

THE UNIVERSITY of EDINBURGH

Edinburgh Research Explorer

Thermodynamics of thermally-driven adsorption compression

Citation for published version:

Santori, G & Luberti, M 2016, 'Thermodynamics of thermally-driven adsorption compression', *Sustainable Materials and Technologies*, vol. 10, pp. 1-9. https://doi.org/10.1016/j.susmat.2016.09.001

Digital Object Identifier (DOI):

10.1016/j.susmat.2016.09.001

Link:

Link to publication record in Edinburgh Research Explorer

Document Version: Peer reviewed version

Published In: Sustainable Materials and Technologies

General rights

Copyright for the publications made accessible via the Edinburgh Research Explorer is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy The University of Edinburgh has made every reasonable effort to ensure that Edinburgh Research Explorer content complies with UK legislation. If you believe that the public display of this file breaches copyright please contact openaccess@ed.ac.uk providing details, and we will remove access to the work immediately and investigate your claim.



1 Thermodynamics of thermally-driven adsorption compression

2 Giulio Santori^a*, Mauro Luberti^a

*corresponding author: Dr. Giulio Santori, g.santori@ed.ac.uk, tel: +4401316519043, fax: +4401316506551

4 ^a The University of Edinburgh, School of Engineering, Institute for Materials and Processes, Sanderson

5 Building, The King's Buildings, Mayfield Road, EH9 3JL, Edinburgh, Scotland, UK 6

7 Abstract

8 Adsorption compressors are an emerging technology used to compress a gas stream with low grade heat 9 which is of interest for the next long term extra-planetary bases. An analysis of thermodynamics of multicomponent gas/vapour mixtures compression with a thermally-driven adsorption bed is reported. In this 10 unit a multicomponent stream is firstly adsorbed and secondly compressed by heating a bed at closed 11 volume. The analysis is based on adsorbed solution theory applied to closed vessels where the composition 12 13 and pressure of the bulk gas phase depends on temperature and volume of the whole system, leading to an 14 isochoric-isothermal flash problem. Analysis of both an ideal and non-ideal adsorption compressor shows that the ideal approach is conservative, resulting in lower compression ratios at higher energy consumption. 15

16

19

Keywords: Isothermal flash; Isochoric flash; Adsorbed solution theory; Common tangent plane; Solidcompressor; Adsorption thermodynamics;

20 1. Introduction

The recent development of highly selective adsorption materials have enabled their integration in a number of traditional technologies, resulting in advanced processes demanding a sensibly lower energy consumption

delivered at lower temperatures than their traditional counterparts [1-3]. A partial list of cases is included in

Table 1. All these new technologies include a fundamental component that is the adsorption bed, which can

be structured in numerous different shapes depending on the regeneration strategy [4-7].

25 26

Table 1: Emerging adsorption technologies

Sector	Traditional technology	Adsorptive technology	Ref.
Refrigeration	Electrically-driven vapour	Adsorption refrigerators	[8-10]
& air conditioning	compression system	& heat pumps	[]
Drying	Electrical dishwasher	Adsorption dishwasher	[11]
Desalination	Multi-effect distillation	Adsorption desalination	[12]
CO ₂ removal	Absorption	Temperature and/or vacuum swing	[13-15]
	*	adsorption	

27

Adsorption beds are often operated as open systems where a single or multicomponent feed flows through the unit. However, some emerging technologies need to operate adsorption beds as closed systems. This is the case of the carbon dioxide removal and compression system, currently operating on the International Space Station and still under development to serve the next long term Mars or lunar bases [16-18].

Adsorption is the favourite method to produce a compressed gas in space applications because it does not have moving parts, requiring virtually no maintenance, and does not generate vibrations. The carbon dioxide removal system is the upstream process of the Sabatier reaction, which will be used in extra-terrestrial bases for the production of water. As demonstrated elsewhere [19, 20], the standard operating conditions to drive the Sabatier reaction towards higher CO₂ conversion are temperature ranging between 250-400°C and pressure of reagents greater than 130 kPa. So, compression of CO₂ from 20 kPa (Mars atmosphere) to 130 kPa (Sabatier reaction pressure) is required, resulting in a minimum compression ratio of 6.5.

In this technology, compression is performed through a change in the bed temperature at closed volume. Theoperational steps of an adsorption compressor can be scheduled as:

Adsorption step: the adsorption bed is kept isothermal and is fed with the inlet stream at constant
 pressure and temperature.

43 2) Heating step: the bed is heated at constant volume. The new equilibrium state after heating results in
44 a partition of the components between bulk gas phase and adsorbed phase at a bulk gas phase pressure higher
45 than the initial pressure during the feeding. The pressure increase is mainly because of the release of material
46 from the adsorbed phase to the bulk gas phase. The composition and the level of pressure in the bulk gas
47 phase depend on the final temperature, volume available for the bulk gas phase, mass of adsorbent and total

47 phase depend on the final temperature, volum48 amount of moles in the enclosure.

49 3) Supply step: the outlet valve is open with a decrease in pressure and a variable flow of material is50 discharged.

51 Thus, adsorption compressors are thermally-driven compression systems adopting adsorption materials and 52 operating according to temperature swing adsorption processes. An early thermodynamic analysis for 53 adsorption of pure fluids has been presented in [21, 22] but no thermodynamic basis has been provided for

54 the adsorption compression of a multicomponent mixture to date and some units such as the air revitalization

55 module of the International Space Station (ISS) have been designed on the basis of simplifying assumptions,

56 refining the performance in successive, trial and error steps. The air revitalization unit of the ISS uses

- 57 zeolites 5A and 13X which adsorb significant amounts of carbon dioxide and nitrogen. This aspect has been 58 addressed, at the design stage, by oversizing the mass of required material with a safety factor [22, 23]. The 59 present work provides a tool for consistent thermodynamic investigation of adsorption compression of 60 multicomponent gas mixtures.
- 61 The fundamental calculation for adsorption compression is the isochoric-isothermal flash (VT flash). VT
- flash in bulk gas/adsorbed phase equilibria has been already formulated by minimization of the Helmholtz
 energy of the system [24]. In this work we show that the well-known Rachford-Rice system of equations can
 be also applied to adsorption equilibrium after appropriate modifications.

For sake of clarity, at first an ideal ternary mixture in equilibrium with an ideal adsorbed solution [25] is evaluated. The aim is to provide the thermodynamic framework for the simple ideal case. Secondly, a nonideal ternary mixture is considered using the Soave-Redlich-Kwong (SRK) equation of state for the description of the bulk gas phase and a Gibbs excess model for the adsorbed phase. This is aimed to show the differences between an ideal and non-ideal formulation.

70 Since no experimental multicomponent equilibrium data are presently available for this problem, the 71 thermodynamic consistency of the results is validated through the common tangent plane approach for

adsorption derived in [26]. The last section of this work is devoted to the performance analysis of ideal and

73 non-ideal compressors.

