 200. These systems have been chosen because of the lack of experimental data in the open literature concerning the system of interest (biogas with H₂S / aqueous solution of sodium hydroxide). Kerosol is a paraffin, characterized by a low surface tension and high viscosity as seen in Table 37. "200" refers to its boiling point (200 °C).

The differences in liquid surface tension, density and viscosity between water and Kerosol 200 allow comparison of models for different conditions, highlighting the effects on pressure drop and liquid holdup.

The experimental data were retrieved from the work of Erasmus [100].

Table 37. Physical properties of the systems tested [100]

Component	Density [kg/m ³]	Viscosity [kg/m.s]	Surface tension [N/m]
Air	0.81	18.10 ⁻⁶	-
Water	1000	0.001	71.2 x 10 ⁻³
Kerosol 200	763	2.31 x 10 ⁻³	23.9 x 10 ⁻³

The type of packing used for this comparison is Flexipac® 350Y. This packing is different with respect to the one used in the BioGNVAL pilot plant (Montz® B1-420), but no literature data are available for this last. The dimensions for the packing Flexipac® 350Y are outlined in Table 38 and the relative constants used by Billet and Schultes are shown in Table 28.

Table 38. Dimensions of Flexipac® 350Y [100]

Property	Value
Void fraction	0.985
Corrugation angle	45 °
Corrugation base	15.5 mm
Corrugation side	11.5 mm
Crimp height	8.4 mm
Height of element	265 mm

5.3.1. Pressure drop and liquid holdup

In Fig. 39, the experimentally determined pressure drop and liquid holdup over Flexipac® 350Y [100] are compared to the results obtained with the models using an Air – Water system.

Fig. 39.a shows that SRP and Billet and Schultes models are accurate in predicting the pressure drop in preloading region ($F_c < 1.9$). The Delft model predicts the correct shape of the pressure drop curve, but compared to experimental data, the results obtained are not realistic.

Although the results are not accurate, Fig. 39.b shows that the model by Billet and Schultes is the best in predicting liquid holdup in a structured packed column. The Delft model assumes that the liquid holdup is not influenced by the gas velocity, which explains the constant shape of the curve. The modified Billet and Schultes model shown in Fig. 39 will be presented in section 5.4.

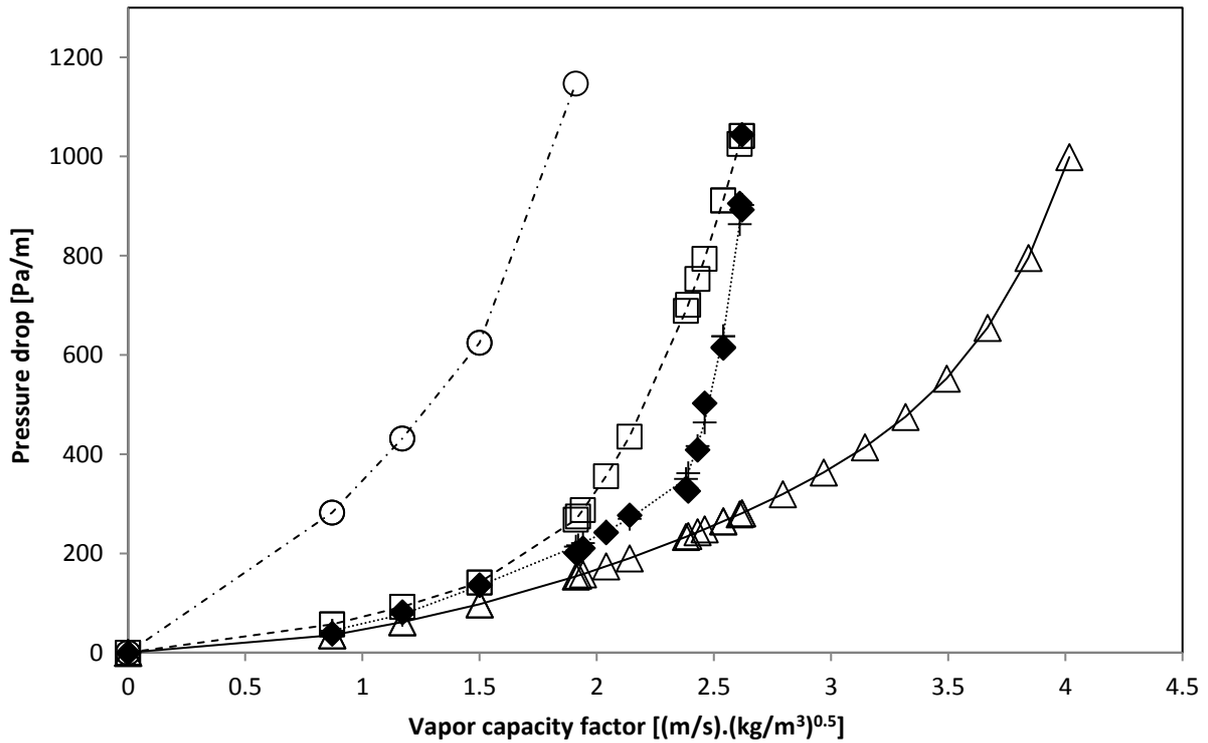


Fig. 39.a. Pressure drop evaluation for liquid load $u_L = 20.5 \text{ m/h}$

Models: (— Δ —) Billet & Schultes ; (- \square -) SRP ; (- \circ -) Delft ; (...+) Billet & Schultes modified "Section 5" ; Experimental values: (\blacklozenge) [100]

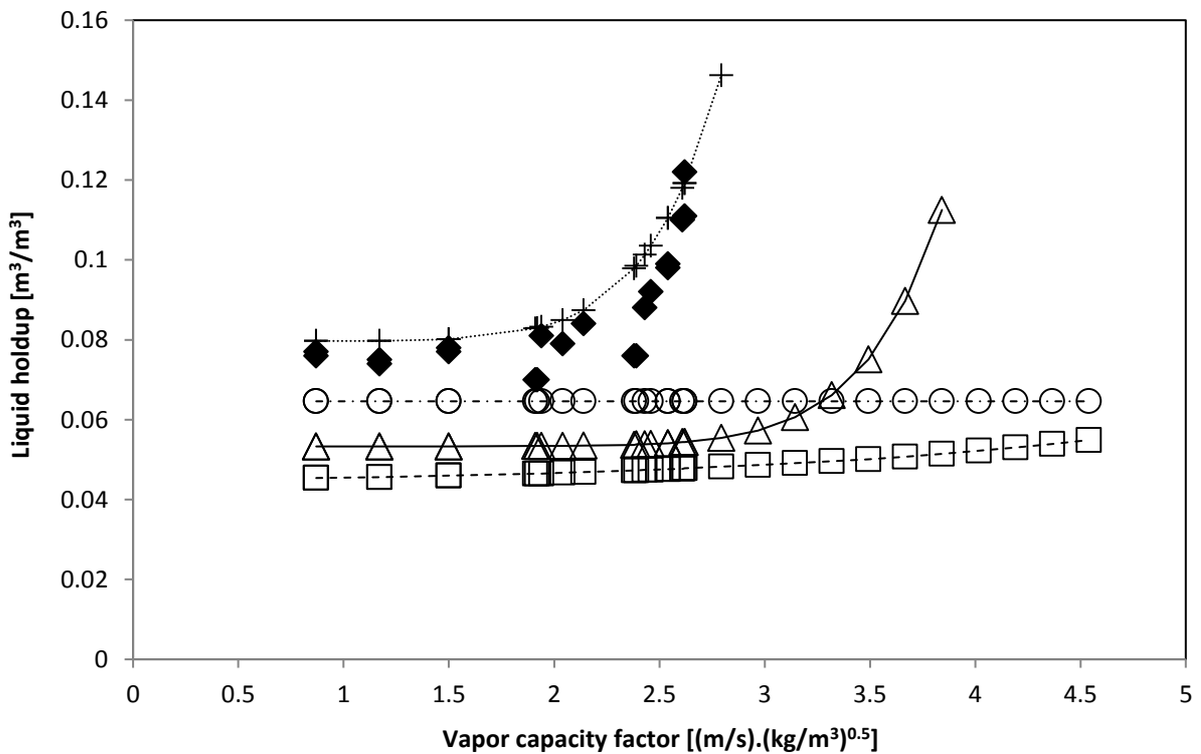


Fig. 39.b. Liquid holdup evaluation for liquid load $u_L = 20.5 \text{ m/h}$

Models: (— Δ —) Billet & Schultes ; (- \square -) SRP ; (- \circ -) Delft ; (...+) Billet & Schultes modified "Section 5" ; Experimental values: (\blacklozenge) [100]

The average absolute deviations between predictive models (Billet & Schultes, SRP and Delft) and experimental results for pressure drop and liquid holdup are shown in Table 39.

Table 39. Deviation between predictive models and experimental data

Model	Average Absolute Deviation AAD [%]	
	Pressure drop	Liquid holdup
Billet and Schultes	36	35
SRP	41	44
Delft	481	22
Modified Billet and Schultes	6	10

5.3.2. Effective interfacial area

In a packed column, the gas and the liquid phases are brought into contact and exchange mass and energy across their common interfacial area. The effective interfacial area accounts for the dead area that does not actively take part in the mass transfer process.

Fig. 40 shows the results of the effective interfacial area obtained with the three models, and compared to experimental data.

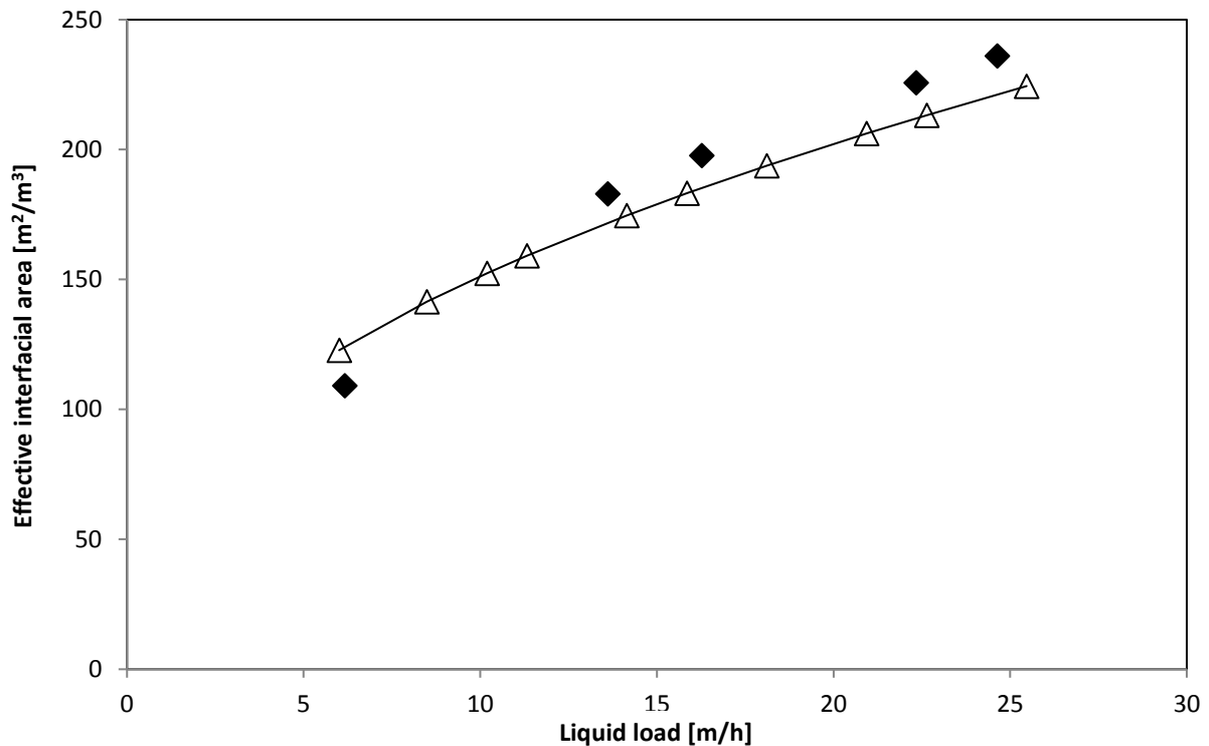


Fig. 40.a. Prediction of effective interfacial area by Billet and Schultes model for the system Air / Kerosol 200

Models: (—△—) Billet & Schultes ; (◆) Experimental values [100]

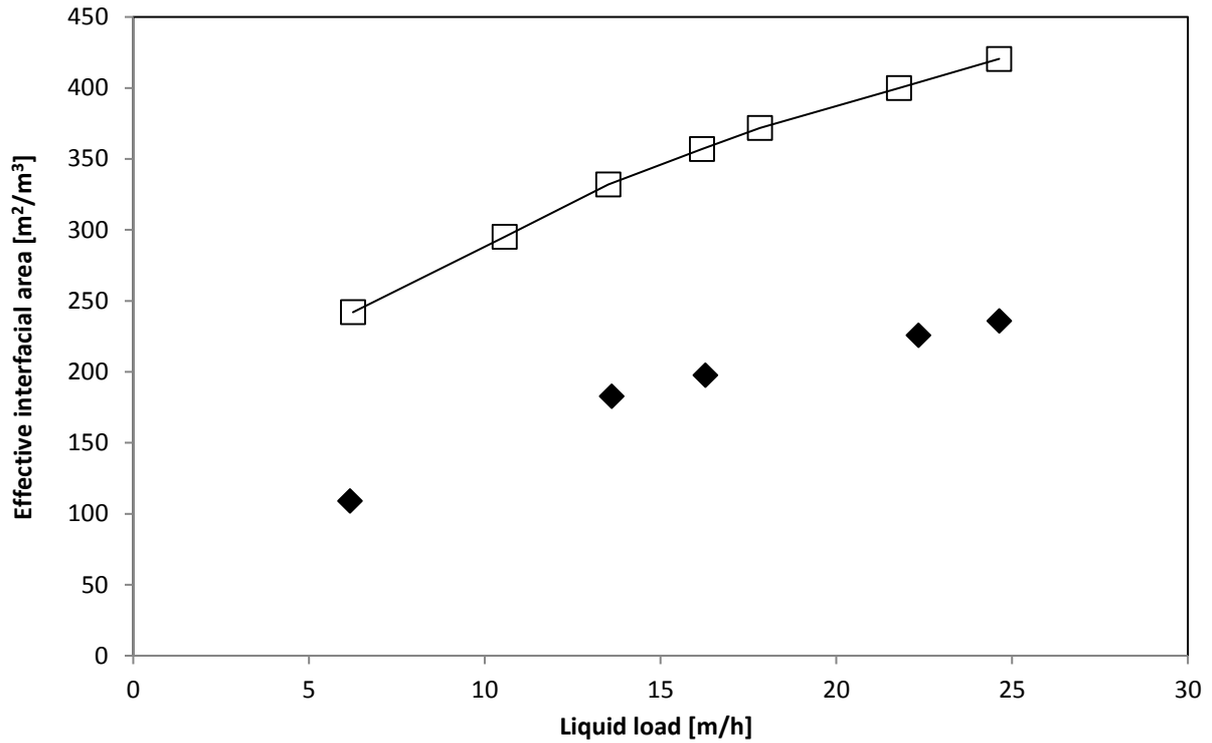


Fig. 40.b. Prediction of effective interfacial area by SRP model for the system Air / Kerosol 200

Models: (—□—) SRP ; (◆) Experimental values [100]

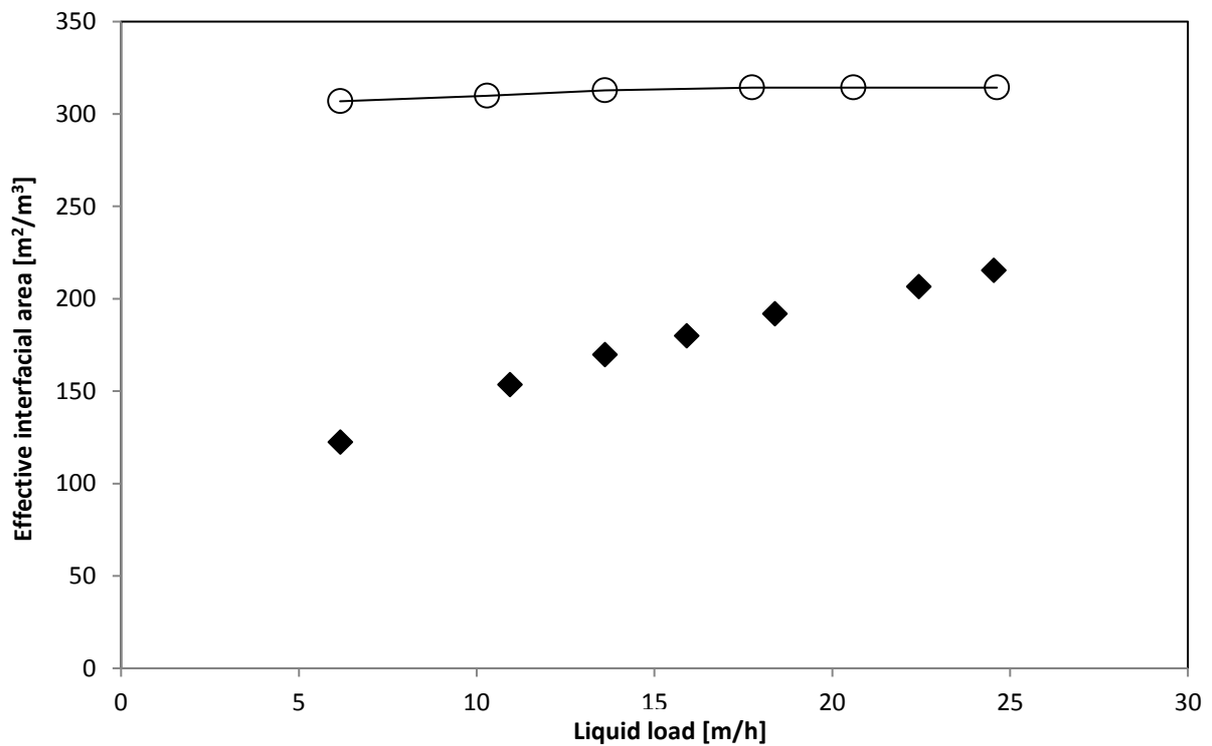


Fig. 40.c. Prediction of effective interfacial area by Delft model for the system Air / Kerosol 200

Models: (—○—) Delft ; (◆) Experimental values [100]

Fig. 40 shows that most models overpredict the effective interfacial area.

The Delft model assumes that the liquid load does not influence the effective interfacial area which presents 90 % of the overall specific area of Flexipac® 350 Y as seen in Fig. 40.c.

Compared to the Delft model, the SRP model predicts the right slope of the curve. However, for liquid loads above 16 m.h⁻¹, the predicted effective interfacial area becomes larger than the packing specific surface.

The Billet and Schultes model [95] is accurate in predicting the effective interfacial area.

The evaluation of the three models shows that the Billet and Schultes model predicts hydrodynamic parameters more accurately than SRP and Delft models. Therefore, the model by Billet and Schultes is retained for the further study.

5.4. Changes made to Billet and Schultes model and results

The Billet and Schultes model was developed for random packings, then it was extended to cover a limited number of commercially available structured packings.

To make this model more realistic and more accurate in predicting hydrodynamic parameters for structured packings, some constants and exponents defined according to experimental observations and used in the correlations were modified as function of liquid load and density. The constants and exponents to modify were selected following a sensitivity analysis. The values of the constants and exponents have been optimized by minimization of an objective function based on the deviations between modelling and experimental results. The modifications made to Equations (31), (36), (43) and (44) are shown in Tables 40, 41 and 42 for liquid holdup and pressure drop. These equations are reminded below by highlighting the modified constants.

$$h_{L,pl} = C_1 \left(\frac{\mu_L a^2 u_L}{\rho_L g} \right)^{\frac{1}{3}} \left(\frac{a_h}{a} \right)^{\frac{2}{3}} \quad (79)$$

$$h_L = h_{L,pl} + (C_2 h_{L,Fl} - h_{L,pl}) \left(\frac{u_V}{u_{V,Fl}} \right)^{n_1} \quad (80)$$

$$\psi'_L = \psi_L f_w = C_p \left(\frac{h_L}{h_{L,lp}} \right)^{n_2} \exp \left(\frac{Re_L}{200} \right) \left(\frac{64}{Re_V} + \frac{1,8}{Re_V^{0,08}} \right) \left(\frac{\varepsilon - h_L}{\varepsilon} \right)^{n_3} \quad (81)$$

$$\frac{dP}{dz} = C_3 \psi_L \frac{f_w a}{(\varepsilon - h_L)^3} \frac{F_c^2}{2} \frac{1}{K} \quad (82)$$

In order to improve predictions, Equations (31), (36), (43) and (44) were slightly modified based on the experimental results of Erasmus [100], but using only three values of liquid load ($u_L = 6$ m/h, $u_L = 20.5$ m/h and $u_L = 35.5$ m/h) for regression set. The modifications made to equations are colored in red.

