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Flexible CO₂ capture for open-cycle gas turbines via vacuum-pressure swing adsorption: a model-based assessment

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Abstract:

As energy systems require flexible and responsive power generators to combat network imbalances, CO₂ post-combustion capture (PCC) technologies need to be capable of transient operation. However, currently only amine absorption has been investigated for its efficacy in Flexible-PCC.

Within this study we develop and validate a vacuum-pressure swing adsorption (VPSA) process model, capable of processing 33.8 kg/s of exhaust flow from a small-scale open-cycle gas turbine (OCGT). The Flexible response scenario is based on realistic load changes of OCGTs during a 5-hour period. To handle the size of flow the system is split into two identical two bed four step VPSA processes, both using Zeolite 13X as the adsorbent material. Included in the Flexible-VPSA operation is the start-up, ramping, and shutdown procedures. Flexible-VPSA showed minute deviations in CO₂ purity and recovery rate, and despite the specific energy demand increasing, results show no technical limitations to transient operation. The Flexible VPSA simulations are compared against the benchmark MEA solvent, with both technologies performing similarly when processing highly transient flue gas flowrate. Thus, VPSA is potentially an attractive alternative technology for Flexible-PCC.

Key Words:

CO₂ Adsorption, Dynamic Modelling, Post-combustion capture, Flexible Operation, VPSA

1. Introduction

With the realistic constraints (peak demand, system inertia and capacity reserve requirements) put on the energy system due to an increased renewable capacity [1], the use of Carbon Capture Utilisation and Storage (CCUS) for fossil-based power generation focusses on flexible operation [2]. System flexibility is crucial in counteracting fluctuations in supply and demand, balancing the network and maintaining security of supply [3]. However, the majority of Flexible-CCUS literature revolves around post-combustion capture (PCC) via amine absorption [4, 5, 6, 7, 8], as this is considered the benchmark CO₂ capture technology [9].

Physical adsorption is a promising alternative and is advantageous because no new bond is formed, enabling the possibility of pressure, temperature or vacuum swing regeneration at a much lower cost compared to amine-absorption. Several physical adsorbents have been extensively researched: zeolites, metal organic frameworks (MOFs), silicas, and carbonaceous material [10, 11, 12, 13, 14]. Ben-Mansour et al. [13] provide an extensive review of physical adsorption and discuss various regeneration processes. The study highlights fixed bed pressure swing adsorption (PSA) as an economic option because of its comparatively simple application, low energy demand, and low investment cost. For PCC vacuum swing adsorption (VSA) is attractive as the feed stream is at atmospheric (or slightly elevated) pressure. However, PSA/VSA systems suffer scalability issues and multiple beds are required to deliver the specifications for downstream compression equipment [9]. With the relatively low thermal regeneration temperatures, and the availability of waste heat from the

flue gas, Zanco et al. [15] suggests the more effective option would be temperature swing adsorption (TSA). In the literature high purities and recovery rates can be obtained by combining processes, i.e. pressure-temperature swing adsorption (PTSA), vacuum-pressure swing adsorption (VPSA) or vacuum-temperature swing adsorption (VTSA) [16, 17, 18, 19, 20].

Only a small number of studies have presented pilot-scale data for CO₂/N₂ adsorption systems, shown in Table 1. Wang et al. [18] achieved >90% capture and >95% purity with a two-unit VPSA system. The energy consumption is a sum of the power demand from blowers and pumps in the pilot facility. The on-site Watt metres measured a power demand of 2.44–2.65 MJ/kgCO₂. Lower than the 3.6–4.0 MJ/kgCO₂ for amine-absorption using the benchmark 30 wt.% monoethanolamine [9]. All the studies shown in Table 1 use fixed columns packed with an adsorbent material. Several studies have highlighted the potential for fluidised beds [21, 22, 23] and moving beds [24, 25, 26]. However, as the majority of CO₂ adsorption studies focus on fixed columns, this is the equipment investigated in this study. Table 2 is a compilation of simulation-based studies investigating CO₂/N₂ separation for PCC. As can be seen in Table 1 and Table 2, Zeolite 13X is the most studied post-combustion CO₂ capture adsorbent, due to its high CO₂/N₂ selectivity and high adsorption capacity at low CO₂ partial pressure [26]. As a result, Zeolite 13X is the adsorbent material investigated in this study.

Jiang et al. [27] compared VPSA, TSA and TVSA configurations at lab-scale and found VPSA to be more effective due to a lower energy consumption and higher CO₂ productivity. VPSA systems have been investigated from lab- to pilot-scale [28, 29, 30, 18, 27, 31] and usually comprise of two stages. The first stage captures >90% of the CO₂, and the second stage is used to increase the purity to >95% [32]. Luberti et al. [33] provides the design and simulation of a rapid VPSA process for PCC on a 10 MW_{th} biomass CHP plant. To handle the high flowrates and overcome scalability issues of adsorption process, in the 1st stage the flue gas stream is split into two parallel two-bed VPSA units, to capture >90% CO₂. The 2nd stage, another two-bed VPSA unit, is used for purification to obtain >95% CO₂.

Rui et al. [34] analysed CO₂ capture from flue gas using VPSA under unstable feed concentrations. The model-based study incorporated PID control strategies (closed-loop and open-loop feedback control) to adjust adsorption step duration for product quality control during variable feed concentration. To the authors knowledge, adsorption technologies for Flexible-CCUS have not previously been investigated.

In our previous study we highlighted the effects of transient open-cycle gas turbine (OCGT) operation on an amine-PCC system [7]. This study aims at investigating VPSA systems for quick response OCGTs (see Figure 1), under the same transient flue gas input to enable a comparison with the benchmark PCC technology. Within this Flexible-VPSA study:

- Development and validation of a single column process model for a VPSA system for CO₂ capture from flue gas using Zeolite13X.
- Designing and scaling the process model for a small-scale OCGT.
- Analysis of system flexibility, i.e., under transient flue gas conditions.
- Comparison against the benchmark amine-PCC system.

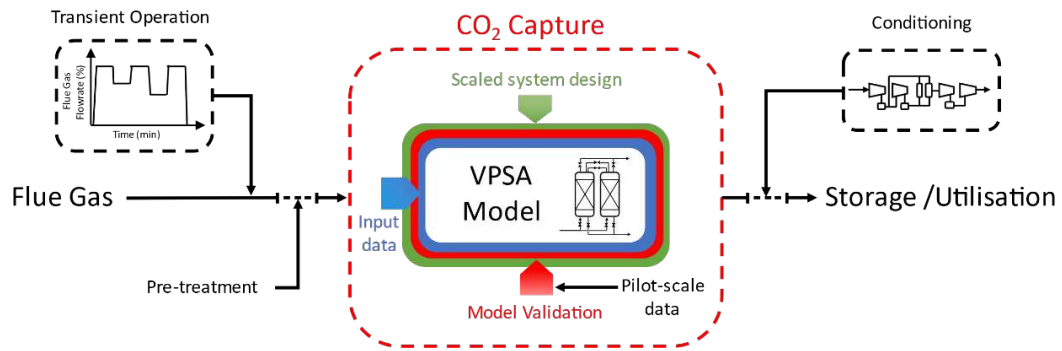


Figure 1: Study overview looking at VPSA CO₂ capture from transient flue gas, only the aspects within the red dotted border are investigated within this study.

Table 1: Pilot-scale studies of CO₂ adsorption from flue gas.

Source	System	Configuration	CO ₂ Feed (%)	Feed Flowrate	Recovery (%)	Purity (%)	Energy Consumption
[35]	PTSA	First stage is a PTSA unit, second stage is a PSA unit, using a Ca-X type zeolite. Feed comes from a dehumidifier unit attached to Yokosuka Thermal Power Station (coal and oil fired)	11.5	1000 N m ³ /h	90	99	560 kWh/t-CO ₂
[36]	TSA	Single bed TSA system with simulated flue gas. Lab scale comparison of Zeolite 13 X and 5A	10	20 N m ³ /min	13X=65 5A=83	13X=94 5A=98	8.8 MJ/kgCO ₂ 6.4 MJ/kgCO ₂
[37]	VSA	Three bed VSA system using Zeolite 13X, evaluating the performance of 6 and 9 step cycles	8-22	66-115 L/min	6 step = 60-80 9 step = 60-70	6 step = 82-83 9 step = 90-95	6 step = 4-8 kW/tCO ₂ 9 step = 6-10 kW/tCO ₂
[29]	VPSA	Three bed seven step VPSA unit removes CO ₂ from an existing coal-fired power station using Zeolite 13X APG	15	32.1-45.9 Nm ³ /h	79	85	2.37 MJ/ kgCO ₂
[18]	VPSA	The first unit is a 3 bed 8 step VPSA system using Zeolite 13X APG, the second unit is a 2 bed 6 step VPSA unit with activated carbon beads.	15-17	35.5-37.0 Nm ³ /h	90.20	95.60	2.44 MJ/kgCO ₂
[38]	VPSA	Single bed four step VSA with simulated flue gas feed for the basic system. Two bed four step VSA for the light product pressurisation (LPP) system. Both using Zeolite 13X	15	1000 SLPM	Basic: 95.9±1 LPP: 94.8±1	Basic: 86.4±5.6 LPP: 89.7±5.6	Basic: 472.2±36.7 kWh /tonne CO ₂ LPP: 475±36.7 kWh /tonne CO ₂
[39]	VPSA	Dual-Reflux VPSA in four fixed beds using two different activated carbon adsorbents within a single bed. Using a flue gas from a 460MW CFB boiler.	11.1	100 N m ³ /h	8 step: 42.7 9 step:30.2	8 step: 78.4 9 step: 89	8 step: 764 kWh/MgCO ₂ 9 step: 978 kWh/MgCO ₂

Table 2: Compilation of studies presenting mathematical models of various adsorption-based PCC systems.