74

75 2. Ideal isothermal-isochoric flash for adsorption

The ideal case includes an ideal bulk gas phase in equilibrium with an ideal adsorbed solution. The pressure and compositions in the new equilibrium state can be calculated considering the usual method based on the Rachford-Rice equations system [27], once a number of additional conditions arising from the adsorbed solution theory are provided [28]. The resulting system of equations for NC components is:

80
$$\sum_{i}^{NC} \frac{z_i (k_i - 1)}{1 + \frac{G}{F} (k_i - 1)} = 0$$
(1)

81
$$x_i = \frac{z_i}{1 + \frac{G}{F}(k_i - 1)}$$
 (2)

82
$$G = \frac{P_{bulk}V_{void}}{RT}$$
(3)

$$k_i = \frac{P_i^0}{P_{bulk}} \tag{4}$$

$$84 y_i P_{bulk} = P_i^0 x_i (5)$$

85
$$\frac{m_{ads}}{N} = \sum_{i=1}^{NC} \left(\frac{x_i}{n_i}\right)$$
(6)

86
$$\psi_i = \int_{0}^{n_i} n_i d(\ln P_i) \quad i = 1, 2, NC$$
 (7)

87
$$\psi_i = \psi_{eq}$$
 $i = 1, 2, ..., NC$ (8)

88
$$F - (G + N) = 0$$
 (9)

Eq. (1) is the original Rachford-Rice solving equation; eq. (3) is the ideal gas equation of state for the calculation of the amount of moles in the bulk gas phase. Eqns (4-8) follow the ideal adsorbed solution

theory [27, 28]. Eq. (9) is the overall mass balance. Assuming that the adsorbent occupies all the volume
available in the vessel, the gas mixture volume is calculated by:

93
$$V_{void} = \frac{m_{ads}}{\rho_b} \left(\varepsilon_b + (1 - \varepsilon_b) \varepsilon_p \right)$$
(10)

94 Usually the parameters involved in eq. (10) are omitted in equilibrium measurements. This work assumes the

95 values reported in Table 2. Although the values of Table 2 do not perfectly correspond to the experimental

96 equilibrium data used for single isotherm parameters regression, they lay in the average range commonly

97 assumed for this kind of materials.

98 The system of eqns (1-9) can be reduced in a straightforward way to only two equations by substitution of 99 variables, holding the final variables P_{bulk} and reduced grand potential ψ_{eq} . The equivalent two equation

100 system includes eq. (1) and eq. (9).

Table 2. Ausorption materials and	beus pro	perces)		
	m _{ads}	ϵ_{b}	ε _p	$ ho_{b}$	Ref
	[kg]			$[\text{kg m}^{-3}]$	
Zeolite 5A	1	0.39	0.50	687	[29]
Activated Carbon Norit R1 Extra	1	0.35	0.84	420	[30]
Zeolite 13X	1	0.37	0.60	641	[31]

Table 2: Adsorption materials and beds properties

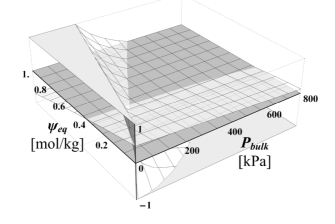
102 Eqns (1-9) can be effectively solved by the Newton numerical method without any issue on the estimation of 103 the best initial guess. That is because the system has always only one solution in the physically meaningful 105 domain $P_{bulk}>0$ and $\psi_{eq}>0$. In order to show this feature for the ideal problem, the Nitrogen/Oxygen/Argon 106 ternary system adsorption on zeolite 5A has been considered, using the Langmuir isotherm with parameters 107 reported in Table 3.

Table 3: Parameters of the Langmuir isotherm model for Nitrogen, Oxygen and Argon on zeolite 5A at 298.55 K. Equilibrium data are from [32].

2,0.00 11			
	Pressure Range	q_s	b
	[kPa]	[mol kg ⁻¹]	[kPa ⁻¹]
$N_2(1)$	34.1-442.9	2.114	0.001756
$O_2(2)$	29.3-404.8	2.313	0.000524
Ar (3)	35.5-438.7	3.348	0.000314

109

Fig. 1 illustrates the solution of the system composed by eq. (1) and eq. (9). In the physically meaningful region, the two functions intersect only in one point. The same equilibrium compositions are confirmed by the common tangent plane of Gibbs energy of mixing (Fig. 2). The presence of a common tangent plane of the Gibbs energy of mixing at the equilibrium point of Table 4 validates the thermodynamic consistency of the solution. In Table 4 it can be noted that while the bulk gas phase composition is very different from the feed, being enriched by oxygen and argon which are the less strongly adsorbed components, the adsorbed phase composition is close to the feed, being slightly richer in nitrogen over the other components.



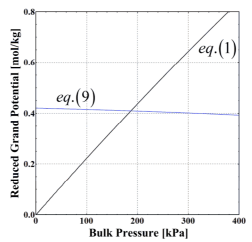


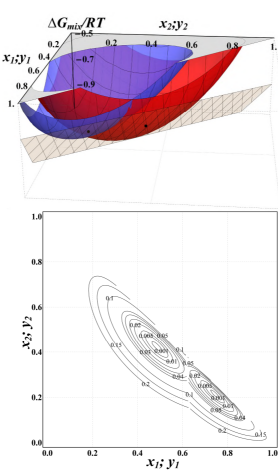


Figure 1: Surfaces representing the left hand side functions of eq. (1) and eq. (9) (top). The horizontal plane passing through (0, 0, 0) defines the contour curves corresponding to eq. (1) and eq. (9) (bottom). eq. (1) and eq. (9) intersect in only one point, highlighting that the solution is necessary and sufficient.

Table 4: Solution for the ideal VT flash calculation at the conditions: $V_{void} =$

$1.021 \ 10^{-9} \ \text{m}^3; F = 0.45 \ \text{moles}; T = 298.55 \ \text{K}$						
Component	$\mathbf{z}_{\mathbf{i}}$	P _{bulk}	Ψ_{eq}	Xi	P_i^0	yi
		[kPa]	$[mol kg^{-1}]$		[kPa]	
$N_{2}(1)$	0.70	186.80	0.410	0.744	122.03	0.486
$O_2(2)$	0.25			0.214	370.54	0.425
Ar (3)	0.05			0.042	398.20	0.089

123



125 $x_l; y_l$ 126Figure 2: Common tangent plane of the Gibbs energy of mixing at the equilibrium point of Table 4 (top).127Contour plot of the distance function between Gibbs energy of mixing and common tangent plane at the128equilibrium point of Table 4 (bottom).

130 **3.** The non-ideal isothermal-isochoric flash

In the non-ideal formulation fugacity and activity coefficients respectively for the bulk gas phase and the adsorbed phase are considered [33]. Based on the experimental measurement reported in [34], the ternary mixture Methane/Nitrogen/Carbon Dioxide on activated carbon Norit R1 Extra at 298 K and high pressure is analysed. Single component adsorption experimental data were fitted using the Unilan isotherms, considering the absolute adsorbed amount versus fugacity, resulting in the parameters of Table 5.

136

Table 5: Parameters of the Unilan isotherm model for Methane, Nitrogen and Carbon Dioxide on activated carbon Norit R1 Extra at 298 K. Experimental data are from [34].