Table 40. Changes made to calculate liquid holdup

Equations to modify	New equations
(79)	<p>For liquid density > 900 kg/m³</p> $h_{L,pl} = [628.4 * (u_L * 3600)^{-0.929}]^{\frac{1}{3}} \left(\frac{\mu_L a^2 u_L}{\rho_L g} \right)^{\frac{1}{3}} \left(\frac{a_h}{a} \right)^{\frac{2}{3}}$
	<p>For liquid density ≤ 900 kg/m³</p> $h_{L,pl} = \text{Exp}(-0.0057 * u_L * 3600 + 1.2074) \left(\frac{\mu_L a^2 u_L}{\rho_L g} \right)^{\frac{1}{3}} \left(\frac{a_h}{a} \right)^{\frac{2}{3}}$
(80)	<p>For liquid density > 900 kg/m³</p> $h_L = h_{L,pl} + (1,3 h_{L,Fl} - h_{L,pl}) \left(\frac{u_V}{u_{V,Fl}} \right)^{10}$
	<p>For liquid density ≤ 900 kg/m³</p> $h_L = h_{L,pl} + (\text{Exp}[-0.0006 * (u_L * 3600)^2 - 0.031 * u_L * 3600 + 0.3736] * h_{L,Fl} - h_{L,pl}) \left(\frac{u_V}{u_{V,Fl}} \right)^{[-0.0215 * (u_L * 3600)^2 + 0.8927 * u_L * 3600 + 5.3358]}$

Table 41. Changes made to calculate Pressure drop for liquid density less than 900 kg.m⁻³

Equations to modify	New equations
(81)	<p>Preloading region</p> $\psi'_L = \psi_L f_w = C_p \left(\frac{h_L}{h_{L,lp}} \right)^{0.3} \exp \left(\frac{Re_L}{200} \right) \left(\frac{64}{Re_V} + \frac{1,8}{Re_V^{0,08}} \right) \left(\frac{\varepsilon - h_L}{\varepsilon} \right)^{1.5}$
	<p>Loading & Flooding regions</p> $\psi'_L = \psi_L f_w = C_p \left(\frac{h_L}{h_{L,lp}} \right)^{0.6} \exp \left(\frac{Re_L}{200} \right) \left(\frac{64}{Re_V} + \frac{1,8}{Re_V^{0,08}} \right) \left(\frac{\varepsilon - h_L}{\varepsilon} \right)^{0.5}$
(82)	<p>Preloading region</p> $\frac{dP}{dz} = \text{Exp}(0.0149 * u_L * 3600 + 0.0829) \psi_L \frac{f_w a}{(\varepsilon - h_L)^3} \frac{F_c^2}{2} \frac{1}{K}$
	<p>Loading & Flooding regions</p> $\frac{dP}{dz} = 0.002 * (u_L * 3600)^2 - 0.0534 * u_L + 1.76 \psi_L \frac{f_w a}{(\varepsilon - h_L)^3} \frac{F_c^2}{2} \frac{1}{K}$

Table 42. Changes made to calculate Pressure drop for liquid density higher than 900 kg.m⁻³

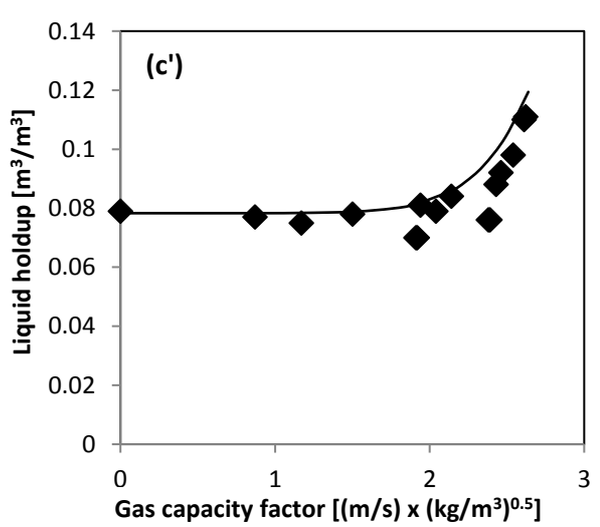
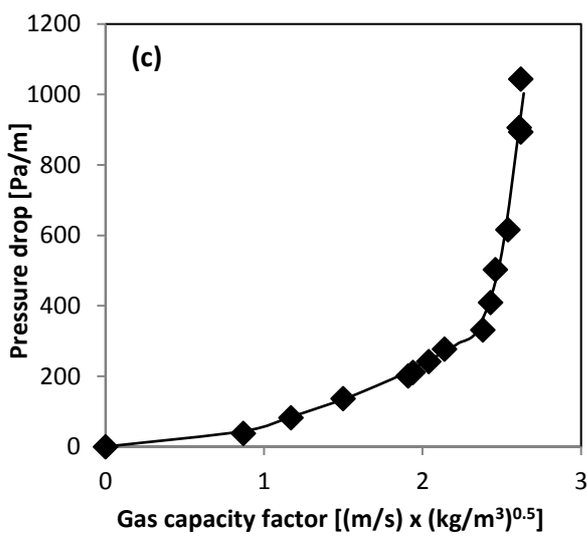
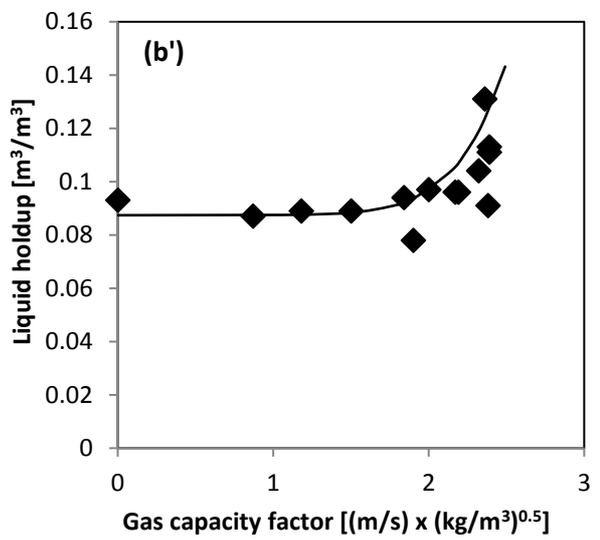
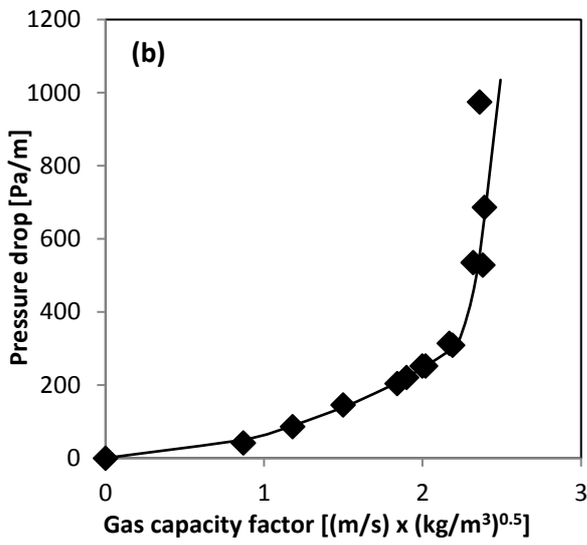
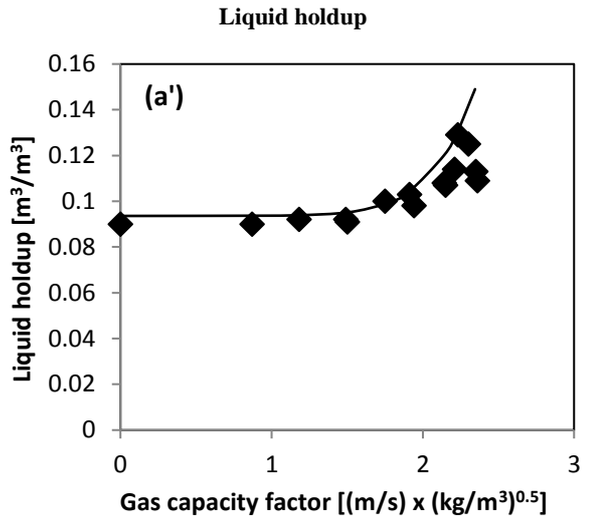
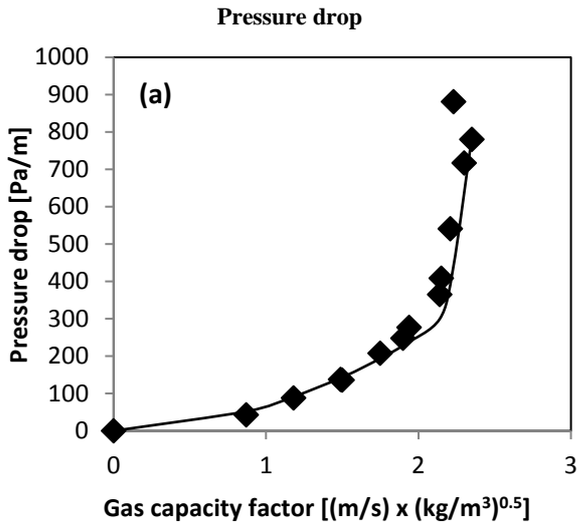
Equation to modify	New equations
(81)	<p>Preloading region</p> $\psi'_L = \psi_L f_w = C_p \left(\frac{h_L}{h_{L,lp}} \right)^{0.3} \exp \left(\frac{Re_L}{200} \right) \left(\frac{64}{Re_V} + \frac{1,8}{Re_V^{0,08}} \right) \left(\frac{\varepsilon - h_L}{\varepsilon} \right)^{1.5}$
	<p>Loading region (For $h_L > h_{L,lp}$)</p> $\psi'_L = \psi_L f_w = C_p \left(\frac{h_L}{h_{L,lp}} \right)^5 \exp \left(\frac{Re_L}{200} \right) \left(\frac{64}{Re_V} + \frac{1,8}{Re_V^{0,08}} \right) \left(\frac{\varepsilon - h_L}{\varepsilon} \right)^{1.2}$
	<p>Loading region (For $h_L \leq h_{L,lp}$)</p> $\psi'_L = \psi_L f_w = C_p \left(\frac{h_L}{h_{L,lp}} \right)^{0.3} \exp \left(\frac{Re_L}{200} \right) \left(\frac{64}{Re_V} + \frac{1,8}{Re_V^{0,08}} \right) \left(\frac{\varepsilon - h_L}{\varepsilon} \right)^3$
	<p>Flooding region</p> $\psi'_L = \psi_L f_w = C_p \left(\frac{h_L}{h_{L,lp}} \right)^{0.6} \exp \left(\frac{Re_L}{200} \right) \left(\frac{64}{Re_V} + \frac{1,8}{Re_V^{0,08}} \right) \left(\frac{\varepsilon - h_L}{\varepsilon} \right)^{1.5}$
(82)	<p>Preloading region</p> $\frac{dP}{dz} = 1.2 \psi_L \frac{f_w a}{(\varepsilon - h_L)^3} \frac{F_c^2}{2} \frac{1}{K}$
	<p>Loading region</p> $\frac{dP}{dz} = 1.5 \psi_L \frac{f_w a}{(\varepsilon - h_L)^3} \frac{F_c^2}{2} \frac{1}{K}$
	<p>Flooding region</p> $\frac{dP}{dz} = 1.4 \psi_L \frac{f_w a}{(\varepsilon - h_L)^3} \frac{F_c^2}{2} \frac{1}{K}$

The refined model is then compared to an extended range of experimental data retrieved also from the work of Erasmus [100] in order to validate the new model.

Comparisons to validate the modified model were made at various liquid loads and using two different systems: Air – Water and Air – Kerosol 200. Results of liquid holdup and pressure drop of the two systems over Flexipac® 350Y are presented in Fig. 41 and 42.

The same conditions used to evaluate the three models (Type of packing: Flexipac® 350Y, system: Air – Water, Liquid load: 20.5 m/h) are used again in order to evaluate the new model and compare it to the other ones and to the experimental data. In Fig. 39, the experimentally determined pressure drop and liquid holdup [100] are compared to the results obtained with all the models including the new one.

Table 39 presents the deviations between predictive models (Billet & Schultes, SRP and Delft), the modified Billet and Schultes model and experimental results for pressure drop and liquid holdup.



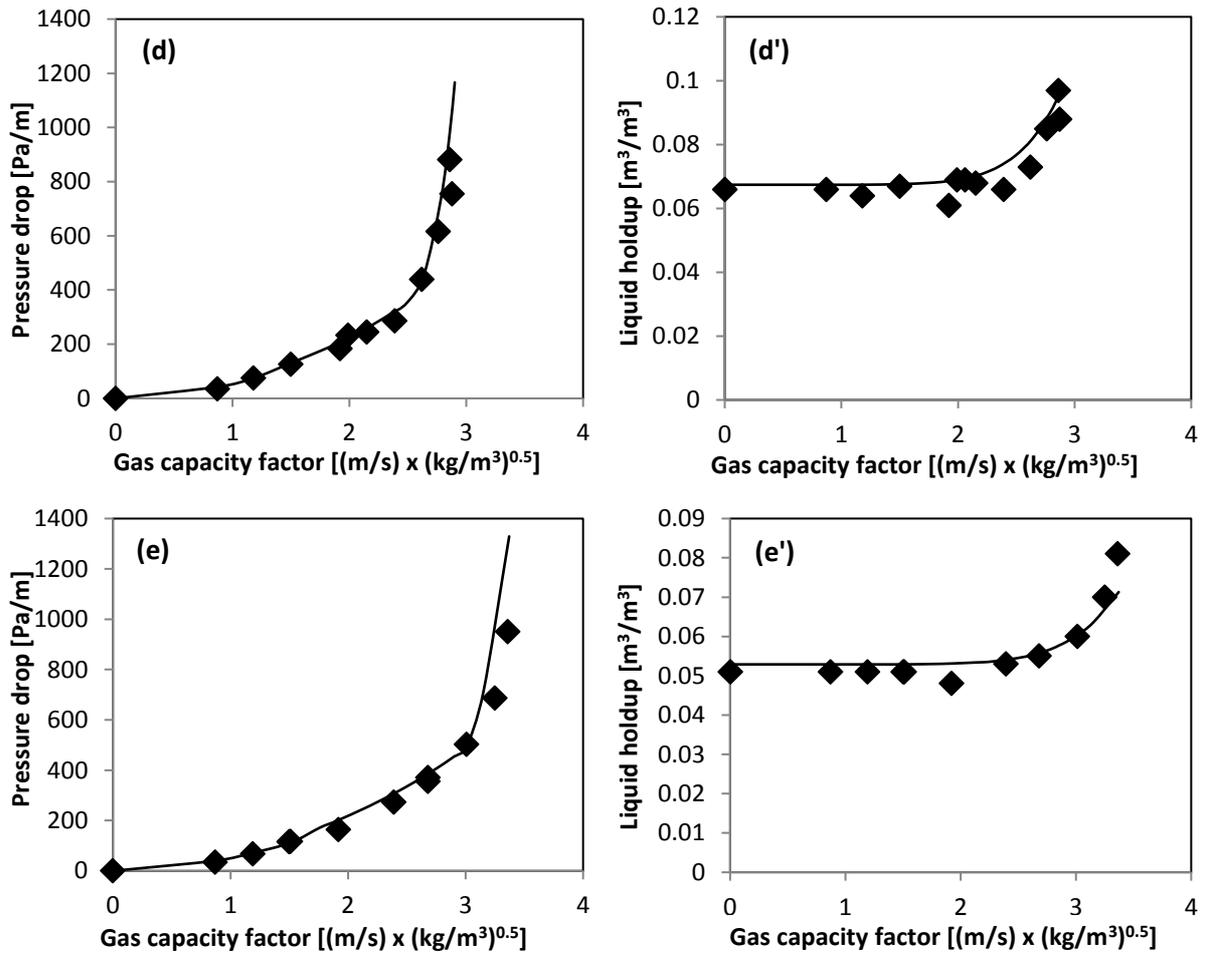
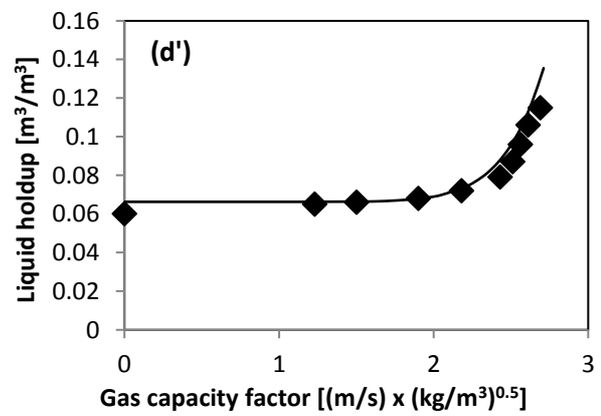
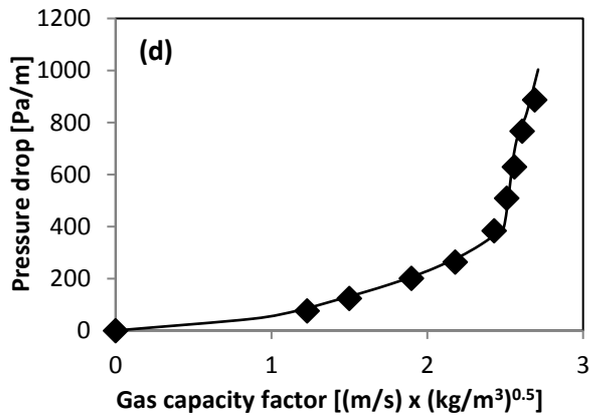
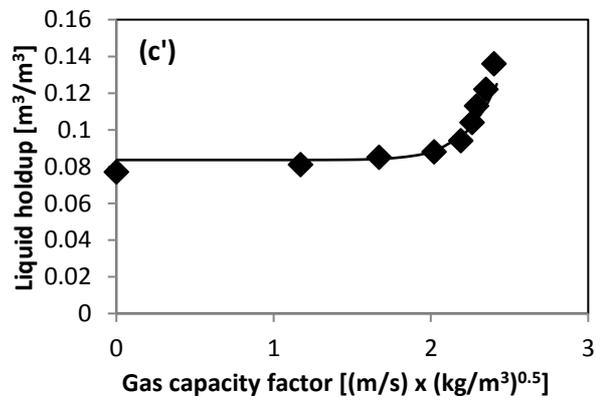
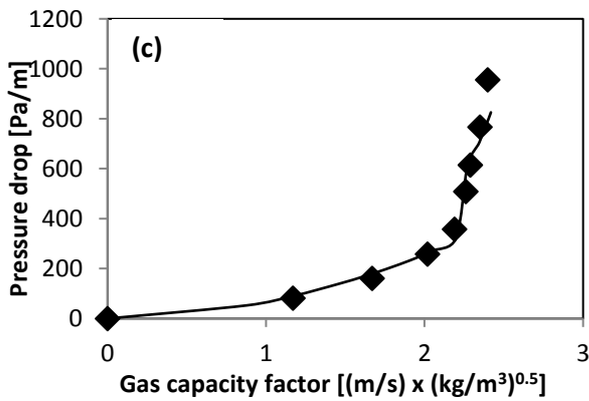
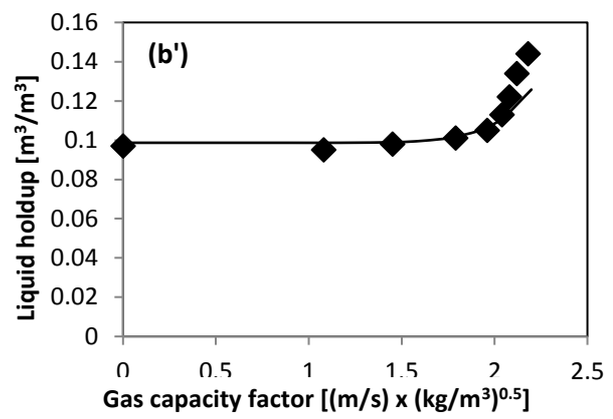
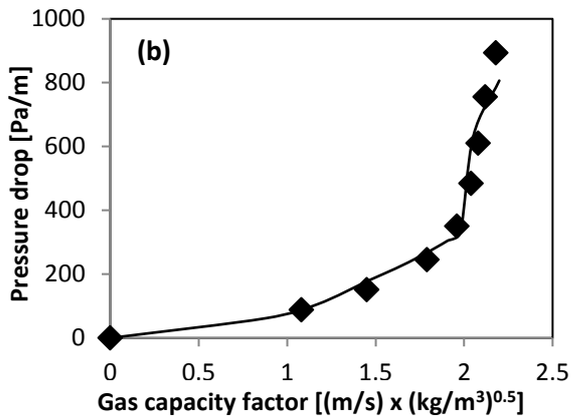
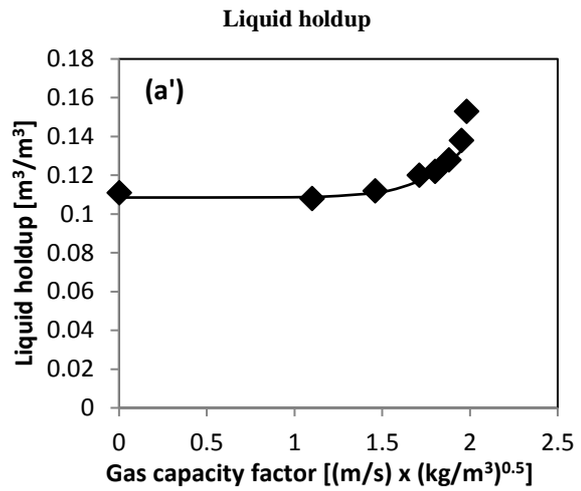
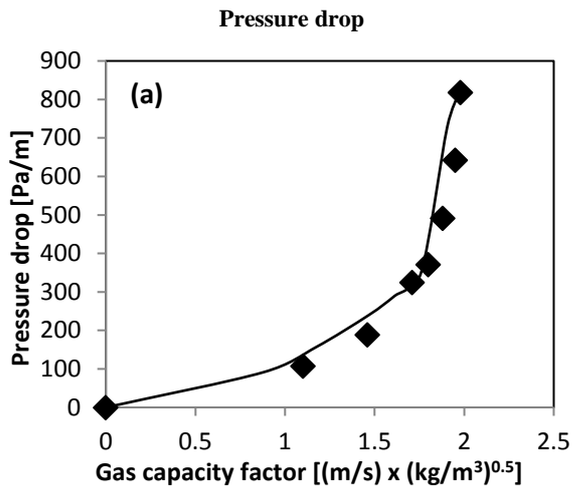


Fig. 41. Liquid holdup and pressure drop with an Air – Water system using Flexipac® 350Y packing

Models: (—) Billet & Schultes modified; (◆) Experimental values [100]

Liquid loads: (a), (a') $u_L = 35.6$ m/h; (b), (b') $u_L = 28.8$ m/h; (c), (c') $u_L = 20.5$ m/h; (d), (d') $u_L = 12.9$ m/h; (e), (e') $u_L = 6$ m/h



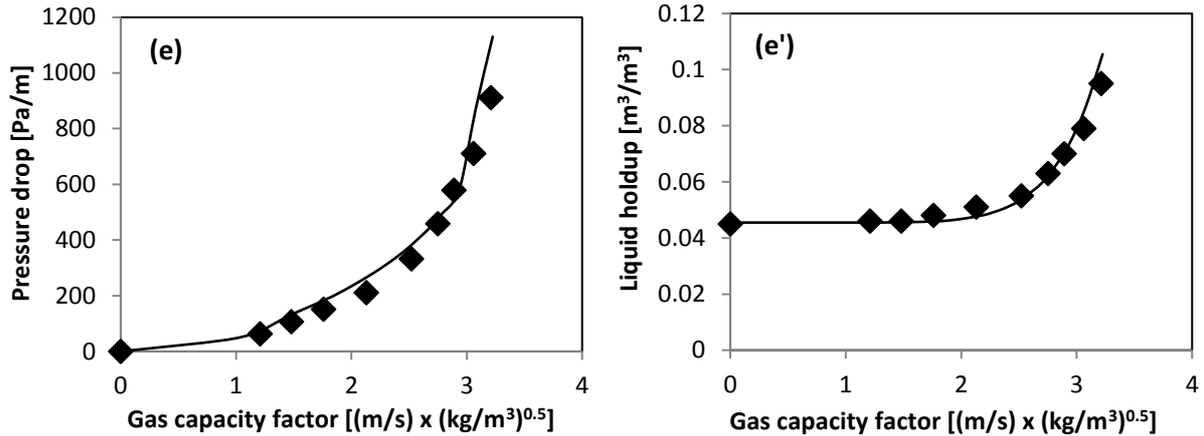


Fig. 42. Liquid holdup and pressure drop with an Air – Kerosol 200 system using Flexipac® 350Y packing

Models: (—) Billet & Schultes modified ; (♦) Experimental values [100]

Liquid loads: (a), (a') $u_L = 35.6$ m/h ; (b), (b') $u_L = 28.8$ m/h ; (c), (c') $u_L = 20.6$ m/h ; (d), (d') $u_L = 12.7$ m/h ; (e), (e') $u_L = 6.1$ m/h

Statistical deviation between experimental data and the modified model results are presented in Table 43 for both systems.