Source	Model					Adsorbent(s) Investigated	System
	Mass Transfer	Flow	Isotherm	Heat Transfer	Pressure Drop		
[40]	LDF	Axially dispersed plug flow regime	Langmuir	Thermal equilibrium	Negligible Pressure Drop	Activated Carbon and Zeolite 13X	PSA
[41]	LDF	Ideal plug flow	Langmuir	Isothermal conditions	Negligible Pressure Drop	Zeolite 13X, 5A, 4A	DAC PSA
[42]	LDF	Axially dispersed plug flow regime	Langmuir	Non-equilibrium	Ergun	Potassium promoted hydrotalcite	PSA
[43]	LDF	Ideal plug flow	Extended Langmuir	Non-isothermal Adiabatic conditions	Negligible Pressure Drop	Zeolite 13X	PSA
[44]	LEM	Axially dispersed plug flow regime	Langmuir	Non-isothermal	Negligible Pressure Drop	Zeolite 13X	VSA
[45]	LDF	Axially dispersed plug flow regime	Dual-site Langmuir	Non-isothermal	Ergun	Zeolite 13X	Fractionated VPSA
[46]	LDF	Axially dispersed plug flow regime	Dual-site Langmuir	Thermal equilibrium	Ergun	Zeolite 13X	PSA
[16]	LDF	Axially dispersed	Toth	Local thermal equilibrium	Darcy	Zeolite 5A	TSA
[47]	LDF	Axially dispersed plug flow regime	Toth	Non-equilibrium	Ergun	Activated Carbon	PSA
[48]	LDF	Axially dispersed plug flow regime	Toth	Non-equilibrium	Ergun	Zeolite 13X	PSA
[49]	LDF	Axially dispersed plug flow regime	Toth	Thermal equilibrium	Ergun	Zeolite 13X	PSA
[50]	LEM	Axially dispersed plug flow regime	IAS theory using dual-site Langmuir-Freundlich isotherm	Isothermal conditions	Negligible Pressure Drop	Zeolite 13X, Mg-MOF-74, MOF-177, CubTTri, BeBtB and Co(BDP)	Breakthrough analysis
[51]	LEM	Axially dispersed plug flow regime	IAS theory using dual-site Langmuir-Freundlich isotherm	Isothermal conditions	Negligible Pressure Drop	Zeolites (MFI, JBW, AFX, NaX) and Metal Organic	PSA

						Frameworks (MgMOF-74, MOF-177, CuBTTri-mmen)	
[52]	LDF	Axially dispersed plug flow regime	Langmuir	Thermal equilibrium	-	Ceca 13X, Alcan AA320-AP and Alcan 650 PCA	TPSA
[30]	LDF	Axially dispersed	Virial	Non-equilibrium	Ergun	Activated Carbon Beads	VPSA
[17]	Bi-LDF	Axially dispersed plug flow regime	Modified multisite Langmuir	Thermal equilibrium	Ergun	Zeolite 13X-APG	VSA, TSA, VTSA
[53]	LDF	Axially dispersed plug flow regime	Multi-site Langmuir	Isothermal conditions	Ergun	activated carbon honeycomb monolith with zeolite 13X particle	ESA
[54]	LDF	Axially dispersed	Langmuir	Thermal equilibrium	Darcy	Zeolite 13X	VSA
[38]	LDF	Axially dispersed	Langmuir	Thermal equilibrium	Darcy	Zeolite 13X	VSA
[55]	LDF	Axially dispersed plug flow regime	Multi-site Langmuir	Thermal equilibrium	Ergun	Zeolite 5A and activated carbon	PSA
[56]	LDF	Axially dispersed	Toth	Non-equilibrium	Ergun	Zeolite 13X, Amine-functionalised Activated Carbon, Lewatit VP OC 1065	VSA
[57]	LDF	Axially dispersed	IAS theory using the Toth isotherm model	Thermal equilibrium	Ergun	Microporous Biochar	Ternary breakthrough analysis
[58]	LDF	Axially dispersed	IAS theory using Toth for CO ₂ and N ₂ , CMMS theory for H ₂ O	Thermal equilibrium	Ergun	Microporous Biochar	Ternary breakthrough analysis
[33]	LDF	Axially dispersed plug flow regime	Langmuir	Thermal equilibrium	Ergun	Zeolite 13X	Rapid VPSA
[59]	LDF	Axially dispersed	Dual-site Langmuir	Thermal equilibrium	Ergun	Zeolite 13X and Mg-MOF-74	VPSA
[60]	LDF	Axially dispersed	IAS theory using Toth for CO ₂ and N ₂ ,	Non-equilibrium	Darcy	Activated Carbon Honeycomb	TSA

CMMS theory for H ₂ O							
[61]	quasi-second order mass transfer	Axially dispersed plug flow regime	Toth	Isothermal conditions	Ergun	Activated Carbon and Zeolite 13X	VSA
[62]	LDF	Axially dispersed plug flow regime	Toth	Non-equilibrium	Ergun	Activated Carbon	PSA
[63]	LDF	2-D Mesh of structured quadrilateral elements	Dual-site Langmuir for CO ₂ & N ₂ , Toth for H ₂ O	Thermal equilibrium	Ergun	Zeolite 13X and Mg-MOF-74	2D and 3D breakthrough analysis
[64]		Axially dispersed	Extended Langmuir	Non-isothermal and adiabatic	Karman-Kozeny	Zeolite MS13X and activated carbon honeycomb	T/ESA
[65]	LDF	2-D Mesh of structured quadrilateral elements	Toth model for Zeolite 13X and the dual-site Langmuir model for Mg-MOF-74	Thermal equilibrium	Ergun	Zeolite 13X and Mg-MOF-74	Breakthrough analysis
[66]	LDF	Axially dispersed plug flow regime	Langmuir	Thermal equilibrium	Darcy	Silica Gel	VSA
[26]	LDF	Axially dispersed	Extended Virial Isotherm	Non-equilibrium	Ergun	Zeolite 13X	MBTSA
[67]	LDF	Axially dispersed plug flow regime	Langmuir-Freundlich	Thermal equilibrium	Ergun	Nanostructured Zeolite pellets	VPSA
[68]	LDF	Axially dispersed	dual-site Langmuir isotherm	Thermal equilibrium	Ergun	Zeolite 13X	VSA
[27]	LDF	Axially dispersed	Extended Langmuir	Non-equilibrium	Ergun	Zeolite 13X	VPSA, TSA, TVSA
[69]	LDF	Axially dispersed	Extended Langmuir	Non-equilibrium	Ergun	Composite adsorption material	VPSA
[70]	LDF	Axially dispersed	Sips	Thermal-equilibrium	Ergun	Zeolite-geopolymer sorbent	TSA-PSA

Note: CMMS is cooperative multimolecular sorption, DAC is direct air capture, IAS is ideal adsorption solution, LDF is linear driving force, and LEM is the local equilibrium model.

2. Adsorption modelling

Throughout the literature, models are usually one-dimensional and axially dispersed, assuming plug flow regime and negligible temperature, pressure, and concentration variation in the radial domain. Li et al. [62] provides a literature review of mathematical models of carbon capture by adsorption (CCA). Figure 2 illustrates the composition of physical adsorption model. The material balance over the fixed bed includes two important sub-models: adsorption kinetics (rate of mass transfer between gas and solid) and adsorption equilibrium (isotherm model). Table 2 highlights mathematical models in the literature specific for CO₂/N₂ adsorption. The Linear Driving Force (LDF) model is the most commonly used mechanism to describe mass transfer between the fluid and adsorbent, as it considers internal diffusional and external convection [71]. For CCA several isotherm models are commonly used in the literature, as highlighted in Table 2, usually an extension of the Langmuir isotherm model (dual-site Langmuir, Freundlich, Langmuir-Freundlich, BET, Sips, Toth) which account for multicomponent mixtures. The Ideal Adsorption Solution (IAS) theory has also been used, utilising pure component adsorption isotherms and enabling different isotherm models to be used for each component [72, 57, 58, 73].