	Pressure Range	Fugacity Range	q_{s}	b	S
	[kPa]	[kPa]	[mol kg ⁻¹]	[kPa ⁻¹]	
$CH_{4}(1)$	0-5753	0-5219	24.537	2.873 10 ⁻⁵	5.220
$N_2(2)$	0-5958	0-5934	21.187	1.403 10 ⁻⁵	4.759
$CO_2(3)$	0-6000	0-4239	34.824	5.833 10 ⁻⁵	5.086

137

138 **3.1 Bulk gas phase model**

139 The eqns (3-7) must be modified introducing fugacity for the description of the bulk gas phase and activity 140 coefficients for the adsorbed phase. The SRK [35, 36] equation of state has been used for the evaluation of 141 the fugacity coefficients.

142
$$Z^3 - Z^2 + Z(A - B - B^2) - AB = 0$$
 (11)

143
$$\ln\left(\varphi_{i,bulk}\right) = \frac{b_i}{b} (Z-1) - \ln\left(Z-B\right) - \frac{A}{B} \left(2\frac{a_i^{1/2}}{a^{1/2}} - \frac{b_i}{b}\right) \ln\left(1 + \frac{B}{Z}\right)$$
 (12)

Eq. (11) is the fugacity coefficient of component i in the bulk gas phase from SRK equation of state. The correlations for the calculation of the parameters of eq. (11) and (12) are reported elsewhere [37]. The interaction parameters are neglected in all of the following calculations and the number of total moles of the mixture in the bulk phase (G) is derived by the compressibility factor Z as follows:

148
$$G = Z \frac{P_{bulk} V_{void}}{RT}$$
(13)

149150 **3.2** Excess Gibbs energy model for the adsorbed phase

151 The adopted activity coefficient model is the ABC equation [38]. At constant temperature, it proposes the 152 following expression for the excess Gibbs energy:

153
$$g_{ex} = \sum_{i=1}^{NC} \sum_{j=1}^{NC} A_{0,ij} x_i x_j \left(1 - e^{-C_{ij} \psi} \right)$$
(14)

154 Its application requires the preliminary regression on the experimental data of binary systems of the binary 155 interaction parameters $A_{0,ij}$ and C_{ij} at constant temperature. The results of the regression on the binary 156 experimental data in [34] are reported in Table 6.

157

Table 6: Parameters of the ABC equation for Methane, Nitrogen and Carbon Dioxide binary systems on activated carbon Norit R1 Extra at 298 K. Experimental data are from [34].

	Fugacity range	A _{0,ij}	C _{ij}
Components (i/j)	[kPa]	[kJ mol ⁻¹]	$[\text{kg mol}^{-1}]$
$CH_4/N_2(1/2)$	150.9-5578.7	-1.726	10.060
CH_4/CO_2 (1/3)	97.7-5342.1	0.474	0.0162
$N_2/CO_2(2/3)$	107.9-4459.2	1.017	14.699

(15)

159 The activity coefficients are defined as: 2(2k-1)

160
$$RT \ln (\gamma_i) = \frac{\partial (Ng_{ex})}{\partial n_i} \Big|_{T, \psi, n_j}$$

161 where N is the total amount of moles adsorbed. According to the definition of eq. (15), for a ternary system,

162 the activity coefficients result in:

¹⁵⁸

$$\ln (\gamma_{1}) = \frac{\left(e^{-(C_{12}+C_{13}+C_{23})\psi}\right)}{RT(x_{1}+x_{2}+x_{3})^{2}} \begin{pmatrix} A_{0,12}e^{(C_{13}+C_{23})\psi}\left(e^{C_{12}\psi}-1\right)x_{2}(x_{2}+x_{3})+\\ +A_{0,23}\left(e^{(C_{12}+C_{13})\psi}-e^{(C_{12}+C_{13}+C_{23})\psi}\right)x_{2}x_{3}+\\ +A_{0,13}e^{(C_{12}+C_{23})\psi}\left(e^{C_{13}\psi}-1\right)(x_{2}+x_{3})x_{3} \end{pmatrix}$$

$$\ln (\gamma_{2}) = \frac{\left(e^{-(C_{12}+C_{13}+C_{23})\psi}\right)}{RT(x_{1}+x_{2}+x_{3})^{2}} \begin{pmatrix} A_{0,12}e^{(C_{13}+C_{23})\psi}\left(e^{C_{12}\psi}-1\right)x_{1}(x_{1}+x_{3})+\\ +A_{0,13}\left(e^{(C_{12}+C_{23})\psi}-e^{(C_{12}+C_{13}+C_{23})\psi}\right)x_{1}x_{3}+\\ +A_{0,23}e^{(C_{12}+C_{23})\psi}\left(e^{C_{23}\psi}-1\right)(x_{1}+x_{3})x_{3} \end{pmatrix}$$

$$\ln (\gamma_{3}) = \frac{\left(e^{-(C_{12}+C_{13}+C_{23})\psi}\right)}{RT(x_{1}+x_{2}+x_{3})^{2}} \begin{pmatrix} A_{0,13}e^{(C_{13}+C_{23})\psi}\left(e^{C_{13}\psi}-1\right)x_{1}(x_{1}+x_{2})+\\ +A_{0,12}\left(e^{(C_{13}+C_{23})\psi}-e^{(C_{12}+C_{13}+C_{23})\psi}\right)x_{1}x_{2}+\\ +A_{0,23}e^{(C_{12}+C_{13})\psi}\left(e^{C_{23}\psi}-1\right)(x_{1}+x_{2})x_{2} \end{pmatrix}$$

$$(18)$$

To take into account the non-idealities both in adsorbed and bulk gas phase, eqns. (6) and (7) are now replaced by eqns (19) and (20):

168
$$\psi_i = \int_{0}^{f_{i,ads}} n_i d\left(\ln f_i\right) \quad i = 1, 2,NC$$
 (19)

169 Eq. (19) is an integral function of each single isotherm. In the ideal case it is fitted directly on the 170 experimental pressures, in the non-ideal the isotherms have to be fitted on the fugacity, modifying the 171 experimental pressure with an equation of state (in this case SRK).

172
$$\frac{m_{ads}}{N} = ex + \sum_{i=1}^{NC} \frac{x_i}{n_i}$$
 (20)

173 where the excess contribution to the total number of adsorbed moles *ex* is given by:

174
$$ex = \frac{\partial \left(g_{ex} / RT\right)}{\partial \psi} \bigg|_{T,x} = \sum_{i=1}^{NC} \sum_{j=1}^{NC} \left(\frac{A_{0,ij}C_{ij}}{RT} x_i x_j e^{-C_{ij}\psi}\right)$$
(21)

The two contributions on the right hand side of eq. (20) are respectively the contribution of the non-idealities in the adsorbed phase according to the ABC equation and the contribution considering ideal adsorption of a non-ideal gas mixture. In fact, in eq. (20), n_i is the adsorption isotherm of the pure component i with parameters regressed on fugacities instead of pressures [28].