Table 43. Statistical deviation between the modified model and experimental data for pressure drop and liquid holdup predictions

Air / Water System				
Liquid load [m/h]	Pressure drop		Liquid holdup	
	AAD [%]	MAD [%]	AAD [%]	MAD [%]
35.6	10	19	7	11
28.8	5	22	2	6
20.5	5	18	6	16
12.9	8	22	4	12
6	9	21	7	12
Air / Kerosol 200 System				
Liquid load [m/h]	Pressure drop		Liquid holdup	
	AAD [%]	MAD [%]	AAD [%]	MAD [%]
35.6	9	26	3	11
28.8	7	20	3	12
20.6	6	13	4	8
12.7	5	12	6	9
6.1	12	23	4	11

After validation of the modified model, it was used to predict pressure drop on a real structured packing column used for the removal of H_2S from biogas.

The results between experimental data obtained from BioGNVAL pilot plant and the refined model are shown in Table 44. The specific constant C_p for pressure drop over Montz B1-420 packing was set to 0.14 by fitting it on experimental data.

The difference between the two results could be explained by pressure drop in the piping which does not contain packing.

Table 44. Comparison between modified correlations and experimental data for the prediction of pressure drop in a structured packing column

No. of point	L [kg/h]	V [kg/h]	$(\Delta P)_{exp}$ [Pa]	$(\Delta P)_{modified\ model}$ [Pa]	$(\Delta P)_{original\ model}$ [Pa]	Absolute value of relative deviation between experimental and modified model [%]	Absolute value of relative deviation between experimental and original model [%]
1	818	89.7	304.4	289.5	200.0	4.89	34.29
2	809	90.2	310.4	292.2	201.6	5.88	35.07
3	809	90.9	312.3	296.5	204.3	5.06	34.58
4	809	91.2	312.4	298.8	205.8	4.36	34.13
5	850	91.8	321.2	307.8	211.1	4.18	34.28
6	854	92.6	325.5	313.3	214.5	3.74	34.09
7	870	93.0	323.9	318.6	217.6	1.63	32.81
8	821	93.8	325.6	318.0	217.7	2.33	33.13
9	797	94.9	326.2	322.6	220.8	1.10	32.31
10	855	95.4	345.0	334.0	227.1	3.16	34.15

5.5. Conclusion

This chapter evaluated three semi-empirical models for prediction of hydrodynamic parameters used for an industrial application concerning biogas purification: Billet and Schultes, SRP and Delft. Flexipac® 350Y structured packing was considered here. Its capacity is closely related to hydrodynamics and mass transfer characteristics. That is why, the performances of these hydrodynamic models were investigated and compared based on existing experimental data, and the choice was finally made on the model of Billet and Schultes.

The correlations of this model were improved in order to develop an accurate prediction of hydrodynamic parameters in a structured packing column.

This model allows to precisely predicting the key hydrodynamic parameters: liquid holdup, pressure drop, effective interfacial area and especially the two transition points: loading and flooding points.

The results of pressure drop using the modified model were compared to those obtained on BioGNVAL pilot plant. Good agreement was obtained with experimental data. It is wise to note that this model may lose generality with varying applications but for the activities of interest, it wins precision. Therefore, this predictive model is ideal to predict accurately the three operating regions of a small scale structured packing column used for biogas or natural gas applications. It would allow the design of structured packing columns without the need of experimental data collected on a pilot plant. The operative conditions of the existing columns could also be optimized using the modified model to operate at full capacity.

After the hydrodynamic study, it will be interesting to investigate the thermodynamic properties in order to accurately predict the efficiencies of H₂S separation from biogas. This study based on simulations using the chemical process optimization software Aspen Plus® V8.0 will be presented in the following section.

Chapter 6: Comparison of experimental and simulation results for the removal of H₂S from biogas by means of sodium hydroxide in structured packed columns

Résumé :

Ce travail est basé sur des simulations utilisant le simulateur de procédés Aspen Plus V8.0 et les résultats sont comparés à des données expérimentales issues de la littérature et du démonstrateur « BioGNVAL ». Le modèle « Rate-based » a été considéré lors des simulations afin de déterminer l'efficacité de séparation pour différentes conditions opératoires. Ce modèle a été adopté car les résultats expérimentaux ont montré une concentration constante de NaOH dans la phase liquide en fonction du temps. Une approche γ/ϕ est employée pour décrire l'équilibre vapeur – liquide : le modèle Electrolyte Non-Random Two-Liquid « ENRTL » est utilisé pour représenter les non-idéalités de la phase liquide, alors que l'équation d'état de Redlich-Kwong est utilisée pour le calcul des propriétés de la phase vapeur. Afin d'étudier de façon réaliste l'efficacité des colonnes à garnissage utilisant de la soude comme solvant pour l'élimination sélective de l'H₂S, le modèle thermodynamique a été vérifié et validé en se basant sur différentes données expérimentales disponibles. Pour une estimation rigoureuse de la solubilité des gaz dans le solvant, les constantes de Henry obtenues ont été comparées aux données expérimentales issues du rapport de recherche RR-48 du Gas Processors Association [105] ainsi qu'à une équation semi-empirique proposée par Harvey [106]. Des propriétés physiques comme les capacités thermiques des composants purs, utilisées dans les calculs de transfert de matière et de chaleur ont été aussi vérifiées. L'équation développée par Aly et Lee [107] a été adoptée pour calculer les capacités calorifiques et les résultats obtenus ont été comparés aux données expérimentales fournies par Elliott et Lira [108]. Le calcul de la masse volumique de la phase liquide dans Aspen Plus a été effectué en fonction de la composition massique de NaOH dans la soude à 25 °C. Les résultats obtenus ont été vérifiés grâce aux données expérimentales de Herrington et al. [109]. Concernant la viscosité de la phase liquide, un modèle correctif adapté pour les électrolytes appelé modèle de « Jones-Dole » [110] a été appliqué dans Aspen Plus. Les paramètres de ce modèle correctif ont été optimisés pour les ions HCO₃⁻, Na⁺ et CO₃²⁻ en utilisant respectivement les données expérimentales de viscosité des systèmes KHCO₃ – H₂O [112], NaOH – H₂O [113] et K₂CO₃ – H₂O [114]. Ce modèle est remplacé par celui de Breslau et Miller [111] dans le cas où la concentration des électrolytes dépasse 0,1 mol/l. Les résultats de ces modèles ont été comparés aux données expérimentales de Klochko et Godneva [115]. Pour le calcul de la tension superficielle de la phase liquide, le modèle de Onsager-Samaras a été considéré dans Aspen Plus et les résultats de ce dernier ont été vérifiés en se basant sur les données expérimentales présentées dans le travail de Gel'perin et al. [116]. Les réactions chimiques impliquées ont été spécifiées. Elles sont toutes à l'équilibre chimique, à l'exception de la réaction entre le CO₂ et l'ion hydroxyde (OH⁻) qui est cinétiquement contrôlée. Pour les réactions d'équilibre, les constantes d'équilibres ont été calculées en s'appuyant sur des données d'Edwards et al. [118] sauf pour les réactions entre H₂S et OH⁻ et entre HS⁻ et OH⁻ où les constantes d'équilibres relatives ont été calculées en utilisant l'énergie libre de Gibbs faute de données dans la littérature. Pour la réaction cinétique entre CO₂ et OH⁻, l'équation d'Arrhenius a été considérée dans Aspen Plus où le facteur de fréquence et l'énergie d'activation ont été extraits du travail de Pinsent et al. [119]. Les résultats des constantes de vitesse obtenues en fonction de la température ont été vérifiées à l'aide des résultats expérimentaux de Faurholt [120]. Après la validation des paramètres physico-chimiques, les simulations ont été réalisées en utilisant les mêmes conditions employées dans le démonstrateur « BioGNVAL » comme présenté dans le tableau suivant.

Diamètre / Hauteur / Type du garnissage	0,15 m / 2,354 m / Flexipac 500Y Métal
Composition molaire du biogaz	CH ₄ (60 %) / CO ₂ (39,997 %) / H ₂ S (30 ppm)
Température / Pression / Débit du biogaz à l'entrée	9 °C / 1 atm / 85 Nm ³ /h
Température / Pression / débit du liquide à l'entrée	4 °C / 1 atm / 420 kg/h

Ces simulations ont permis de montrer l'influence de certains paramètres sur l'abattement de H₂S, comme la concentration de NaOH dans l'eau, le débit de la phase liquide et la température. Ces simulations ont aussi montré la sélectivité de la solution sodée dans l'élimination du H₂S car la réaction entre H₂S et l'ion hydroxyde est très rapide comparée à celle entre CO₂ et OH⁻. La sélectivité a donc été atteinte grâce au temps de contact très court entre les deux phases dans la colonne.

Le modèle thermodynamique utilisé dans les simulations a été validé avec des données de la littérature. L'influence de la température ainsi des conditions chimiques et hydrodynamiques dans l'absorption du H₂S a été étudiée. Les résultats des simulations ont ensuite été comparés aux résultats obtenus sur le démonstrateur « BioGNVAL ». Cette comparaison a montré que les deux résultats sont en bon accord permettant une prédiction réaliste des efficacités de séparation de H₂S du biogaz.

6.1. Introduction

This chapter was treated at an international stay conducted within the "Group on Advanced Separation Processes & GAS Processing" at Politecnico di Milano. This group is led by the Professor Laura Pellegrini who is an expert internationally recognized in process simulation and using the software Aspen Plus®.

Purification of biogas particularly requires the removal of hydrogen sulfide, which negatively affects the operation and viability of equipment especially pumps, heat exchangers and pipes, causing their corrosion. Several methods described in chapter 2 are available to eliminate hydrogen sulfide from biogas. Herein, reactive absorption in structured packed column by means of chemical absorption in aqueous sodium hydroxide solutions is considered. This study is based on simulations using Aspen Plus™ V8.0, and comparisons are done with data from BioGNVAL pilot plant treating 85 Nm³/h of biogas which contains about 30 ppm of hydrogen sulfide. The rate-based model approach has been used for simulations in order to determine the efficiencies of separation for different operating conditions. To describe vapor-liquid equilibrium, a γ/ϕ approach has been considered: the Electrolyte Non-Random Two-Liquid (ENRTL) model has been adopted to represent non-idealities in the liquid phase, while the Redlich-Kwong equation of state has been used for the vapor phase. In order to validate the thermodynamic model, Henry's law constants of each compound in water have been verified against experimental data. Default values available in Aspen Plus™ V8.0 for the properties of pure components as heat capacity, density, viscosity and surface tension have also been verified. Reactions involved in the process have been studied rigorously. Equilibrium constants for equilibrium reactions and the reaction rate constant for the kinetically controlled reaction between carbon dioxide and the hydroxide ion have been checked. Results of simulations of the pilot plant purification section show the influence of low temperatures, concentration of sodium hydroxide and hydrodynamic parameters on the selective absorption of hydrogen sulfide.

6.2. Aspen Plus® simulations

The aim of these simulations in Aspen Plus™ V8.0 is to study realistically the effectiveness of a structured packed column which uses sodium hydroxide as a chemical solvent for the selective removal of hydrogen sulfide.

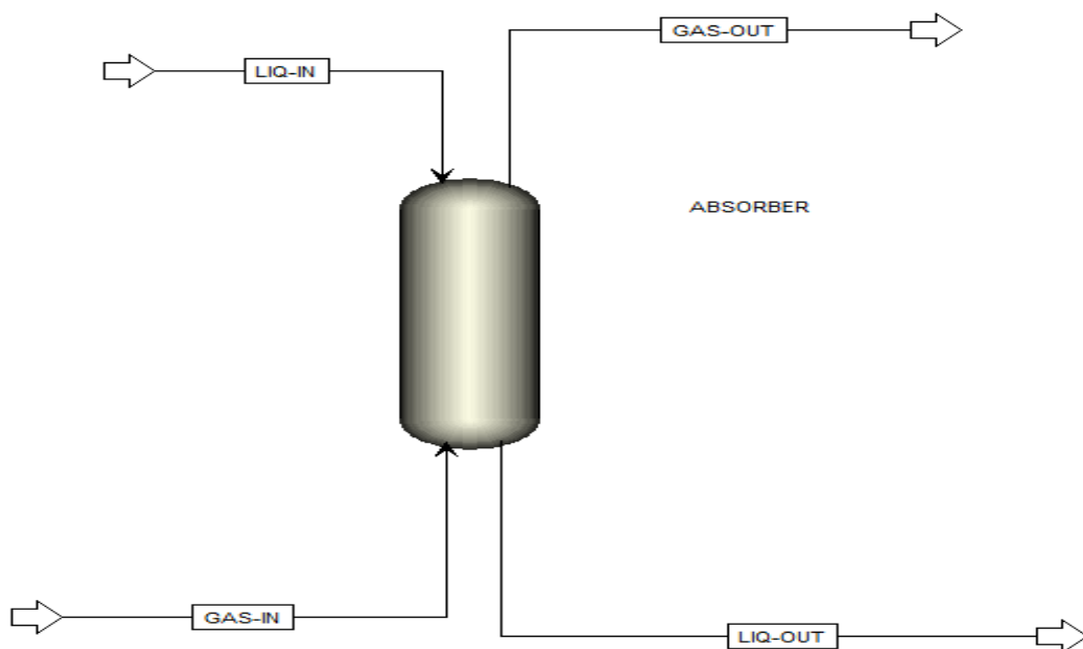


Fig. 43: Flowsheet of the absorption process simulated using Aspen Plus®

Unlike amines, sodium hydroxide is not regenerable but it is very effective in removing low contents of H₂S [36].

Although the liquid solution is recycled, experimental data showed that the NaOH consumption is quite constant. NaOH concentration can be assumed to be constant over time in the liquid phase. This assumption justifies the use of the model "Rate-based" for the study.

The rate-based modeling approach is realistic compared to the traditional equilibrium-stage modeling approach that has been employed extensively in the process industries over the decades. The rate-based models assume that separation is caused by mass transfer between the contacting phases, and use the Maxwell-Stefan theory to calculate mass transfer rates [102].

Conversely, the equilibrium-stage models assume that the contacting phases are in equilibrium with each other, which is an inherent approximation because the contacting phases are never in equilibrium in a real column.

The rate-based modeling approach has many advantages over the equilibrium-stage modeling approach. The rate-based models represent a higher fidelity, more realistic modeling approach and the simulation results are more accurate than those attainable from the equilibrium-stage models [103].

The Electrolyte Non-Random Two-Liquid model proposed by Chen and Song [104] is used for calculating the liquid phase properties, while the Redlich-Kwong equation of state is used to calculate the vapor phase properties.

This model is verified and validated using various experimental data from literature.

6.2.1. Validation of the Temperature-Dependent Henry's constant for CH₄ - H₂O, CO₂ - H₂O and H₂S - H₂O systems

For a rigorous estimation of gas solubility in the solvent, the Henry's constants obtained with Aspen Plus® should be verified by experimental data.

The Henry's constants based on the mole fraction scale are taken from Aspen Plus® databanks for the gaseous components (CH₄, CO₂ and H₂S) with water.

The temperature dependence of the Henry's constants used by Aspen Plus® is represented by:

$$\ln H_{ij} = a_{ij} + \frac{b_{ij}}{T} + c_{ij} \ln T + d_{ij}T + \frac{e_{ij}}{T^2} \quad (83)$$

Coefficients a_{ij}, b_{ij}, c_{ij}, d_{ij} and e_{ij} are summarized in Table 45 for each system.

Table 45. Coefficients used by Aspen Plus® to calculate Henry's constant

Component i	CH ₄	CO ₂	H ₂ S
Component j	H ₂ O	H ₂ O	H ₂ O
Low temperature [°C]	1.85	-0.15	-0.15
High temperature [°C]	79.85	226.85	149.85
a _{ij}	183.7811	159.1997	346.625
b _{ij}	-9111.67	-8477.711	-13236.8
c _{ij}	-25.0379	-21.957	-55.0551
d _{ij}	0.0001434	0.00578	0.05957
e _{ij}	0	0	0

Henry's constants obtained with Aspen Plus® were compared to experimental data from the research report RR-48 of the Gas processors Association [105].

Results of Henry's constants were also compared to the semi-empirical Equation (84) proposed by Harvey [106].

$$\ln H_{ij} = \ln P_{s,j} - \frac{a_{ij}}{T^*} + b_{ij} \frac{(1 - T^*)^{0.355}}{T^*} + c_{ij} e^{1-T^*} T^{*(-0.41)} \quad (84)$$

Where $P_{s,j}$ is the vapor pressure of the component j , and T^* is the reduced temperature. They are calculated respectively using the expressions (85) and (86) for water.

$$P_{s,water} = \text{Exp} \left(73.649 - \frac{72582}{T} - 7.3037 \ln T + 4.1653 \cdot 10^{-6} T^2 \right) \quad (85)$$

$$T^* = \frac{T}{T_{c,water}} \quad (86)$$

Where $T_{c,water}$ is the critical temperature of water.

Coefficients a_{ij} , b_{ij} and c_{ij} used by Harvey [106] in Equation (84) are summarized in Table 46 for each system.

Table 46. Coefficients used by Harvey to calculate Henry's constants

Component i	CH ₄	CO ₂	H ₂ S
Component j	H ₂ O	H ₂ O	H ₂ O
a_{ij}	11.01	9.4234	5.7131
b_{ij}	4.836	4	5.3727
c_{ij}	12.52	10.32	5.4227

The obtained results are similar, with no major differences. Fig. 44, 45 and 46 show the adequacy of the results for the systems CH₄ – H₂O, CO₂ – H₂O and H₂S – H₂O respectively.

The average absolute deviation is equal to 1.2 % for CH₄ – H₂O system, 1.9 for CO₂ – H₂O system and 7.8 % for H₂S – H₂O system.

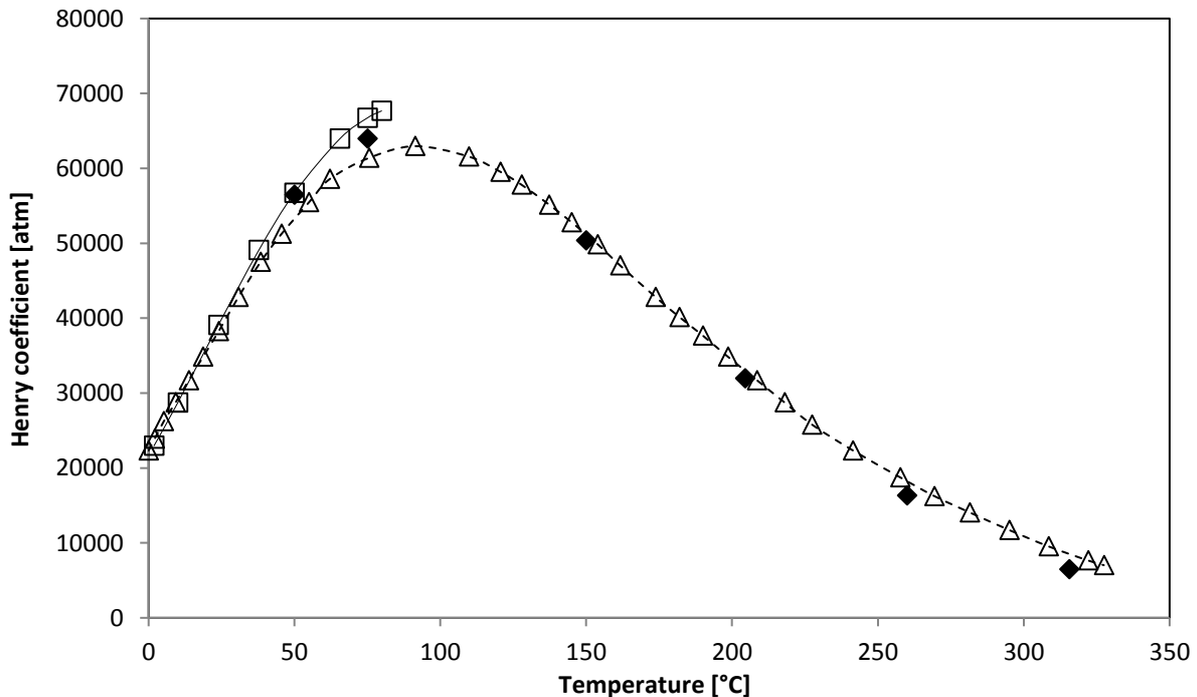


Fig. 44: Henry coefficients for CH₄ – H₂O system

(—□—) Aspen Plus®; (-Δ-) Harvey equation [106]; (◆) Experimental values [105]

