

PROSIM DAC APPLICATION EXAMPLE

TSA PROCESS

PROPANE ADSORPTION ON AN ACTIVATED CARBON

INTEREST OF THIS EXAMPLE

This example deals with a TSA process (Thermal Swing Adsorption) in which propane is adsorbed on an activated carbon. The thermal regeneration of the activated carbon is done by a hot nitrogen stream. This process is modeled within ProSim DAC, ProSim's dynamic simulation software dedicated to gas adsorption columns.

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Please note that this use case is only an example and should not be used for other purposes. Although this example is based on an actual case, it may not be considered as typical nor are the data used always the most accurate available. ProSim shall have no responsibility or liability for damages arising out of or related to the use of the results of calculations based on this example.

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1. PROCESS MODELING

1.1. Process description

ProSim DAC is a dynamic simulation software based on mass and enthalpy balances dedicated to gas-solid adsorption columns. The adsorption and the regeneration steps can be modeled. The available types of regeneration are: thermal regeneration (co- or counter-current), steam regeneration (co- or counter-current) and pressure regeneration. The TSA (Thermal Swing Adsorption), PSA (Pressure Swing Adsorption) or VTSA (Vacuum Thermal Swing Adsorption) processes can be represented. A management system of events allows going from one step to an other depending on criterions specified by the user (duration, breakthrough ratio, concentration...). An end simulation time or a number of cycles can be defined. The characteristics (temperature, pressure, flow rate, compositions) of the feeds (flow to be purified, regenerating flows...) can be constant or time dependent. The main models of adsorption isotherm (Langmuir, Toth, Dubinin-Radushkevich, Langmuir extended, Sips...) are available. The mass and heat transfer coefficients can be specified by the user or calculated by ProSim DAC. A data base of activated carbons and of adsorption isotherms of VOCs on activated carbons is available. An economical balance of the regeneration steps can be performed to compare different solutions.

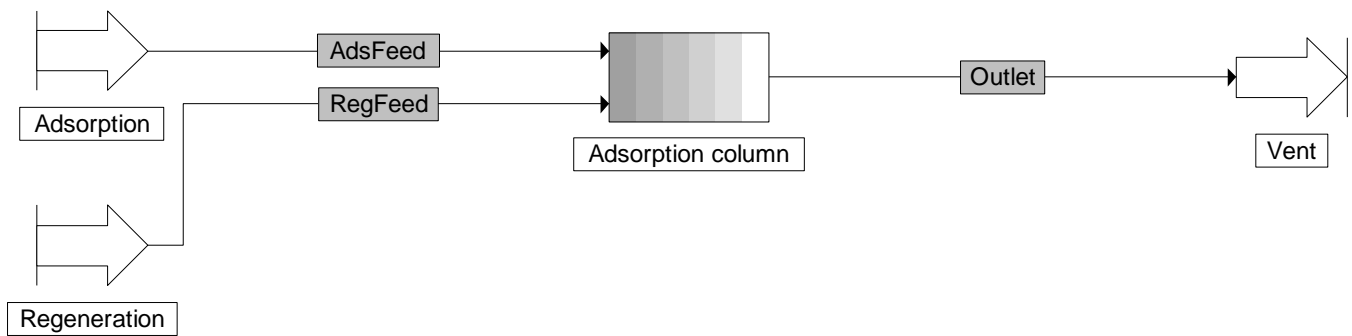
In this example, propane is adsorbed on a Witco JCX activated carbon. The regeneration of the adsorbent is done by a hot pure nitrogen stream (TSA process). The regeneration flow is counter-current compared to the adsorption flow.

Two inlet streams are used in this example: the "AdsFeed" stream and the "RegFeed" stream. During the adsorption step only the "AdsFeed" stream is used. This stream is the gas stream to be purified (i.e. in which the propane has to be removed). During the regeneration step only the "RegFeed" stream is used. This stream is the hot nitrogen stream used to regenerate the bed of adsorbent. The sequence of the steps is done automatically thanks to the definition of events ending the different steps.

The parameters of the column (size, thermal behavior...), the adsorption isotherms, the type of the cycle (TSA, VTSA, PSA), the step events (breakthrough level, temperature, concentration...) are defined within the "Adsorption column" unit operation module.

This example is adapted from [SCH88].

1.2. Process flowsheet



1.3. Compounds

The compounds used in the simulation, their chemical formula and CAS numbers are shown in the following table. Their pure component properties are extracted from the standard data base provided with ProSim DAC [ROW11].

Compound	Chemical formula	CAS number
Propane	C_3H_8	74-98-6
Nitrogen	N_2	7727-37-9

1.4. Thermodynamic model

The adsorption and the regeneration are done at a moderate pressure level (around 2.5 atm) and at temperature below 100°C. Thus the "Ideal" thermodynamic profile is selected in the Simulis Thermodynamics calculator.