180 **3.3.** Non-ideal flash calculation

181 The k_i factors for the non-ideal case are no longer only function of pressures but also of the compositions 182 because of the presence of the fugacity coefficient in the bulk gas phase ($\varphi_{i,bulk}$), the activity coefficient of the 183 adsorbed phase (γ_i) and the fugacity of the pure component i in adsorbed phase ($f_{i,ads}$). Therefore k_i factors 184 become:

185
$$\frac{y_i}{x_i} = \frac{f_{i,ads}\gamma_i}{\varphi_{i,bulk}P_{bulk}}$$
(22)

186 This feature makes the system of equations larger and more nonlinear than the ideal case. Furthermore, in 187 order to have consistency between the number of equations and the number of variables, for a three 188 components mixture, three isofugacity condition equations and conservation of moles equation for one of the 189 components must be included.

190 The Newton method can again effectively solve this problem, providing an initial guess reasonably close to 191 the actual solution, e.g. assuming a value of 1 for fugacity and activity coefficients. After some sensitivity 192 analysis, it has been noticed that the convergence to a feasible solution is affected by the initial value of P_{bulk}

193 more than the initial value of the other variables.

Assuming the parameters of Tables 5 and 6, the results of the calculation for the non-ideal high pressure case are reported in Table 7. Carbon dioxide, being the most strongly adsorbed component, exhibits a higher

196 concentration in the adsorbed phase compared to the feed; an opposite trend can be seen for methane and 197 nitrogen components. The result is again validated testing the presence of the common tangent plane of the

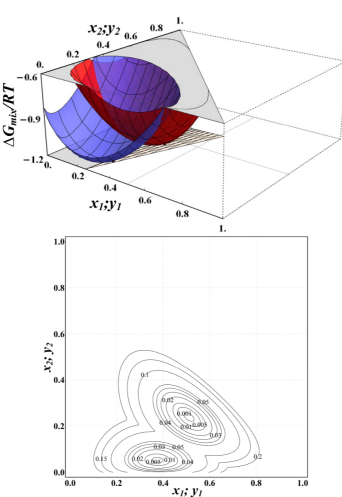
198 Gibbs energy of mixing surface at the equilibrium (Fig. 3).

199

Table 7: Solution for the non-ideal VT flash calculation at the conditions: $V_{void} = 2.133 \ 10^{-3} \ m^3$; F = 10 moles; T = 298 K

Component	zi	P _{bulk}	ψ_{eq}	Xi	γ_i	$f_{i,ads}$	yi	$\phi_{i,bulk}$
		[kPa]	[kg/mol]			[kPa]		
$CH_4(1)$	0.4	2837.29	14.336	0.373	0.980	3784.88	0.511	0.511
$N_2(2)$	0.1			0.052	0.970	13724.31	0.239	0.239
$\overline{CO_2}(3)$	0.5			0.575	1.029	1055.05	0.250	0.250





201

202

206

Figure 3: Common tangent plane of the Gibbs energy of mixing locating at the equilibrium point calculated
 in Table 7 (top). Contour plot of the distance function between Gibbs energy of mixing and common tangent
 plane at the equilibrium point of Table 7 (bottom).

207 4. Enthalpy of adsorption

The thermal energy required for compression includes both enthalpy of desorption and sensible heat. In case of multicomponent mixture, the enthalpy of desorption ΔH_{ads} [kJ] can be calculated following the derivation described in [38]:

211
$$\Delta H_{ads} = m_{ads} \sum_{i=1}^{NC} \left(\int_{0}^{n_{i}^{o}} \Delta \overline{h}_{i} \, dn_{i} \right)$$
(23)

where $\Delta \bar{h_i}$ is the differential enthalpy of adsorption [kJ mol⁻¹] of the ith component of the mixture. For nonideal adsorbed solutions and non-ideal bulk gases, the differential enthalpy is:

$$\Delta \overline{h}_{i} = \Delta h_{i}^{o} + RT^{2} \left(\frac{\partial \ln \gamma_{i}}{\partial T} \right)_{\psi,x} + \left[\frac{1}{n_{i}^{o}} + \left(\frac{\partial \ln \gamma_{i}}{\partial \psi} \right)_{T,x} \right]$$

$$214 \times \left[\frac{\sum_{j} x_{j} G_{j}^{o} n_{j}^{o} \left(\Delta \overline{h}_{j}^{o} - \Delta h_{j}^{o} \right) + RT^{2} \left(\frac{\partial (ex)}{\partial T} \right)_{\psi,x}}{\sum_{j} x_{j} G_{j}^{o} - \left(\frac{\partial (ex)}{\partial T} \right)_{T,x}} \right]$$

$$i=1,2,...NC$$
(24)

215 which, for an ideal gas adsorbed ideally, becomes:

216
$$\Delta \overline{h_i} = \Delta h_i^o + \frac{1}{n_i^o} \left(\frac{\sum_j x_j G_j^o n_j^o \left(\Delta \overline{h_j^o} - \Delta h_j^o \right)}{\sum_j x_j G_j^o} \right)$$
 i=1,2,...NC (25)

217 The terms composing eq. (24) are:

218
$$G_i^o = \frac{1}{\left(n_i^o\right)^2} \left(\frac{\partial \ln n_i^o}{\partial \ln f_i}\right)_T$$
 i=1,2,...NC (26)

Eq. (24) is a function of the pure component molar integral enthalpy of adsorption [kJ mol⁻¹], which neglecting the Poynting term, becomes:

221
$$\Delta h_i^o = \frac{1}{n_i^o} T^2 \frac{\partial}{\partial T} \left(\frac{\Omega_i}{T}\right)_{f_i}$$
(27)

where Ω [kJ kg⁻¹] is the grand potential of adsorption. The other variable in eq. (24) is the pure component differential enthalpy [kJ mol⁻¹]:

224
$$\Delta \overline{h}_{i}^{o} = RT^{2} \left(\frac{\partial \ln f_{i}}{\partial T} \right)_{n_{i}}$$
(28)

Assuming that the specific heats of the gases are constant in the operating range of temperatures and the specific heats in adsorbed and bulk gas phase are identical [39], the sensible heat can be derived as follows:

227
$$\Delta H_{sens} = \left[m_{ads} c_{p,ads} + \sum_{i}^{NC} \left(n_{i,bulk} + n_{i,ads} \right) c_{p,i} \right] \left(T_{heating} - T_{ads} \right)$$
(29)

228 where $c_{p,ads}$ is the specific heat capacity of the adsorbent assumed 0.858 kJ kg⁻¹ K⁻¹ [11], $c_{p,i}$ is the bulk gas 229 phase specific heat capacity of the ith component of the mixture [kJ mol⁻¹ K⁻¹] as reported in [40]. $n_{i,bulk}$ is the 230 number of moles of component i in the bulk gas phase [mol] and $n_{i,ads}$ is the number of moles of component i 231 in the adsorbed phase [mol], T_{ads} is the adsorption temperature which is kept at 298 K and $T_{heating}$ is the 232 heating temperature.

234 5. Ideal adsorption compression

The case of a solid compressor is presented where 1 kg of zeolite 13X adsorbs from a three components feed mixture of Nitrogen, Oxygen, Carbon Dioxide with composition (0.7897,0.2099, 0.0004) mole fraction at 101.325 kPa, 298 K in accordance to the ideal adsorbed solution theory, with adsorption isotherms parameters reported in Table 8.