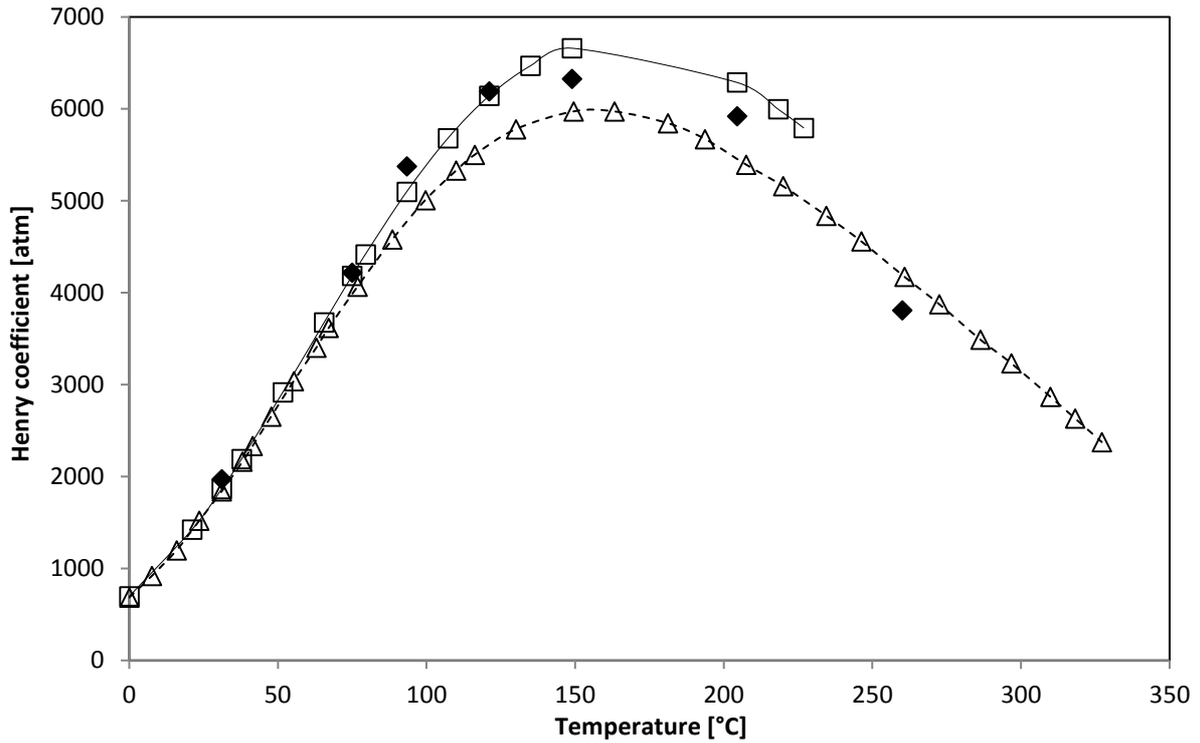


Fig. 45: Henry coefficients for CO₂ – H₂O system

(—□—) Aspen Plus®; (- -Δ- -) Harvey equation [106]; (◆) Experimental values [105]

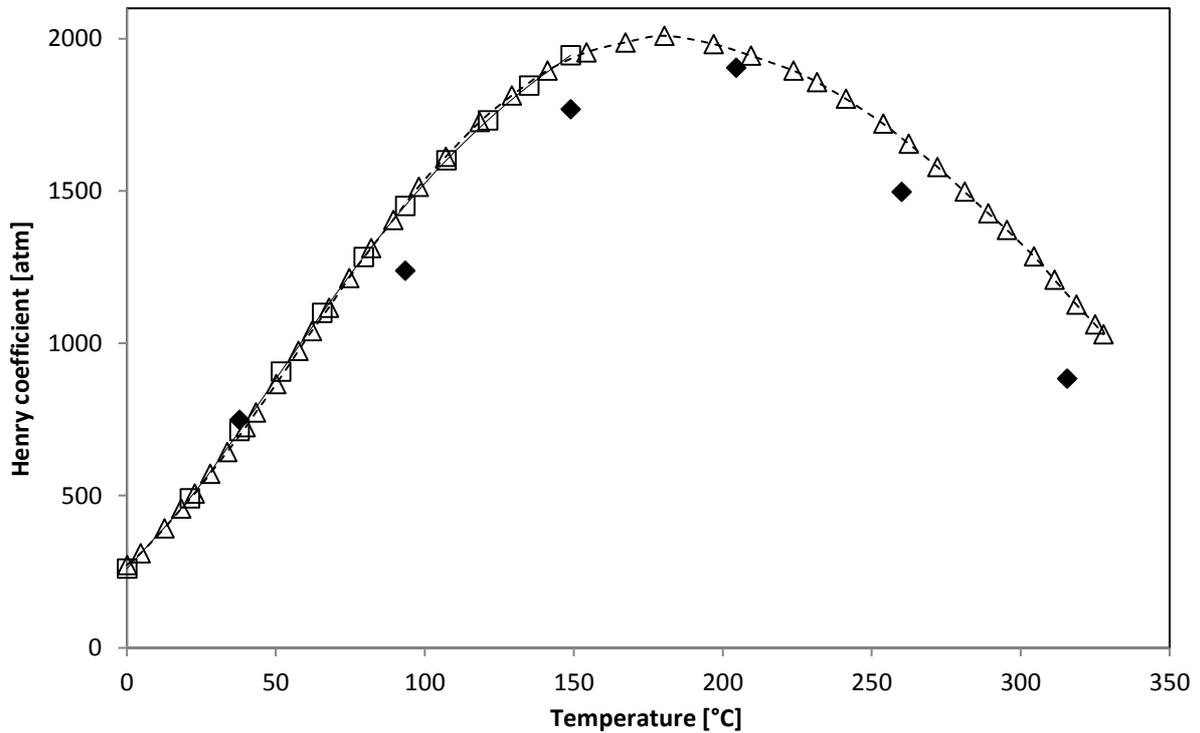


Fig. 46: Henry coefficients for H₂S – H₂O system

(—□—) Aspen Plus®; (- -Δ- -) Harvey equation [106]; (◆) Experimental values [105]

6.2.2. Validation of heat capacity for carbon dioxide

Some physical properties as heat capacities of pure components used for heat and mass transfer modelling were also checked. The ideal gas heat capacity equation (87) developed by Aly and Lee [107] is used for Aspen Plus® simulations. Fig. 47 shows for example the comparison of results obtained with the adopted model [107] in Aspen Plus® with experimental data [108].

$$C_p^{*ig} = C_{1i} + C_{2i} \left(\frac{C_{3i}/T}{\sinh(C_{3i}/T)} \right)^2 + C_{4i} \left(\frac{C_{5i}/T}{\cosh(C_{5i}/T)} \right)^2 \quad (87)$$

The values of the constants of equation (87) used to calculate the heat capacities of carbon dioxide are listed in Table 47.

Table 47. Values of the constants used by Equation (87) to calculate the heat capacity of carbon dioxide

Constants	Values
C_{1i}	29.37
C_{2i}	34.54
C_{3i}	1428
C_{4i}	26.4
C_{5i}	588

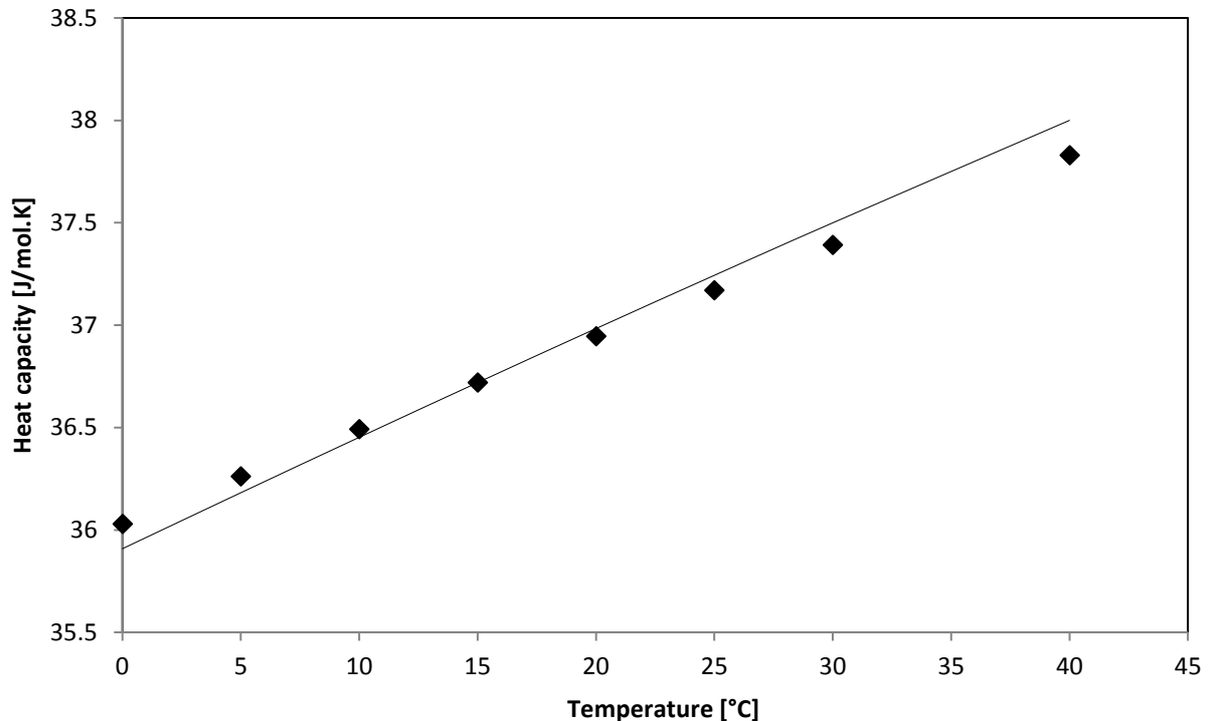


Fig. 47: Comparison between model and experimental data for heat capacity for carbon dioxide

(—) Aspen Plus® ; (♦) Experimental values [108]

The deviations between experimental and calculated results of heat capacity of carbon dioxide are presented in Fig. 48.

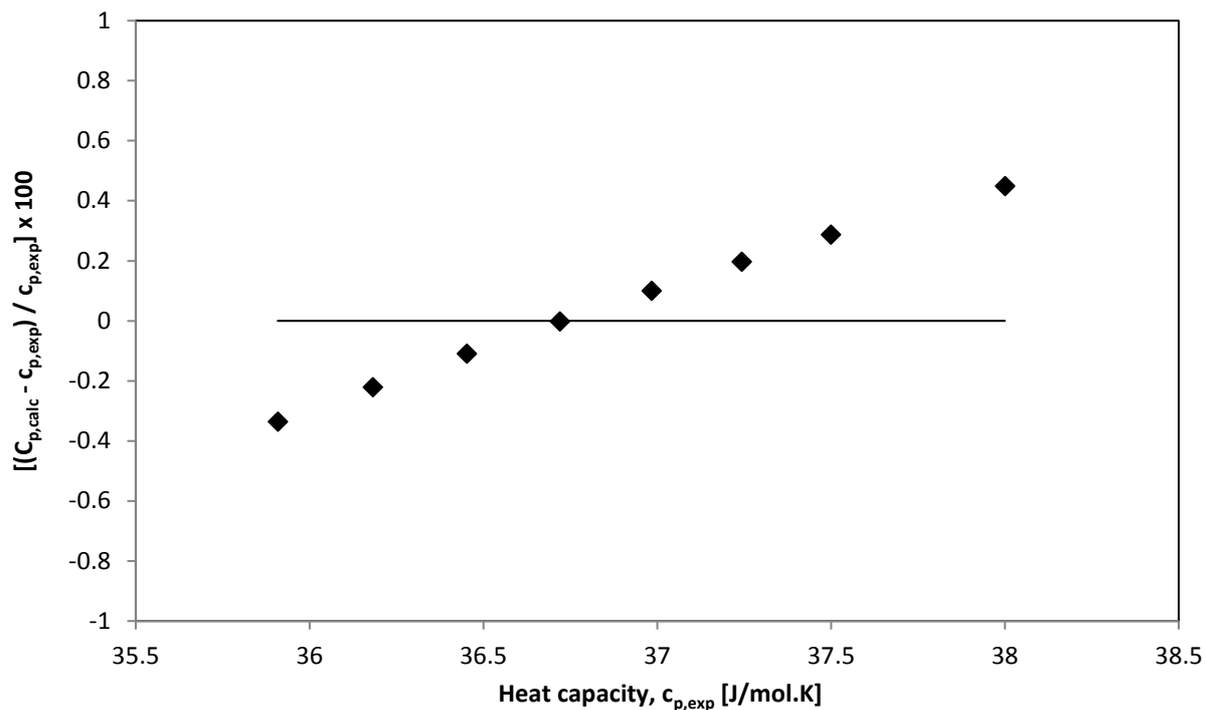


Fig. 48: Deviation between experimental and calculated results of carbon dioxide heat capacity

6.2.3. Validation of liquid density of NaOH – H₂O

The calculations of the density of the liquid phase have been verified depending on the mass fraction of sodium hydroxide at 25 ° C.

The results obtained with Aspen Plus™ are in good agreement with experimental data [109] as shown in Fig. 49.

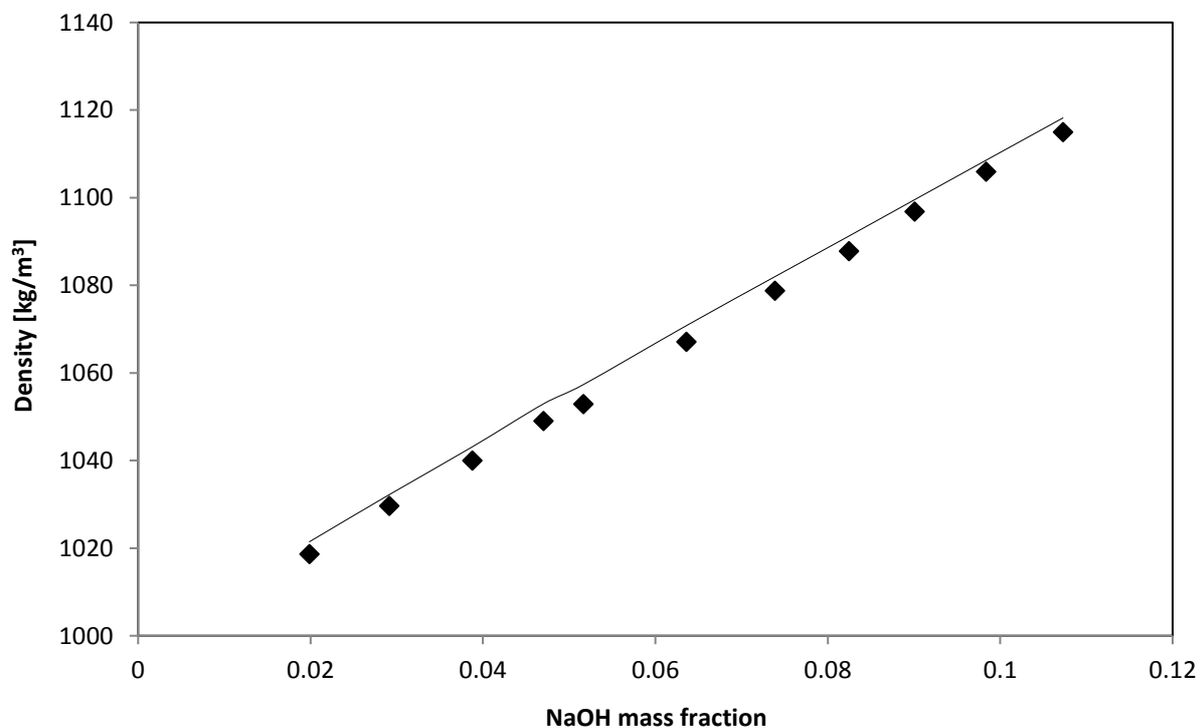


Fig. 49: Comparison between model and experimental data for liquid density of NaOH – H₂O

(—) Aspen Plus® ; (♦) Experimental values [109]

The deviations between experimental and calculated results of liquid density are depicted in Fig. 50.

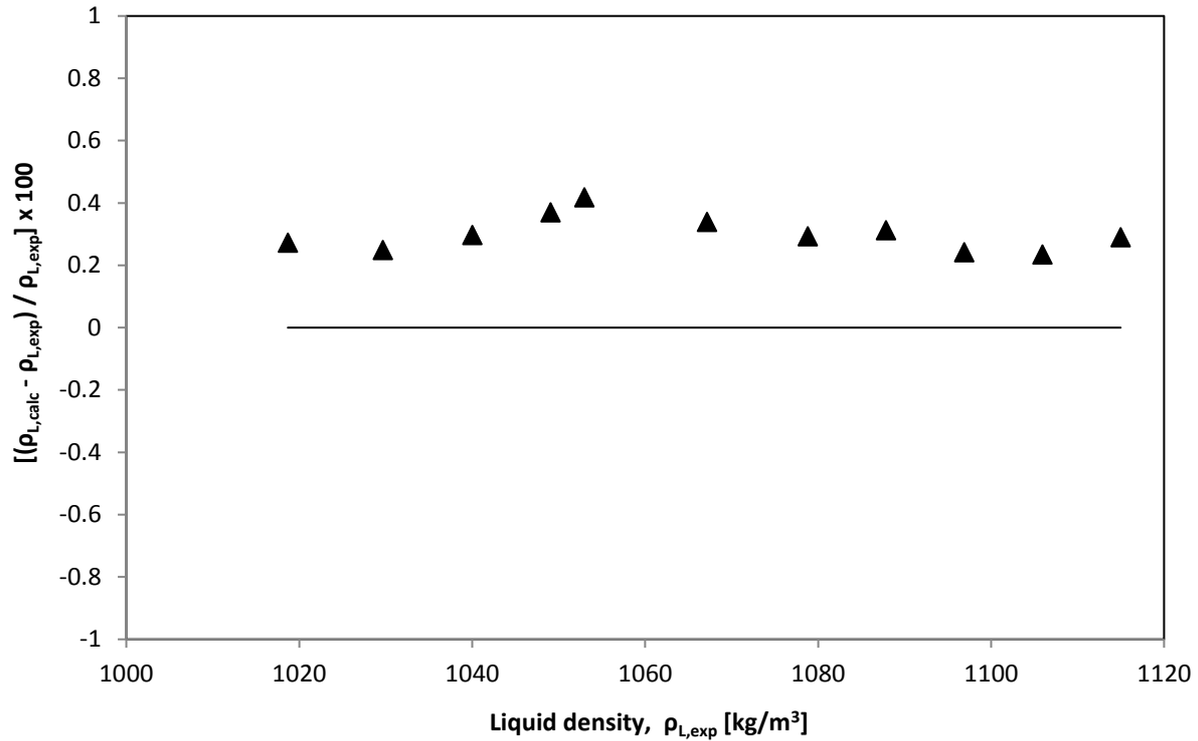


Fig. 50: Deviation between experimental and calculated results of liquid density

6.2.4. Validation of liquid viscosity of NaOH – H₂O

For the liquid viscosity, a corrective model for electrolytes called "Jones-Dole" is applied in Aspen Plus®. This model uses the mass fraction of the solvent in the liquid phase. This model is presented in Equation (88) [110].

$$\mu^l = \mu_{solv} \left(1 + \sum_{ca} \Delta\mu_{ca}^l \right) \quad (88)$$

Where μ_{solv} is the viscosity of the liquid solvent mixture, calculated using the Andrade model and $\Delta\mu_{ca}^l$ is the contribution to the viscosity correction due to apparent electrolyte ca (cation-anion).

In Aspen Plus®, the ENRTL model calculates the viscosity of the liquid solvent mixture by default using the modified Andrade Equation (89).

$$\ln \mu^l = \sum_i f_i \ln \mu_i^{*l} + \sum_i \sum_j (k_{ij} f_i f_j + m_{ij} f_i^2 f_j^2) \quad (89)$$

Where f_i is by default the mole fraction of the component i . k_{ij} and m_{ij} are binary parameters, they allow accurate representation of complex liquid mixture viscosity. k_{ij} and m_{ij} are given respectively by Equations (90) and (91).

$$k_{ij} = a_{ij} + \frac{b_{ij}}{T} \quad (90)$$

$$m_{ij} = c_{ij} + \frac{d_{ij}}{T} \quad (91)$$

When the electrolyte concentration exceeds 0.1 M, Aspen Plus® uses the Equation (92) of Breslau and Miller instead of that of Jones and Dole [111].

$$\Delta\mu_{ca}^l = 2.5 V_e c_{ca}^a + 10.05 V_e (c_{ca}^a)^2 \quad (92)$$

Where V_e is the effective volume. It is given by Equation (92).

$$V_e = \frac{B_{ca} - 0.002}{2.6} \quad \text{For salts involving univalent ions} \quad (93.a)$$

$$V_e = \frac{B_{ca} - 0.011}{5.06} \quad \text{For other salts} \quad (93.b)$$

Where B_{ca} is calculated using Equation (94).

$$B_{ca} = (b_{c,1} + b_{c,2} T) + (b_{a,1} + b_{a,2} T) \quad (94)$$

c_{ca}^a is the concentration of apparent electrolyte ca. It is calculated using Equation (95).

$$c_{ca}^a = \frac{x_{ca}^a}{V_m^l} \quad (95)$$

Where x_{ca}^a is the mole fraction of the apparent electrolyte ca and V_m^l is the molar volume of the liquid mixture calculated by the Clarke model.

The electrolyte correction model parameters were improved for the ion HCO_3^- using $\text{KHCO}_3 - \text{H}_2\text{O}$ viscosity data [112].

The regression of parameters for Na^+ ion was performed with viscosity data of $\text{NaOH} - \text{H}_2\text{O}$ system [113].

For ion CO_3^{2-} , parameters have been optimized considering experimental data for the $\text{K}_2\text{CO}_3 - \text{H}_2\text{O}$ system [114].

For other ions, values provided by Aspen Plus® database were used.

Fig. 51 shows the fit between experimental data and Aspen Plus® results for the viscosity of the liquid phase as a function of the mass fraction of sodium hydroxide.

Data of this comparison are carried out at a temperature of about 25 °C.

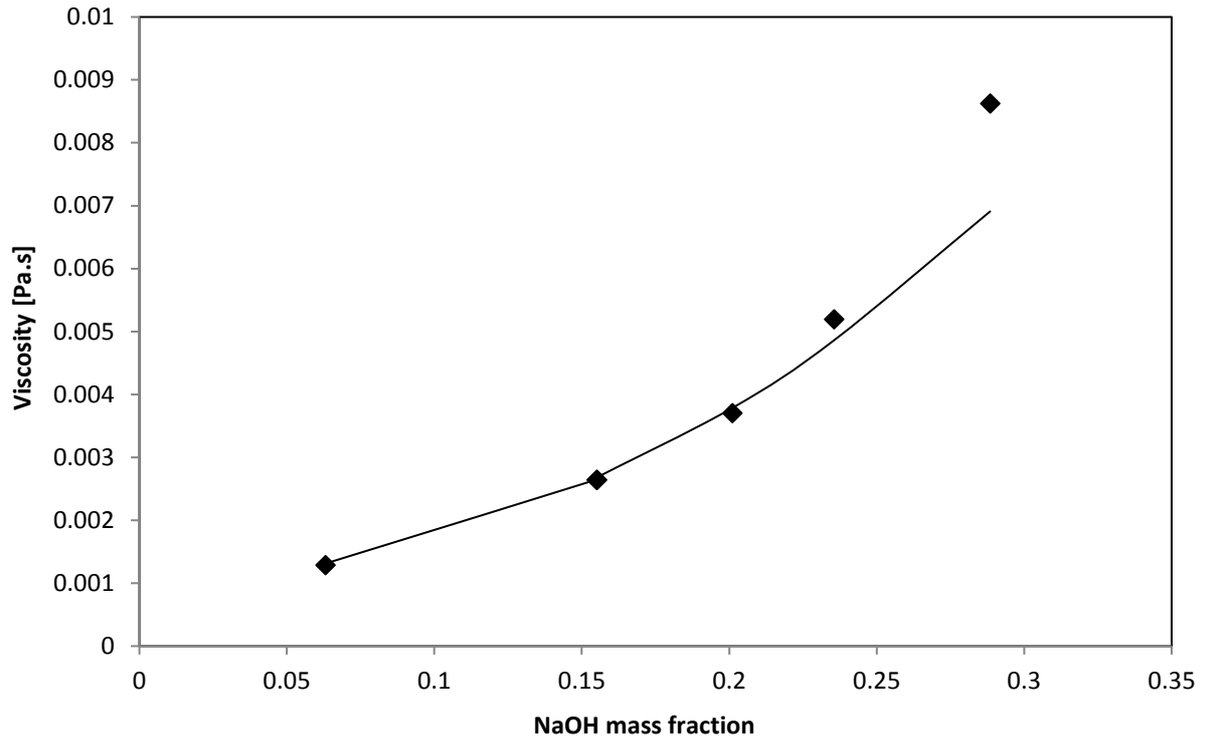


Fig. 51: Comparison between model and experimental data for liquid viscosity of NaOH – H₂O

(—) Aspen Plus® ; (◆) Experimental values [115]

The deviations between experimental and calculated results of liquid viscosity are shown in Fig. 52.

