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# 1 Modelling, Simulation and Analysis of Intensified

# 2 Regenerator for Solvent based Carbon Capture using

**3 Rotating Packed Bed Technology** 

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# 10 ABSTRACT

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6 7

Intensified regenerator/stripper using rotating packed bed (RPB) for regeneration of 11 rich-MEA solvent in post-combustion CO<sub>2</sub> capture with chemical absorption process 12 was studied through modelling and simulation in this paper. This is the first 13 systematic study of RPB regenerator through modelling as there is no such 14 publication in the open literature. Correlations for liquid and gas mass transfer 15 coefficients, heat transfer coefficient, liquid hold-up, interfacial area and pressure 16 drop which are suitable for RPB regenerator were written in visual FORTRAN as 17 subroutines and then dynamically linked with Aspen Plus<sup>®</sup> rate-based model to 18 replace the default mass and heat transfer correlations in the Aspen Plus<sup>®</sup>. The 19 model now represents intensified regenerator/stripper. Model validation shows good 20 21 agreement between model predictions and experimental data from literature. Process analyses were performed to investigate the effect of rotor speed on the 22 regeneration efficiency and regeneration energy (including motor power). The rotor 23 speed was varied from 200 to 1200 rpm, which was selected to cover the validation 24 25 range of rotor speed. Impact of reboiler temperature on the rate of CO<sub>2</sub> stripping was also investigated. Effect of rich-MEA flow rate on regeneration energy and 26 regeneration efficiency was studied. All the process analyses were done for wide 27 range of MEA concentration (32.6 wt%, 50 wt% and 60 wt%). Comparative study 28 between regenerator using packed column and intensified regenerator using RPB 29 30 was performed and the study shows a size reduction of 9.691 times. This study indicates that RPB process has great potential in thermal regeneration application. 31

Keywords: Post-combustion CO<sub>2</sub> capture, MEA solvent, Process Intensification (PI),
 Rotating Packed Bed (RPB), Process Modelling, Process simulation

# 34 **1** Introduction

## 35 1.1 Background

Environmental concern has posed many questions as to the impact of greenhouse gas to those changes currently noticed in world climate and the future dangers that

will be expected if mitigation measures are not put in place. Combustion of coal and 38 petroleum accounts for the majority of the anthropogenic CO<sub>2</sub> emissions. Petroleum 39 is mostly used as a transportation fuel for vehicles while coal is used mostly for 40 electricity generation, for instance about 85.5% of coal is used for electricity 41 generation in 2011 in the UK [1]. Albo et al. [2] stated that among the greenhouse 42 gases, CO<sub>2</sub> contributes to more than 60% of global warming. Statistics from World 43 Metrological Organisation (WMO) showed the amount of CO2 in the atmosphere 44 reached 393.1 ppm in 2012. The WMO report also showed that the amount of CO<sub>2</sub> in 45 the atmosphere has increased on average by 2 ppm per year for the past 10 years. 46 Recent report by CO2-Earth [3] shows that as at 8 April 2017 CO<sub>2</sub> atmospheric 47 concentration stood at 407.78 ppm, this increased atmospheric concentration of CO<sub>2</sub> 48 affects the radiative balance of the earth surface [4]. 49

In order to meet the set target of 50% emission reduction by 2050 as compared to 50 the level of 1990 as proposed by Intergovernmental panel on climate change (IPCC) 51 [5], carbon capture and storage (CCS) is an important option for that target to be 52 53 achieved. The International Energy Agency (IEA) [6] identifies CCS as a significant and low-cost option in fighting climate change. The most matured CO<sub>2</sub> capture 54 technology is post-combustion CO<sub>2</sub> capture (PCC) based on chemical absorption as 55 reported in Mac Dowell et al. [7] which is also believed to be a low-risk technology 56 and promising near-term option for large-scale CO<sub>2</sub> capture. 57

58 PCC for coal-fired power plants using conventional packed columns has been 59 reported by many authors. Dugas [8] carried out pilot plant study of PCC in the context of fossil fuel-fired power plants. Lawal et al. [9-11] carried out dynamic 60 modelling and process analysis of CO<sub>2</sub> absorption for PCC in coal-fired power 61 plants. In all these studies, one of the identified challenges to the commercial roll-out 62 63 of the technology has been the high capital and operating costs which has an unavoidable impact on electricity cost. Systematic study of 64 aqueous monoethanolamine (MEA)-based CO2 capture process looking at the techno-65 economic assessment of the MEA process and its improvements was reported by Li 66 67 et al. [12]. Oh et al. [13] study energy minimization of MEA-based CO<sub>2</sub> capture process it was found that Flue gas splitting gives a significant reduction of energy 68 consumption. Solvent performance comparison for a large scale pulverized coal 69 power plant was reported by Sharifzadeh et al. [14]. Hanak et al. [15] reported 70 efficiency improvements for the coal-fired power plant retrofit with CO<sub>2</sub> capture plant 71 72 using chilled ammonia process showing efficiency penalty reduced to 8.7% Also Zhao et al. [16] using mixed solvent for 650 MW power plant reported that the net 73 power efficiency penalty was reduced from 9.13% to 7.66%. Approaches such as 74 heat integration, inter-cooling among others could reduce the operating cost slightly. 75 76 However, they limit the plant flexibility and will make operation and control more difficult [17]. Process intensification (PI) has the potential to meet this challenge [18-77 78 20].

Study of intensified absorber was reported in Joel et al [21,22] and Agarwal et al. 79 [23]. Joel et al [21] reported 12 times volume reduction for absorber if using RPB 80 technology as compared to packed column. Results from Agarwal et al. [23] 81 indicated 7 times volume reduction when using RPB as compared to conventional 82 packed column. The study by Joel et al. [21] uses aqueous MEA solvent while 83 Agarwal et al. [23] uses diethanolamine (DEA) as solvent. This is the main reason for 84 the differences in size reduction since faster reaction rate means shorter residence 85 86 time and slower reaction rate means longer residence time required for the same capture rate. Jassim et al. [24] and Cheng et al. [25] reported experimental studies 87 on intensified regenerator using RPB. Zhao et al. [26] study the mass transfer 88 performance of CO<sub>2</sub> capture in rotating packed bed and Chamchan et al. [27] 89 compared RPB and PB absorber in pilot plant. 90

Figure 1 is a typical process flow diagram of an intensified regenerator using RPB for solvent regeneration. The flowsheet was used by Jassim *et al.* [24] and Cheng *et al.* [25] for experimental study. One of the operational benefits of using RPB is its ability to be operated at higher gas and/or liquid flow rates owing to the low tendency of flooding compared to that in the conventional packed bed [28]. Another benefit of using RPB is its better self-cleaning, avoidance of blocking in the system, and being unaffected by a moderate disturbance in its orientation [29].