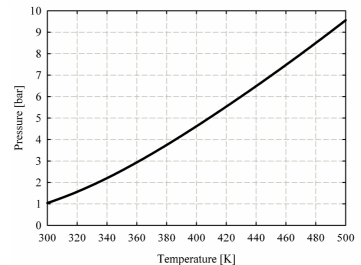
239

233

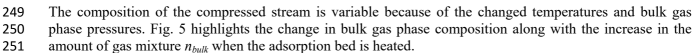
Table 8: Langmuir, Dual Site Langmuir and Toth isotherm parameters of the ideal and non-ideal adsorption zeolite 13X compressor

-		Isotherm	q_{s1}	b ₀₁	ΔH_1	q_{s2}	b ₀₂	ΔH_2	t	Ref
			[mol/kg]	[kPa ⁻¹]	[kJ mol ⁻¹]	$[mol kg^{-1}]$	[kPa ⁻¹]	[kJ mol ⁻¹]		
	N_2	Langmuir	4.725	3.616 10 ⁻⁷	-19.59					[41]
	O_2	Langmuir	4.650	1.452 10 ⁻⁵	-7.80					[41]
	CO_2	Dual site Langmuir	3.573	$1.681 \ 10^{-8}$	-35.00	1.789	3.812 10 ⁻⁹	-49.26		[42]
	C_2H_4	Toth	2.21	5.22 10 ⁻⁶	-21.4				1.75	[43]
210-	C_2H_6	Toth	2.72	1.13 10 ⁻⁷	-36.3				0.97	[43]

- 241 After adsorption, the bed is heated up at closed volume. The significant amount of material moving from the
- 242 adsorbed phase to the bulk gas phase causes a pressure increase with a magnitude depending on the heating 243 temperature (Fig. 4).



Temperature [K]
 Figure 4: Evolution of the bulk gas phase pressure heating the bed at different temperature levels for 1 kg of
 zeolite 13X compressing a stream of Nitrogen, Oxygen, Carbon Dioxide adsorbed at 101.325 kPa, 298 K and
 composition (0.7897, 0.2099, 0.0004) mole fraction.



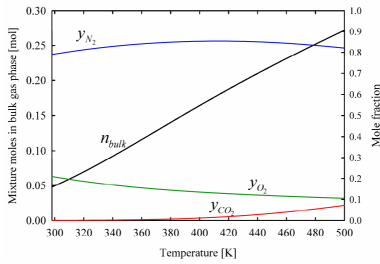


Figure 5: Evolution of the gas mixture moles (left) and component mole fraction (right) at different heating
 temperatures for 1 kg of zeolite 13X compressing a stream of Nitrogen, Oxygen, Carbon Dioxide adsorbed at
 101.325 kPa, 298 K and composition (0.7897, 0.2099, 0.0004) mole fraction.

- 256
- Fig. 6 shows that the dominant part in the total energy required for compression is the sensible heat, accounting, in this particular case, for an average of 91% of the total energy.

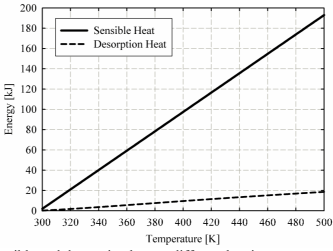


Figure 6: Evolution of sensible and desorption heat at different heating temperatures for 1 kg of zeolite 13X
 compressing a stream of Nitrogen, Oxygen, Carbon Dioxide adsorbed at 101.325 kPa, 298 K and
 composition (0.7897, 0.2099, 0.0004) mole fraction

263

264 6. Non-ideal adsorption compression

When adsorption cannot be described as ideal, ideal and non-ideal interpretations can results in significant 265 266 performance discrepancies. In this section a gas mixture of Carbon Dioxide/Ethylene/Ethane adsorbed on zeolite 13X is considered. This gas mixture shows non-ideal behaviour in gas phase already at moderate 267 pressure and ambient temperature and non-idealities in adsorbed phase on zeolite 13X. As measured in [30 268 269 38], ABC equation excess model can describe correctly the equilibrium of a non-ideal adsorbed phase of Carbon Dioxide/Ethylene/Ethane on zeolite 13X with the binary interaction parameters of Table 9. The bulk 270 gas phase, especially for Ethane and Ethylene, require the specification of an equation of state for the 271 calculation of fugacity coefficient in the range of operating conditions. Accordingly SRK equation of state is 272 used to describe the thermodynamic behaviour of the bulk gas phase. 373

Unitary systems on .	Zeonie 15A. Experimer		[58].
Components	A_{ij}	\mathbf{B}_{ij}	\mathbf{C}_{ij}
(i/j)	$[kJ mol^{-1}]$	$[kJ mol^{-1} K^{-1}]$	$[kg mol^{-1}]$
$CO_2/C_2H_6(1/3)$	-10.0	0.01917	0.110
$CO_2/C_2H_4(1/2)$	-6.5	0.01450	0.030
$C_2H_4/C_2H_6(2/3)$	-4.5	0.00437	0.067
Note: temperature	dependent interaction	parameters are	considered according to

Table 9: Parameters of the ABC equation for Carbon Dioxide, Ethylene and Ethane binary systems on zeolite 13X. Experimental data are from [38].

275

Note: temperature dependent interaction parameters are considered according to $A_{0,ij} = A_{ij} + T B_{ij}$

276 In the first operational step, the adsorption compressor is loaded at 101.325 kPa and 298 K with a multicomponent gas mixture having composition (0.05, 0.15, 0.8) mole fraction. Table 10 reports the 277 278 distribution of the components in bulk gas phase and adsorbed phase at this initial adsorption step. The non-279 ideal approach predicts more amount of initial gas in the vessel than the ideal approach. The main difference is in the carbon dioxide adsorbed. This is because both the binary interaction parameters of carbon dioxide. 280 which is the most strongly adsorbed compound, are higher in magnitude than the interaction parameters of 281 282 ethane/ethylene, making carbon dioxide equilibrium more sensible to the non-ideal approach. However, total amount of moles in the vessel does not change significantly because carbon dioxide is present in small 283 284 amount in the feeding mixture.

- 285
- 286 287
- 288
- 289

- 291
- 292

compressor after a	adsorption at 101.32.	5 kPa and 298 I	x for 1 kg of zeo	me 13A.
	Carbon Dioxide	Ethylene	Ethane	Total
Ideal case				
Bulk gas phase	0.002	0.007	0.038	0.048
Adsorbed phase	0.459	0.041	2.183	2.683
Total	0.461	0.048	2.221	2.731
Non-ideal case				
Bulk gas phase	0.002	0.007	0.038	0.047
Adsorbed phase	0.756	0.045	2.079	2.880
Total	0.758	0.052	2.117	2.927

Table 10: comparison of the amount of moles between ideal and non-ideal compressor after adsorption at 101.325 kPa and 298 K for 1 kg of zeolite 13X.