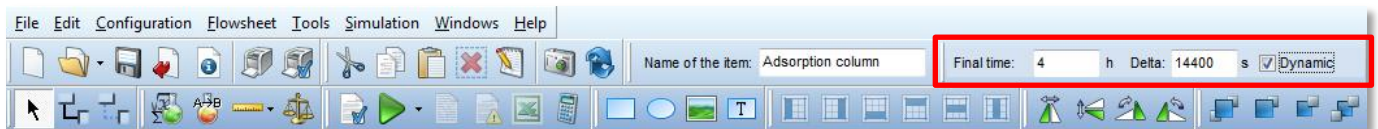
1.5. Operating parameters

1.5.1. Process feed

	Propane stream	Hot nitrogen stream
Temperature (°C)	23	100
Pressure (mmHg)	1 882	1 849
Total flow rate (kmol/h)	0.09396	0.10332
Mole fractions		
Dichloromethane	0.0106	0
Nitrogen	0.9894	1

1.5.2. Simulation time

The “Final time” is the real working duration of the process (adsorption + thermal regeneration in this example). The “Delta” is the duration during which the “Adsorption column” module is working. As only one module is present, the “Final time” and the “Delta” should be identical. These two values are specified in the icon bar of ProSim DAC:



Simulation time	Value
Final	4 h
Delta	14 400 s

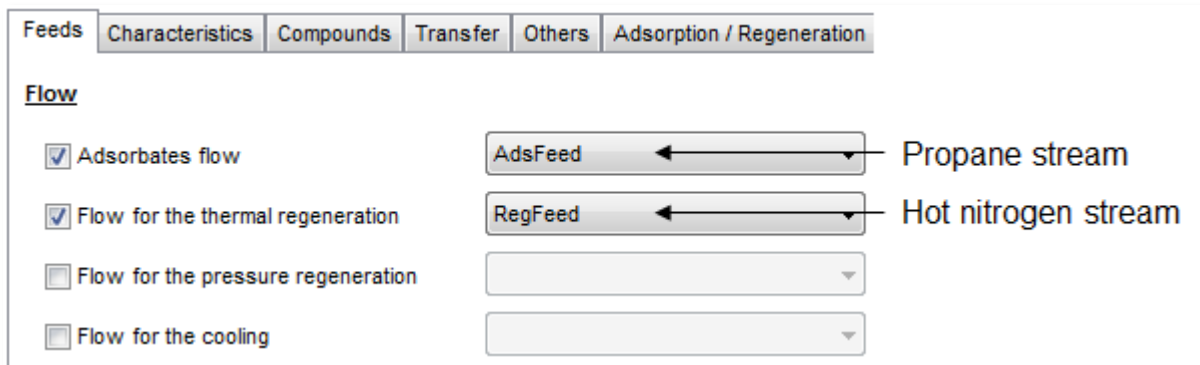
1.5.3. Adsorption column

1.5.3.1. Feeds

Four feeds can be used in ProSim DAC:

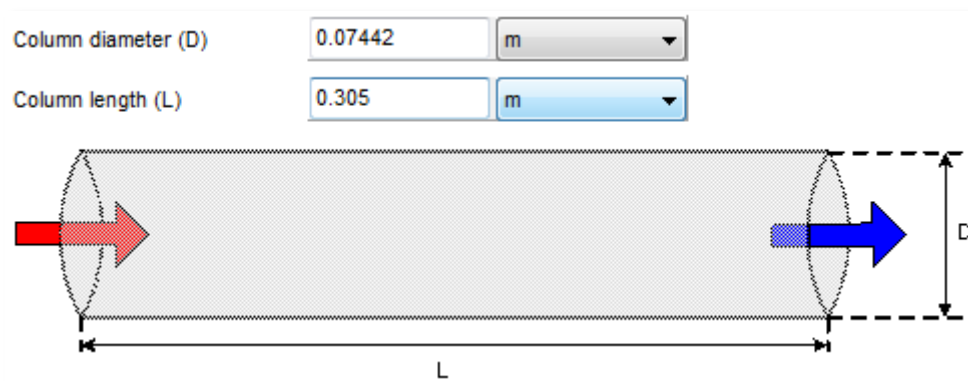
- ✓ Adsorbates flow: Flow to be purified during the adsorption step
- ✓ Flow for the thermal regeneration: Flow used during the thermal regeneration (hot nitrogen, steam...)
- ✓ Flow for the pressure regeneration: Flow used during the pressure regeneration (low pressure nitrogen...)
- ✓ Flow for the cooling: Flow used to cool the column at the end of the thermal regeneration step

Only the two first feeds are used in this example as shown on the screen shot below.



1.5.3.2. Column characteristics

The column used is a lengthwise flow column which characteristics are shown on the following screen shot. The dimensions to be specified are the ones of the adsorbent bed.



Transvers flow columns can also be modeled in ProSim DAC.

1.5.3.3. Thermal behavior

The following thermal heat exchanges can be modeled in ProSim DAC:

- ✓ Heat exchange in the bed: "Given heat duty without wall transfer"
This possibility allows to model an adiabatic working or a given heat duty for each step working (heat exchanger in the adsorbent bed).
- ✓ Wall heat transfer + thermal heat exchange in the bed: "Given heat duty and wall transfer"
The heat exchange is done by the wall of the column. By default, the wall temperature is constant during the time and along the length of the column. To go beyond this assumption, the option "take into account thermal inertia of the column wall" has to be activated. It is possible to add a given heat duty (by default zero) in the adsorbent bed for each step.