98 99

Figure 1 Schematic diagram of an RPB regenerator

100

# 101 Nomenclature

- a effective interfacial area (m<sup>2</sup>/m<sup>3</sup>)
- *a<sub>i</sub>* activity of species *i* in a solution
- at total specific surface area of packing (m<sup>2</sup>/m<sup>3</sup>)
- $a_w$  wetted area per unit volume (m<sup>2</sup>/m<sup>3</sup>)

$a'_p$	parameter for Chen et al. [24] and Chen [25] correlations for liquid and gas film
	mass transfer coefficients (= 3000 m <sup>2</sup> /m <sup>3</sup> )
С	width of wire mesh packing opening (mm)
$C_i^l$	concentration of component <i>i</i>
$Cp_i$	heat capacity for component <i>i</i>
d	wire diameter of wire mesh packing (mm)
D	column diameter (m)
$D_G$	diffusivity of gas (m <sup>2</sup> /s)
$D_L$	diffusivity of liquid (m²/s)
$E_j$	activation energy (kJ/mol)
dp	packing size (m)
G	volumetric gas flow rate (m <sup>3</sup> /s)
$G^m$	Gas molar flowrate (kmol/s)
g <sub>c</sub>	gravitational acceleration or acceleration due to centrifugal field (m <sup>2</sup> /s)
g₀	characteristic acceleration value (100 m <sup>2</sup> /s)
Н	height of packing (m)
h <sub>G</sub>	gas phase specific molar enthalpy (J/kmol)
$h_L$	liquid phase specific molar enthalpy (J/kmol)
$h_{g/l}$	interfacial heat transfer coefficient (W/m <sup>2</sup> K)
$\Delta H_r$	heat of desorption of CO <sub>2</sub> ( <i>J/kmol</i> )
$\Delta H_{vap}$	heat of vaporisation of H <sub>2</sub> O ( <i>J/kmol</i> )
k <sub>G</sub>	gas film mass transfer coefficient (m/s)
$K_G^a$	overall mass transfer coefficient (1/s)
$k_j^o$	pre-exponential factor (kmol/m <sup>3</sup> .s)
$k_L$	liquid film mass transfer coefficient (m/s)
L	Liquid mass flowrate per tangential area (kg/m²/s)
L <sup>m</sup>	Liquid molar flowrate ( <i>kmol/s</i> )
MEA	Monoethanolamine
N <sub>i</sub>	molar fluxes for component <i>i</i> ( $kmol/m^2 s$ )
P <sub>motor</sub>	motor power (kilowatts)
$Q_L$	volumetric flow rate of liquid (m <sup>3</sup> /s)
r	radial position (m)
R <sub>c</sub>	ideal gas constant (J kmol <sup>-1</sup> K <sup>-1</sup> )
$r_j$	reaction rate for reaction <i>j</i>
rxn <sub>i</sub>	reaction rate of component <i>i</i> , ( <i>kmol/m<sup>3</sup>/s</i> )

r <sub>i</sub>	inner radius of the RPB (m)
$r_o$	outer radius of the RPB (m)
$r_s$	radius of the stationary housing of the RPB (m)
Т	temperature (K)
t <sub>res</sub>	residence time (s)
u <sub>l</sub>	superficial liquid velocity (m/s)
$u_g$	superficial gas velocity (m/s)
Uo	characteristic superficial liquid velocity (1cm/s)
V	volume of the liquid films in the RPB (m <sup>3</sup> )
V <sub>i</sub>	volume inside the inner radius of the RPB = $\pi r_i^2 Z$ (m <sup>3</sup> )
Vo	volume between the outer radius of the bed and the stationary housing =
	$\pi (r_s^2 - r_0^2) Z$ (m <sup>3</sup> )
V <sub>t</sub>	total volume of the RPB = $\pi r_s^2 Z$ (m <sup>3</sup> )
<i>x</i> <sub>i</sub>	Component molar fraction in liquid phase
<i>Y</i> <sub>i</sub>	Component molar fraction in gas phase
$y_{CO_{2,in}}$	mole fraction of $CO_2$ in inlet gas stream
$y_{CO_{2,out}}$	mole fraction of CO2 in outlet gas stream
Ζ	axial height of the RPB (m)

#### 102 Greek letters

 $\alpha_{ij}$  reaction order of species *i* in reaction *j* 

- $\varepsilon$  porosity of packing, m<sup>3</sup>/m<sup>3</sup>
- $\in_L$  liquid holdup ( $m^3/m^3$ )
- $\mu$  viscosity (Pa.s)
- $\rho_L$  liquid density (kg/m<sup>3</sup>)
- $\rho_G$  gas density (kg/m<sup>3</sup>)
- $\sigma$  liquid surface tension (N/m)
- $\sigma_c$  critical surface tension (N/m)
- $\sigma_w$  surface tension of water (N/m)
- $v_L$  kinematic liquid viscosity (m<sup>2</sup>/s)
- $v_G$  kinematic gas viscosity (m<sup>2</sup>/s)
- $\omega$  angular velocity (rad/s)

## 103 Dimensionless groups

 $Fr_L$ liquid Froude number  $(u_l^2 a_t/g_c)$  $Gr_G$ gas Grashof number  $\left(d_p^3 g_c/v_G^2\right)$ 

- $Gr_L$  liquid Grashof number  $\left(d_p^3 g_c / v_L^2\right)$
- $Re_G$  gas Reynolds number  $(u_g/a_tv_G)$
- $Re_L$  liquid Reynolds number  $(u_l/a_tv_L)$
- $Sc_L$  liquid Schmidt number  $(v_L/D_L)$
- $We_L$  liquid Webber number  $(u_l^2 \rho_L / a_t \sigma)$
- $\varphi$  theoretical probability of liquid uncaptured by fibers  $(c^2/(d+c)^2))$

104 1.2 *Motivation* 

Over 8,000 tonnes of CO<sub>2</sub> per day will be released from a typical 500 MWe 105 advanced supercritical coal fired power plant operating at 46% overall net power 106 plant efficiency (LHV basis) [30]. This huge volume of flue gas will require big column 107 size. Lawal et al. [9] reported dynamic modelling study of a 500 MWe sub-critical 108 coal-fired power plant using the packed column (i.e. conventional technology). From 109 the study, one regenerator of 17m in packing height and 9 m in diameter will be 110 needed for regeneration of rich-MEA solvent. This huge packed column will mean 111 higher capital and operating costs, therefore a technological option leading to smaller 112 equipment size is very important. Kothandaraman et al. [31] reported that in 113 conventional packed tower majority (approximately 62%) of the energy consumed 114 during the CO<sub>2</sub> capture process was used for the solvent regeneration, therefore it is 115 necessary to look for technological options that will reduce this energy requirement. 116