304

294 The excess part of the total adsorbed amount (eq. (20)) makes the compressor performance also dependent of

the excess thermodynamic potentials which are inverse functions of temperature. This results in a decrease of

the excess adsorbed amount with increasing temperatures. Thus, the non-ideal approach introduces a higher sensitivity of equilibrium with temperature. This is illustrated in Fig. 7, where in the non-ideal compressor

the release of gas from the adsorbed phase to the bulk gas phase is larger than the ideal compressor, resulting

eventually in higher equilibrium pressures.

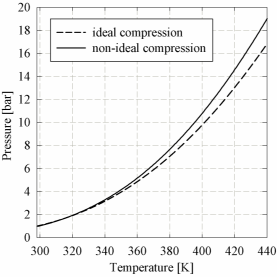


Figure 7: Compression of a gas mixture of Carbon Dioxide/Ethylene/Ethane with 1 kg of zeolite 13X and
 (0.05,0.15,0.80) feed composition, adsorbed at 101.325 kPa, 298 K. Pressure profile against heating
 temperature in the ideal (dashed line) and non-ideal (solid line) cases.

In Fig.8, the change in the activity coefficients over temperature promotes increasingly higher pressures.
 Among the three components, carbon dioxide shows the steepest trend, spanning from 0.55 at 298 K to 0.92
 at 440 K. Fugacity coefficients vary in smaller proportion. In this case ethane has the larger variation ranging
 between 0.992 at 298 K and 0.96 at 340 K.

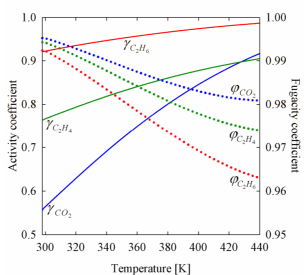
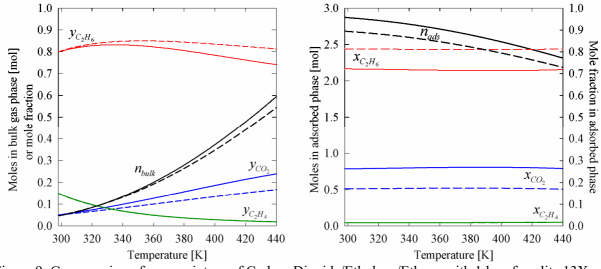




Figure 8: Compression of a gas mixture of Carbon Dioxide/Ethylene/Ethane with 1 kg of zeolite 13X and (0.05, 0.15, 0.80) feed composition, adsorbed at 101.325 kPa, 298 K. Activity and fugacity coefficients profiles against heating temperature.

Compositions in bulk gas phase and adsorbed phase are illustrated in Fig. 9, where adsorbed phase compositions keep essentially constant throughout the heating range, while bulk gas phase compositions visibly change. Also a significant difference is observed in the adsorbed phase compositions of carbon dioxide and ethane between ideal and non-ideal approach already in the adsorption step. The feature of the non-ideal approach of predicting higher pressures depends also on this initial difference.



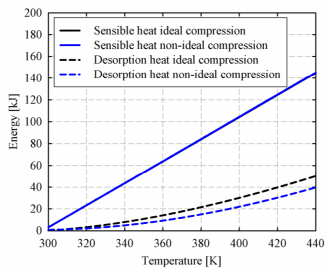
319 320

Figure 9: Compression of a gas mixture of Carbon Dioxide/Ethylene/Ethane with 1 kg of zeolite 13X and
 (0.05,0.15,0.80) feed composition, adsorbed at 101.325 kPa, 298 K. Comparison between compositions, bulk
 gas phase amount and adsorbed amount against heating temperature in the ideal (dashed line) and non-ideal
 (solid line) cases.

324

Fig. 9 shows also that in the bulk gas phase, at temperatures <320 K (<2 bar), the ideal and non-ideal approaches provide the same results, while in the adsorbed phase the two approaches result in significant discrepancies already at low pressure. This is due to non-idealities in the adsorbed phase.

Energy consumption for the two cases is depicted in Fig. 10, where the difference in sensible energy is negligible because it is dominated by the adsorption material heat capacity more than by the adsorbed amount. A difference between ideal and non-ideal desorption heat can be observed already at temperatures >320 K. This difference increases up to 10.5 kJ at 440 K. This results in a lower total energy consumption for the non-ideal compression than ideal compression.



333

Figure 10: Evolution of sensible and desorption heat at different heating temperatures for 1 kg of zeolite 13X
 compressing a stream of Nitrogen, Oxygen, Carbon Dioxide. Comparison between ideal and non-ideal
 compression.

Besorption heat is dominated by the pure component molar integral enthalpies (Δh_i^0) which are similar for the ideal and non-ideal approaches. The second major contribution to the desorption heat consists of the term $RT^2 (\partial ln\gamma_i/\partial T)_{\psi,x}$ which is negligible for all gases except that ethane. This builds the difference between ideal compressor and non-ideal compressor in terms of energy consumption. These results show that considerations based on ideal adsorbed solution theory lead to conservative assessment of the compressor performance, with lower pressure generated at higher energy consumption.

345 5. Conclusion and outlook

346 Adsorption compressor is an emerging technology which is currently under development for the next long term Mars and lunar missions. Thermodynamics of this system is based on the solution of the isochoric-347 348 isothermal flash problem for bulk gas/adsorbed phases. A system composed by Rachford-Rice equation and additional equations embedding the iso-reduced grand potential condition enables the determination of the 349 compressed state. The flash problem has been solved in both cases respectively for two ternary mixtures and 350 351 validated through its thermodynamic consistency testing the presence of the common tangent plane to the Gibbs energy of mixing at the equilibrium compositions. In this way the applicability of the Rachford-Rice 352 equation is extended to the adsorbed solution theory. This procedure has been applied for the characterisation 353 of one zeolite 13X compressor working ideally with a Nitrogen/Oxygen/Carbon Dioxide gas mixture and 354 non-ideally with a Carbon Dioxide/Ethylene/Ethane mixture. In the latter non-ideal mixture, calculations 355 356 based on ideal approach proved to be more conservative compared to the non-ideal compressor performance. The application of the proposed thermodynamic framework is at the basis of adsorption compressor 357 performance calculation. The precise design of the single unit and of multi-compressor systems [44] can be 358 359 reached by embedding the described approach of the equilibrium in the wider dynamic description of the compressor. This impacts directly on the development of sustainable systems that capture, compress and 360 concentrate gases in extremely dilute conditions such as planetary and extra-planetary systems for 361 362 greenhouse gases removal and compression [45, 46].

363

364 Acknowledgements

The research leading to these results has received funding from the European Union Seventh Framework
 Programme (FP7/2007-2013) under grant agreement n°630863.