In this example, the thermal transfer is done by the wall without any direct heating/cooling in the bed of adsorbent:

Thermal behavior	
Heat transfer	Given heat duty and wall transfer
Wall temperature	23°C

1.5.3.4. Adsorbent bed characteristics

Bed characteristics	
Void ratio	0.44

1.5.3.5. Adsorbent characteristics

The density of the particles takes into account of the intra-particulate porosity.

Particles characteristics	
Diameter	3.2 mm
Density	827 kg/m ³
Specific heat	0.2 cal/g/K
Surface/Volume ratio	1 320 m ² /m ³

1.5.3.6. Measures conditions for concentration results

The user can specify the temperature and pressure conditions he wants to calculate the volumetric concentrations in the gas phase. Indeed, the density is then necessary and this one, particularly in the case of a gas, is sensitive to the temperature and the pressure. In this example, the volumetric concentrations in the gas phase are evaluated at the temperature and pressure conditions of the column outlet.

Measures conditions	Value
Conditions	Column

1.5.3.7. Initialization

It's necessary to define the state of the column before its first use. In this example, it's assumed that the column is inerted with nitrogen at temperature and pressure levels corresponding to the flow to be purified.

Initialization	Value
Type	Supplied by the user
Pressure	1 882 mmHg
Temperature	23°C
Molar fractions	
Propane	0
Nitrogen	1

The screen shot below shows the information of the previous paragraphs (from paragraph 1.5.3.2 to paragraph 1.5.3.7 included) specified in the module interface: "Parameters" tab, "Characteristics" sub-tab.

The screenshot displays the software interface with the following settings:

- Columns:** Identification, Parameters, Scripts, Report, Streams, Profiles, Notes, Advanced parameters
- Sub-tabs:** Feeds, Characteristics, Compounds, Transfer, Others, Adsorption / Regeneration, Financial results, Prints, Parameters
- Column Section:**
 - Column type: Lengthwise flow colmun
 - Heat transfer: Given heat duty and wall transfer
 - Wall temperature: 23 °C
- Adsorbent Section:**
 - Bed void ratio: 0,44 m³/m³
 - Particles diameter: 0.0032 m
 - Particles density: 827 kg/m³
 - Specific heat of the solid: 0,2 cal/g/K
 - Particle surface/volume ratio: 1320 m²/m³
- Measures conditions (T,P):** Conditions: Column
- Initialization Section:**
 - Initialization type: Supplied by user
 - Initial pressure: 1882 mmHg
 - Initial temperature: 23 °C
 - Initial molar fractions table:

1	PROPANE	0
2	NITROGEN	1
 - Summation: 1,0000

1.5.3.8. Adsorption isotherms and enthalpies

The Langmuir model is selected for the adsorption isotherm of propane:

$$q_i = \frac{\left[q_{m0} \exp\left(\frac{q_{m1}}{T}\right) \right] \left[K_0 \exp\left(\frac{K_1}{T}\right) \right] P_i}{1 + \left[K_0 \exp\left(\frac{K_1}{T}\right) \right] P_i}$$

The linear model is selected for the adsorption isotherm of nitrogen:

$$q_i = \left[K_0 \exp\left(\frac{K_1}{T}\right) \right] P_i$$

The values of the parameters with the corresponding units are displayed in the table below.

Propane		Nitrogen	
Isotherm	Langmuir	Isotherm	Linear
q_{m0} (mol/kg)	0.93589302	K₀ (mol/kg/atm)	0.000988
K₀ (atm⁻¹)	0.50456976	K₁ (K)	1752
q_{m1} (K)	425.905386		
K₁ (K)	1299.44563		

The adsorption phenomenon being exothermic, the values of the enthalpies of adsorption are negatives.

Adsorption enthalpy	Propane	Nitrogen
Type	Given	
ΔH_{ads} (cal/mol)	-9 500	-3 840

This information has to be specified in the “Parameters” tab, “Compounds” sub-tab:

The screenshot shows the 'Parameters' tab for 'Compounds' in the software. It is divided into two panels for 'PROPANE' and 'NITROGEN'.
 For **PROPANE**:
 - **Adsorption enthalpy**: Enthalpy of adsorption is set to 'Given', Adsorption heat is -9500 cal/mol.
 - **Adsorption isotherm**: Correlation is set to 'Langmuir'. The equation $q_i = \frac{\left[q_{m0} \exp\left(\frac{q_{m1}}{T}\right) \right] \left[K_0 \exp\left(\frac{K_1}{T}\right) \right] P_i}{1 + \left[K_0 \exp\left(\frac{K_1}{T}\right) \right] P_i}$ is displayed. Parameters: q_{m0} = 0.93589302 mol/kg, K₀ = 0.50456976 atm⁻¹, q_{m1} = 425.905386 K, K₁ = 1299.44563 K.
 For **NITROGEN**:
 - **Adsorption enthalpy**: Enthalpy of adsorption is set to 'Given', Adsorption heat is -3840 cal/mol.
 - **Adsorption isotherm**: Correlation is set to 'Linear isotherm'. The equation $q_i = K_0 \exp\left(\frac{K_1}{T}\right) P_i$ is displayed. Parameters: K₀ = 0.000988 mol/kg/atm, K₁ = 1752 K.