## 117 1.3 Novel contributions of the paper

This is the first systematic study on RPB regenerator through modelling as there is 118 no such publication in the open literature. There are four novel aspects in this paper: 119 (a) A new first principle model for intensified regenerator using RPB was developed 120 which was implemented in Aspen Plus® rate-based model by replacing different 121 correlations for mass transfer, interfacial area and liquid hold-up. Steady state 122 validation of the intensified regenerator is performed using experimental data from 123 Jassim et al. [24] and Cheng et al. [25]. (b) Process analysis of the intensified 124 regenerator involving different process scenarios were carried out to gain insights for 125 process design and operation. These process scenarios are: (i) the impact of 126 rotational speed on the regeneration efficiency and regeneration energy at fixed 127 intensified regenerator size and rich-MEA flow rate was studied: (ii) the effect of rich-128 MEA solvent flow rate on the regeneration efficiency and regeneration energy was 129 explored; (iii) the effect of reboiler temperature on regeneration efficiency and 130 regeneration energy was explored. (c) Comparative study was performed between 131 132 intensified regenerator using RPB and conventional regenerator using packed bed. It was found that there is 9.69 times reduction in size under the same conditions which 133 means decrease in equipment capital cost. (d) The study were done over wide range 134

of MEA concentrations (32.6 wt%, 50 wt% and 60 wt%) and the energy consumption
 per ton of CO<sub>2</sub> is within the range of conventional packed column.

# 137 2 Model Development

Model for intensified regenerator using RPB does not exist in any commercially 138 available model library (including Aspen Plus®). To model intensified regenerator 139 using RPB, the default mass/heat transfer correlations in the Aspen Plus<sup>®</sup> rate-based 140 model have to be replaced with subroutines written in Intel® visual FORTRAN. The 141 new model now represents an intensified regenerator using RPB. The new steady 142 state model is still developed based on two-film theory Joel et al. [21]. The 143 correlations include: liquid phase mass transfer coefficient given by Chen et al. [32]. 144 gas-phase mass transfer coefficient given by Chen [33], interfacial area correlation 145 estimated by Luo et al. [34] and liquid hold-up correlation given by Burns et al. [35]. 146 Dry pressure drop expression was used since it accounts in an additive manner of 147 the drag and centrifugal forces, the gas-solid slip and radial acceleration effect [36]. 148

149 2.1 *Main governing equations* 

The main governing equations include material and energy balance equations.
Momentum balance is reflected in the pressure drop relation presented in Section
2.7.

# 153 *2.1.1 Gas and liquid phase material balances*

Assuming steady state conditions, material balances for gas and liquid phase in the RPB is described by Equations 1 and 2. Due to flow directions in RPB, the equations are based on numerical discretisation in the radial direction. Also, the term  $2\pi rZ$  is described preferably as tangential area to differentiate it from cross sectional area as it varies from section to section along the radial direction.

159 Material balances for gas phase:

160 
$$0 = \frac{1}{2\pi rZ} \frac{\partial (G^m y_i)}{\partial r} - aN_i$$
(1)

161 Material balances for liquid phase:

162 
$$0 = -\frac{1}{2\pi rZ} \frac{\partial (L^m x_i)}{\partial r} + aN_i + \epsilon_L rxn_i$$
(2)

The equation includes component molar flow balances (represented by the derivative terms) across each radial segment per tangential area, interfacial molar fluxes  $(aN_i)$  and liquid phase reaction rate  $(\in_L rxn_i)$ . By this, reactions are deemed to occur only in the liquid phase and ionic components therefore only exist in the liquid phase. Energy balances for gas and liquid phase is given by Equations 3 and 4 respectively [37]

171 Energy balances for gas phase:

172 
$$0 = \frac{1}{2\pi rZ} \frac{\partial (G^m h_G)}{\partial r} - ah_g (T_l - T_g) - Q_G^{Loss}$$
(3)

173 Energy balances for liquid phase:

174 
$$0 = -\frac{1}{2\pi r Z} \frac{\partial (L^m h_L)}{\partial r} - a \left( h_{g/l} \left( T_l - T_g \right) - \Delta H_r N_{CO_2} - \Delta H_{vap} N_{H_2O} \right) - Q_L^{Loss}$$
(4)

The equations include interfacial heat transfer,  $h_{g/l}(T_l - T_g)$ , heat released due to CO<sub>2</sub> desorption from the loaded MEA solvent,  $\Delta H_r N_{CO_2}$  and heat released or absorbed due to H<sub>2</sub>O condensation or vaporization,  $\Delta H_{vap}N_{H_2O}$ , [17]. Due to the relatively higher temperature of the stripper compared to ambient condition, heat losses ( $Q_G^{Loss}$  and  $Q_L^{Loss}$ ) are also taken into account.

180 2.2 Physical property

Electrolyte Non-Random-Two-Liquid (ElecNRTL) activity coefficient model in Aspen 181 Plus<sup>®</sup> was used to describe the vapour-liquid equilibrium, the chemical equilibrium 182 and the physical properties of the system. The equilibrium constants for reactions 5-183 9 are calculated from the standard Gibbs free energy change, the equilibrium 184 reactions are assumed to occur in the liquid film and kinetic reactions equations and 185 parameters are obtained from AspenTech [38]. The electrolyte solution chemistry 186 which is used in property calculation is modelled with chemistry model and all the 187 ionic reactions are assumed to be in chemical equilibrium as shown in Equations 5-9 188 [38]. 189

190	Equilibrium	$2H_20 \leftrightarrow H_30^+ + 0H^-$	(5)
-----	-------------	---------------------------------------	-----

191 Equilibrium  $CO_2 + 2H_2O \leftrightarrow H_3O^+ + HCO_3^-$  (6)

192 Equilibrium 
$$HCO_3^- + H_2O \leftrightarrow H_3O^+ + CO_3^{2-}$$
 (7)

193 Equilibrium 
$$MEAH^+ + H_2O \leftrightarrow MEA + H_3O^+$$
 (8)

194 Equilibrium  $MEACOO^- + H_2O \leftrightarrow MEA + HCO_3^-$  (9)

195 Kinetic reactions used for the intensified stripping calculation is specified by 196 Equations 10-13 in the reaction part of the regenerator model in the Aspen Plus.