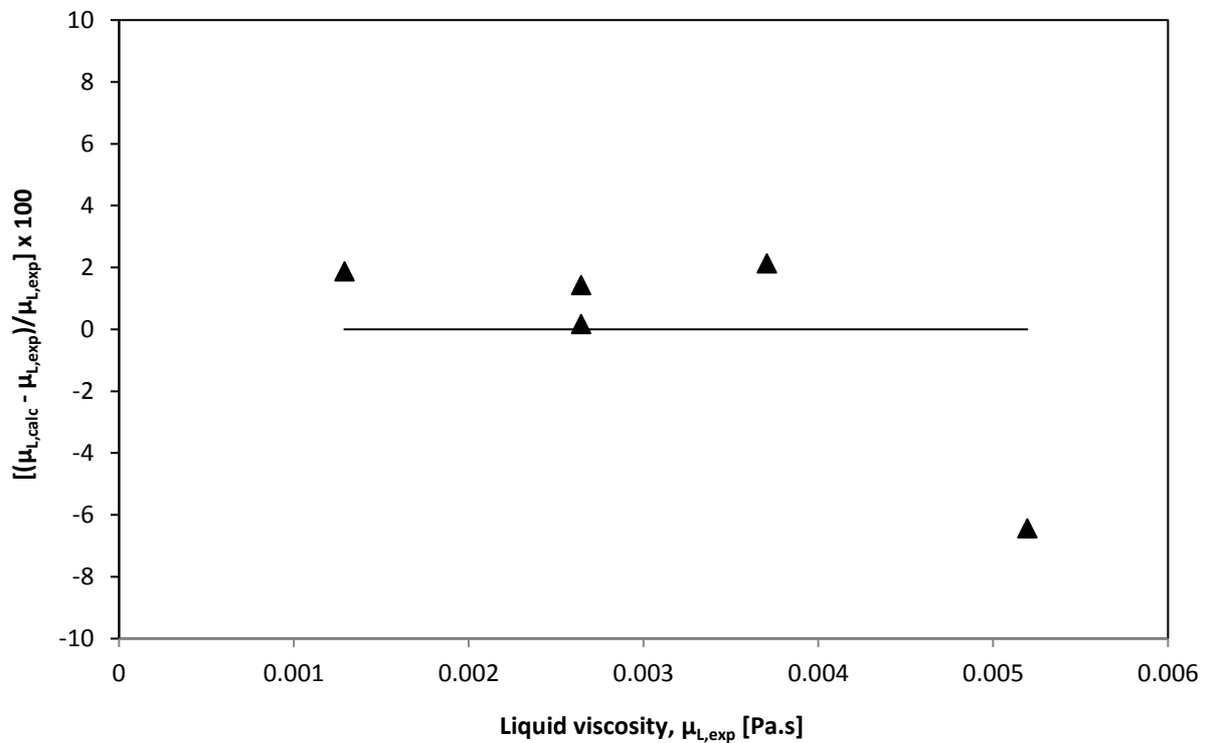


Fig. 52: Deviation between experimental and calculated results of liquid viscosity

6.2.5. Validation of surface tension of NaOH - H₂O

To calculate the liquid phase surface tension, Aspen Plus® uses the model of Onsager-Samaras presented by Equation (96). Results obtained were compared to experimental data found in literature, as shown in Fig. 53 [116].

$$\sigma = \sigma_{solv} + \sum_{ca} x_{ca}^a \Delta\sigma_{ca} \quad (96)$$

Where σ_{solv} is the surface tension of the solvent mixture calculated using the General Pure Component Liquid Surface Tension Model [117] and $\Delta\sigma_{ca}$ is the contribution to the surface tension correction due to apparent electrolyte ca , calculated using Equation (97).

$$\Delta\sigma_{ca} = \frac{80}{\epsilon_{solv}} c_{ca}^a \log \left[\frac{1.13 \times 10^{-13} (\epsilon_{solv} T)^3}{c_{ca}^a} \right] \quad (97)$$

Where ϵ_{solv} is the dielectric constant of the solvent mixture.

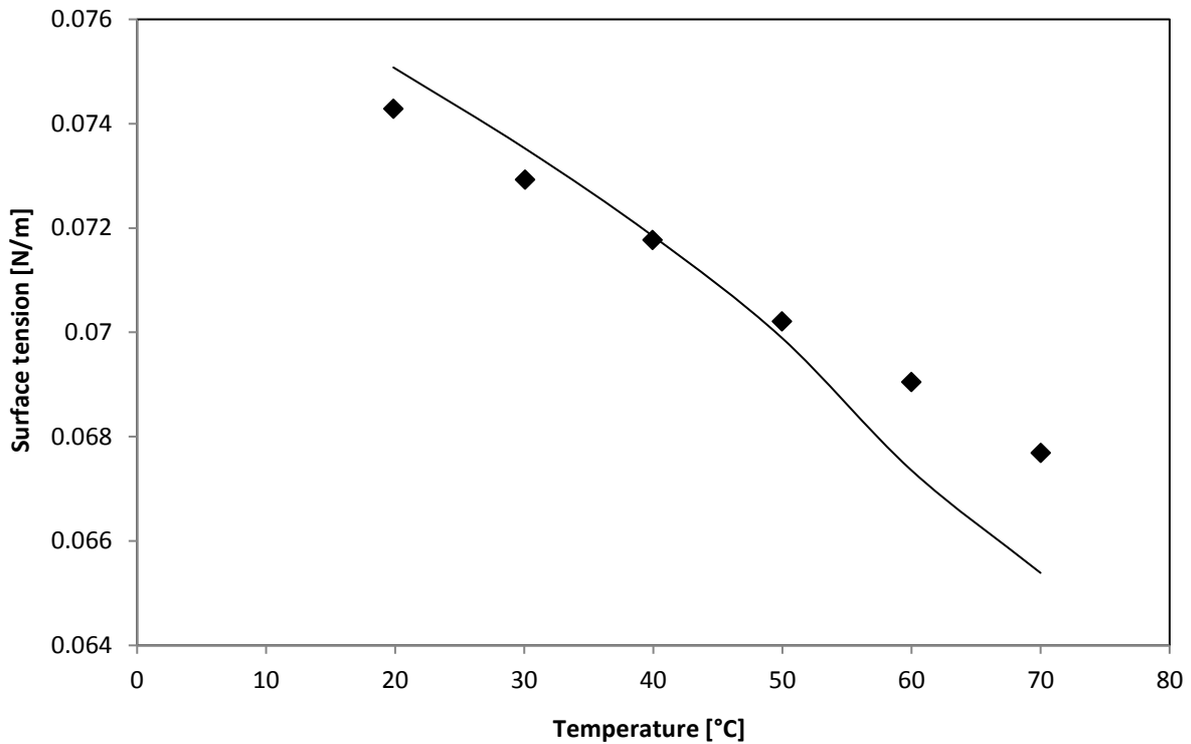


Fig. 53: Comparison between model and experimental data for liquid phase surface tension of 5 wt% NaOH aqueous solution

(—) Aspen Plus® ; (♦) Experimental values [116]

The deviations between experimental and calculated results of surface tension are presented in Fig. 54.

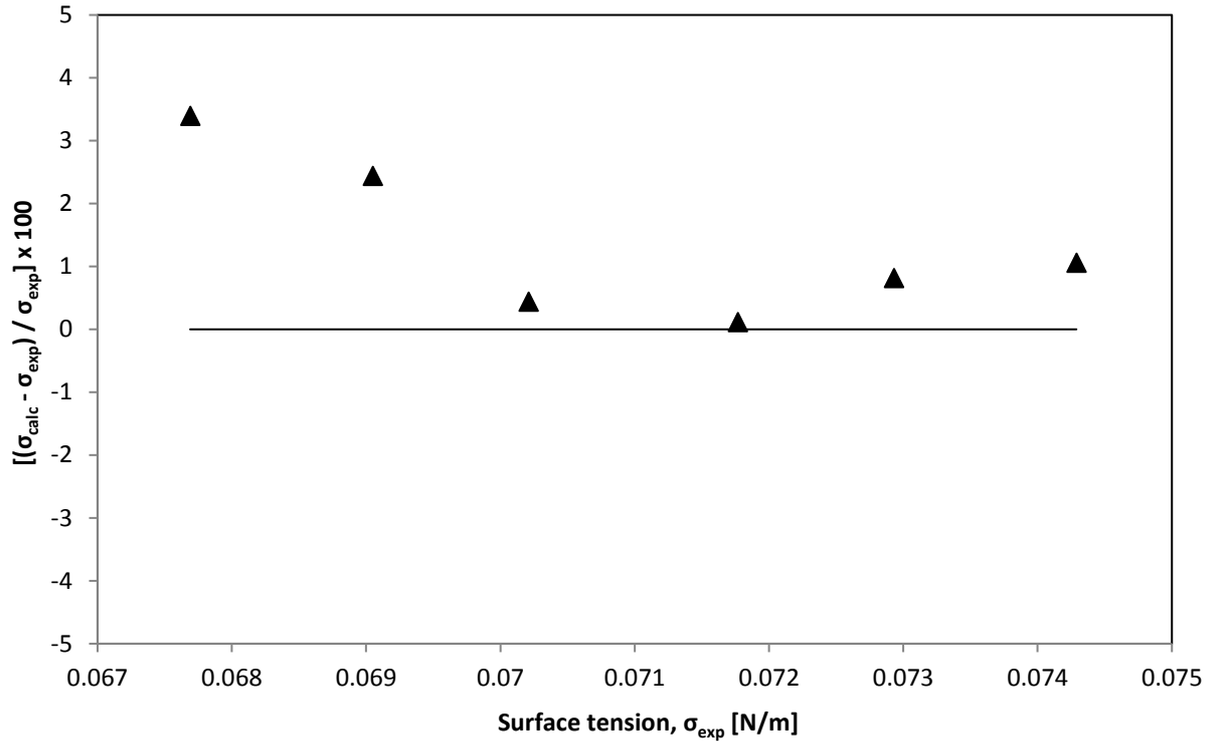
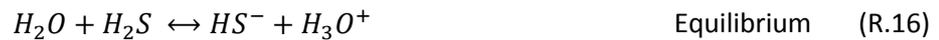


Fig. 54: Deviation between experimental and calculated results of surface tension

6.2.6. Validation of chemical parameters

All the reactions involved in the process have been specified. These reactions are assumed to be in chemical equilibrium. Only the reaction between carbon dioxide and hydroxyl ion is kinetically controlled. The reactions defined in Aspen Plus® are presented in the following expressions.



Chemical equilibrium constants for the instantaneous reversible reactions are calculated using the following Expression (98) employed by Aspen Plus®.

$$\ln K_{eq} = A + \frac{B}{T} + C \ln T \quad (98)$$

The calculation of the temperature-dependent equilibrium constants requires the knowledge of coefficients A, B and C, which were taken from the work of Edwards et al [118]. Coefficients used for Reactions (R.13) to (R.17) are presented in Table 48.

Table 48. Coefficients used in the calculation of the equilibrium constant

Reaction	A	B	C
R.13	231.456	-12092.1	-36.7816
R.14	132.899	-13445.9	-22.4773
R.15	216.05	-12431.7	-35.4819
R.16	214.582	-12995.4	-33.55471
R.17	-9.74	-8585.47	0

These coefficients are unavailable in literature for Reactions (R.18) and (R.19). Therefore, equilibrium constants of these two reactions were calculated by Aspen Plus® using Gibbs free energies. Results for Reaction (R.18) were verified using equilibrium constants of Reactions (R.14) and (R.16) by using Equation (99). Fig. 55 shows the results obtained.

$$K_{eq,R.6} = \frac{K_{eq,R.4}}{K_{eq,R.2}} \quad (99)$$

A temperature-dependent expression was proposed in order to define coefficients A, B and C for Reaction (R.18).

The values of the defined coefficients are presented in Table 49.

Table 49. Coefficients used in the calculation of the equilibrium constant of Reaction (R.6)

Reaction	A	B	C
R.18	147	-1930	-21.15

The same method has been adopted in order to validate the equilibrium constant for Reaction (R.19).

For kinetic-controlled reactions (R.24) and (R.25), the power law expression (100) is adopted by Aspen Plus®.

$$r = k e^{(-E/RT)} \prod_{i=1}^N C_i^{a_i} \quad (100)$$

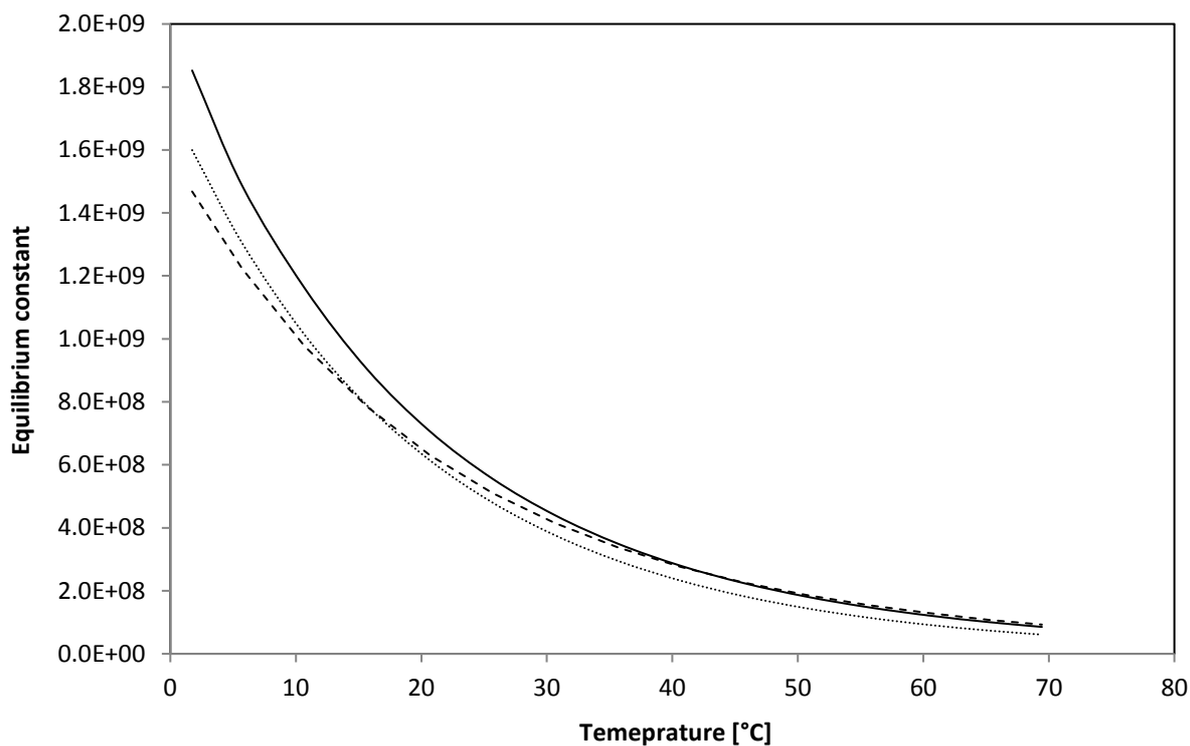


Fig. 55: Comparison of results of equilibrium constant for reaction R.18

(—) Aspen Plus®; (- - -) Equation (99); (.....) Proposed equation

k and E parameters are given in Table 50 for reactions (R.24) and (R.25), knowing that the concentration is based on the molarity.

Table 50. Parameters k and E for kinetic-controlled reactions [119]

Reaction	k	E [cal/mol]
R.24	4.32E+13	13249
R.25	2.83E+17	29451

The reaction rate constants for the kinetic reaction between CO₂ and OH⁻ have been verified against experimental data [120]. Fig. 56 shows the good agreement between the results.

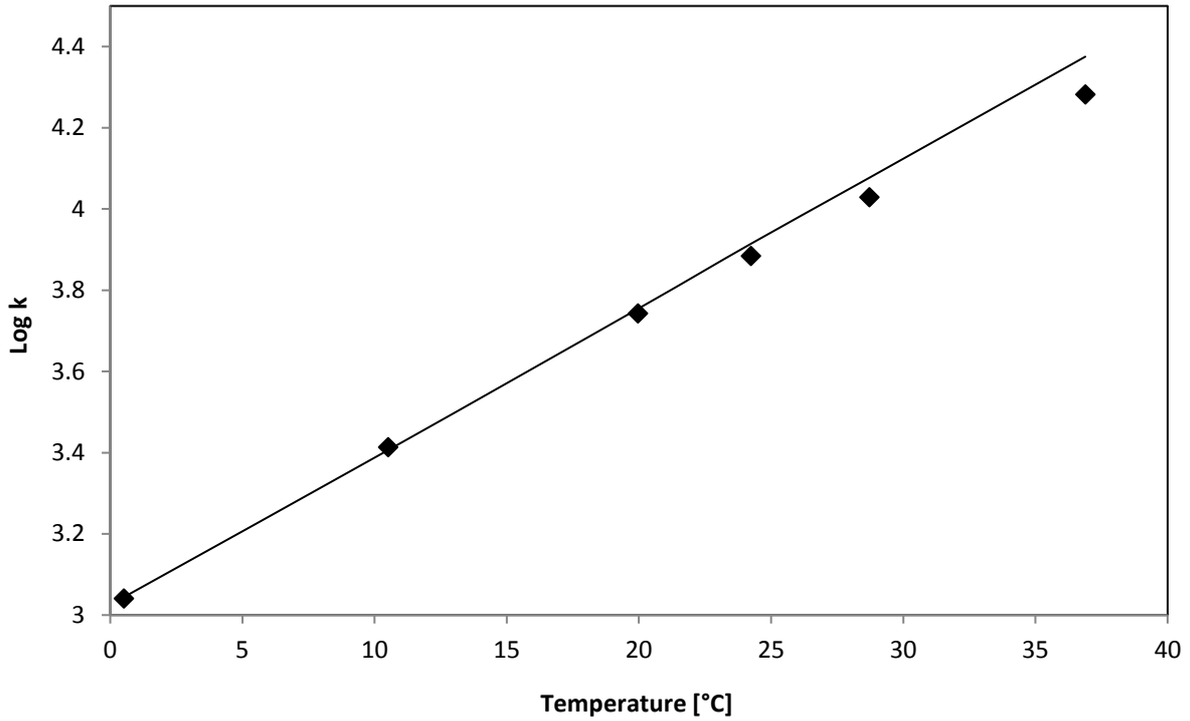


Fig. 56: Comparison of results of reaction rate constant for reaction R.24

(—) Aspen Plus® ; (◆) Experimental values [120]

The deviations between Arrhenius equation (88) and experimental results are depicted in Fig. 57.

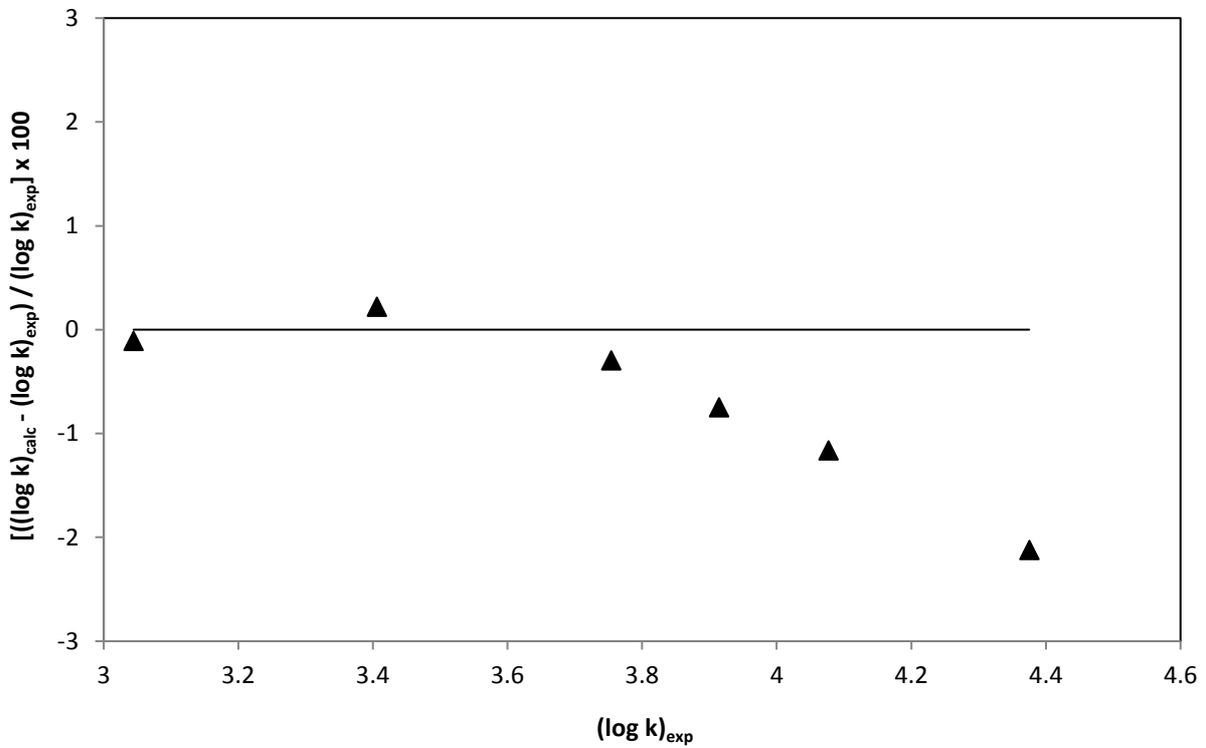


Fig. 57: Deviation between experimental and calculated results of reaction rate constant (R.24)

6.2.7. Simulation results

After validation of physicochemical parameters, Aspen Plus® simulations of the packed absorption column of the pilot plant have been performed using the Rate-based model.