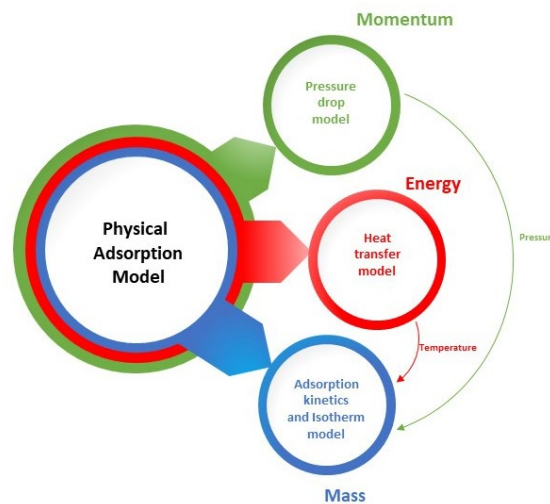


Figure 2: Physical adsorption model components

The energy balance describes the heat transfer between the gas and solid particles, as well as the column wall. Mathematic models in the literature vary in complexity of their consideration of heat transfer. Non-isothermal and non-adiabatic models that account for heat transfer between the gas and solid adsorbent, are classified as non-equilibrium models. Those that consider the energy flux between the gas and solid adsorbent as negligible are in thermal equilibrium [62]. The momentum balance considers the pressure drop in the column due to resistances in gas flow, modelled using the Blake-Kozeny equation (linear-laminar flow), the Ergun equation (non-linear turbulent flow) or the Darcy equation [71].

2.1. Process simulation

The process model used in this work (single column used for validation and multi-column used for large-scale dynamic analysis) is one dimensional and axially dispersed, with fluid flow described through the plug flow model, i.e., no radial variation in temperature, pressure, and concentration. The fluid phase material balance for component i through the packed bed is represented by the axial plug flow model [72]:

$$-D_L \frac{\partial^2 C_i}{\partial z^2} + \frac{\partial}{\partial z} (v_s C_i) + \frac{\partial C_i}{\partial t} + \left(\frac{1 - \varepsilon}{\varepsilon} \right) \frac{\partial \bar{q}_i}{\partial t} = 0, \quad \forall i \in \text{Components}, \quad \forall z \in (0, L) \quad 1$$

Where D_L is the axial dispersion coefficient [m^2/s], C_i is the fluid component concentration [kmol/m^3], z is the distance along the axial direction [m], L is the bed length [m], v_s is superficial fluid velocity [m/s], and ε is the bed voidage. Assuming axial flow is from the feed to the product side of the adsorption column, the boundary conditions are:

$$\frac{F^{in} w_i^{in}}{A_{bed}} = v_s C_i - \varepsilon D_L \frac{\partial C_i}{\partial z}, \quad \forall i \in \text{Components}, \quad z = 0 \quad 2$$

$$\varepsilon D_L \frac{\partial C_i}{\partial z} = 0, \quad \forall i \in \text{Components}, \quad z = L \quad 3$$

Where F^{in} is the molar flowrate entering the bed [kmol/s], w^{in} is the component molar fraction [kmol/kmol], and A_{bed} is the bed cross-sectional area [m^2]. The mass transfer rate is described through the LDF model [74]:

$$\frac{\partial \bar{q}_i}{\partial t} = k(q_{eq,i} - \bar{q}_i), \quad \forall i \in \text{Components} \quad 4$$

Where \bar{q}_i is the averaged absorption amount of species i [kmol/kg], $q_{eq,i}$ is the equilibrium absorption amount of species i [kmol/kg], and k is the mass transfer coefficient [$1/\text{s}$]. The dual-site Langmuir isotherm model was chosen as this is commonly used in the literature [68, 59, 75, 76, 38, 45]. The dual-site Langmuir isotherm is non-competitive as N_2 adsorption is low and does not significantly affect the adsorption amount of CO_2 [77], it calculates the equilibrium adsorption amount [38]:

$$q_{eq,i} = \frac{q_{sb,i} b_i C_i}{1 + b_i C_i} + \frac{q_{sd,i} d_i C_i}{1 + d_i C_i}, \quad \forall i \in \text{Components}, \quad \forall z \in (0, L) \quad 5$$

$$b_i = b_0 e^{(-\Delta H_{b,i}/RT)} \quad 6$$

$$d_i = d_0 e^{(-\Delta H_{d,i}/RT)} \quad 7$$

Where $q_{sb,i}$ is the maximum equilibrium adsorption amount of species i on site 1 [$\text{mol}/\text{kg}_{\text{ads}}$], $q_{sd,i}$ is the maximum equilibrium adsorption amount of species i on site 2 [$\text{mol}/\text{kg}_{\text{ads}}$], b_i and d_i are equilibrium absorption constants [m^3/kmol], b_0 and d_0 are pre-exponential constants [K], $\Delta H_{b,i}$ and $\Delta H_{d,i}$ are the heat of adsorptions at each site [J/mol], R is the universal gas constant [$\text{J}/\text{mol}/\text{K}$], and T is the temperature [K]. The equilibrium parameters b_i and d_i , in equations 6 and 7 respectively, follow the Van't Hoff equilibrium temperature dependence equation [72]. The momentum balance/pressure drop along each column is expressed through the Ergun equation [78]:

$$-\frac{\partial P}{\partial z} = 150 \frac{\mu_g (1 - \varepsilon)^2}{\varepsilon^3 d_p^2} v_s + 1.75 \frac{(1 - \varepsilon)}{\varepsilon^3 d_p} \rho_g v_s^2, \quad \forall z \in (0, L) \quad 8$$

Where P is the pressure [bar], μ_g is the dynamic viscosity [$\text{Pa}\cdot\text{s}$], d_p is the pellet particle diameter [m] and ρ_g is the fluid phase mass density [kg/m^3]. The model is non-isothermal and non-adiabatic, accounting for heat transfer between the bed and column wall using thermal inertia properties of the column (i.e. thermal conductivity and specific heat capacity), excluding heat transfer between the solid adsorbent and gas flow, i.e. thermal equilibrium. The fluid phase energy balance [79]:

$$\frac{\partial U_{bed}}{\partial t} = \varepsilon \frac{\partial}{\partial z} \left(\rho_T D_L \frac{\partial H_f}{\partial z} \right) - \frac{\partial (v_s \rho_T H_f)}{\partial z} + \frac{1}{1000} \frac{\partial}{\partial z} \left(\lambda_e \frac{\partial T}{\partial z} \right) + - \frac{4h_W}{d_b} \frac{1}{1000} (T_f - T_w), \quad \forall z \in (0, L) \quad 9$$

Where U_{bed} is the energy per unit volume of the bed [kJ/m^3], H_f is the bulk fluid phase mass specific enthalpy [kJ/kg], ρ_T is the total mass density of the fluid phase [kg/m^3], λ_e is the

effective axial thermal conductivity [W/m²/K], h_w heat transfer coefficient between the gas and column wall [W/m²/K], d_b is the bed diameter [m], T_f is the fluid phase temperature [K] and T_w is the bed wall temperature [K]. Assuming axial flow is from the feed to the product side of the adsorption column, the boundary conditions are:

$$\frac{F^{in}H_f^{in}}{A_{bed}} = v_s\rho_T H_f - \varepsilon D_L \rho_T \frac{\partial H_f}{\partial z} - \frac{1}{1000} \lambda_e \frac{\partial T}{\partial z}, \quad \forall z \in (0, L) \quad 10$$

$$\frac{\partial T}{\partial z} = 0, \quad z = L \quad 11$$

Where F^{in} is the mass flowrate entering the bed [kg/s], and H_f^{in} is the mass specific enthalpy of the entering fluid on the feed side of the adsorption bed [kJ/kg]. Taking into consideration the thermal inertia and conduction within the bed wall, the energy balance for the metal wall is [79]:

$$\rho_w C_{p,w} A_w \frac{\partial T_w}{\partial t} = A_w \lambda_w \frac{\partial^2 T_w}{\partial z^2} + h_w \frac{4}{d_b} (T_f - T_w) + h_{w,a} \frac{4(d_b + 2l_w)}{d_b^2} (T_w - T_a), \quad 12$$

$$\forall z \in (0, L)$$

Where ρ_w is the bed wall mass density [kg/m³], $C_{p,w}$ is the bed wall specific heat capacity [kJ/kg/K], A_w is the bed wall cross-sectional area [m²], T_a is the ambient temperature [K], λ_w is the wall material thermal conductivity [W/m/K], $h_{w,a}$ heat transfer coefficient between the column wall and external environment [W/m²/K] and l_w is the wall thickness [m].