197 Kinetic 
$$CO_2 + OH^- \rightarrow HCO_3^-$$
 (10)

198 Kinetic 
$$HCO_3^- \rightarrow CO_2 + OH^-$$
 (11)

199 Kinetic 
$$MEA + CO_2 + H_2O \rightarrow MEACOO^- + H_3O^+$$
 (12)

200 Kinetic MEACOO<sup>-</sup> + 
$$H_3O^+ \rightarrow MEA + CO_2 + H_2O$$
 (13)

Power law expression Equation 14 is used for the rate-controlled reactions. The kinetic parameters for reactions in Equations 10-13 were listed in Table 1

203 
$$r_{j} = k_{j}^{o} exp\left(-\frac{E_{j}}{R_{c}}\left[\frac{1}{T} - \frac{1}{298.15}\right]\right) \prod_{i=1}^{N} a_{i}^{\alpha_{ij}}$$
(14)

Table 1 Constants for power law expressions for the absorption of CO<sub>2</sub> by MEA [39]

Reaction No.	kj⁰ (kmol∕m³.s)	<i>E<sub>j</sub></i> , kJ/mol
10	1.33e+17	55.38
11	6.63e+16	107.24
12	3.02e+14	41.2
13	6.56e+27	95.24

205

#### 206 2.3 Liquid phase mass transfer coefficient

An expression was introduced by Tung and Mah [40] based on penetration theory to describe the liquid mass transfer behaviour in the RPB.

209 
$$\frac{k_L d_p}{D_L} = 0.919 \left(\frac{a_t}{a}\right)^{1/3} Sc_L^{1/2} Re_L^{2/3} Gr_L^{1/6}$$
 (15)

This correlation was developed without considering the Coriolis force or the effect of the packing geometry. This is why there is a need for an alternative correlation for liquid phase mass transfer coefficient.

Chen *et al.* [32] developed liquid phase mass transfer correlation considering the end effect and packing geometry. The correlation was found to be valid for different sizes of the RPBs and for viscous Newtonian and non-Newtonian fluids. Because of these advantages, Equation 16 is selected for calculating the liquid phase mass transfer coefficient, and also findings from Joel *et al.* [22] suggested the use of Equation 16 because of it smaller error prediction.

220 
$$\frac{k_L a d_p}{D_L a_t} \left( 1 - 0.93 \frac{V_o}{V_t} - 1.13 \frac{V_i}{V_t} \right) = 0.35 S c_L^{0.5} R e_L^{0.17} G r_L^{0.3} W e_L^{0.3}$$
221 
$$\left( \frac{a_t}{a'_p} \right)^{-0.5} \left( \frac{\sigma_c}{\sigma_w} \right)^{0.14}$$
(16)

222

#### 223 2.4 Gas phase mass transfer coefficient

Onda *et al.* [41] correlation for calculating gas-side mass transfer coefficient (Equation 17) was developed for conventional packed column. Sandilya *et al.* [42] suggested that the gas rotates like a solid body in the rotor because of the drag force caused by the packing, which means that gas-side mass transfer coefficient should
be similar to that in a conventional packed column, but the end effect and packing
effect were not considered, this makes the authors to select Equation 18 proposed
by Chen [33] instead of Equation 17.

231 
$$k_G = 2.0(a_t D_G) R e_G^{0.7} S c_G^{1/3} (a_t d_p)^{-2}$$
 (17)

Chen [33] presented local gas-side mass transfer coefficient correlation using twofilm theory for RPB. Equation 18 for calculating the gas phase mass transfer coefficient was used in the model because it accounts for the effect of rotation of the RPB.

236 
$$\frac{k_G a}{D_G a_t^2} \left( 1 - 0.9 \frac{V_o}{V_t} \right) = 0.023 R e_G^{1.13} R e_L^{0.14} G r_G^{0.31} W e_L^{0.07} \left( \frac{a_t}{a_p'} \right)^{1.4}$$
(18)

237

238 2.5 Total gas-liquid interfacial area

Total gas-liquid interfacial area correlation for conventional packed column was developed by Onda *et al.* [41] as shown in Equation 19. It can be modified to account for the effect of rotation of the bed but because it is not originally designed for RPB and also it was not designed for different types of packing, Equation 20 developed by Luo *et al* [34] was selected.

244 
$$\frac{a}{a_t} = 1 - exp\left[-1.45\left(\frac{\sigma_c}{\sigma}\right)^{0.75} Re_L^{0.1} W e_L^{0.2} F r_L^{-0.05}\right]$$
(19)

Luo *et al.* [34] studied gas-liquid effective interfacial area in an RPB considering different types of packing, also taking into account the effect of fibre diameter and opening of the wire mesh.

248 
$$\frac{a}{a_t} = 66510Re_L^{-1.41}Fr_L^{-0.12}We_L^{1.21}\varphi^{-0.74}$$
(20)

249 2.6 Liquid hold-up

Liquid holdup correlation given by Burns *et al.* [35] was used. The correlation is based on data obtained through measurement of electrical resistance across sections of an RPB. The study showed that the liquid hold-up is approximately inversely proportional to the local packing radius and is largely independent of gas flow up to the flooding point and also liquid viscosity has only a weak influence on hold-up [35].

256 
$$\in_L = 0.039 \left(\frac{g_c}{g_o}\right)^{-0.5} \left(\frac{u_l}{u_o}\right)^{0.6} \left(\frac{v_L}{v_o}\right)^{0.22}$$
 (21)

257 
$$g_o = 100 \text{ m s}^{-2}, \quad U_o = 1 \text{ cm s}^{-1}, \quad v_o = 1 \text{ cS} = 10^{-6} \text{ m}^2 \text{ s}^{-1}$$

$$258 u_l = \frac{Q_L}{2\pi rZ} (22)$$

259 2.7 Dry pressure drop expression

Semi-empirical dry pressure drop expression was given by Llerena-Chavez and 260 Larachi [36]. The correlation was developed based on Ergun-type semi-empirical 261 relationships in which the gas-slip and radial acceleration effects, the laminar and 262 inertial drag effects and the centrifugal effect were aggregated additively to form the 263 pressure drops correlation in the RPB [36]. 264

(23)

265 
$$\Delta P_{Packed bed} = \frac{150(1-\varepsilon)^2 \mu}{d^2 \varepsilon^3} \left(\frac{G}{2\pi Z}\right) \ln \frac{r_o}{r_i} + \frac{1.75(1-\varepsilon)\rho}{d\varepsilon^3} \left(\frac{G}{2\pi Z}\right)^2 \left(\frac{1}{r_i} - \frac{1}{r_o}\right) + \frac{1}{2}\rho\omega^2(r_o^2 - r_i^2) + F_c$$
(23)

where  $F_c$  is a corrective function given as: 267

268 
$$F_c = \varepsilon (a - G + (b + \omega^c)G^2)$$
(24)

a, b, and c are fitting parameters given as: 269

 $a = -0.08 \ m^3/s$   $b = 2000(rpm)^c$ 270 c = 1.22

#### 2.8 Power consumption by RPB stripper motor 271

272 The amount of power consumed by motor for rotating RPB absorber and stripper is 273 calculated using the correlation proposed by Singh et al. [43]. The correlation was used to account for all the frictional losses and also the power required for 274 accelerating the liquid entering the packing bed to the rotational speed at the outer 275 radius. It is important to note that frictional losses are highly dependent upon the 276 design of the machine and cannot be predicted without advance knowledge of the 277 design (i.e., type of bearings, direct or pulley drive, etc.) [43]. 278