367368 Nomenclature

369	c _{p,ads}	Specific heat capacity of the adsorben material [kJ kg ⁻¹ K ⁻¹]
370	c _{p,i}	Specific heat capacity of component i [kJ mol ⁻¹ K ⁻¹]
371	ex	Excess contribution to the total number of adsorbed moles [kg mol ⁻¹]
372	F	Total number of moles in the feed [mol]

373	$\mathbf{f}_{i,ads}$	Fugacity of pure component i at the system temperature and pressure [kPa]
374	G	Total number of moles in the bulk gas phase [mol]
375	g _{ex}	Excess Gibbs energy [kJ mol ⁻¹]
376	k _i	Equilibrium constant of component i
377	m _{ads}	Mass of adsorbent [kg]
378	Ν	Total number of moles in the adsorbed phase [mol]
379	NC	Number of components participating in the adsorption
380	n _i	Specific amount adsorbed of component i [mol kg ⁻¹]. It is calculated directly from adsorption
381	isotherm using	; either P_i^0 (ideal adsorbed phase) or adsorbed phase fugacity $f_{i,ads}$ (non-ideal adsorbed phase).
382	n _{i,ads}	Amount of adsorbed moles of component i [mol] in adsorbed phase phase;
383	n _{i,bulk}	Amount of adsorbed moles of component i [mol] in bulk gas phase;
384	P _{bulk}	Pressure of the mixture in the bulk gas phase [kPa]
385	$\mathbf{P_i}^0$	Bulk pressure of the component i at a given reduced grand potential ψ_i [kPa]
386	R	Universal gas constant [kJ mol ⁻¹ K ⁻¹]
387	Т	Equilibrium temperature [K]
388	T_{ads}	Temperature of the adsorption bed during adsorption [K]
389	Theating	Temperature of the bed after heating [K]
390	V _{void}	Volume occupied by the bulk gas phase (dead volume) [m ³]
391	Xi	Molar fraction of the component i in the adsorbed phase
392	y _i	Molar fraction of the component i in the bulk gas phase
393	Zi	Molar fraction of the component i in the feed
394	Ζ	Compressibility factor
395	ΔH_{ads}	Enthalpy for adsorption or desorption [kJ];
396	ΔH_{sens}	Sensible thermal energy and its bulk gas and adsorbed phases [kJ];
397	$\Delta ar{ extbf{h}}_{ extbf{i}}$	Differential enthalpy of adsorption [kJ mol ⁻¹] for component i in the mixture;
398	$\Delta {ar{ extbf{h}}_{ extbf{i}}}^{0}$	Differential enthalpy of adsorption [kJ mol ⁻¹] for pure component i;
399	Δh_i^0	Integral enthalpy of adsorption [kJ mol ⁻¹] for pure component i;
400		
401	Greek letters	
402	γ _i	Activity coefficient of component i
403	ε _b	Bed porosity
404	ε _p	Particle porosity
405	$\rho_{\rm b}$	Bed bulk density [kg m ⁻³]
406	φ _{i,bulk}	Fugacity coefficient of component i in the bulk gas phase
407	Ψeq	Reduced grand potential at equilibrium [mol kg ⁻¹]
100	N/	Peduced grand notantial of component i [mal ka ⁻¹]

- 408 ψ_i Reduced grand potential of component i [mol kg⁻¹]

411 **References**

- [1] N. Augustine, J. Ling, P. Xiao, P.A. Webley, Y. Zhai, CO₂ Capture by Temperature Swing Adsorption:
 Use of Hot CO₂-Rich Gas for Regeneration. Industrial Engineering Chemistry Research 55 (2016) 703-713.
- [2] G. Maggio, L.G. Gordeeva, A. Freni, Yu.I. Aristov, G. Santori, F. Polonara, G. Restuccia, Simulation of
 a solid sorption ice-maker based on the novel composite sorbent "lithium chloride in silica gel pores".
 Applied Thermal Engineering 29 (2009) 1714-1720.
- [3] J.C. Atuonwu, X. Jin, G. van Straten, H.C. van Deventer Antonius, J.B. van Boxtel, Reducing energy
 consumption in food drying: Opportunities in desiccant adsorption and other dehumidification strategies.
 Procedia Food Science 1 (2011) 1799-1805.
- [4] L. Bonaccorsi, P. Bruzzaniti, L. Calabrese, A. Freni, E. Proverbio, G. Restuccia, Synthesis of SAPO-34
 on graphite foams for adsorber heat exchangers. Applied Thermal Engineering 61 (2013) 848-852.
- 426 [5] A. Freni, L. Bonaccorsi, L. Calabrese, A. Caprì, A. Frazzica, A. Sapienza, SAPO-34 coated adsorbent 427 heat exchanger for adsorption chillers. Applied Thermal Engineering 82 (2015) 1-7.
- 428

414

418

422

- [6] Y.Y. Li, S.P. Perera, B.D. Crittenden, J. Bridgwater, The effect of the binder on the manufacture of a 5A
 zeolite monolith. Powder Technology 116 (2001) 85-96.
- [7] G. Li, R. Singh, D. Li, C. Zhao, L. Liu, P.A. Webley, Synthesis of biomorphic zeolite honeycomb
 monoliths with 16000 cells per square inch. Journal of Materials Chemistry 19 (2009) 8372-8377.
- [8] R.E. Critoph, Y. Zhong, Review of trends in solid sorption refrigeration and heat pumping technology.
 Proceedings of the Institution of Mechanical Engineers 219 (2005) 285-300.
- 437

449

453

456

459

462

466

469

472

476

479

482

431

434

[9] G. Santori,S. Santamaria, A. Sapienza, S. Brandani, A. Freni, A., A stand-alone solar adsorption
refrigerator for humanitarian aid. Solar Energy 100 (2014) 172-178.

- [10] G. Santori, A. Sapienza, A. Freni, A dynamic multi-level model for adsorptive solar cooling. Renewable
 Energy 43 (2012) 301-312.
- [11] G. Santori, A. Frazzica, A. Freni, M. Galieni, L. Bonaccorsi, F. Polonara, G. Restuccia, Optimization
 and testing on an adsorption dishwasher. Energy 50 (2013) 170-176.
- [12] J.W. Wu, M.J. Biggs, P. Pendleton, A. Badalyan, E.J. Hu, Experimental implementation and validation
 of thermodynamic cycles of adsorption-based desalination. Applied Energy 98 (2012) 190-197.
- [13] L. Joss, M. Gazzani, M. Hefti, D. Marx, M. Mazzotti, Temperature Swing Adsorption for the Recovery
 of the Heavy Component: An Equilibrium-Based Shortcut Model. Industrial and Engineering Chemistry
 Research 54 (2015) 3027-3038.
- [14] E.M. Mattox, J.C. Knox, D.M. Bardot, Carbon dioxide removal system for closed loop atmosphere
 revitalization, candidate sorbents screening and test results. Acta Astronautica 86 (2013) 39-46.
- [15] R.J. Kay, D. El Sherif, Carbon Dioxide Removal Assembly Performance Comparison. SAE Technical
 paper 2009-01-2431.
- 460 [16] L. Mulloth, J. Finn, A Solid-State Compressor for Integration of CO₂ Removal and Reduction
 461 Assemblies. SAE Technical Paper 2000-01-2352.
- [17] L.M. Mulloth, J.E. Finn, Air quality systems for related enclosed spaces: spacecraft air, in: D. Barceló,
 A.G. Kostianoy (Eds), The Handbook of Environmental Chemistry, Springer-Verlag, Berlin, 2005, part H,
 pp. 383-404.
- [18] E. Seedhouse, Lunar Outpost. The Challenges of Establishing a Human Settlement on the Moon, PraxisPublishing Ltd, Chichester, UK, 2009.
- [19] H.J. Strumpf, C.Y. Chin, G.R. Lester, S.T. Homeyer, Sabatier Carbon Dioxide Reduction System for
 Long-Duration Manned Space Application, SAE Technical Paper 911541.
- [20] J.C. Knox, M. Campbell, L.A. Miller, L. Mulloth, M. Varghese, B. Luna, Integrated Test and Evaluation
 of a 4-Bed Molecular Sieve, Temperature Swing Adsorption Compressor, and Sabatier Engineering
 Development Unit, SAE Technical Paper 2006-01-2271
- 477 [21] R.P. Hoover, P.C. Wankat, Gas compression using temperature swing adsorption. Separation Science478 and Technology 37 (2002) 3187-3199.
- 480 [22] J. R. Moate, Temperature swing adsorption compression and membrane separations, PhD thesis,
 481 Vanderbilt University, 2009.
- [23] J.R. Moate, M.D. LeVan, Temperature swing adsorption compression: Effects of nonuniform heating on
 bed efficiency. Applied Thermal Engineering 30 (2010) 658-663.