1.5.3.9. Material transfer

The following options are available for the material transfer:

- ✓ Transfer coupled in gas and solid phase
- ✓ Material transfer resistance (“linear driving force” model in gas phase and/or in solid phase, the material transfer coefficient needed could be specified by the user or calculated by ProSim DAC. It’s also possible to neglect the material transfer resistance.

In this example, the material transfer resistance has been taken into account in the two phases and the necessary coefficients are calculated by ProSim DAC.

	Value
Material transfer	
Type	Gas and solid transfer
Gas material transfer	
Type	kf calculated, Petrovic-Thodos
Solid material transfer	
Type	kf calculated

1.5.3.10. Thermal transfer

It’s possible to take into account the enthalpy balance in ProSim DAC or to ignore it. An isothermal working can be modeled if the enthalpy balance is not taken into account. If the enthalpy balance is taken into account, the gas-adsorbent and the gas-wall heat transfer coefficients are necessary. They can be calculated, as in this example, or specified.

Thermal transfer	Value
Enthalpy balances	Taken into account
Gas – Adsorbent	Calculated (Satterfield)
Gas – Wall	Calculated (Leva)

The screen shot below shows the information of the previous paragraphs 1.5.3.9 and 1.5.3.10 specified in the module interface: "Parameters" tab, "Transfer" sub-tab.

Identification	Parameters	Scripts	Report	Streams	Profiles	Notes	Advanced parameters	
Feeds	Characteristics	Compounds	Transfer	Others	Adsorption / Regeneration	Financial results	Prints	Parameters
Material transfer		Thermal transfer						
Transfer type	Gas and solid transfer	<input checked="" type="checkbox"/> Enthalpy balances ?						
Gas material transfer		Gas-adsorbent						
Gas transfer type	kf calculated, Petrovic-Thodos	Calculated (Satterfield)						
		Gas-wall						
		Calculated (Leva)						
Solid material transfer		Wall thermal inertia						
Solid transfer type	kf calculated	<input type="checkbox"/> Take into account thermal inertia of the column wall						
		Mass (wall)						
		0 kg						
		Specific heat (wall)						
		0 cal/g/K						

1.5.3.11. Adsorption thermodynamics

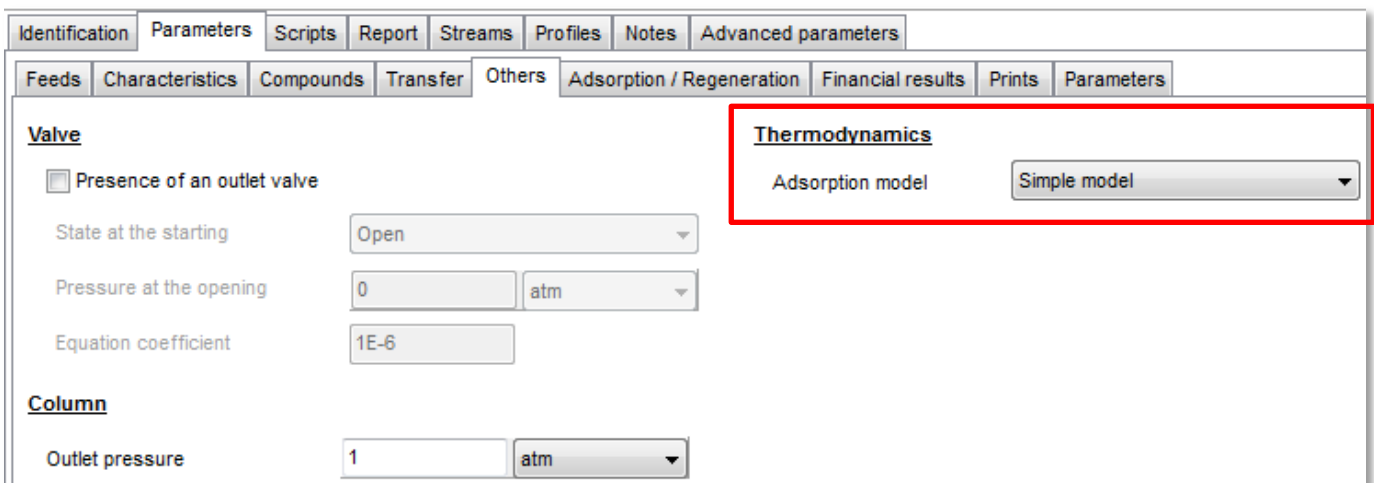
The adsorption thermodynamic models available are:

- ✓ Simple model: This model doesn't take into account the influence of the compounds on each other's. The adsorbed quantity of a given compound is directly given by its adsorption isotherm specified by the user.
- ✓ IAS or RAS: This model takes into account the influence of the compounds on each other's. The IAS model assumes an ideal adsorbed phase. The RAS model is used if an activity coefficient model is selected in the thermodynamic profile of the Simulis Thermodynamics calculator.
- ✓ Langmuir extended: This model takes into account the influence of the compounds on each other's. It has to be used with pure compound adsorption isotherms of the type "Langmuir extended".