279 
$$P_{motor} = 1.2 + 1.1 \times 10^{-3} \rho_L r_o^2 \omega^2 Q_L$$
 (25)

## 280 2.9 *Modelling and simulation methodology*

The procedure used in this paper for modelling and simulation of the RPB is shown

in **Figure 2** and summary of the model parameters and correlations were presented

# 283 in **Table 2**



284

285

Figure 2 Methodology used in this paper [21,22]

Table 2 Summary of model parameters

Parameters	Correlations or values		
	Jassim <i>et al</i>	Cheng et al	
Model geometry			
ri (m)	0.156	0.076	
r <sub>o</sub> (m)	0.398	0.160	
h (m)	0.025	0.020	
surface area of the packing per unit	2132	803	
volume of the bed (m <sup>2</sup> /m <sup>3</sup> )			
Liquid phase mass transfer coefficient	See E	Equation 16	
Gas phase mass transfer coefficient	See E	Equation 18	
Total gas-liquid interfacial area	See E	Equation 20	
Liquid holdup	See E	Equation 21	
Dry pressure drop expression	See E	Equation 23	
Motor power	See E	Equation 25	

#### 287 3 Model Validation

#### 288 3.1 Model validation using experimental data from Jassim et al. [24]

The experimental data used for the model validation was obtained from Jassim *et al.* [24]. From their experiments, rich-MEA concentration of 32.9 wt%, 35.7 wt%, 30.8 wt%, 57.4 wt% and 52 wt% were selected for the validation study. The equipment specifications and process input conditions for the validation study are shown in **Tables 3 and 4.** The study was done under two different rotor speeds 800 rpm and 1000 rpm.

Description	Value
RPB outer diameter	0.398 m
RPB inner diameter	0.156 m
RPB axial depth	0.025 m
Packing specific surface area	2132 m²/m³
Packing porosity	0.76

Table 3 RPB stripper packing specifications used by Jassim et al. [24]

#### Table 4 Input process conditions for Run 1 to Run 5 [24]

	Runs				
	Run 1	Run 2	Run 3	Run 4	Run 5
Rotor speed (RPM)	800	800	800	1000	1000
Rich-MEA temperature (°C)	67.100	69.000	70.000	57.200	58.400
Rich-MEA pressure (kPa)	101.325	101.325	101.325	101.325	101.325
Rich-MEA flow rate (kg/s)	0.200	0.200	0.400	0.400	0.200
Rich-MEA composition (wt. %) H <sub>2</sub> O CO <sub>2</sub> MEA	58.116 8.984 32.900	54.013 10.287 35.700	61.536 7.664 30.800	25.142 17.458 57.400	32.895 15.105 52.000
Rich-MEA CO2 loading (mol CO2 /mol MEA)	0.3790	0.3999	0.3454	0.4221	0.4030

297

#### Table 5 Simulation results compared to experimental data [24] for Run 1 to Run 5

	Runs				
	Run 1	Run 2	Run 3	Run 4	Run 5
Rotor speed (RPM)	800	800	800	1000	1000
Experimental measurement					
Lean-MEA CO <sub>2</sub> loading (mol/mol)	0.321	0.329	0.329	0.403	0.334
Model prediction					
Lean-MEA CO <sub>2</sub> loading (mol/mol)	0.316	0.295	0.298	0.355	0.320
Relative error (%)	1.558	10.334	9.422	11.911	4.192

299

Model validation is shown in **Table 5** which gives percentage error prediction of not more than 12 % on the lean-MEA CO<sub>2</sub> loading. The lean-MEA CO<sub>2</sub> loading was

302 evaluated on mole basis as shown in Equation 26.

303	Loading =	Moles of all CO <sub>2</sub> carrying species _	$[CO_2] + [HCO_3^-] + [CO_3^{2-}] + [MEACOO^-]$	(26)
		Moles of all MEA carrying species	$[MEA] + [MEA^+] + [MEACOO^-]$	(20)

Jassim *et al.* [24] didn't include experimental results on reboiler duty, therefore the authors cannot compare model predictions with experimental tests.

306 3.2 Model validation based on experimental data from Cheng et al. [25]

Cheng et al [25] carried out experimental study on the thermal regeneration of alkanolamines solutions in a RPB using 30 wt% MEA aqueous solution loaded with CO<sub>2</sub> and a CO<sub>2</sub> - loaded aqueous solution consisting of 20 wt% diethylenetriamine and 10 wt% piperazine. For the purpose of this study, experimental data with 30 wt% MEA aqueous solution was used for model validation. RPB stripper specifications and process input conditions for the model are shown in **Tables 6 and 7**.

Table 6 RPB stripper packing specifications used by Cheng et al [25]

Description	Value
RPB outer diameter	0.160 m
RPB inner diameter	0.076 m
RPB axial depth	0.020 m
Packing specific surface area	803 m²/m³
Packing porosity	0.960

Table 7 Input process conditions for different reboiler temperature [25]

Variable	Reboiler Temperature			
Vanable	105 °C	115 °C	120 °C	
Rotor speed (RPM)	900	900	900	
Rich-MEA temperature (°C)	96.6	97	97	
Rich-MEA pressure (kPa)	202.65	202.65	202.65	
Rich-MEA flow rate (mL/min)	400	400	400	
Rich-MEA CO <sub>2</sub> loading (mol CO <sub>2</sub> /mol MEA)	0.484	0.484	0.484	

Table 8 Simulation results compared to experimental data [25]

Variable		Reboiler Temperature (°C		
		105	115	120
	Experimental measurement	0.418	0.340	0.271
Lean Loading	Model prediction	0.423	0.367	0.289
(mol CO <sub>2</sub> /mol MEA)	Relative error (%)	1.132	8.054	6.848
	Experimental measurement	0.620	0.900	1.240
	Model prediction	0.629	0.989	1.383
Reboiler duty (kW)	Relative error (%)	1.487	9.951	11.498

Model results compared with the experimental data from Cheng *et al.* [25] shown in **Table 8** indicates a good agreement with relative error on lean loading of less than 9% and reboiler duty percentage error of less than 12% for different reboiler temperatures.