Process simulation tools such as Aspen Adsorption [80, 57, 58, 64] and gPROMS ProcessBuilder [81, 82], have been used to effectively simulate, analyse and optimise CO₂/N₂ adsorption. In this study, the model is developed in gPROMS ProcessBuilder utilising the physical properties package Multiflash™. Adsorption processes are inherently dynamic; therefore, the process models must include valves to control the flow in and out of the adsorption bed. For this study each valve is considered adiabatic, where the mass flowrate (F in kg/s) and pressure drop (Δp) are related through:

$$F = C_v x \Delta p \quad 13$$

As the model is pressure driven, the flow coefficient (C_v in kg/s/kPa) for each valve is calculated to give the required pressure drop during each stage of the simulation. The timings of the valve stem position (x) are defined using a scheduling unit. This controls which valves are open at any given time, allowing for step management and cyclic behaviour.

The energy demand comes from the compressors needed to elevate the bed pressure and the vacuum pumps needed to depressurise the bed to desorb the CO₂. For real gas compression and evacuation power consumption [W], Nikolaidis et al. [32] used:

$$Power_{comp} = F_{in} T_{in} R \frac{1}{n_c} \left(\frac{\gamma}{\gamma - 1} \right) \left[\left(\frac{P_{out}}{P_{atm}} \right)^{\frac{\gamma-1}{\gamma}} - 1 \right] \quad 14$$

$$Power_{vac} = F_{in} T_{in} R \frac{1}{n_v} \left(\frac{\gamma}{\gamma - 1} \right) \left[\left(\frac{P_{atm}}{P_{vac}} \right)^{\frac{\gamma-1}{\gamma}} - 1 \right] \quad 15$$

Where F_{in} is the inlet flowrate [mol/s], T_{in} is the inlet temperature [K], R is the universal gas constant [J/mol/K], P_{out} is the compressors discharge pressure [Pa], P_{atm} is atmospheric pressure [Pa], and P_{vac} is the vacuum pressure draw [Pa]. The ratio of specific heat capacities (γ), also known as the adiabatic constant, is assumed to be 1.4. Within the literature the isentropic efficiency (n_c and n_v) for N₂/CO₂ systems is 0.72 [54]. To compare and evaluate

CCA technologies two important metrics are used: recovery rate (difference between the mass of CO₂ entering and exiting the process) and the purity (end CO₂ composition). Alongside these metrics are two important performance indicators: the specific energy demand (E_T) calculated using equation 16 and the Productivity (P_r) calculated using equation 17.

$$E_T[\text{kWh}/\text{tCO}_2] = \frac{\text{Total Energy Demand [kWh]}}{\text{Mass of CO}_2 \text{ Captured [ton]}} \quad 16$$

$$P_r[\text{mol}/\text{m}^3 \text{ Adsorbent}/\text{s}] = \frac{\text{CO}_2 \text{ Captured [mol]}}{\text{Volume of Adsorbent [m}^3] \times \text{total cycle time [s]}} \quad 17$$

2.2. Single column validation

In order to evaluate the validity of the process model, a single column is validated against pilot scale data from Krishnamurthy et al. [38]. The pilot study investigated CO₂/N₂ VPSA on the commercial sorbent Zeolite 13X. The feed contains 15 mol.% CO₂ and 85 mol.% N₂ at 1000 SLPM. The column specifications are shown in Table 3.

Table 3: Single column specifications [38]

Parameter	Value
Column internal diameter (m)	0.3
Column packing height (m)	0.867
Mass of adsorbent (kg)	41
Particle size (mm)	1.6-2.6
Bed Voidage	0.428

Run 1 in [38] is chosen for its high CO₂ purity (94.7±1.05%) and high CO₂ recovery (85.4±4.52%). The four-step Skarstrom cycle VPSA system includes: (i) pressurisation to P_H using the feed gas, (ii) adsorption at high pressure with product valve open, (iii) forward blowdown to depressurise the column to P_i , and (iv) reverse evacuation to recover CO₂ at P_L (see Figure 3). The step duration and pressure specifications are shown in Table 4.

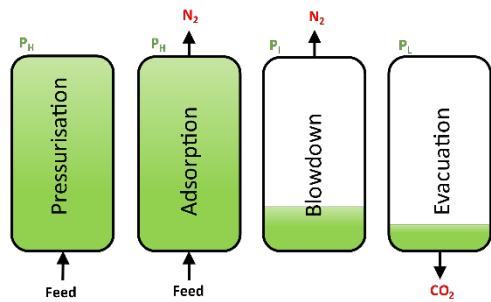


Figure 3: Four-step VPSA cycle

Table 4: Step duration and pressure specifications [38]

Step	Time (s)	Pressure (level)	Pressure (bar)
Pressurisation	20	P_H	1.5
Adsorption	60	P_H	1.5
Blowdown	150	P_i	0.07
Evacuation	310	P_L	0.025

The gravimetric equilibrium isotherm data is shown in Table 5. Zeolite 13X is a Faujssite type zeolite, and [83] determined the thermal conductivity to be 2.28 (W/m/K). Mass transfer coefficients for N₂ and CO₂ on Zeolite 13X are taken as 2.733 s⁻¹ and 0.065 s⁻¹ respectively, and the axially dispersion co-efficient is 0.23 m²/s [48]. Table 6 highlights the other key process parameters necessary to calculate the mass and heat transfer along the adsorption column.

Table 5: Dual-site Langmuir isotherm parameters [76]

Parameter	Symbol	N ₂	CO ₂
Equilibrium adsorption amount for site 1 (mol/kg)	$q_{sb,i}$	5.84	3.09
Equilibrium adsorption amount for site 2 (mol/kg)	$q_{sd,i}$	0	2.54
Pre-exponential constant for site 1 (m ³ /mol)	b_0	2.50x10-6	8.65x10-7
Pre-exponential constant for site 2 (m ³ /mol)	d_0	0	2.63x10-8
Heat of adsorption for site 1 (J/mol)	$-\Delta H_{b,i}$	-1580000	-36641.21
Heat of adsorption for site 2 (J/mol)	$-\Delta H_{d,i}$	0	-35690.66

Table 6: Mass and heat transfer parameters [76]

Parameter	Symbol	Value
Pellet bulk density (kg/m ³)	-	1130
Pellet void fraction (m ³ /m ³)	-	0.35
Specific heat capacity of fluid (J/mol/K)	-	30.7
Specific heat capacity of adsorbent (J/kg/K)	-	1070
Wall thickness (mm)	l_w	17.5
Wall density (kg/m ³)	ρ_w	7800
Heat transfer coefficient between bed and wall (W/m ² /K)	h_w	8.6
Heat transfer coefficient between wall and environment (W/m ² /K)	$h_{w,a}$	2.5
Specific heat capacity of the column wall (J/kg/K)	$C_{p,w}$	502
Thermal conductivity of the column wall (W/m/K)	λ_w	16
Universal gas constant (m ³ Pa/mol/K)	R	8.314

Figure 4 compares the simulations adsorption bed pressure and temperature against the pilot results (Run 1) from Krishnamurthy et al. [38]. The simulation exhibits an almost identical pressure increase during the pressurisation step (t=0 to 20s), using 1000 SLPM flowrate containing 85 mol.% N₂ and 15 mol.% CO₂. During the adsorption (t=20 to 80s) and blowdown (t=80 to 230s) steps, similar pressure profiles are seen between the simulation and pilot results, the pressure drop rate is slightly greater in the pilot test, however, both results produce a bed pressure of 0.07 bar once the blowdown step is complete. Small variations between the results could be due to data extraction errors, as the pilot experiments performance indicators are for the 300th cycle; however, individual error bars are not available for the dynamic pilot results. Included in Figure 4 is the feed and product header temperatures, within Krishnamurthy et al [38] the temperature profile is not given for a single cycle; however, they have been included to compare with the full-scale VPSA simulations.