- 485
 486 [24] V.F. Cabral, M. Castier, F.W. Tavares, Thermodynamic equilibrium in systems with multiple adsorbed
 487 and bulk phases. Chemical Engineering Science 60 (2005) 1773-1782.
 488
 - [25] A.L. Myers, J.M. Prausnitz, Thermodynamics of mixed-gas adsorption. AIChE Journal 11 (1965) 121 127.
 - 492 [26] G. Santori, M. Luberti, S. Brandani, Common tangent plane in mixed-gas adsorption. Fluid Phase
 493 Equilibria 392 (2015) 49-55.
 - [27] H.H. Rachford Jr., J.D. Rice, Procedure for Use of Electronic Digital Computers in Calculating Flash
 Vaporization Hydrocarbon Equilibrium. Journal of Petroleum Technology 195 (1952) 327-328.
 - 498 [28] O. Talu, A. Myers, Rigorous Thermodynamic Treatment of Gas Adsorption. AIChE Journal 34 (1988)499 1887-1893.
 - [29] F.V.S. Lopes, C.A. Grande, A.M. Ribeiro, J.M. Loureiro, O. Evaggelos, V. Nikolakis, A.E. Rodrigues.
 Adsorption of H₂, CO₂, CH₄, CO, N₂ and H₂O in Activated Carbon and Zeolite for Hydrogen Production.
 Separation Science and Technology 44 (2009) 1045-1073.
 - 505 [30] S.K. Henninger, M. Schicktanz, P.P.C. Hugenell, H. Sievers, H.M. Henning, Evaluation of methanol
 506 adsorption on activated carbons for thermally driven chillers part I: Thermophysical characterisation.
 507 International Journal of Refrigeration 35 (2012) 543-553.
 - [31] P. Xiao, J. Zhang, P. Webley, G. Li, R. Singh, R. Todd, Capture of CO₂ from flue gas streams with
 zeolite 13X by vacuum-pressure swing adsorption. Adsorption 14 (2008) 575-582.
 - [32] H. Verelst, G.V. Baron, Adsorption of oxygen, nitrogen, and argon on 5A molecular sieve. Journal of
 Chemical and Engineering Data 30 (1985) 66-70.
 - [33] O. Talu, I. Zwiebel, Multicomponent Adsorption Equilibria of Nonideal Mixtures. AlChE Journal 32
 (1986) 1263-1276.
 - 518 [34] F. Dreisbach, R. Staudt, J.U. Keller, High Pressure Adsorption Data of Methane, Nitrogen, Carbon
 519 Dioxide and their Binary and Ternary Mixtures on Activated Carbon. Adsorption 5 (1999) 215-227.
 520
 - [35] S. Gamba, G.S. Soave, L.A. Pellegrini, Use of normal boiling point correlations for predicting critical
 parameters of paraffins for vapour-liquid equilibrium calculations with the SRK equation of state. Fluid
 Phase Equilibria 276 (2009) 133-141.
 - [36] G. Soave, Equilibrium constants from a modified Redlich-Kwong equation of state. Chemical
 Engineering Science 27 (1972) 1197-1203.
 - [37] O. Redlich, J.N.S. Kwong, On the Thermodynamics of Solutions. V. An Equation of State. Fugacities of
 Gaseous Solutions. Chemical Reviews 44 (1949) 233-244.
 - 531 [38] F.R. Siperstein, A.L. Myers, Mixed Gas Adsorption. AIChE Journal 47 (2001) 1141-1159.
 - [39] K.S. Walton, M.D. LeVan, Adsorbed-Phase Heat Capacities: Thermodynamically Consistent Values
 Determined from Temperature-Dependent Equilibrium Models. Industrial and Engineering Chemistry
 Research 44 (2005) 178-182.
 - 537 [40] E.W. Lemmon, M.O. McLinden, M.L. Huber, NIST Standard Reference Database 23, NIST
 538 Thermodynamic Properties of Refrigerants and Refrigerant Mixtures Database (REFPROP) Version 9.0.
 539 Gaithersburg: National Institute of Standards and Technology (2010)

491

494

497

500

504

508

511

514

517

524

527

530

- [41] M. L. Zanota, H. Heymans, F. Gilles, B.L. Su, M. Frère, G. De Weireld, Adsorption Isotherms of Pure
 Gas and Binary Mixtures of Air Compounds on Faujasite Zeolite Adsorbents: Effect of Compensation
 Cation. Journal of Chemical and Engineering Data 55 (2010) 448-458.
- 544
 545 [42] F.V.S. Lopes, C.A. Grande, A.M. Ribeiro, V.J.P. Vilar, J.M. Loureiro, A.E. Rodrigues, Effect of Ion
 546 Exchange on the Adsorption of Steam Methane Reforming Off-Gases on Zeolite 13X. Journal of Chemical
 547 and Engineering Data 55 (2010) 184-195.
- [43] M.A. Granato, V.D. Martins, J.C. Santos, M. Jorge, A.E. Rodrigues, From molecules to processes:
 molecular simulations applied to the design of simulated moving bed for ethane/ethylene separation. The
 Canadian Journal of Chemical Engineering 92 (2014) 148-155.
- 553 [44] G. Santori, Open-cycle adsorption compression train, Sorption Friends meeting, Milazzo, Italy, 2015.
- [45] M.-C. Ferrari, D. Friedrich, G. Santori, S. Brandani, Design of a small scale air capture system, FOA11:
 11th International Symposium on the Fundamentals of Adsorption, Baltimore, MD, United States, 2013.
- 558 [46] C. Charalambous, M.-C. Ferrari, S. Brandani, G. Santori, Thermally-driven adsorption, concentration 559 and purification of highly dilute gases, Heat Powered Cycle Conference 2016, Nottingham, UK, 2016.
- 560

552

554