In summary, the model has predicted all experimental data reasonably well with not more than 12% error prediction, the model developed can then be used to carry out process analysis in order to study the system behaviour when there is a change in some variables.

## 324 4 Process Analysis

With the validated models, we carried out process analysis to explore the effect of 325 rich-MEA flow rate, rotor speed and reboiler temperature on (a) the regeneration 326 efficiency calculated based on loading (Equation 27) and calculated based on 327 328 amount of CO<sub>2</sub> in rich-MEA and lean-MEA solvent (Equation 28), (b) the regeneration energy (with and without motor power) expressed in Equations 29 and 329 30 respectively. But the electricity power consumed by motor is high grade while the 330 steam power in the reboiler is a low grade, therefore for the two powers to be 331 332 comparable, efficiency loss needs to be accounted for by multiplying the motor

power with 2.5 (i.e. assuming 40% thermal efficiency in converting thermal heat into 333 electricity. The reason is that solvent-based carbon capture can be used in different 334 scenarios such as coal-fired subcritical power plants, coal-fired supercritical power 335 plants, gas-fired power plants and other industrial manufacturing plants. 40% is a 336 good assumption for these different scenarios). Equations 31 and 32 were used to 337 estimate the solvent residence time in the RPB with the assumption that wetted area 338 per unit volume is equal to specific surface area of the packing [44]. The RPB 339 stripper used for the process analysis has the following packing geometry: outer 340 radius = 0.371 m; inner radius = 0.152 m; axial depth of packing = 0.167 m; packing 341 void fraction = 0.76; packing specific surface area =  $2,132 \text{ m}^2/\text{m}^3$ . 342

343 Regeneration eff. 1 = 
$$\left(\frac{\text{Rich CO}_2 \text{ loading} - \text{Lean CO}_2 \text{ loading}}{\text{Rich CO}_2 \text{ loading}}\right) \times 100$$
 (27)

Regeneration eff. 2 = 
$$\left(\frac{\text{Amount of } CO_2 \text{ in Rich } - \text{Amount of } CO_2 \text{ in Lean}}{\text{Amount of } CO_2 \text{ in Rich}}\right) \times 100$$
 (28)

Regeneration energy (without motor power) = 
$$\frac{\text{Reboiler duty}}{\text{Mass of CO}_2 \text{ desorbed}}$$
 (29)

Regeneration energy (with motor power) = 
$$\frac{(\text{Reboiler duty} + P_{motor} \times 2.5)}{\text{Mass of CO}_2 \text{ desorbed}}$$
(30)

347 Residence time 
$$(t_{res}) = \frac{V}{Q_L}$$
 (31)

348 Where

349 
$$V = \left(\frac{3v_L Q_L}{2\pi Z a_w \omega^2}\right)^{1/3} (a_w 2\pi Z) \left[r_o^{4/3} - r_i^{4/3}\right]$$
(32)

350  $Q_L$  = liquid volumetric flowrate (m<sup>3</sup>/s)

351

#### 4.1 Effect of rich solvent flow rate on regeneration efficiency and energy

#### 353 4.1.1 Justification for case study

Rich-MEA solvent flow rate not only has influence on the amount of CO<sub>2</sub> that will be stripped off from the regenerator, but also has relationship with the reboiler duty. Therefore study on the right quantity of rich-MEA solvent coming into the regenerator of fixed or given size is necessary.

#### 358 *4.1.2* Setup of the case study

For this study, the process input conditions are shown in Table **9** with the rich-MEA flow rate varying from 0.2 kg/s to 0.8 kg/s. Here the rich-MEA loading is kept constant (i.e. 0.482 mol CO<sub>2</sub>/mol MEA). Also in this study lean-MEA loading and reboiler temperature are the two outputs parameters that were maintained at 0.3178 mol CO<sub>2</sub>/mol MEA and 120 °C respectively. Reboiler temperature is maintained at 120 °C by controlling the back pressure regulator to the stripper.

Variable	Case 1	Case 2	Case 3
Rich-MEA temperature (°C)	104	104	104
Rich-MEA pressure (kPa)	202.650	202.650	202.650
Rich-MEA flow rate (kg/s)	0.2 - 0.8	0.2 - 0.8	0.2 - 0.8
Rich-MEA composition (wt. %)			
H <sub>2</sub> O	56.072	32.027	18.559
CO <sub>2</sub>	11.328	17.530	21.010
MEA	32.600	50.443	60.431
Rich loading (mol CO <sub>2</sub> /mol MEA)	0.482	0.482	0.482
Reboiler temperature (°C)	120	120	120
Rotor speed (RPM)	1000	1000	1000

365 Table 9 Process inputs

#### 366

#### 367 4.1.3 Results and discussion

Figure 3a (using Equation 27) and Figure 3b (using Equation 28) show a constant 368 regeneration efficiency as the rich-MEA flow rate increases, this is because the rich-369 370 MEA loading is the same for all the cases and the lean-MEA loading which is one of the output is controlled at the same value of 0.3178 mol/mol. Figure 4a presents the 371 amount of CO<sub>2</sub> strip-off from the stripper as the rich-MEA flow rate increases. The 372 graph shows an increase in the amount of CO<sub>2</sub> desorbed as the rich-MEA flow rate 373 increases. This is attributed to increase in droplet flow regime. As liquid flowrate 374 increases, the liquid breaks up more readily as they enter the rotating packing 375 forming more droplets due to their higher velocity. Studies by Chambers and Walls 376 [45] already showed that droplet flow regime in RPBs generally favours better mass 377 transfer performance than the film flow regime. It is not surprising then that CO<sub>2</sub> 378 desorption rate noticeably increases (Figure 4a) as liquid flowrate increases. Figure 379 **4a** also shows that higher MEA concentration gives higher CO<sub>2</sub> desorption rate. This 380 is due to their higher loading capacity, which means the amount of CO<sub>2</sub> absorbed is 381 more. Therefore, under similar conditions then, desorption rate from more 382 383 concentrated MEA solution is expected to be more.

**Figure 4b** shows firstly that the residence time decreases with increasing liquid flowrate. To understand this, referring to Equations 31 and 32, it is seen that the residence time can be related to flowrate as follows (assuming other parameters in Equations 31 and 32 remain constant):

388 
$$t_{res} = KQ_L^{-\frac{2}{3}}$$
 (33)

With K (= constant), this simply shows that increasing flowrate will result to lower residence time. In physical terms, this can be further explained by acknowledging that liquid velocity increases with flowrate. Higher liquid velocity means that delay within the system is less and this ultimately means lower residence time.