Details of the column used in the experiments, as well as the description of the given gas and liquid inlets are defined in Table 51.

The type of packing used for this comparison is Flexipac® 350Y. This packing is different with respect to the one used in the pilot BioGNVAL (Montz® B1-420), but no literature data are available for this last.

Table 51. Details of the simulated process

Packed column	
Diameter [m]	0.15
Type of the packing / Size / Material / Vendor	Flexipac / 500Y / Metal / KOCH
Packing height [m]	2.354
Gas inlet	
Temperature [°C]	9
Pressure [atm]	1
Mass flow rate [kg/h]	90
Volume flow rate [m ³ /h]	77
Molar composition	CH ₄ (60 %) CO ₂ (39.997 %) H ₂ S (30 ppm)
Liquid inlet	
Temperature [°C]	4
Pressure [atm]	1
Mass flow rate [kg/h]	420
Composition	Water with NaOH (0.5 g/l)

When designing a packed column, it is desired to minimize the flow of liquid to reduce the consumption of water and the energy needed by the pump for its circulation. However, the flow must allow the absorption of H₂S and reduce its content to less than 1 ppm. This is a very important parameter since it has an influence on the thermodynamic and hydrodynamic conditions.

In 2007, Sanchez et al. showed that with a fixed air flow, an increase in liquid flow rate will improve the velocity of transfer [121].

Fig. 58 obtained using Aspen Plus® demonstrates that increasing the liquid flow improves the absorption of H₂S. However, a mass flow rate of 240 kg/h is sufficient to eliminate the totality of hydrogen sulfide with a minimal pressure loss.

As shown in Fig. 58, chemical conditions strongly influence the transfer percentage. The absorption rate increases with the concentration of sodium hydroxide in the liquid phase.

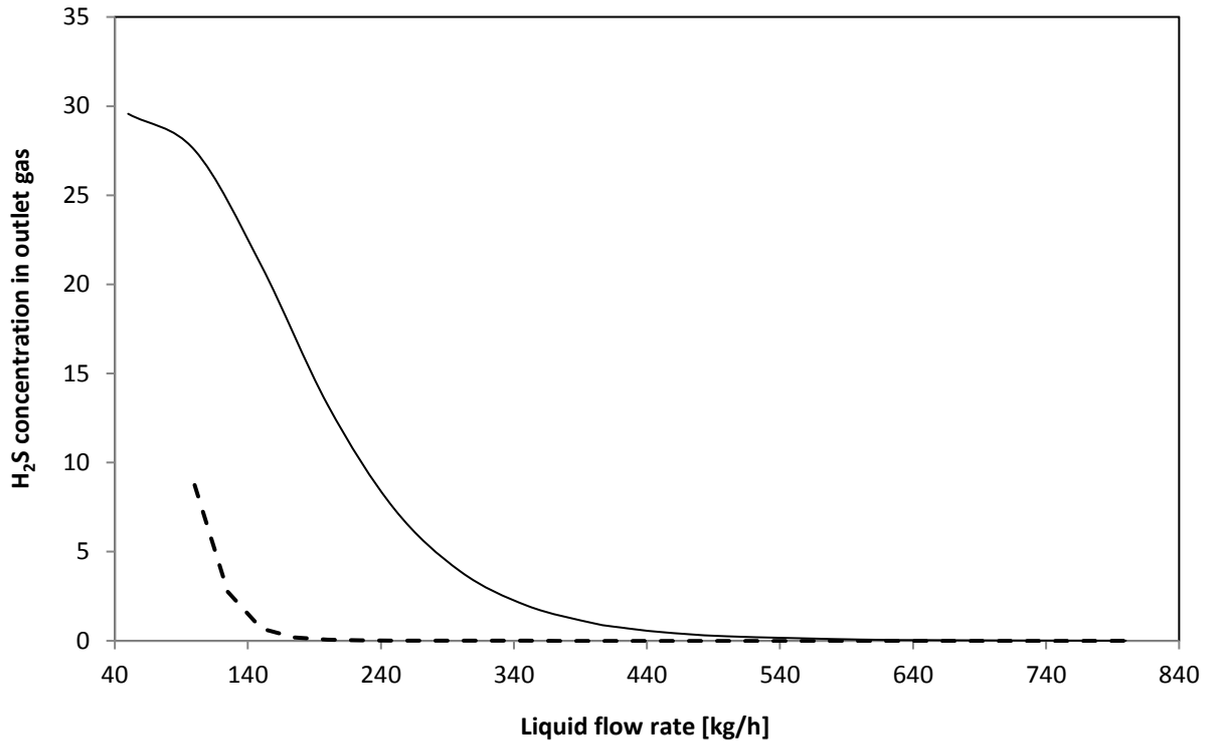


Fig. 58: Influence of the liquid flow on the absorption of hydrogen sulfide

Sodium hydroxide concentrations: (—) 0.5 g/l ; (- - -) 1.9 g/l

Table 52 shows that the increase in liquid flow causes a rise in pressure drop. This increase is limited, and shows that this parameter does not depend too much on the liquid flow.

The results obtained on the BioGNVAL pilot plant were compared to those from the modified Billet and Schultes model. These results confirm the precision of this model for activities of interest.

Table 52. Influence of liquid flow rate on the pressure drop

No. of point	L [kg/h]	V [kg/h]	$(\Delta P)_{exp}$ [Pa]	$(\Delta P)_{modified\ model}$ [Pa]	Absolute value of relative deviation between experimental and modified model [%]
1	1185	74.9	232	234	0.86
2	1026	77.1	217	229	5.4
3	852	74.5	204	200	1.9
4	766	73.0	180	187	3.8
5	686	75.0	182	191	5.1
6	423	77.0	204	186	8.7
7	329	78.9	211	190	9.8

The dependence between the pressure drop and the gas flow rate is much more important as shown in Fig. 59, because the gas has the continuous phase in the packed column.

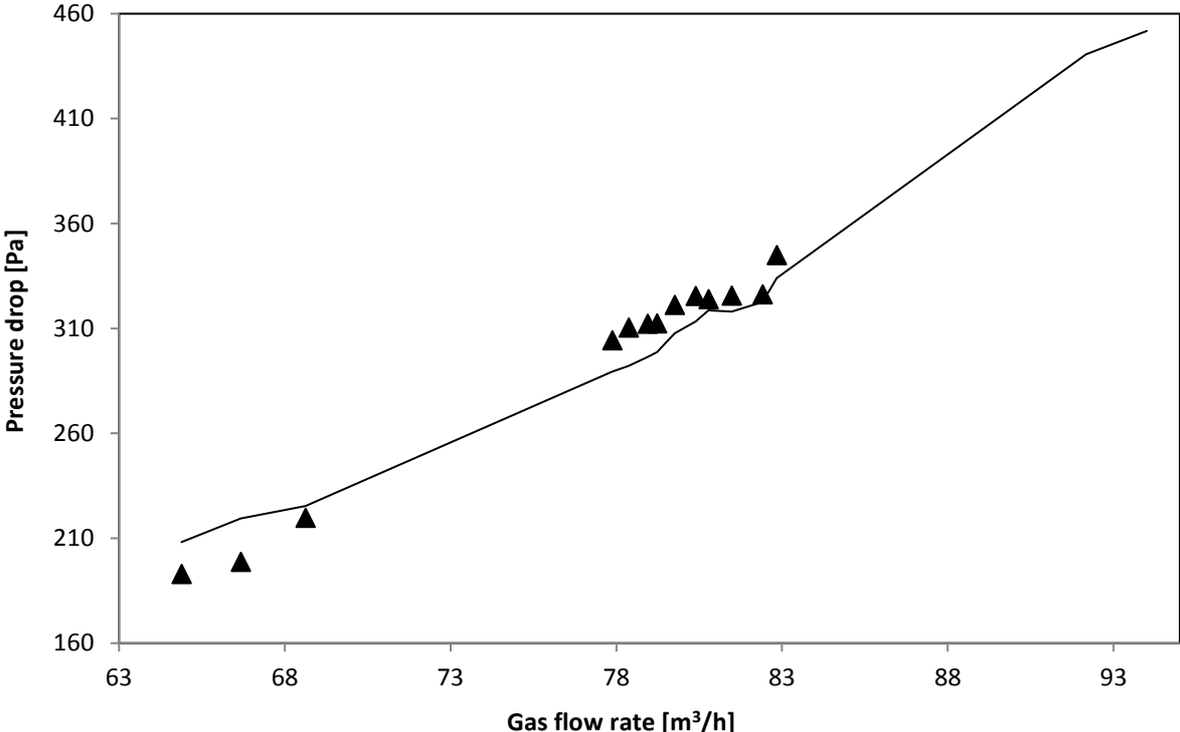


Fig. 59: Influence of gas flow rate on the pressure drop

(—) Modified Billet and Schultes model ; (♦) Experimental values [BioGNVAL, 2015]

The deviations between experimental and calculated results of pressure drop are presented in Fig. 60.

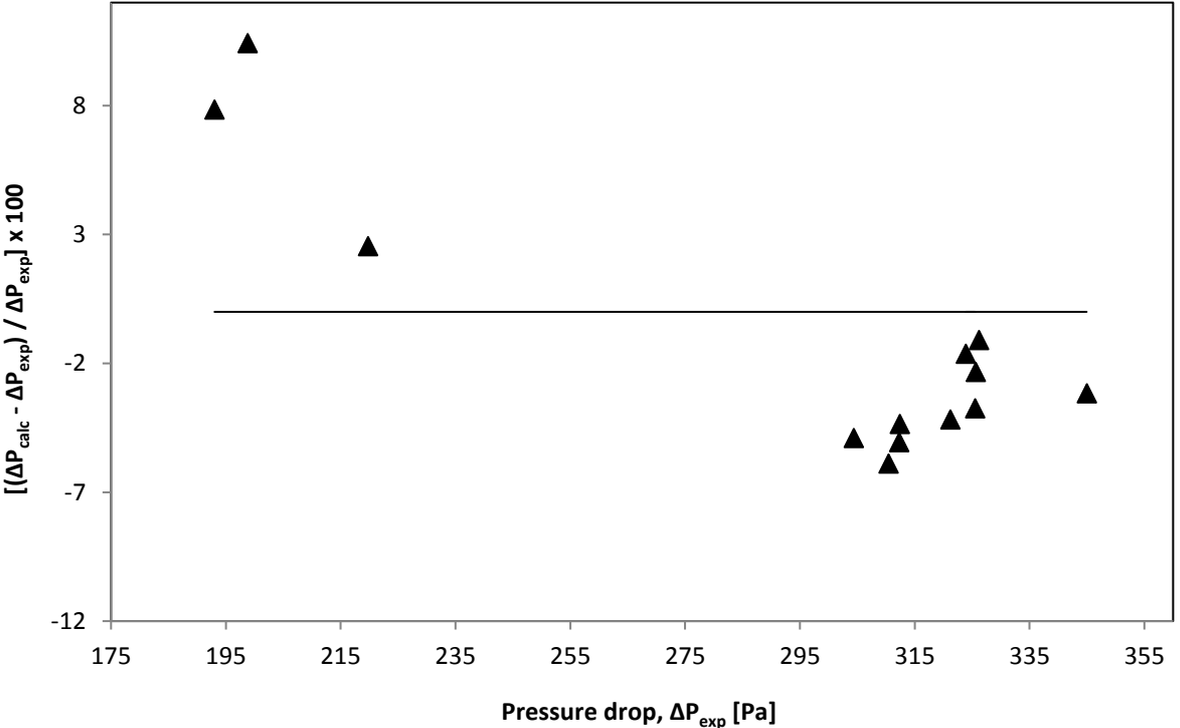


Fig. 60: Deviation between experimental and calculated results of pressure drop

When hydrogen sulfide is absorbed into a sodium hydroxide solution, it can react directly with hydroxyl ions by a proton transfer reaction as seen in Reaction R.18. Compared to the diffusion phenomena, this reaction is extremely rapid and can be considered instantaneous.

Since hydrogen sulfide is absorbed more rapidly than carbon dioxide by aqueous sodium hydroxide solutions, partial selectivity can be attained when both gases are present as seen in Fig. 61. Selectivity is favored by short gas-liquid contact times and low temperatures [36].

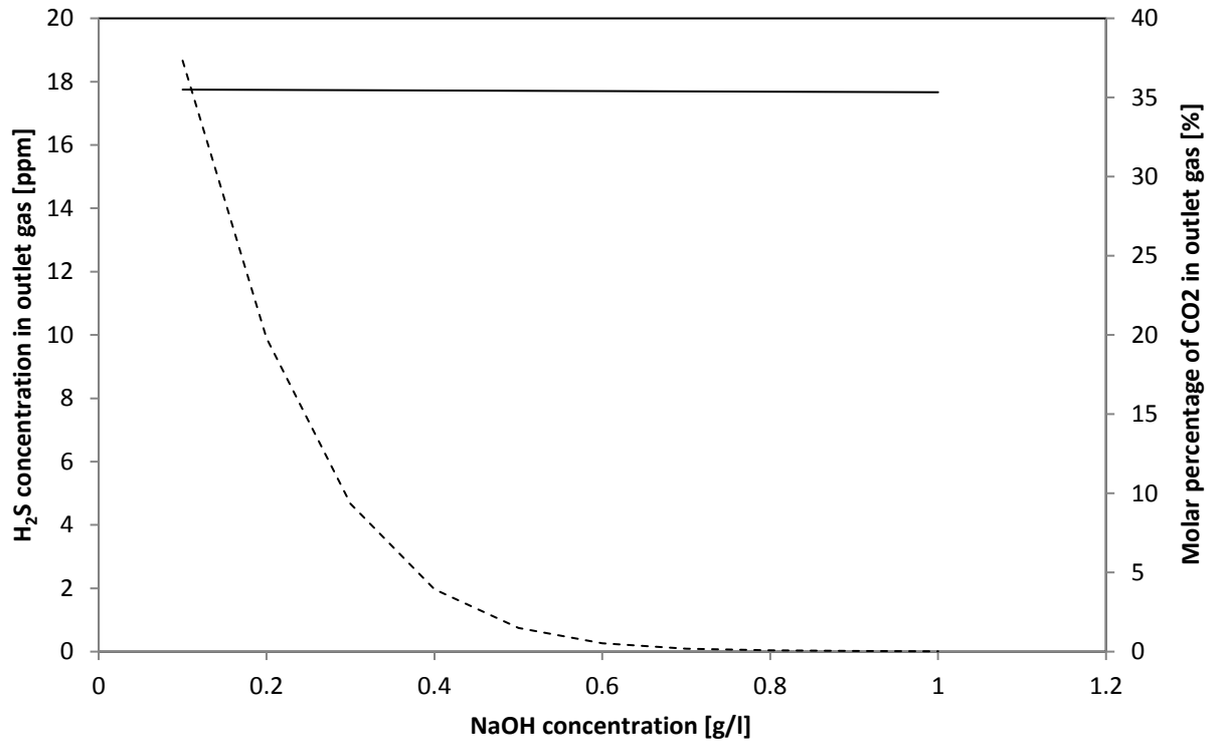


Fig. 61: Influence of the concentration of NaOH in the removal of H₂S and CO₂

Compound: (—) CO₂; (- - -) H₂S

A key parameter affecting the overall performances of the absorption unit is the temperature, since it affects physicochemical properties (such as the solubility of acidic compounds in the aqueous phase, according to the Henry's law) and the chemical reactions in the liquid phase.

Fig. 62 shows a good agreement between the results of Aspen Plus® simulations and those obtained from the pilot plant.

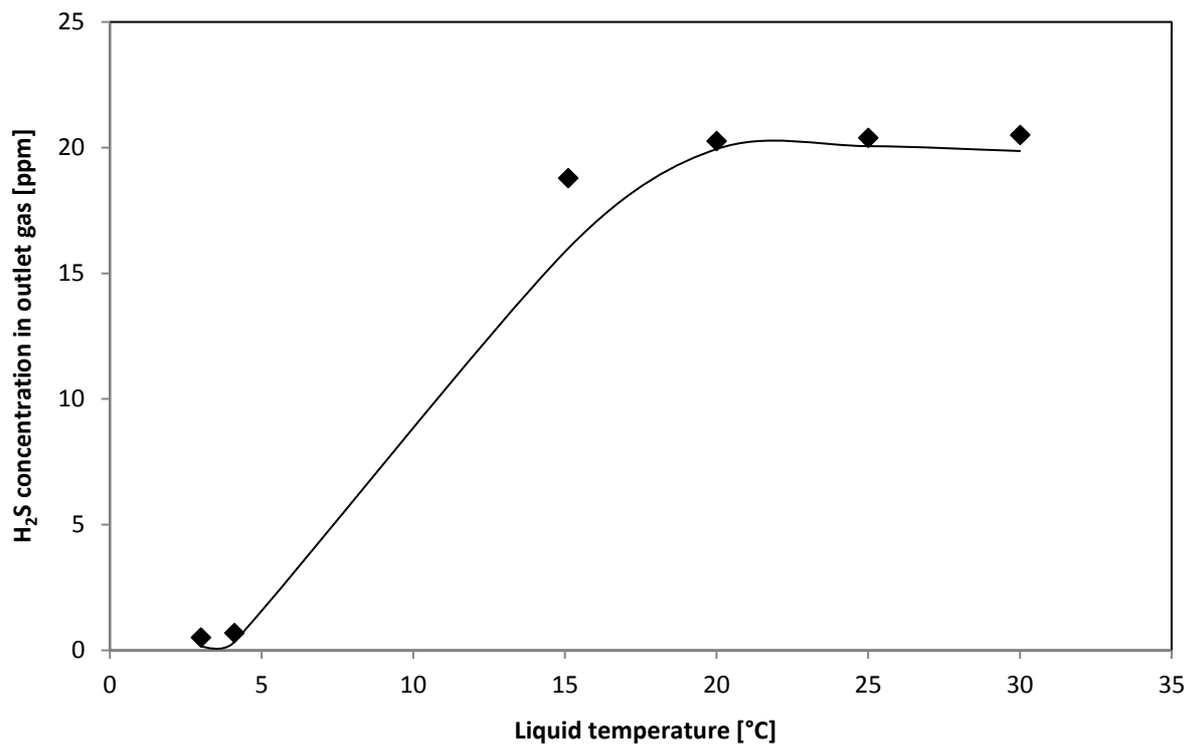


Fig. 62: Influence of the temperature of liquid in the absorption of hydrogen sulfide

(—) Aspen Plus® ; (♦) Experimental values [BioGNVAL, 2015]

The deviations between experimental and calculated results of hydrogen sulfide concentration at the outlet of the column are depicted in Fig. 63.

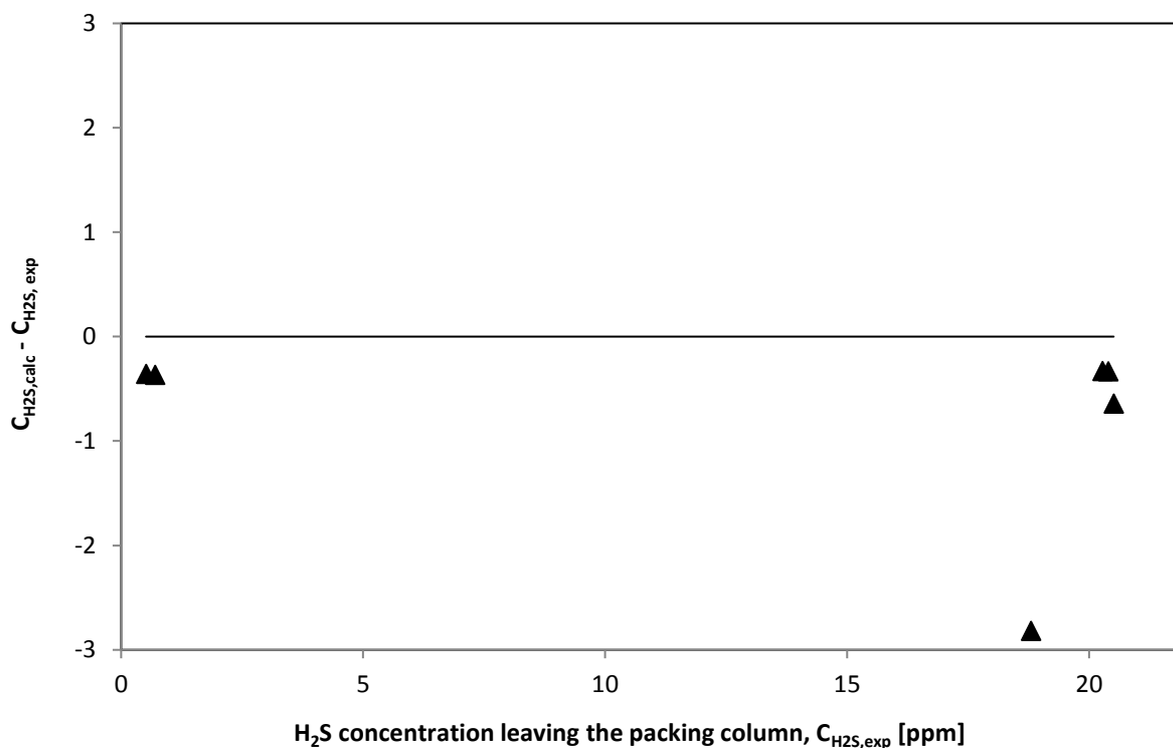


Fig. 63: Deviation between experimental and calculated results of H₂S concentration leaving the packing column

6.3. Conclusion

The thermodynamic model used in the simulations (Electrolyte NRTL) was validated with experimental data from the literature.

Simulations were performed in order to study the influence of temperatures, chemical and hydrodynamic parameters on H₂S absorption.

The simulation results were compared to experimental data obtained on the BioGNVAL pilot plant.

The comparison was successful and shows that the two results are in good agreement.

The model allows predicting realistically the separation efficiencies of H₂S in biogas.

The simulation results confirm the observations made on the demonstrator. The NaOH aqueous solution is effective for the removal of H₂S in a packing column. The removal efficiency reaches values higher than 99.5 % throughout the operation period of the demonstrator. The use of NaOCl is important to prevent the accumulation of H₂S in the aqueous solution by creating an irreversible reaction.

From a practical point of view, the use of hazardous substances (NaOH and NaOCl) requires an operator to manipulate them, which complicates the commercialization of the technology. Furthermore, salt precipitation may occur and can cause blockage of pumps and heat exchangers.

Despite the advantages of absorption technology, this system will be bypassed in order to test another promising technology that requires less financial means and which does not use hazardous chemical products. This technology is adsorption using activated carbon. It will be discussed in the next chapter.