In this example, the simple model is selected.

Adsorption thermodynamics	Value
Adsorption model	Simple model

The adsorption thermodynamic model is specified in the "Parameters" tab, "Others" sub-tab. The left part of this sub-tab is used to specify an optional valve at the column outlet during the adsorption step.



1.5.3.12. Sequence

The “Adsorption/Regeneration” sub-tab of the “Parameters” tab allows to select the type of the cycle to simulate between the five available:

1. Adsorption only
2. Adsorption + thermal regeneration
3. Adsorption + pressure regeneration
4. Adsorption + pressure regeneration + thermal regeneration
5. Adsorption + thermal regeneration + pressure regeneration

The cycle “Adsorption + thermal regeneration” is used in this example to model a TSA (Thermal Swing Adsorption) process as shown in the following figure.

The screenshot shows the 'Parameters' tab for 'Adsorption / Regeneration'. The 'Sequence type' dropdown menu is open, showing five options: 'Adsorption only', 'Adsorption + thermal regeneration' (selected), 'Adsorption + pressure regeneration', 'Adsorption + pressure regeneration + thermal regeneration', and 'Adsorption + thermal regeneration + pressure regeneration'. Below the dropdown, there are sections for 'Adsorption', 'Thermal regeneration', and 'End of simulation', each with 'Parameters...' and 'Events...' buttons. To the right, a flowchart illustrates the cycle: a box labeled 'A' (Adsorption) leads to a circle labeled 'EVA' (Stop adsorption events), which leads to a box labeled 'RT' (Thermal regeneration), which leads to a circle labeled 'EVT' (Stop thermal regeneration events), which then loops back to the 'A' box. A legend on the right identifies these symbols: 'A: Adsorption', 'EVA: Stop adsorption events', 'RT: Thermal regeneration', and 'EVT: Stop thermal regeneration events'.

Optional parameters can be defined for each step ("Parameters..." buttons):

- ✓ Adsorption:
 - Wall temperature specific to this step
 - Heat duty exchanged in the bed
- ✓ Thermal regeneration:
 - Co- or counter-current regeneration
 - Bed and/or wall preheating of the column
 - Cooling of the column at the end of the step
 - Temporization of the column at the end of the step
 - Heat duty exchanged in the bed

Each step is ended by an event. The first event reached starts the next step. The available events are the following ones ("Events..." buttons):

- ✓ Adsorption
 - Step duration
 - Breakthrough rate of a compound at a given position in the column
 - Gas phase concentration of a compound at a given position in the column
 - Solid phase concentration of a compound at a given position in the column
 - Maximal temperature reached at a given position in the column
 - Maximal pressure reached at a given position in the column
- ✓ Thermal or pressure regeneration
 - Step duration
 - Gas phase concentration of a compound at a given position in the column
 - Solid phase concentration of a compound at a given position in the column
 - Maximal temperature reached at a given position in the column
 - Maximal pressure reached at a given position in the column
 - Amount produced of a compound at a given position in the column
- ✓ Simulation end
 - End simulation time (§ 1.5.2) or number of cycles
 - Total amount produced of a compound at a given position in the column
 - Maximal temperature reached at a given position in the column
 - Maximal pressure reached at a given position in the column
 - Total amount treated of a compound

In this example, the regeneration of the bed of adsorbent is a thermal regeneration done by a hot nitrogen stream. This stream flows counter-currently compared to the adsorption flow. The default parameters are kept for the two steps (adsorption and thermal regeneration). These two steps and the simulation are stopped by a duration.

	Value
Sequence	
Type	Adsorption + Thermal regeneration
Adsorption	
Parameters	Default values
Events	Duration = 7 200 s
Thermal regeneration	
Parameters	Default values
Events	Duration = 7 200 s
End of simulation	
Events	End simulation time

1.5.3.13. Financial balance

ProSim DAC performs a financial balance on the regeneration steps. If the user is interested, it's possible to modify the parameters default values in the "Financial results" sub-tab of the "Parameters" tab.

Identification	Parameters	Scripts	Report	Streams	Profiles	Notes	Advanced parameters	
Feeds	Characteristics	Compounds	Transfer	Others	Adsorption / Regeneration	Financial results	Prints	Parameters
<u>Thermal regeneration costs</u>								
Cost of regeneration flow	<input type="text" value="0.2"/>	€/Nm ³						
Cost of regeneration flow heating	<input type="text" value="100"/>	€/MWh						
Cost of bed preheating	<input type="text" value="120"/>	€/MWh						
<u>Pressure regeneration costs</u>								
Cost of electricity	<input type="text" value="100"/>	€/MWh						
Power of vacuum pump	<input type="text" value="0"/>	kcal/h						

1.5.3.14. Prints

ProSim DAC offers different options for the printing of the calculation results. The parameters specified for this example are listed in the table below.