The other result shown in **Figure 4b** is increase in residence time as MEA solution concentration increases. This is due to increase in solution density (i.e. 1062.784kg/m<sup>3</sup> for 32.6 wt%, 1162.062 kg/m<sup>3</sup> for 50.443 wt%, 1209.465 kg/m<sup>3</sup> for 60.431 wt%) and viscosity (i.e. 0.000681235 N.s/m<sup>2</sup> for 32.6 wt%, 0.000990415 N.s/m<sup>2</sup> for 50.443 wt%, 0.00125367 N.s/m<sup>2</sup> for 60.431 wt%).



Figure 3 Effect of rich-MEA flow rate on regeneration efficiency (a) using Equation 27 (b) using Equation 28



401 Figure 4 Effect of rich-MEA flow rate on (a) CO<sub>2</sub> desorbed (b) Residence time

It can be observed from Figure 5a,b that the regeneration energy increases with 402 403 increase in rich-MEA flow rate for Cases 2 and 3 (with and without motor power). For Case 1 (without motor energy) the regeneration energy is fairly constant as the rich-404 MEA flowrate increases as shown in Figure 5a. This is because the percentage 405 increase in the reboiler duty shown in **Figure 6** is same as the percentage in CO<sub>2</sub> 406 407 desorbed (Figure 4a) (i.e. 75.103% increase in reboiler duty and 74.973% increase in amount of CO<sub>2</sub> desorbed for Case 1) while for Case 2 there is 77.383% increase 408 in reboiler duty and 75.003% increase in amount of CO<sub>2</sub> desorbed and Case 3 has 409 80.703% increase in reboiler duty, 74.986% increase in amount of CO<sub>2</sub> desorbed. 410 The lowest regeneration energy obtained from the study is at flow rate of 0.2 kg/s 411 rich-MEA. For Case 3 the regeneration energy is 5.17 GJ/ton CO<sub>2</sub> (without motor 412 energy) and 5.44 GJ/ton CO<sub>2</sub> (with motor energy). The highest regeneration energy 413 for Case 2 is at rich-MEA flow rate of 0.8 kg/s and the regeneration energy is 5.17 414 GJ/ton CO<sub>2</sub> (without motor energy) and 5.31 GJ/ton CO<sub>2</sub> (with motor energy). For 415 Case 1 (with motor power) regeneration energy decreases with increase in rich-MEA 416 flow. 417

The percentage increase in regeneration energy (i.e. including motor energy) when rich-MEA flowrate increases from 0.2 kg/s to 0.8 kg/s is 5.74% and 20.18% for Cases 2 and 3 respectively, while for Case 1 the regeneration energy decreases by 5.72%.

Looking at **Figures 5a,b and 6**, one may wonder why higher MEA concentration solvent is preferred for the RPB technology when the energy consumption is higher. It should be noted that here it is for stripper only. For a closed loop process including RPB Absorber and RPB stripper, the recycling solvent flowrate would be much lower for higher concentration solvent for the same capture efficiency in the absorber. Thus the energy consumption will be lower.





0 \_\_\_\_\_\_0.1

0.2

0.3





0.5

Rich MEA flow rate (kg/s)

0.6

0.7

0.8

0.9

0.4

434



Figure 7 Effect of Rich-MEA flow rate on (a) mass transfer rate (b) heat transfer rate

Figure 7a shows how the CO<sub>2</sub> mass transfer rate increases as the rich-MEA flow 436 rate increases. Mass transfer is generally defined as transfer from either vapour to 437 liquid or liquid to vapour. Here CO<sub>2</sub> is transferred from liquid to vapour, therefore 438 negative sign appears. Figure 7a shows higher mass transfer rate for high MEA 439 concentration this is because more CO<sub>2</sub> has been dissolve in it, therefore the rate of 440 CO<sub>2</sub> desorbed will be higher than the lower MEA concentration. In Figure 7b heat 441 transfer rate increases with increase in rich-MEA flow rate, the negative sign indicate 442 443 transfer from liquid to vapour. The studies on CO<sub>2</sub> mass transfer rate and the heat transfer rate look at mass and heat transfer from the inner to outer radius of the RPB 444 excluding the mass and heat transfer in the condenser and reboiler. 445

To further account for why there is an increase in the regeneration energy as shown 446 in Figures 5a,b, the heat duty requirement in the reboiler is divided into three 447 different parts: (i) Sensible heat to raise the temperature of the rich-MEA stream in 448 the reboiler; (ii) Heat of reaction to reverse the absorption reaction and release CO<sub>2</sub>; 449 (iii) Heat of vapourisation to maintain the driving force for transfer of CO<sub>2</sub> from liquid 450 phase to gas phase. Figures 8a,b and 9 show how the heat of vapourisation, 451 sensible heat and the heat of reversible reaction increases with increase in rich-MEA 452 flow rate respectively. Figure 8a shows that Heat of vapourisation is higher for high 453 MEA concentration than lower MEA concentration this is due to the difference in their 454 vapour composition (i.e. Case 1 has vapour composition of 0.25% H<sub>2</sub>O, 97.13% 455 MEA and Case 2 has 0.70% H<sub>2</sub>O, 97.64% MEA and Case 3 has 1.49% H<sub>2</sub>O, 456 97.85% MEA). Sensible heat for the three Cases is almost the same this is because 457 the rich-MEA flow rate coming into the stripper is same and the specific heat 458 capacity is relatively same and the difference in the specific heat capacity is counter 459 balanced by the temperature differences. Figure 9 shows that heat of reversible 460 reaction increases with increase in concentration this is as a result more energy 461 needed to break the CO<sub>2</sub> and MEA bonds and because of the decrease in the 462 amount of free CO<sub>2</sub> as the MEA concentration increases. Figure 10 shows how the 463

reboiler duty requirement was divided for a fixed rich-MEA flowrate of 0.8 kg/s at three different MEA concentrations. This shows that the heat of reaction increases from 47%, to 65% and 74% when rich-MEA concentration increases from 32.9 wt%, to 50.443 wt% and 60.431 wt%. This is consistent with amount of CO<sub>2</sub> stripped as presented in Figure 4 (a).