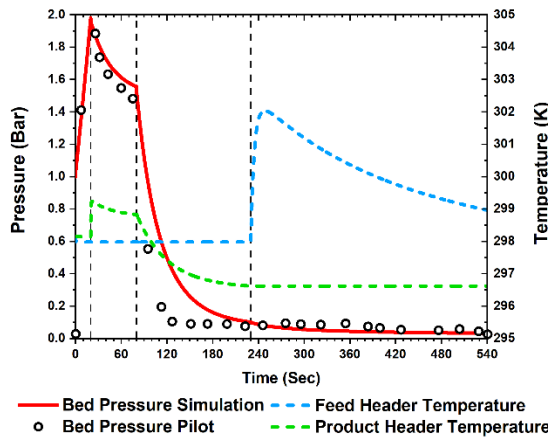


Figure 4: Adsorption bed pressure and temperature profiles

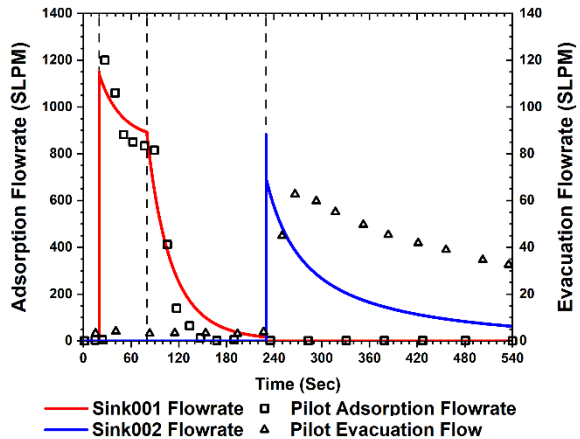


Figure 5: Flowrate profiles for the adsorption/blowdown and evacuation steps

Two important key performance indicators for CO₂ capture technologies are the recovery rate and CO₂ purity. The recovery rate is defined as the percentage difference between the CO₂ flow entering and exiting the system [5]; in this case, the flowrates (Figure 5) and compositions (Figure 6) are used to calculate the amount of CO₂ captured (84.30%), which lies within the bounds (85.4 ± 4.52%) specified in Krishnamurthy et al. [38]. The purity of the end product during the evacuation step in the simulation is 89.56% (mol/mol), 5.14% points lower than the pilot facility (94.7 ± 1.05%). The simulation's predicted purity is not only a function of bed dynamics, in particular the adsorption rate calculated through the isotherm model, it is also affected by the valve stem position and the flowrate through the evacuation valve. As multiple factors affect the purity it is difficult to align all the input parameters simultaneously in order to improve the results. Also, as the model is pressure driven it is vital the pressure profile is as accurate as possible. Therefore, the evacuation valve and flowrate cannot be further adjusted (to increase the purity), as this would prevent the pressure level from decreasing to the 0.025 bar goal.

Figure 5 compares the flowrates exiting the adsorption column, and the composition profiles during the adsorption, blowdown and evacuation steps are shown in Figure 6. The adsorption flowrate exiting the top of the column (mainly composed of N₂) exhibits a similar profile to the bed pressure. The pilot evacuation profile shows a smaller decrease in flowrate compared to the simulation, due to a rapid pressure drop from the intermediate pressure (0.07 bar) to the low pressure (0.025 bar) used to desorb the captured CO₂.

The pilot results productivity was 1.4 tCO₂/m³/day, the simulation result is 15% lower at 1.18 tCO₂/m³/day. The overall power demand (for the flue gas blower and vacuum pumps) is 934.05 kWh/tCO₂, 74.26% higher than the pilot run power demand (510.5 ± 25.5 kWh/tCO₂). If the efficiency of the pumps is set at 100% the overall power demand is 672.52 kWh/tCO₂ and 25.47% higher than the pilot test. The accuracy is further improved if the ratio of specific heat capacities is closer to that of CO₂ (1.28 [84]), specifically during the evacuation step due to the higher CO₂ concentration, the power demand is 611.79 kWh/tCO₂ and 14.14% higher than the upper bound of the pilot test. Highlighting the need for variable specific heat capacity ratios when determining overall pump power requirement.

During the adsorption step the average adsorption amount for CO₂ and N₂ is 3.569 mmol/g and 0.464 mmol/g, respectively, comparable to sources in the literature looking specifically at Zeolite 13X [85, 86]. Even though the CO₂ concentration leaving the column is lower than expected, the accuracy of the capture rate and the adsorption capacity shows the fidelity of the overall process model.

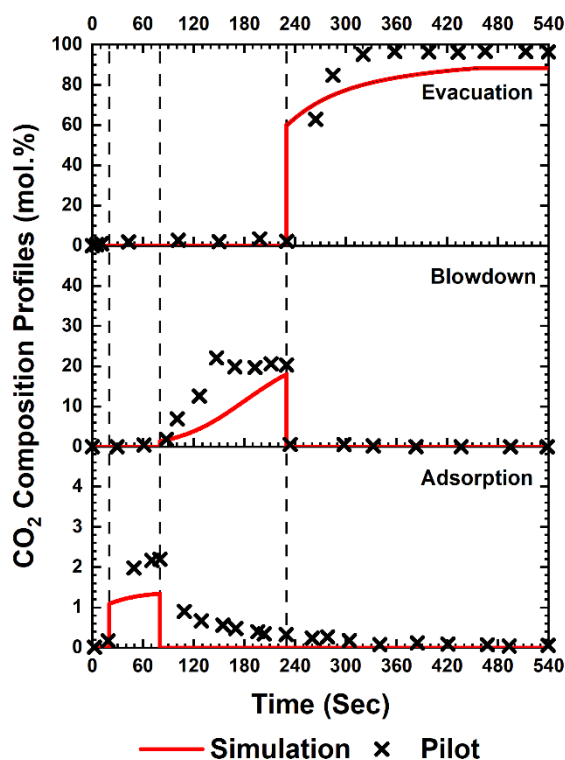


Figure 6: Comparison of CO₂ composition profiles during the Adsorption, Blowdown and Evacuation steps

3. Large-scale system design

Large-scale CCA systems require multiple beds and multiple stages for continuous separation of the desired component. A limitation for large-scale vacuum-adsorption systems is the ability of industrial machinery to deliver the vacuum pressure necessary for desorption [87], i.e., the vacuum pressures achieved in lab/pilot-scale systems may not be achievable at large-scale [33]. Therefore, the large-scale VPSA system used in this study is based on Luberti et al. [33], where the 1st stage is split into two identical two-bed VPSA units to handle the large inlet flowrate. The process topology for this studies adsorption CO₂ capture system is shown in Figure 7. In Shen et al. [30], Wang et al. [17], and Luberti et al. [33] a secondary stage in series is used to ensure the end CO₂ purity is above 95%; however, in this study the purity is already >95% meaning the second stage is not required. In Figure 7, the scheduling unit dictates the valve stem position (x in Equation 13), i.e., how open the valve is. The valve flow coefficient (C_V in Equation 13) is specified to give an adequate pressure drop over each valve.

Assuming the power generation source is a 10 MWe open-cycle gas turbine (OCGT), 33.8 kg/s of flue gas is produced with 4.27 vol.% CO₂ (6.78 wt.% or 4.42 mol.%) [7]. Each parallel train processes 16.9 kg/s of flue gas (FGS1 and FGS2 on Figure 7). The process simulation uses the same isotherm (Table 5) and mass and heat transfer parameters (Table 6) as the validation study. In order to process the large flue gas flowrate, the column properties are scaled such that each adsorption column has 11.15 m packing height, 4.80 m bed diameter, 2 mm particle size, and the bed voidage is 0.428. These column properties ensured the scaled design has the same adsorbent surface area to volume ratio (3000 m²/m³) as the pilot study.

The scheduling unit dictates the valve stem positions, within Figure 7 the scheduling table highlights each valves stem position during each of the two bed four step operations, i.e.,

Press – Bb denotes the first bed is in pressurisation mode and the second bed is in blowdown mode.

Also included in Figure 7 is the scheduling table that highlights each valve’s flow co-efficient used in Equation 13 to calculate each valve’s pressure drop. Similarly to the pilot study used for the validation simulation, the scaled-up system design has four process steps with three main target pressures. The step timings and corresponding pressure levels are shown in Table 7. The Finite Volume Method is selected as the numerical discretisation scheme. There are 10 discretisation points per layer (one layer in total) with uniform distribution across the layer. The structural analysis for the Baseline process model showed the total number of model equations is 2,781, with 3,191 process variables (2, 781 unknown and 410 inputs) and 18,327 parameters. The 5-hour operation was simulated in 879 seconds (CPU).