Parameters	Value
Print results files	Frequency = 60 s
Print 3D plots	Not activated
Print output stream with time step of module	Activated
Printing of inlet data	Activated
Type of results	Molar
Inert detection	Not activated

The following screen shot shows this information specified in the “Prints” sub-tab of the “Parameters” tab.

Print

Print results files
 Frequency: 60 s

Print 3D plots
 Frequency: 0,1 h

Print output stream with time step of module

Printing of inlet data

Type of results: Molar

Inert detection
 Threshold: 1E-6 kmol

1.5.3.15. Integration, tolerances and model parameters

In ProSim DAC, the user has access to a set of integration, tolerances and model parameters. Only the parameters shown in the table below are modified compared to the default values. The use of the analytical calculation of the derivatives speeds up the solving of the problem. The model consists in a numerical time integration and on a spatial discretization. In this example, the column is discretized in 30 cells to obtain a precise solving of the problem. The thermal accumulation of the solid has a weak influence in this example. Thus, this option is deactivated to decrease the computational time required.

Parameters	Value
Derivatives	Calculated analytically
Number of discretization cells	30
Thermal accumulation in the solid	Not taken into account
Other parameters	Default values

The following screen shot shows this information specified in the “Parameters” sub-tab of the “Parameters” tab.

2. RESULTS

2.1. Simulation report of the adsorption column

The simulation report of the adsorption column ("Report") presents global results (integrated during the time): initial characteristics of the column, adsorbed quantities, quantities recovered during the regeneration...

The mass of adsorbent is not directly specified in the input data. This value is calculated by ProSim DAC depending on the geometrical characteristics of the bed and the adsorbent characteristics. It's then interesting to verify in the simulation report the mass of adsorbent calculated corresponds to the end wished. For this example, the mass of adsorbent is about 615 g.

COLUMN GENERAL CHARACTERISTICS

Adsorbent mass in the column:	614.416	(G)
Column volume	: 1.32669	(L)
Solid volume	: 0.742946	(L)
Void volume	: 0.583743	(L)

With the operating parameters specified (see § 1.5.3.12 among others), the bed of adsorbent is nearly completely regenerated (number of moles of propane adsorbed close to the number of moles of propane regenerated).

AMOUNT ADSORBED (MOL)

COMPONENT	Cycle N° 1
PROPANE	1.24936
NITROGEN	-6.550613E-03

AMOUNT RECOVERED DURING THERMAL REGENERATION (MOL)

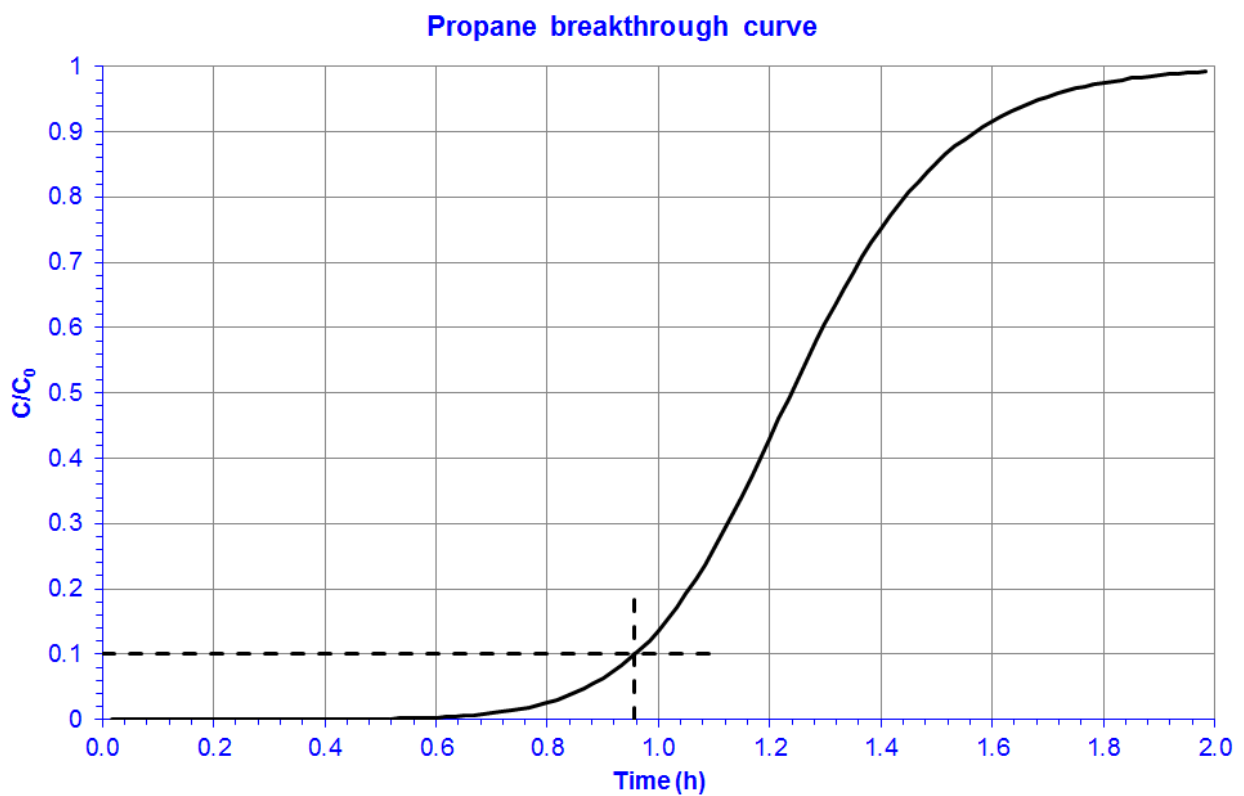
COMPONENT	Cycle N° 1
PROPANE	1.24849
NITROGEN	206.948