Chapter 7: Modeling hydrogen sulfide adsorption onto activated carbon

Résumé :

Le dernier chapitre concerne la modélisation du procédé d'adsorption du H₂S sur charbon actif. Un modèle dynamique a été développé pour modéliser la courbe de percée du système H₂S – charbon actif.

La courbe de percée est utilisée pour décrire l'évolution spatio-temporelle de la concentration de H₂S en phase gazeuse. Ce type de modélisation de la dynamique de la colonne d'adsorption est basé sur la définition des bilans de matière : un bilan de masse de la phase gazeuse où le transfert par convection domine et un bilan massique des particules adsorbantes où le transfert par diffusion domine.

Au cours de cette étude, les différents coefficients de transfert de masse impliqués dans le processus ont été estimés dans les mêmes conditions dans lesquelles les expérimentations sur le démonstrateur "BioGNVAL" ont eu lieu.

Les simulations ont été effectuées seulement pour des concentrations élevées en H₂S car une unité d'adsorption industrielle qui traite de faibles concentrations en H₂S atteint la saturation dans une période de quelques mois (≈ 3 mois). La simulation de cette période avec un pas de temps de 0,01 s exige des temps de calculs énormes qui ne sont pas supportés par la machine de travail. De plus, l'augmentation du pas de temps entraînera la divergence des calculs.

Les résultats obtenus avec le modèle développé devraient être comparés aux données expérimentales afin d'ajuster le coefficient global de transfert de masse et les paramètres d'équilibre donnés par l'isotherme d'adsorption.

7.1. Introduction

This chapter deals with modeling the breakthrough curve in the case of hydrogen sulfide adsorption onto activated carbon. Physical adsorption using activated carbon is a traditional technology widely used for the removal of H₂S. The adsorption efficiency depends on several factors, such as relative humidity, temperature, concentration of H₂S in biogas and characteristics of the activated carbon. To improve efficiency, the activated carbon may be impregnated with sodium hydroxide (NaOH), potassium hydroxide (KOH), sodium carbonate (Na₂CO₃), sodium bicarbonate (NaHCO₃), potassium iodide (KI) or potassium permanganate (KMnO₄).

7.2. Operating conditions of the adsorption column for the removal of hydrogen sulfide

The adsorbent chosen for this study is activated carbon impregnated with a base (NaOH or KOH) dedicated to the elimination of hydrogen sulfide. The selected commercial adsorbent called “*Airpel Ultra DS*” is provided by Desotec® Company. Its properties are shown in Table 53.

Table 53. Activated carbon properties used for this study [53]

Properties	Unit	Value
Shape	-	Pellet or cylindrical shaped
Particle density	kg/m ³	460
Particle size	mm	4
BET surface area	m ² /g	1134
Micropore volume	cm ³ /g	0.48

The extruded activated carbon was chosen because it causes a lower pressure drop than granular and powder activated carbons.

The micropore volume is an important parameter for the kinetics of hydrogen sulfide adsorption. Compared to the other commercial adsorbents, the “*Airpel Ultra DS*” contains a high volume of micropores.

The hydrogen sulfide adsorption capacity is also influenced by other operating conditions. One of the important parameters is the relative humidity. As seen in Fig. 64 presenting the breakthrough curve of H₂S, the best hydrogen sulfide adsorption capacities are obtained for values of relative humidity between 55 % and 100 % with an optimal value of 85 %.

The breakthrough curve presents the evolution of concentration of the pollutant to be removed (H₂S) as a function of time. It predicts the time required for the saturation of the activated carbon in the adsorption bed.

The adsorption using impregnated activated carbon is improved by injection of a small quantity of oxygen. The amount injected is generally of the order of 4 times the amount of hydrogen sulfide present in the biogas. The oxidation of hydrogen sulfide will make larger molecules to block them in the micropores.

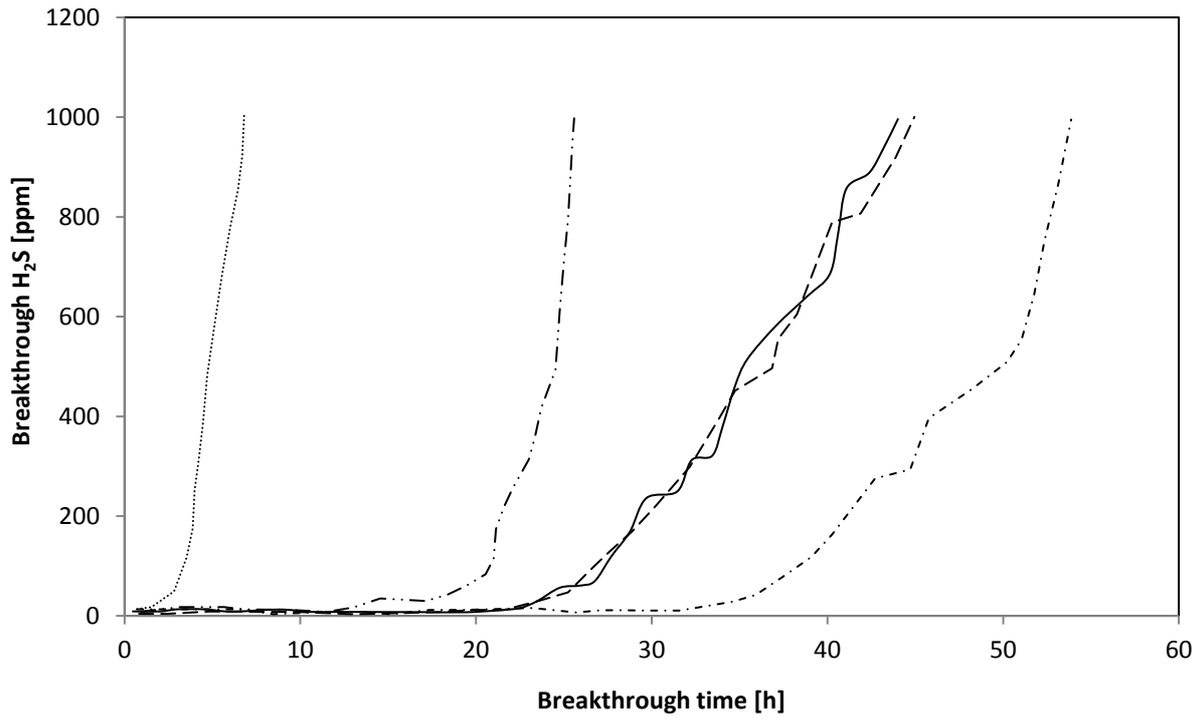


Fig. 64: Influence of relative humidity in the adsorption of hydrogen sulfide using Airpel Ultra DS [53]

Relative humidity: (.....) 0 % ; (-) 33 % ; (- - - -) 58 % ; (-) 85 % ; (—) 100 %

7.3. Breakthrough curve modeling

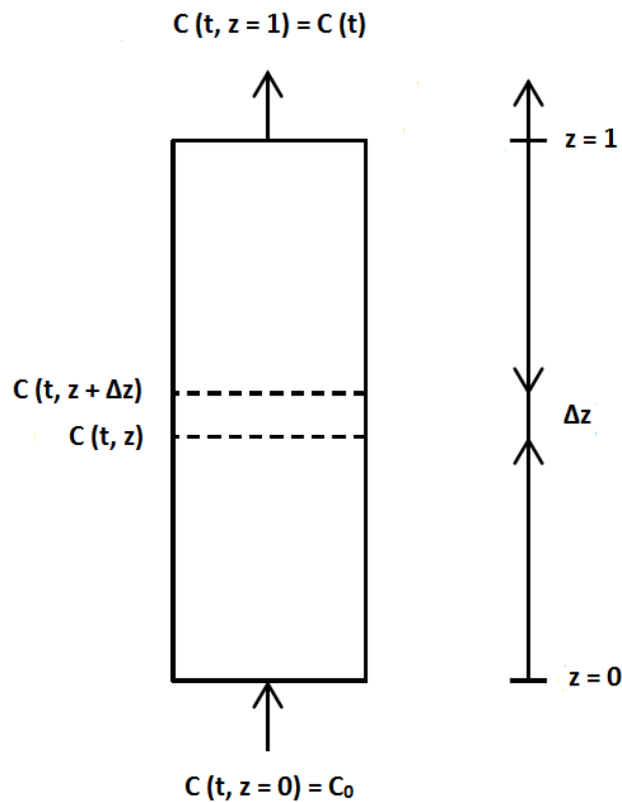


Fig. 65: Schematic representation of an adsorption column [122]

The operating conditions used in the model are the same measured on the BioGNVAL pilot plant, in order to adjust the estimated parameters, such as the overall mass transfer coefficient and the maximum adsorption capacity of activated carbon.

These operating conditions include the temperature and the pressure of the process, the adsorption column dimensions, the relative humidity of biogas and its composition. They are presented in Table 54.

Table 54. Biogas composition and operating conditions of the adsorption process

Properties	Unit	Value
Molar biogas composition	mol%	CH ₄ (60) CO ₂ (37) N ₂ (1.88) H ₂ O (1.11) H ₂ S (0.01) ≈ 100 ppm
Volume flow rate of biogas	Nm ³ /h	85
Inlet pressure	bar	1.103
Inlet temperature	°C	21
Relative humidity	%	50
Adsorption column diameter	m	0.8
Height of the adsorption column	m	1

The adsorption mechanism is governed by different types of mass transfer: external and internal diffusion. That's why, before modeling the breakthrough curve, mass transfer coefficients should be correctly estimated.

7.3.1. Estimation of mass transfer coefficients

The estimation of mass transfer coefficients is an essential step in order to simulate the dynamic behavior of adsorption.

- **External mass transfer coefficient**

The evaluation of the external mass transfer coefficient depends on the H₂S molecular diffusion coefficient, the activated carbon particle size and dimensionless numbers as seen in Equation (101).

$$k_D = \frac{Sh D_{H_2S-Biogas}}{d_p} \quad (101)$$

The particle size is indicated in Table 53.

The Sherwood number (Sh) estimation is based on correlations using Reynolds (Re) and Schmidt (Sc) numbers. Some correlations defining the Sherwood number are presented in Table 55.

Table 55. Sherwood number estimation [123]

Correlations	Application range	References	
$Sh = 2 + 1.1 Re^{0.6} Sc^{\frac{1}{3}}$	$3 < Re < 10^4$	[124]	(102)
$Sh = 2 + 0.6 Re^{0.5} Sc^{\frac{1}{3}}$		[125]	(103)
$Sh = 1.65 Re^{0.49} Sc^{\frac{1}{3}}$		[126]	(104)
$Sh = \left(\frac{1}{\varepsilon}\right) [0.765 (\varepsilon Re)^{0.18} + 0.365 (\varepsilon Re)^{0.614}] Sh^{\frac{1}{3}}$	$Re < 15000$	[127]	(105)
$Sh = 1.85 \left(\frac{1-\varepsilon}{\varepsilon}\right)^{\frac{1}{3}} Re^{\frac{1}{3}} Sc^{\frac{1}{3}}$	$\left(\frac{\varepsilon}{1-\varepsilon}\right) Re \leq 100$	[128]	(106)
$Sh = 2 \alpha \phi^2 \left[\frac{1 - (1-\varepsilon)^{\frac{1}{3}}}{2(1-\varepsilon)^{\frac{1}{3}}} + \phi \right] \sqrt{\frac{Pe_p}{\pi \varepsilon \tau}}$		[129]	(107)
$Sh = \left[2 + \sqrt{(Sh_{lam}^2 + Sh_{turb}^2)} \right] [1 + 1.5(1-\varepsilon)]$ $Sh_{lam} = 0.644 Re^{0.5} Sc^{\frac{1}{3}}$ $Sh_{turb} = \frac{0.037 Re^{0.8} Sc}{1 + 2.443 Re^{-0.1} (Sc^{\frac{2}{3}} - 1)}$	$1 < Re < 10^4$ $0.6 < Sc < 10^4$	[130]	(108)
$Sh = 1.09 \varepsilon^{-0.66} Re^{\frac{1}{3}} Sc^{\frac{1}{3}}$	$0.0015 \leq \varepsilon Re \leq 55$ $950 < Sc < 70000$	[131]	(109)
$Sh = 0.25 \varepsilon^{-0.31} u_V Re^{0.69} Sc^{\frac{1}{3}}$	$55 \leq \varepsilon Re \leq 1050$ $950 < Sc < 70000$	[131]	(110)
$Sh = 2 + 1.58 Re^{0.4} Sc^{\frac{1}{3}}$	$10^{-3} \leq Re \leq 5.8$	[132]	(111)
$Sh = 2 + 1.21 Re^{0.5} Sc^{\frac{1}{3}}$	$5.8 \leq Re \leq 500$	[132]	(112)
$Sh = 2 + 0.59 Re^{0.6} Sc^{\frac{1}{3}}$	$Re \geq 500$	[132]	(113)

Reynolds and Schmidt numbers are calculated using Equations (114) and (115) respectively.

$$Re = \frac{\rho_V u_V d_p}{\mu_V} \quad (114)$$

$$Sc = \frac{\mu_V}{\rho_V D_{H_2S-Biogas}} \quad (115)$$

In Expression (107) defined by Doytchava et al. [129], α is the available surface coefficient, ϕ is the particle shape factor, τ is the tortuosity and Pe_p is the Peclet number of the particle calculated using Equation (116).

$$Pe_p = \frac{d_p u_V}{D_{H_2S-Biogas}} \quad (116)$$

The H₂S molecular diffusion coefficient is calculated using The Equation (117) proposed by Wilke and Fairbanks [133]. This equation is derived from the theories of Maxwell and Stefan.

$$D_m = \frac{1 - y_{H_2S}}{\frac{y_{CH_4}}{D_{H_2S-CH_4}} + \frac{y_{CO_2}}{D_{H_2S-CO_2}} + \frac{y_{N_2}}{D_{H_2S-N_2}} + \frac{y_w}{D_{H_2S-w}}} \quad (117)$$

Equations (118), (119), (120) and (121) are used to estimate the diffusivities for the different binary systems at low pressure. These equations developed by Slattery and Bird [134] from a kinetic theory are shown in Table 56.

Table 56. Estimation of the binary diffusion coefficients [135]

Binary systems	Correlations	
H ₂ S – CH ₄	$D_{H_2S-CH_4} = \frac{2.745 \times 10^{-4} (P_{c,H_2S} P_{c,CH_4})^{\frac{1}{3}} (T_{c,H_2S} T_{c,CH_4})^{\frac{5}{12}} \left(\frac{1}{M_{H_2S}} + \frac{1}{M_{CH_4}} \right)^{0.5}}{P} \left(\frac{T}{\sqrt{T_{c,H_2S} T_{c,CH_4}}} \right)^{1.823}$	(118)
H ₂ S – CO ₂	$D_{H_2S-CO_2} = \frac{2.745 \times 10^{-4} (P_{c,H_2S} P_{c,CO_2})^{\frac{1}{3}} (T_{c,H_2S} T_{c,CO_2})^{\frac{5}{12}} \left(\frac{1}{M_{H_2S}} + \frac{1}{M_{CO_2}} \right)^{0.5}}{P} \left(\frac{T}{\sqrt{T_{c,H_2S} T_{c,CO_2}}} \right)^{1.823}$	(119)
H ₂ S – N ₂	$D_{H_2S-N_2} = \frac{2.745 \times 10^{-4} (P_{c,H_2S} P_{c,N_2})^{\frac{1}{3}} (T_{c,H_2S} T_{c,N_2})^{\frac{5}{12}} \left(\frac{1}{M_{H_2S}} + \frac{1}{M_{N_2}} \right)^{0.5}}{P} \left(\frac{T}{\sqrt{T_{c,H_2S} T_{c,N_2}}} \right)^{1.823}$	(120.a)
	$D_{H_2S-N_2} = \frac{10^{-7} T^{1.75} \sqrt{\frac{1}{M_{H_2S}} + \frac{1}{M_{N_2}}}}{P \left(V_{m,H_2S}^{1/3} + V_{m,N_2}^{1/3} \right)^2}$	(120.b)
H ₂ S – H ₂ O	$D_{H_2S-H_2O} = \frac{3.64 \times 10^{-4} (P_{c,H_2S} P_{c,H_2O})^{\frac{1}{3}} (T_{c,H_2S} T_{c,H_2O})^{\frac{5}{12}} \left(\frac{1}{M_{H_2S}} + \frac{1}{M_{H_2O}} \right)^{0.5}}{P} \left(\frac{T}{\sqrt{T_{c,H_2S} T_{c,H_2O}}} \right)^{2.334}$	(121)

- **Internal mass transfer coefficient**

The hydrogen sulfide molecules are now located on the surface of the activated carbon. The diffusion mechanisms govern the transport of hydrogen sulfide molecules into the pores of activated carbon. In porous materials, diffusion mechanisms are of four types: molecular diffusion caused by collisions between molecules, Knudsen diffusion caused by collisions of the molecules with the walls

of the pore, surface diffusion caused by the electrostatic forces exerted by the walls on the molecules and Poiseuille diffusion caused by the difference in total pressure across a particle. The equations used to estimate these diffusivities are listed in Table 57.

Table 57. Equations used to estimate internal diffusion coefficients

Type of diffusion mechanism	Equation	
Knudsen diffusion	$D_{Knudsen} = 97 r_{pore} \sqrt{\frac{T}{M_{H_2S}}}$	(122)
Surface diffusion	$D_s = 1.1 \times 10^{-8} e^{\left(\frac{-5.32 T_{B,H_2S}}{T}\right)}$	(123)
Poiseuille diffusion	$D_{Poiseuille} = \frac{P r_{pore}^2}{8 \mu_{H_2S}}$	(124)
Pore diffusion (Global internal diffusion)	$\frac{1}{D_p} = \frac{1}{D_m} + \frac{1}{D_{Knudsen}}$	(125)

The values obtained for the dimensionless numbers and the diffusion coefficients are presented in Table 58.

Table 58. Estimation of dimensionless numbers and diffusion coefficients

Properties	Unit	Value
Reynolds number, Re	-	17
Schmidt number, Sc	-	0.81
Sherwood number, Sh	-	7.66
Peclet number, Pe	-	13.77
Molecular diffusivity	m ² /s	1.35 x 10 ⁻⁵
Knudsen diffusivity	m ² /s	1.84 x 10 ⁻⁷
Surface diffusivity	m ² /s	2.34 x 10 ⁻¹⁰
Poiseuille diffusivity	m ² /s	4.35 x 10 ⁻¹⁰
Pore diffusivity	m ² /s	1.81 x 10 ⁻⁷

As seen in Table 58, the molecular diffusion is the dominant transport mechanism. The Poiseuille diffusivity can be neglected because the pressure drop over a particle is very small [49].

Knudsen diffusion and surface diffusion contribute to global internal diffusion. This contribution is very small compared to the contribution of external diffusion. Therefore, only external diffusion will be considered.

The external mass transfer coefficient calculated using Equation (101) is equal to $0.021 \text{ m}\cdot\text{s}^{-1}$.

7.3.2. Modeling of breakthrough curves

The breakthrough curve is used to describe the spatiotemporal evolution of the concentration of H_2S in the gas phase. This type of modeling the dynamics of the adsorption column is based on the definition of mass transfer balances: a mass balance of the gas phase where the transfer by convection dominates and a mass balance of the adsorbent particles where the diffusion transfer dominates [122].

Several hypotheses govern the flow of the biogas in the absorption column: The properties and the superficial velocity of the biogas are assumed constant throughout the adsorption column and the porosity of the adsorption column is considered uniform [136].

Based on the assumptions outlined, the mass balance of the adsorption column can be equated as seen in Equation (126). This equation has been used by several authors as Ruthven [49], Suzuki [137], Hwang et al. [138], Brosillon et al. [139], Yang [140], Boulinguez [136] and Sigot [122].

$$\frac{-D_L}{L^2} \frac{\partial^2 C}{\partial z^2} + \frac{u}{L} \frac{\partial C}{\partial z} + \frac{\partial C}{\partial t} + \frac{\rho}{\varepsilon} \frac{\partial q}{\partial t} = 0 \quad (126)$$

Where:

$D_L [\text{m}^2\cdot\text{s}^{-1}]$ is the axial dispersion coefficient.

$C [\text{mol}\cdot\text{m}^{-3}]$ is the concentration of H_2S in the biogas.

$L [\text{m}]$ is the length of the adsorption column.

$z [\text{m}]$ is the axial dimension.

$t [\text{s}]$ is the time.

$u [\text{m}\cdot\text{s}^{-1}]$ is the superficial velocity of the biogas.

$\rho [\text{kg}\cdot\text{m}^{-3}]$ is the density of activated carbon.

$\varepsilon [-]$ is the void fraction of the adsorption column.

$q [\text{g}\cdot\text{kg}^{-1}]$ is the amount of H_2S adsorbed per kg of activated carbon.

The mass balance Equation (126) takes into account the axial dispersion, convection, the accumulation of the gas phase and the overall mass transfer by adsorption [136].

To simplify the problem, initially the axial diffusion coefficient is neglected. The mass balance presented by Equation (126) is simplified.

$$\frac{u}{L} \frac{\partial C}{\partial z} + \frac{\partial C}{\partial t} + \frac{\rho}{\varepsilon} \frac{\partial q}{\partial t} = 0 \quad (127)$$

Considering a single transfer resistance material represented by the linear driving force model, the flux transferred may then be expressed relative to the solid phase concentration as shown in Equation (128), or relative to the gas phase concentration as shown in Equation (129).

$$\frac{\partial q}{\partial t} = k_g a_p (q_e - q) \quad (128)$$

$$\frac{\partial q}{\partial t} = k_g a_p \frac{1 - \varepsilon}{\rho} (C - C_e) \quad (129)$$

Where:

k_g [$\text{m}\cdot\text{s}^{-1}$] is global mass transfer coefficient.

a_p [$\text{m}^2\cdot\text{m}^{-3}$] is the external surface of the adsorbent particle per unit volume of adsorbent.

q_e [$\text{g}\cdot\text{kg}^{-1}$] is the amount of H_2S in equilibrium with the gas phase concentration.

C_e [$\text{g}\cdot\text{m}^{-3}$] is the H_2S concentration in the gas phase, in equilibrium with the solid phase.

The adsorption equilibrium between the two phases is defined by means of the Langmuir isotherm Equation (type I).

$$q = \frac{q_m b C_e}{1 + b C_e} \quad (130)$$

Where q_m and b are the Langmuir parameters.

Fig. 66 shows that the equilibrium parameters are sensitive parameters. They have a great influence on the simulation results. These parameters must be adjusted with experimental data.