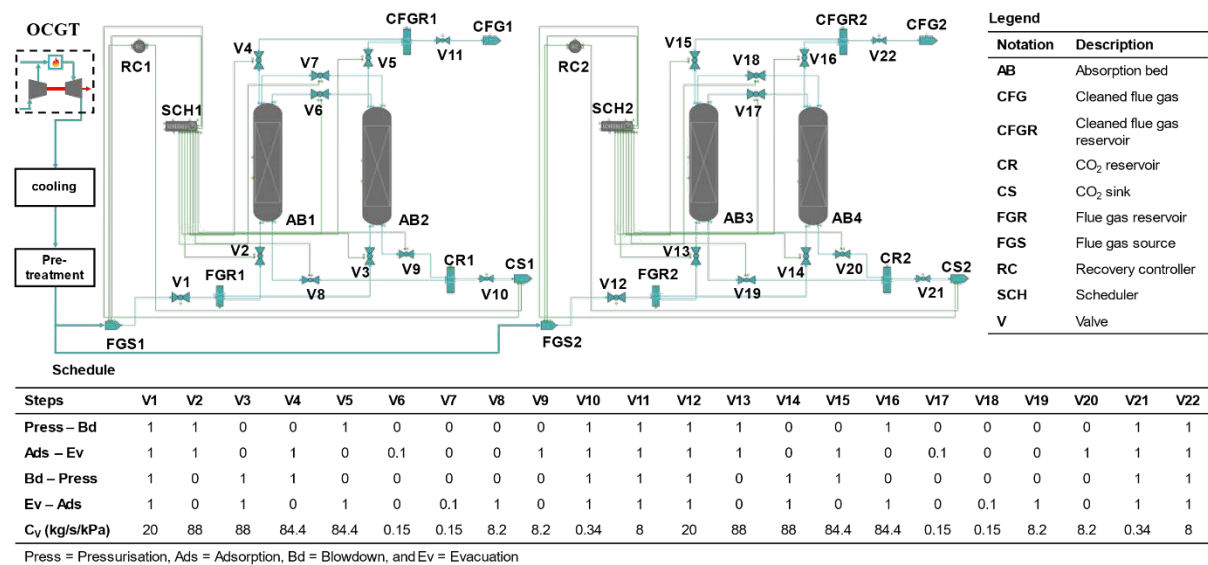


Figure 7: Large-scale VPSA model topology, where the flue gas stream is split into two parallel streams, included in the figure is valve scheduling and each valve’s flow co-efficient

Table 7: Step duration and pressure specification for the large-scale VPSA system

Step	Time (s)	Pressure (level)	Pressure (bar)
Pressurisation	80	P_H	1.5
Adsorption	220	P_H	1.5
Blowdown	80	P_i	0.07
Evacuation	220	P_L	0.025

3.1. Flexible operation

The flexible operation of OCGT’s is based on our previous work [7], which analysed data from the Balancing Mechanism Report Services (BMRS) and industrial gas turbine suppliers, to show the typical operation of OCGT’s in the UK. The results showed the highly transient nature of OCGT power generation, with average operating times around 5 hours, while ramping to different power outputs within the same operating window. Figure 8 is an example OCGT operation for a 10 MWe plant, using a Siemens SGT-400 gas turbine. The OCGT can ramp to different power outputs at a rate of 10% baseload (MWe) per minute, and the start-up and shutdown rates are 12.5% baseload (MWe) per minute [88]. Therefore, the start-up/shutdown time is 8 minutes, the ramp time to 70% load is 3 minutes, and the ramp time to 50% load is 5 minutes. The balancing period in the UK is 30 minutes [89]. Each new load is maintained until the end of the 2nd balancing period before ramping or shutdown is performed, i.e., at 60

minutes the flue gas is ramped to 70% load in 3 minutes, 57 minutes later the flue gas is then ramped back up to full load. As the total cycle operating time is 600 seconds, 31 cycles are required for a 5-hour operation plus the 8-minute shutdown time.

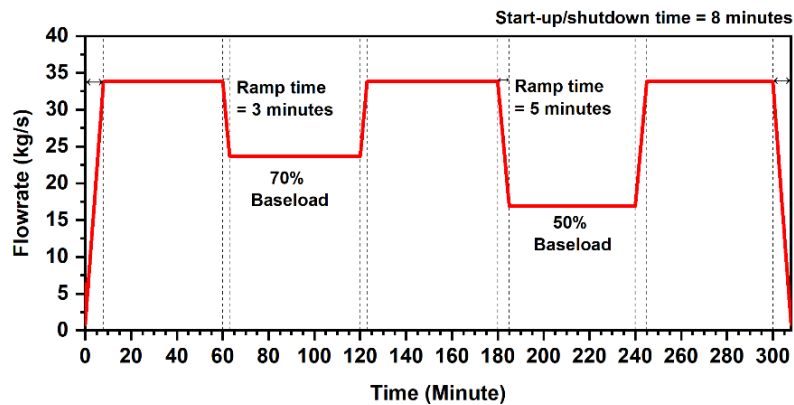


Figure 8: Transient OCGT operation from Wilkes et al. [7]

4. Results

The results section shows the large-scale VPSA operation, during the Baseline (no transient behaviour) and Flexible (transient flue gas input) operating scenarios. The scaled system analysis focusses on the effects transient operation has on the processes key performance indicators (KPI), this study does not look at the optimisation of the scaled VPSA design. Figure 9 shows the flowrates through flue gas source (FGS), cleaned flue gas (CFG) and CO₂ sink (CS) streams, these are highlighted in Figure 7, for both the Baseline and Flexible operating scenario. Each parallel train handles 16.9 kg/s exhaust flow. The flowrate profiles for the CFG

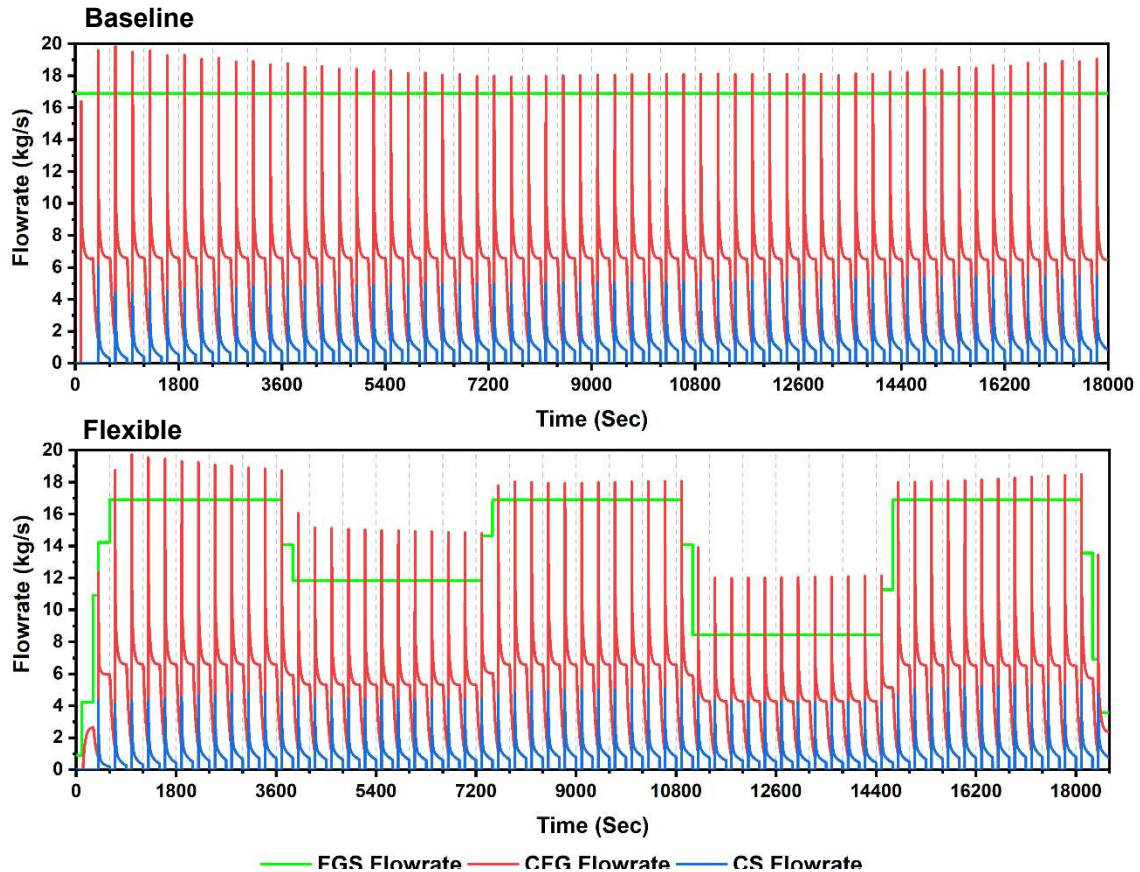


Figure 9: Flowrate profiles for the flue gas source (FGS), cleaned flue gas (CFG) and CO₂ sink (CS) flowrates in each of the parallel VPSA units, during the Baseline and Flexible scenarios.

and CS flows are similar to the pilot study, where there is a sharp initial spike in flow that levels out during each operating step. When one bed is adsorbing the other bed is evacuating, allowing for continuous CO₂ capture. In the Baseline scenario, the system is fed with a constant flue gas flowrate for the entire 5-hour operating period. In the Flexible scenario, the system is fed with a transient flue gas supply. Included in this scenario is the start-up procedure, ramping to 70% and 50% loads, and the shutdown procedure. Due to computational limitations the simulation cannot operate at 0 kg/s feed, therefore, the system starts at 0.9 kg/s and ramps to 16.9 kg/s during start-up in 8 minutes. Similarly, for the shutdown operation, the feed is at 16.9 kg/s and ramps to 0.9 kg/s in 8 minutes.