The negative value of the amount of nitrogen adsorbed corresponds to the mas balance deviation.

2.2. Profiles of the adsorption column

Several profiles (temperatures, pressures, velocities, concentrations, compositions and breakthrough curves) in the adsorption column are available after the simulation from the module edition window ("Profiles" tab).

The following figure shows the breakthrough curve of propane. The breakthrough is assumed to be reached when the concentration at the column outlet is equal to 10% of the inlet concentration. The breakthrough is reached after 57 min (0.96 h). The bed of adsorbent is saturated when the outlet concentration is equal to the inlet one. This is the case after 2 h.



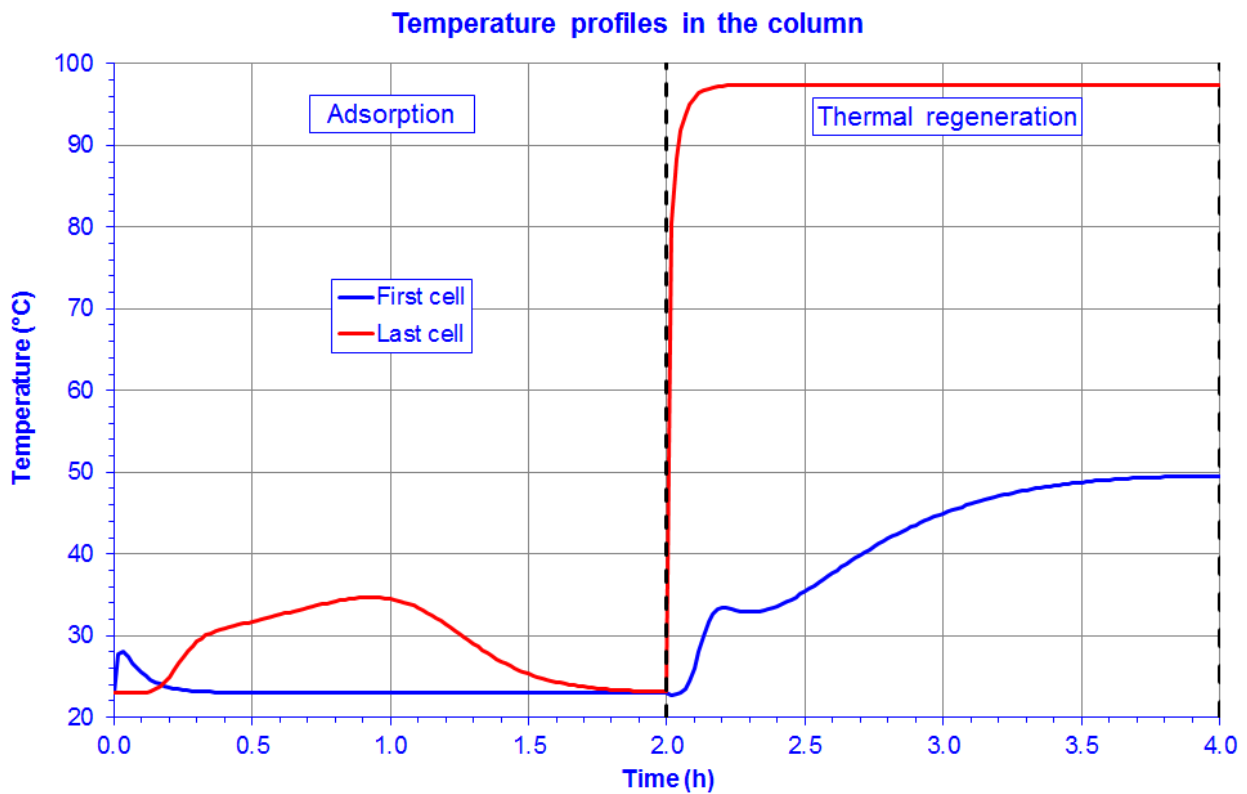
The figure below presents the temperature profile in the first and the last discretization cell. In the case of the adsorption:

- ✓ First cell: Inlet cell of the column
- ✓ Last cell: Outlet cell of the column

As the regeneration is done counter-currently:

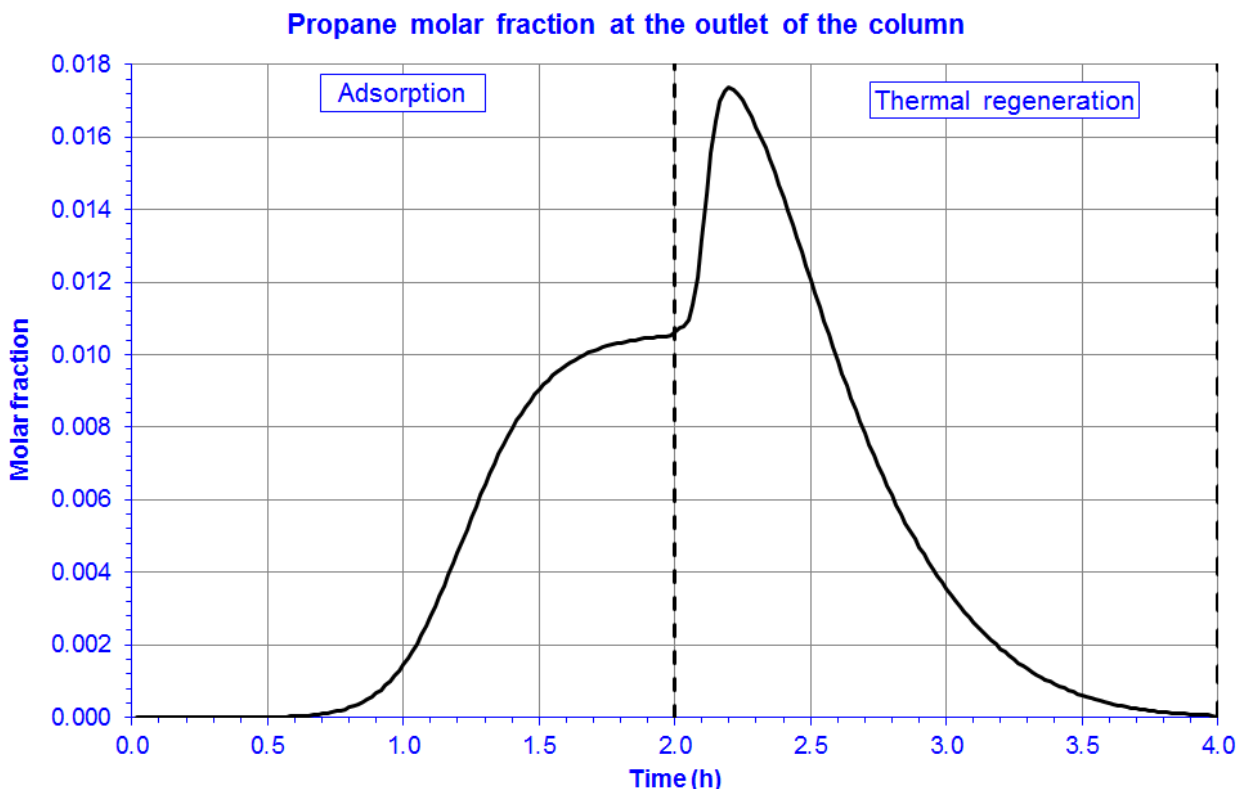
- ✓ First cell: Outlet cell of the column
- ✓ Last cell: Inlet cell of the column

The temperature profiles during the adsorption step show the corresponding temperature peak. During the thermal regeneration the temperature gradient is higher. Indeed the extremity where the hot gas enters (last cell) is rapidly at the thermal equilibrium with the incoming flow (100°C). At the opposite extremity, the temperature increase is less and the equilibrium between the wall temperature (23°C) and the hot gas (100°C) has the time to be reached.



2.3. Profiles of the outlet stream

The following figure shows the evolution of the molar fraction of propane at the column outlet. Once the breakthrough is not reached, the adsorbent plays its role and the propane amount at the column outlet is null. When the breakthrough time is reached, the outlet stream is charged progressively in propane. In this example, the adsorption step is ended at the time corresponding to the saturation of the bed. The bed no more retained the propane and its outlet concentration is equal to its inlet concentration. The counter-current thermal regeneration by a hot nitrogen stream allows to desorb the propane retained in the adsorbent. After a peak corresponding to the propane accumulated in the bed, the propane amount decreases because of the bed regeneration. This step is ended at the time corresponding to a complete regeneration of the bed.



The above profile and other ones (temperature, pressure, flow rate, enthalpy, and compositions) are available for the column outlet stream at the end of the simulation from the outlet stream edition window (“Tabulated results...” button in the “Parameters” tab).

3. REFERENCES

- [ROW11] ROWLEY R.L., WILDING W.V., OSCARSON J.L., GILES N.F., "DIPPR® Data Compilation of Pure Chemical Properties", Design Institute for Physical Properties, AIChE, New York, NY (2011)
- [SCH88] SCHORK J.M., FAIR J.R., "Parametric Analysis of Thermal Regeneration of Adsorption Beds", Ind. Eng. Chem. Res, 27, 457-469 (1988)