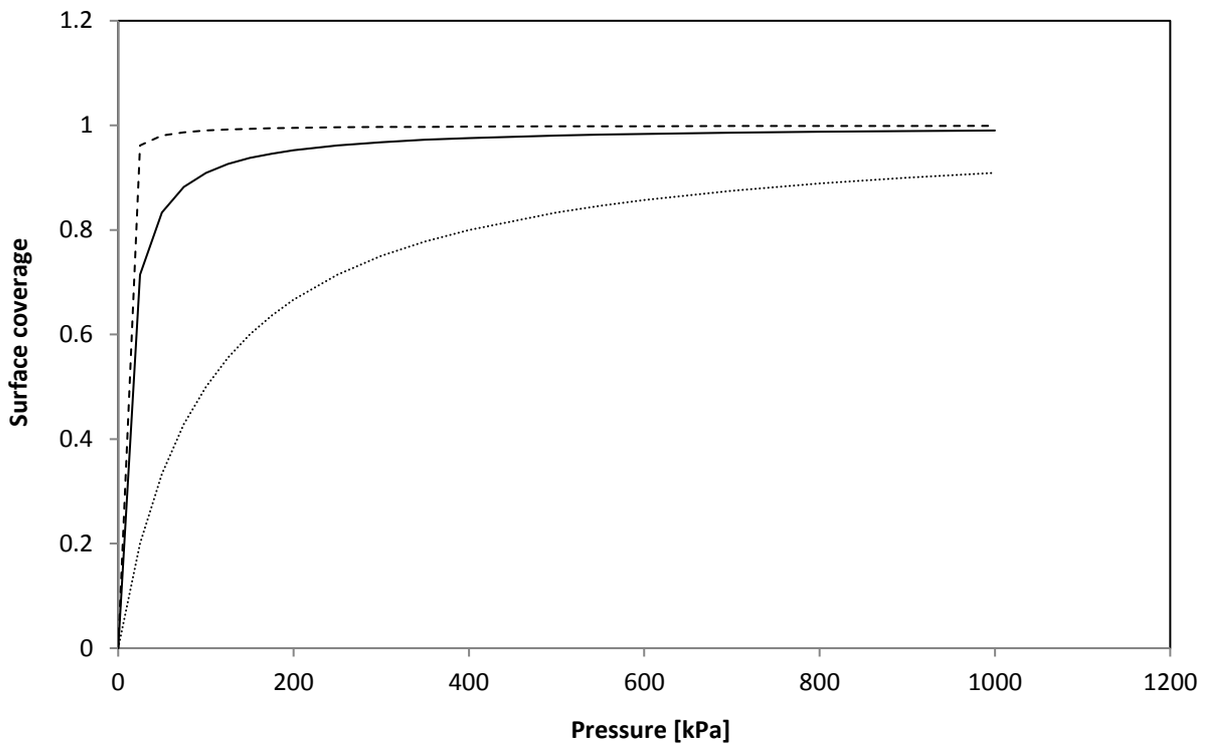


Fig. 66: Influence of Langmuir equilibrium parameters on adsorption sites occupied

(.....) $b = 0.01$; (—) $b = 0.1$; (- - -) $b = 1$

Boundary and initial conditions are summarized in Table 59.

Table 59. Boundary and initial conditions of the system

Condition type	Equation	
Boundary condition – 1	$C(t, z = 0) = C_0$	(131)
Boundary condition – 2	$\left(\frac{\partial C}{\partial z}\right)_{z=1} = 0$	(132)
Initial condition – 1	$C(t = 0, z = 0) = C_0$	(133)
Initial condition – 2	$C(t = 0, z > 0) = 0$	(134)
Initial condition – 3	$q(t = 0, z) = 0$	(135)

By combining the equation (127) and (129), the expression (136) is obtained [122].

$$\frac{u}{L} \frac{\partial C}{\partial z} + \frac{\partial C}{\partial t} + \frac{1 - \varepsilon}{\varepsilon} k_g a_p (C - C_e) = 0 \quad (136)$$

Similarly, the expression (137) is obtained by combining equations (129) and (130).

$$\frac{\partial C_e}{\partial t} = \frac{k_g a_p (1 - \varepsilon)}{\rho q_m b} (1 + b C_e)^2 (C - C_e) \quad (137)$$

To simplify the system of equations (136) and (137), three time constants associated to mass transfer τ_1 , τ_2 and τ_3 [s⁻¹] are highlighted. The system of equation becomes [122]:

$$\frac{1}{\tau_1} \frac{\partial C}{\partial z} + \frac{\partial C}{\partial t} + \frac{1}{\tau_2} (C - C_e) = 0 \quad (138)$$

$$\frac{\partial C_e}{\partial t} = \frac{1}{\tau_3} (1 + b C_e)^2 (C - C_e) \quad (139)$$

Where:

$$\frac{1}{\tau_1} = \frac{u}{L} \quad (140)$$

$$\frac{1}{\tau_2} = \frac{1 - \varepsilon}{\varepsilon} k_g a_p \quad (141)$$

$$\frac{1}{\tau_3} = \frac{k_g a_p (1 - \varepsilon)}{\rho q_m b} \quad (142)$$

In order to solve the system of equations numerically, it is necessary to discretize it by means of Euler method which uses the finite difference quotient. The system of equations thus obtained is given by the Expressions (143) and (144). The boundary and initial conditions are given respectively by Equations (145) and (146) [122].

$$C_i^{j+1} = -\Delta t \left[\frac{1}{\tau_1} \left(\frac{C_i^j - C_{i-1}^j}{\Delta z} \right) + \frac{1}{\tau_2} (C_i^j - C_{e,i}^j) \right] + C_i^j \quad (143)$$

$$C_{e,i}^{j+1} = \frac{\Delta t}{\tau_3} (1 + b C_{e,i}^j)^2 (C_i^j - C_{e,i}^j) + C_{e,i}^j \quad (144)$$

$$C_1^j = C_0 \quad \text{and} \quad C_n^j = C_{n-1}^j \quad (145)$$

$$C_1^1 = C_0 \quad ; \quad C_{i>1}^1 = 0 \quad \text{and} \quad C_{e,i}^1 = 0 \quad (146)$$

After the construction of the discrete form of the analytical mass balance equations, the input parameters for the simulation of the breakthrough curve are listed in Table 60.

Table 60. Input parameters for the simulation of the breakthrough curve

Parameters	Unit	Value
Dimensions of the adsorption column		
Diameter	m	0.8
Height	m	0.8
Biogas composition and flow rate		
Methane	mol%	60
Carbon dioxide	mol%	33 / 30.5 / 28
Hydrogen sulfide	mol%	5 / 7.5 / 10
Nitrogen	mol%	0.89
Water vapor	mol%	1.11
Biogas flow rate	Nm ³ /h	80
Pressure and temperature		
Inlet pressure	bar	1.103
Inlet temperature	°C	21
Langmuir constants		
b	m ³ /g	0.9
Activated carbon maximal capacity	g/kg	110
Mesh		
Number of mesh nodes over the column height	-	20
Time step	s	0.01
Number of time steps	-	2 x 10 ⁶

The simulation was performed for high hydrogen sulfide concentrations (5, 7.5 and 10 mol%) as seen in Table 60. The breakthrough curves obtained are presented respectively in Fig. 67, 68 and 69. An industrial adsorption column with such dimensions (0.8 m in height and 0.8 m in diameter) and treating low concentrations of H₂S (< 100 ppm) reaches saturation after few months (\approx 3 months).

The simulation of such period with a time step of 0.01 s demands enormous time calculations that are not supported by the working machine. In addition, increasing the time step will cause the divergence of calculations.

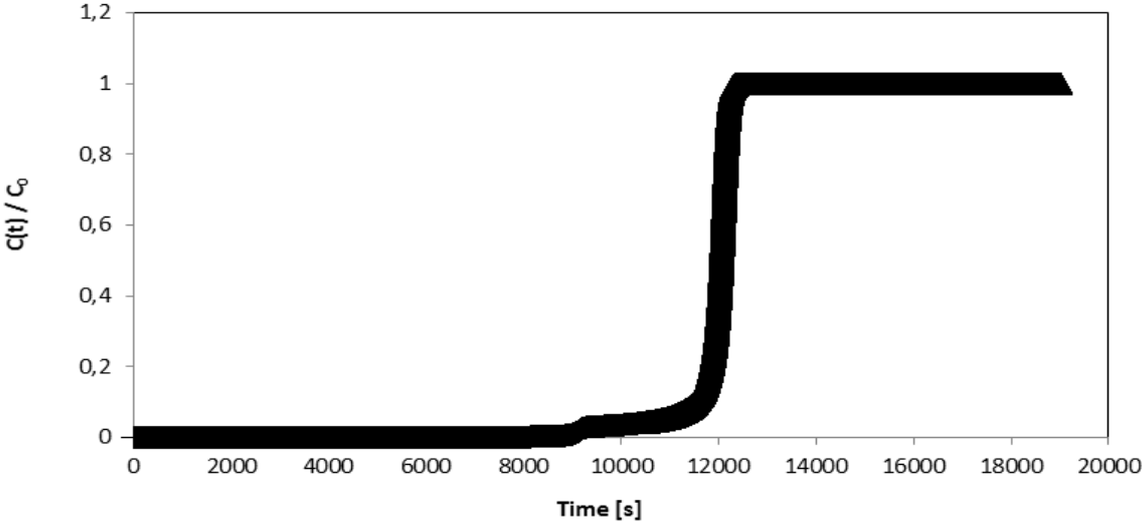


Fig. 67: Breakthrough curve simulated for H₂S molar percentage of 5 %

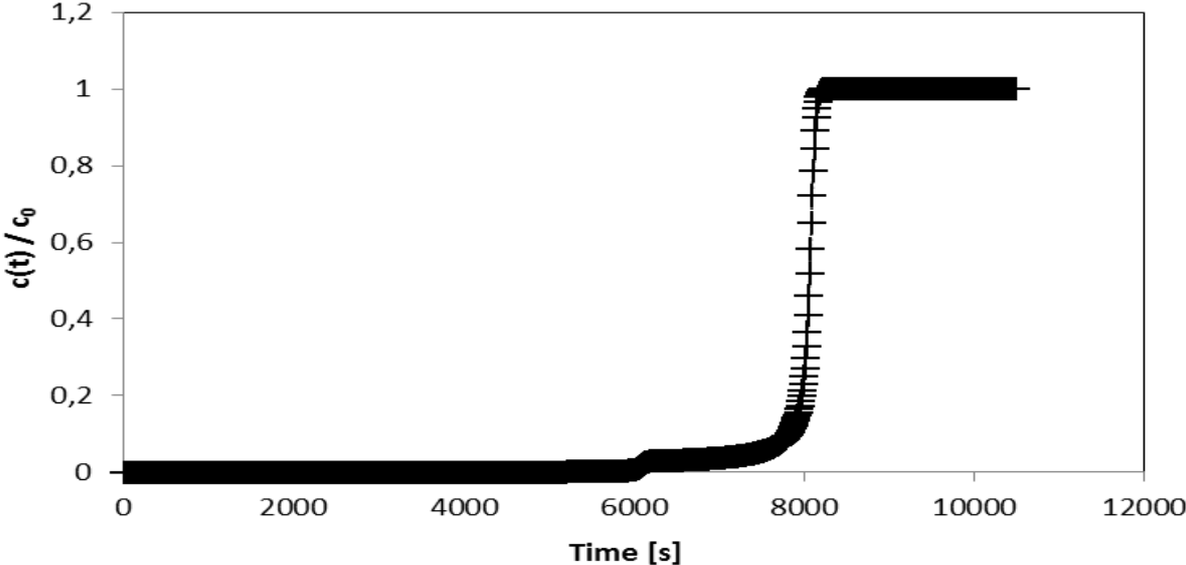


Fig. 68: Breakthrough curve simulated for H₂S molar percentage of 7.5 %

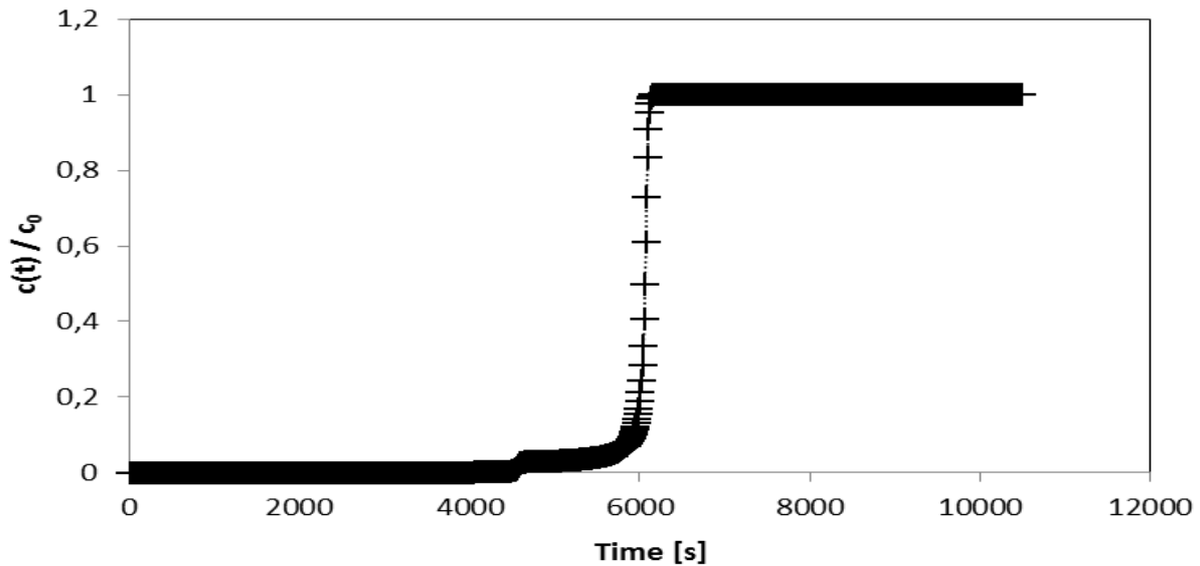


Fig. 69: Breakthrough curve simulated for H₂S molar percentage of 10 %

7.4. Conclusion

Based on the mass balance equations, a dynamic model has been developed to simulate the breakthrough curve for the system H₂S – Activated carbon. To simulate the dynamic behavior of an adsorption column for the removal of low concentrations of H₂S, a powerful calculation tool is necessary.

The results obtained with the model developed should be compared to experimental data in order to adjust the overall mass transfer coefficient and equilibrium parameters given by the Langmuir isotherm. The axial dispersion coefficient should be added in the mass balance equation to evaluate its contribution on the overall mass transfer coefficient.

This study has highlighted the analytical difficulties due to low concentrations of H₂S. It has also showed the importance of the experimental work to develop an accurate model which could be a reliable design tool for biogas purification units.

Conclusions and perspectives

This thesis is part of the innovative project led by the Cryo Pur® society that aims to develop a technology to purify and upgrade biogas in order to be used as a liquid fuel or injected into the natural gas grid. The purpose of this work is to ensure a continuous removal of hydrogen sulfide upstream of the process because of the risks of toxicity, corrosion and odors.

This thesis has allowed drawing scientific and technical conclusions.

From a technical point of view, two desulfurization technologies were tested: chemical absorption in a packing column using sodium hydroxide as an aqueous solvent and adsorption using impregnated activated carbon in a fixed bed.

Both techniques have been experimented on a demonstrator “BioGNVAL” developed by Cryo Pur® Society which treats 85 Nm³/h of real biogas from the sewage treatment plant of Valenton.

The general process is characterized by the following conditions:

- The concentrations of methane and carbon dioxide are about ≈ 60 mol% of CH₄ and ≈ 35 mol% of CO₂.
- The process operates at a pressure slightly higher than the atmospheric pressure to prevent the infiltration of oxygen.
- The desulfurization of biogas must be complete to ensure a high quality of CO₂ and biomethane.
- The concentration of H₂S in the biogas is low and generally varies between 10 and 100 ppm.

Both technologies have shown satisfactory separation efficiency greater than 99.5 %. The bibliographic study has allowed choosing an efficient absorption process that uses sodium hydroxide as a chemical solvent for the selective separation of hydrogen sulfide. This method is based on contacting the biogas to be treated with the aqueous NaOH in a structured packing column.

Despite the advantages of this process, it also has some drawbacks. For example, in the presence of CO₂, the aqueous solution of sodium hydroxide cannot be regenerated because there is formation of sodium carbonate (Na₂CO₃) precipitate. Moreover, when the tanks of NaOH and NaOCl reach their lowest level, they must be filled by an operator who has to handle hazardous products. This operation is an obstacle to the commercialization of the process which must be automated without the repetitive intervention of an operator. Nevertheless for high H₂S concentrations for processes which are continuously operated with personnel, this absorption process can be adopted.

To overcome these drawbacks for fully automated biogas purification equipment, the adsorption technology using impregnated activated carbon was tested. It has achieved similar separation efficiencies to the absorption technology without the use of hazardous products and without the need for an operator to ensure continuous operation of the process. Furthermore, the investment and operating costs of the adsorption process are lower than those of the absorption process.

From a scientific point of view, experimental data collected on the BioGNVAL pilot plant were used to develop a new hydrodynamic model which predicts accurately the key hydrodynamic parameters in a structured packing column: liquid holdup, pressure drop, effective interfacial area and the two transition points: loading and flooding points. It also allows the design and optimization of structured packing columns to operate at full capacity.

This model was developed based on an existing model: Billet and Schultes. Some constants and exponents present in the correlations of this last model were developed following a sensitivity analysis and then optimized based on experimental data to finally implement a new model adapted to predict precisely the hydrodynamic parameters of a small scale structured packing column used for biogas or natural gas applications.

After the hydrodynamic study, simulations using the chemical process optimization software Aspen Plus® V8.0 have been performed in order to determine the efficiencies of separation for different operating conditions.

The rate-based model approach has been used. To describe vapor-liquid equilibrium, a γ/ϕ approach has been considered: the Electrolyte Non-Random Two-Liquid (ENRTL) model has been adopted to represent non-idealities in the liquid phase, while the Redlich-Kwong equation of state has been used for the vapor phase.

In order to improve the simulation of absorption, Henry's law constants of the main biogas components (CH_4 , CO_2 and H_2S) in water, the properties of pure components, equilibrium constants for the equilibrium reactions and the reaction rate constant for the kinetically controlled reaction between carbon dioxide and the hydroxide ion have been verified against experimental data.

After the verification and modification of the thermodynamic model, the results of the simulations were compared to experimental data obtained on the demonstrator "BioGNVAL".

The comparison shows that the two results are in good agreement.

In connection with this work, it would be interesting to study the perspective of incorporating the hydrodynamic model developed, in the database of Aspen Plus.

For adsorption technology, a dynamic model has been developed to simulate the breakthrough curve for the system H_2S – Activated carbon.

During this study, the different mass transfer coefficients involved in the process have been estimated under the same conditions in which the experiments on "BioGNVAL" pilot plant were held.

The results obtained with the model developed should be compared to experimental data in order to adjust the overall mass transfer coefficient and equilibrium parameters given by the adsorption isotherm. Another perspective is to consider the reaction phenomena due to the impregnation of the activated carbon in order to have a prediction tool able to manage the design of adsorption columns. It will be also interesting to reinsert the axial dispersion coefficient in the mass balance equation to evaluate its contribution.

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Résumé

Le biogaz doit être purifié pour devenir un combustible renouvelable. De nombreux traitements actuels ne sont pas satisfaisants car, pour des raisons de coûts les procédés de séparation privilégiés aboutissent souvent au rejet direct ou indirect du sulfure d'hydrogène (H_2S) à l'atmosphère. C'est le cas de la séparation à l'eau sous pression.

Les objectifs de la thèse portent d'abord sur l'étude et la modélisation des méthodes connues de séparation de l'hydrogène sulfuré du méthane. Les concentrations typiques varient de 200 à 5000 ppm et la séparation devra réduire la teneur résiduelle en H_2S à moins de 1 ppm. Parallèlement seront étudiées les méthodes de traitement de H_2S .

Une fois la (ou les) méthode(s) de séparation sélectionnée(s), des essais de validation seront effectués sur un système traitant de l'ordre de $85 \text{ Nm}^3/\text{h}$ de méthane où seront injectées des quantités de H_2S variant entre 1 et 100 ppm.

Cette thèse requiert des modélisations réalistes sous Aspen Plus® ou sous un code équivalent pour établir a priori des efficacités de séparation selon différentes conditions opératoires et en prenant en compte le paramètre température. L'énergie dépensée pour la séparation effective sera un des critères fort de la comparaison, de même que l'économie de matière. Une approche système est indispensable pour étudier la rétroaction de la méthode de valorisation du H_2S sur la ou les méthodes séparatives. A priori c'est aussi l'outil Aspen Plus® ou équivalent qui permettra cette approche système. L'étude du procédé sera menée selon la double approche modélisation et expérimentation, pour l'étude expérimentale des méthodes séparatives, l'échelle du banc sera semi-industrielle et le banc permettra d'étudier les méthodes de séparation jusqu'à -90°C .

Mots Clés

Biogaz, Hydrogène sulfuré, Absorption réactive, Adsorption sur charbon actif, Aspen Plus®, Hydrodynamiques, Basses températures, Pilote BioGNVAL, Carburant pour véhicules ...

Abstract

Biogas must be purified for becoming a renewable fuel. At now, the most part of the purification techniques are not satisfactory because they imply hydrogen sulfide (H_2S) rejection to the atmosphere. One example of these methods is the treatment with high pressure water.

The first objective of the thesis is modeling the conventional methods for separating H_2S from methane. Typical concentrations of H_2S in methane vary from 200 to 5000 ppm. Separation methods must decrease the concentration of H_2S in methane to less than 1 ppm. At the same time, methods for H_2S treatment will be studied.

Once the most appropriated separation methods will be selected, some test will be carried out on a pilot plant capable of treating $85 \text{ Nm}^3/\text{h}$ of methane, where quantities of H_2S ranging from 1 and 100 ppm will be injected. These tests will allow validating the modeling of the separation process.

The thesis work requires simulating the separation process using the software Aspen Plus® or an equivalent one. The effectiveness of different operative conditions will be tested, varying also the parameter temperature. The energy necessary for the separation will be one of the most important criteria for the comparison, as well as the mass consumption of the different fluids involved in the process.

A system approach is fundamental for evaluating the backward effect of the H_2S valorization method on the separation techniques. The process simulator (Aspen Plus® or equivalent) will allow the system approach. The study will involve modeling and experimental parts. The experimental part will be carried out taking advantage of a semi-industrial size test bench, allowing studying the separation methods down to -90°C .

Keywords

Biogas, Hydrogen sulfur, Reactive absorption, Adsorption on activated carbon, Aspen Plus®, Hydrodynamics, Low temperatures, BioGNVAL pilot plant, Vehicle fuel ...