Figure 10 presents the adsorption bed (AB1 and AB2 on Figure 7) pressure profiles during the Baseline and Flexible operating scenarios. The profiles are comparable to the pilot-scale study, see Figure 4. As the models are pressure driven it is important the pressure profiles are accurate. There is a sharp initial increase in pressure (pressurisation step), which then equalises during the adsorption step when the product valves are opened. At t=300 seconds in each adsorption cycle the blowdown valves (V4, V5, V15, and V16 in Figure 7) are opened and the bed pressure drops to 0.07 bar, at t=500 seconds in each cycle the blowdown valve is closed, and the evacuation valve is opened (V8, V9, V19, and V20 in Figure 7), further reducing the bed pressure to 0.025 bar and the CO₂ is desorbed from the bed. In the Flexible scenario the valve coefficient for V6, V7, V17, and V18 is changed from 0.15 to 0.01 kg/s/kPa to ensure an adequate exchange between the adsorption columns during the low load (50%) operation, without this alteration the full operation could not be completed.

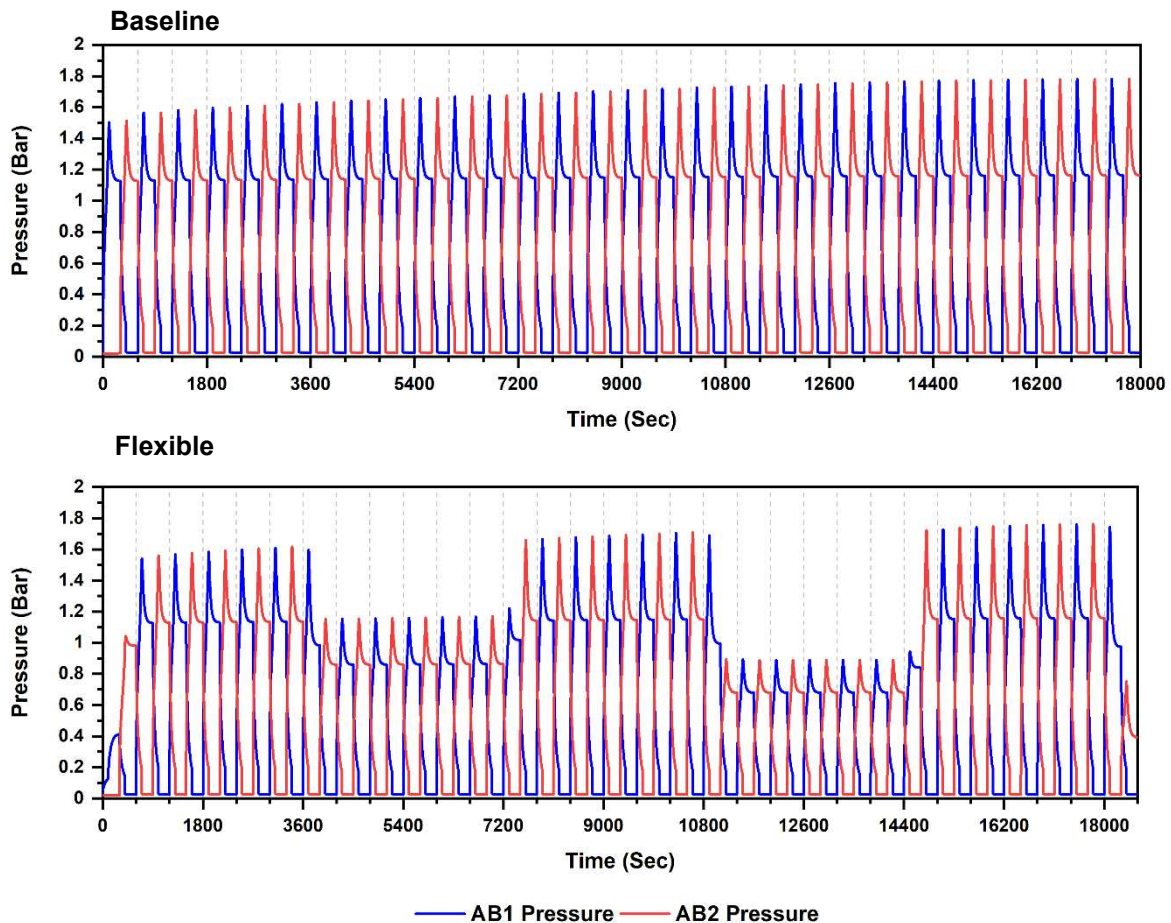


Figure 10: Pressure profiles for adsorption bed 1 (AB1) and adsorption bed 2 (AB2) in each of the parallel VPSA units, during the Baseline and Flexible scenarios.

Over the course of operation there is a gradual increase in the initial pressure spike during the pressurisation step. During the pressurisation step in the first cycle (Baseline operation), the bed pressure (AB1) is elevated to 1.47 bar. Whereas, in the final cycle the bed pressure is elevated to 1.76 bar. This is a result of the increasing column temperature, especially at the bottom of column, shown in Figure 11. The increase in step pressure is also shown in the pilot scale results in Krishnamurthy et al. [38]. At the end of each adsorption cycle the columns equalise to 0.025 bar ready for the next adsorption cycle.

During the Flexible scenario, less flow is going through the system during the start-up, ramping and shutdown periods, therefore the adsorption columns cannot pressurise to 1.5 bar. Hence, the bed pressure follows the same profile as the input (FGS) flowrate. The valve flow coefficients are set input parameters that are used alongside the flowrate to calculate the pressure drop across the valve. Incorporating control systems to modify the flow coefficients during low load operation could promote acceptable pressure increases in the column. However, this would involve developing Model Predictive Control (MPC) strategies, which is beyond the scope of this study. Several studies have investigated optimisation and control strategies for Flexible-MEA systems [90, 91] and flexible gas turbines [92], which have provided valuable information on the potential options available to reduce the specific energy of the processes. Therefore, future work should focus on optimising and developing strategies specifically for Flexible-VPSA systems.

The temperature profile in the adsorption column changes throughout each of the 5-hour operations, Figure 11 and Figure 12 show this transient behaviour for the Baseline and Flexible scenarios, respectively. As the adsorption process releases the heat of adsorption when CO₂ is adsorbed, the temperature increases in the column. In the beginning of the operation, the temperature bulge is located at the bottom of the column; indicating most of the adsorption is taking place within the first 2 m of packing. The temperature bulge transitions towards the centre of the column (between 4 and 8 m) as the operation progresses.

There is a small difference in the rate at which the temperature increases in the first operating cycle, between the Baseline and Flexible operating scenarios. This is due to the start-up procedure lasting 480 seconds; therefore, it is at full-load by the end of the first operating cycle. As less CO₂ is entering the system, less CO₂ is adsorbed onto the zeolite surface and less heat is emitted into the column. There is also a sudden drop in temperature in the Flexible scenario in the bottom of the column between t=10000 to t=14000 seconds, due to decreasing the inlet flowrate to 50% load. Overall, there is a small difference in the temperature increase (<1.5K) at the temperature bulge, highlighting that less CO₂ is being adsorbed in the column.

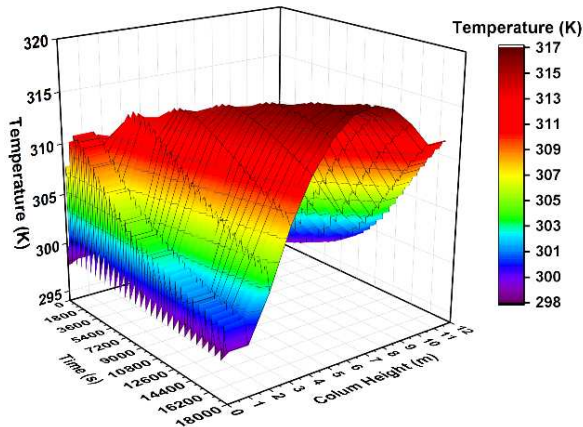


Figure 11: Adsorption bed (AB1) 3-D temperature profile for the Baseline scenario

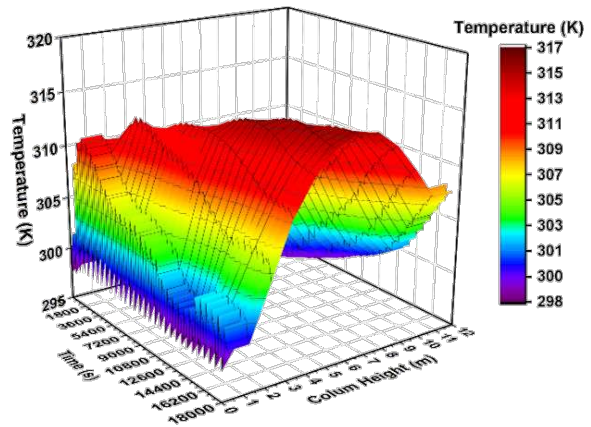


Figure 12: Adsorption bed (AB1) 3-D temperature profile for the Flexible scenario

The dissimilarities between scaled Baseline and pilot KPI results (see Table 8) are related to the scaled system design. The adsorption cycle steps and timings were modified for continuous operation. The timings used in this study enabled the large adsorption columns to pressurise and minimise the inlet FGS velocity, preventing issues such as flow reversal, increased pressure drop, and loss in separation efficiency. Krishnamurthy et al. [38] showed the effect of altering the step duration and blowdown pressure on the recovery rate and purity. Longer adsorption step times increases the purity and decreases the recovery rate, due to the CO₂ front moving further into the column. This is observed in the 3-D temperature profile, see Figure 11. Highlighting, the adsorption timing within this study needs to be longer, however, this is limited by the pressure increase in the bed. Parameter scaling is another important issue: without demonstration scale results it is difficult to ascertain the fidelity of the scaled models [93].

The continuous operation and modified step timings also explain the increase in energy demand. The scaled Baseline results are more than double the pilot scale results, due to constant flue gas processing (increased flow through the FGS blower, CFG vacuum pump and CS vacuum pump), thus requiring more energy. Despite the issues with the scaled design, it is important to see the ramifications transient operation has on the performance of VPSA CO₂ capture.

An issue observed in industrial adsorption processes is complex valve operation [94, 95]. Within this study the valves are operated based on flow co-efficients dictating the pressure drop. Each valve operates simultaneously with the active status constantly changing. The complex partnership between the valves within the system make scaling up the VPSA processes difficult and affects the KPI's. When scaling the VPSA design it is important to consider the adsorption steps and timings, column geometry, process design and valve operation.

The differences between the Baseline and Flexible operating scenarios KPI's can be explained by the non-identical temperature profiles in Figure 11 and Figure 12. In this study, for VPSA there is a small drop in recovery rate (0.03% points). This is lower than all of the MEA CO₂ capture flexible response scenarios (0.73 -1.20% points lower than the steady-state results at 92.48%) shown in Wilkes et al. [7]. There is also a small drop in purity for flexible VPSA system (1.20 % points). Although Wilkes et al. [7] do not specify the end purity, their report states there is also a drop in purity during flexible operation. This will affect downstream compression equipment, so too will the transient operation [96].

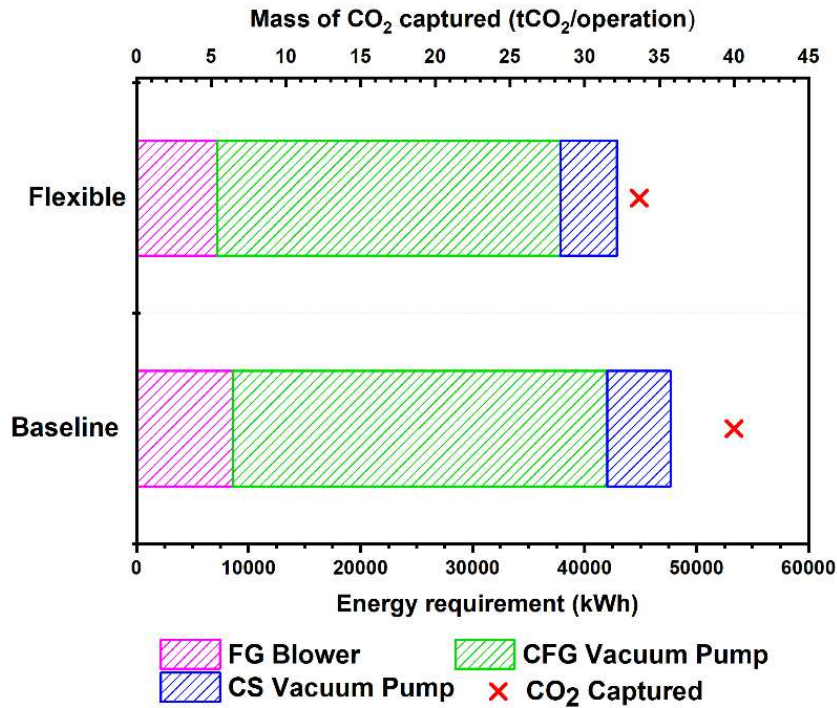


Figure 13: Pump energy requirement and mass of CO₂ captured during the 5-hour operation for the Baseline and Flexible scenarios

Table 8: Key performance indicators (KPI) for the Baseline and Flexible VPSA simulations

KPI	Pilot [38]	Scaled-Baseline	Deviation (%)	Scaled-Flexible	Deviation (%)
Recovery rate (%)	85.40 ± 4.52	97.07	+13.66	97.03	-0.04
Purity (%)	94.7 ± 1.05	80.74	-14.74	79.54	-1.49
Energy demand (kWh/tCO ₂)	510.5 ± 25.5	1191.72	+133.67	1274.04	+6.91
Productivity (tCO ₂ /m ³ ads/day)	1.40 ± 0.07	1.28	-9.02	1.07	-15.85

Figure 13 highlights the FGS blower, CFG vacuum pump and CS vacuum pump power requirements for the Baseline and Flexible scenarios. The majority of the energy demand comes from the CFG vacuum pump used in the adsorption and reverse blowdown steps. This has the largest energy demand because it handles a large amount of flow compared to the CS vacuum pump, and the energy drop is the greatest (approximately 1.5 to 0.07 bar). Despite the Flexible scenario consuming less energy, it has a higher specific energy demand (see Table 8) due to the smaller quantity of CO₂ captured (see Figure 13). The quantity of CO₂ captured during the Baseline (39.99 tCO₂/operation) and Flexible (33.65 tCO₂/operation) operations are comparable to the results in Wilkes et al. [7] (31-37 tCO₂/operation), the slightly higher amount is due to the higher capture rate (92.48 % MEA steady-state results).

Several studies have highlighted the use of MPC strategies for solvent-based [97] and adsorption [90, 98] capture technologies. It has been observed that MPC can improve the capture performance and aid in reducing deviations caused by flexible operation [99, 100]. Neither the VPSA system analysed in this study, nor the MEA system analysed in Wilkes et al. [7] include MPC, as these studies focus on the technical feasibility of transient operation. Future work should incorporate MPC strategies to optimise the performance of dispatchable OCGT-PCC, specifically at low-load operation.

5. Conclusion

As the need for flexible CO₂ captured increases, alternate technologies to amine-based post-combustion capture (PCC) need to be investigated specially for flexible response. Within this study a vacuum-pressure swing adsorption (VPSA) PCC system has been modelled, validated, and scaled up to process flow from a small-scale 10 MWe open-cycle gas turbine (OCGT). The purpose of the study is to assess the ability of a VPSA CO₂ Capture system to handle highly transient flue gas operation, to highlight the technologies applicability in future energy systems that require fast-response dispatchable generators to combat intermittent renewables. The validation study showed the process model accurately simulated the adsorption of CO₂ on the Zeolite 13X sorbent. The amount of CO₂ captured (84.30%) lies within the error bounds ($85.4 \pm 4.52\%$) specified in that pilot-scale study [38].

The full-scale simulations showed the limitations with process scaling for VPSA systems are related to adsorption step timings, column sizing, and valve operation. The aim of the paper is not to provide scaled system design, but to investigate the effects of flexible operation on VPSA CO₂ capture. To the authors knowledge, no studies have investigated Flexible-VPSA, i.e., the effects of transient flue gas production on the operation and performance of VPSA CO₂ capture. Included in the Flexible-VPSA operation is the start-up, ramping, and shutdown procedures. The simulations showed minimal deviation in CO₂ purity (-1.49%) and recovery rate (-0.04%), between the Baseline and Flexible operating scenarios. The decrease in purity and recovery rate during flexible operation is lower than experienced in MEA based Flexible-PCC.

As demonstrated in our earlier study, the energy demand for MEA based Flexible-PCC varies between each of the flexible operating scenarios investigated [7]. When the steam supply to the reboiler (for solvent regeneration) is constant, the energy demand for the process is 4.69 GJ/tCO₂ (1302.78 kWh/tCO₂). When the steam supply is altered in accordance with flue gas changes, the energy demand decreases to 3.95 GJ/tCO₂ (1097.22 kWh/tCO₂). For Flexible-VPSA the energy demand (for the blower and vacuum pumps) is 4.59 GJ/tCO₂ and 16.11% higher than the best Flexible-MEA case. This is due to the fact that no specific energy minimisation scenario was considered in this study. Therefore, future work should focus on developing dedicated operating scenarios and control strategies for a VPSA system, in order to minimise the negative energy effects of transient operation and scale-up. A possible solution is manipulating the scheduling (step time and pressure level) to deliver lower specific energy demands for the pumps during low load operation.

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