469 Therefore, this study will help operators and designers of RPB regenerator to

balance between energy consumption requirement and amount of CO<sub>2</sub> desorbed for

a given Rich-MEA flow rate. 62000 50000 -Case 1\_32.9 wt% 45000 Case 1\_32.9 wt% -Case 2\_50.443 wt% Heat of vapourization (W) 52000 Case 2\_50.443 wt% 40000 Case 3 60.431 wt% Case 3\_60.431 wt% €35000 42000 Sensible heat Se 32000 22000 10000 12000 5000 0 2000 0.3 0.5 0.7 Rich-MEA flow rate (kg/s) 0.1 0.9 0.1 0.5 0.7 0.9 0.3 Rich-MEA flow rate (kg/s) b а

#### Figure 8 Effect of Rich-MEA flow rate on (a) Heat of vapourisation (b) Sensible heat



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471

472



Figure 9 Effect of Rich-MEA flow rate on heat of reversible reaction



- Figure 10 Heat contributions for 0.8 kg/s rich-MEA flowrate at different MEA
  concentration (a) 32.9 wt% MEA (b) 50.443 wt% MEA (c) 60.431 wt% MEA
- 478 4.2 Effect of rotor speed on regeneration efficiency and regeneration energy

## 479 4.2.1 Justification for case study

The higher the rotating speed of the intensified regenerator the higher the energy consumed, therefore it is important to understand the relationship that rotor speed has with rich-MEA solvent flow rate so that the energy requirement for driving the stripper can be reduced with respect to the amount of rich-MEA solvent regenerated.

484 4.2.2 Setup of the case study

For this study, the rotor speed was varied from 200 rpm and 1200 rpm in order to cover the experimental range of rotor speed reported in Jassim *et al* [24] and Cheng et al [25]. Input process conditions for this study are shown in **Table 9** (i.e. Cases 1 and 2). The reboiler temperature, rich-MEA flow rate and rich-MEA loading were kept constant at 120 °C 0.3 kg/s and 0.4823 mol/mol respectively for all the cases.

490 *4.2.3 Results and discussion* 

Figure 11a shows that the regeneration efficiency increases with increase in the 491 rotor speed. The impact of rotor speed on lean-MEA loading is shown in Figure 11b. 492 Though higher rotor speed can produce opposite effect on mass and heat transfer by 493 decreasing the residence time (as shown in Figure 12a) but this effect was counter 494 balanced by the increase in the interfacial area which enhances mass and heat 495 transfer. Burns et al. [46] stated that at higher rotor speed there are more of smaller 496 liquid droplets and thinner liquid films in the packing regions of the bed, which means 497 increase in interfacial area. The set-up in this study is different with the one reported 498 in Section 4.1 where the lean-MEA loading was kept constant but in this study the 499 recovery rate is kept constant for all the MEA concentrations, but the recovery rate 500

changes as the rotor speed increases. Figure 11a shows that regeneration 501 efficiency decreases with increase in concentration this is because the rich-MEA 502 loading is maintained at same value and the recovery rate is fixed at the same value 503 for different rotor speed meaning that the change between rich-MEA loading to lean-504 MEA loading is smaller for higher MEA concentration than lower MEA concentration. 505 The amount of CO<sub>2</sub> desorbed from the stripper increases as the rotor speed 506 increases as shown in Figure 12b. The amount of CO2 desorbed for the two 507 different cases are similar this is due to the model set-up where the recovery rate 508 were maintained at the same value but varied with rotor speed. 509





510

Figure 11 Effect of rotor speed on (a) regeneration efficiency (b) lean loading



512

513 Figure 12 Effect of rotor speed on (a) residence time (b) CO<sub>2</sub> desorbed

Increase in rotor speed decreases the regeneration energy as shown in Figure 13a.
This is because increase in rotor speed leads to more liquid droplet and thin liquid
films to dominate the packing resulting in increase in mass and heat transfer. Also at

higher rotational speed the problem of liquid mal-distribution is overcome leading to 517 higher wetted area which subsequently contributes to improving mass transfer. For 518 all cases, the trend in Figure 13a (without motor energy) shows a drop in the 519 regeneration energy as the rotor speed increases from 200 rpm to 1200 rpm this is 520 521 because of increase in the rate of CO<sub>2</sub> stripped-off (**Figure 12b**). But when energy consumed by the motor is included **Figure 13b**, there is an increase in regeneration 522 energy at rotor speed above 600 rpm for Case 2 and above 400 rpm for Case 1. This 523 is because the motor energy is a function of square of rotor speed. Also Figures 524 13a,b shows that regeneration energy decreases with increase in MEA 525 concentration this is due to smaller difference between rich-MEA loading and lean-526 loading as seen in Figure 11b (i.e. at rotor speed of 600rpm Cases 1 and 2 has 527 lean-MEA loading as an output from the model of 0.2898 mol/mol and 0.354076 528 mol/mol respectively). The average percentage increase in regeneration energy 529 530 when motor power is included is 6.44% and 6.84% for Case 1 and Case 2 respectively. Figure 14 shows how the reboiler duty increases with increase in rotor 531 speed. Case 1 has higher reboiler duty because the difference in rich-MEA to lean-532 533 MEA loading is bigger which means higher reboiler duty, since reboiler duty is related to the difference in lean and rich loading. 534

535 This study will help operator and designer of RPB regenerator in chosen the rotor 536 speed that give lower regeneration energy without compromising the amount of CO<sub>2</sub> 537 desorbed.



539 Figure 13 Effect of rotor speed on regeneration energy (a) without motor energy (b) 540 with motor energy



541



Figure 14 Effect of rotor speed on reboiler duty

543 4.3 Effect of reboiler temperature on regeneration efficiency and energy

## 544 *4.3.1 Justification for case study*

545 Operating intensified regenerator at the right reboiler temperature will lead to good 546 system performance by reducing regeneration energy waste and also operating at 547 relatively high regeneration efficiency.

548 4.3.2 Setup of the case study

549 For this study, the reboiler temperature was varied from 105 to 125 °C. Process input 550 conditions are same as in **Table 9** (i.e. Cases 1 and 2). The rich-MEA flow rate and 551 rich-MEA loading were kept constant at 0.3 kg/s and 0.4823 mol/mol respectively for 552 all the cases.

## 553 4.3.3 Results and discussion

Figure 15a shows that the regeneration efficiency increases with increase in reboiler 554 temperature. The percentage increase in regeneration efficiency as the reboiler 555 temperature increases for Cases 1 and 2 is about the same 9.67% and 9.33% 556 respectively. The model shows that regeneration efficiency for Case 1 which has 557 lower MEA concentration is higher. This is due to lower lean-MEA loading coming 558 out as shown in Figure 15b since at each reboiler temperature for Case 1 and 2, the 559 amount CO<sub>2</sub> desorbed is maintained at around the same value and also more CO<sub>2</sub> 560 was dissolved in higher MEA concentration than the lower MEA concentration in 561 order to have the same rich-MEA loading coming into the stripper. Figure 15b shows 562 that there is a decrease in lean-MEA loading as the reboiler temperature increases. 563 This is because of increase in the amount of CO<sub>2</sub> stripped-off as shown in Figure 16. 564







Figure 15 Effect of reboiler temperature on (a) regeneration efficiency (b) lean loading



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569

Figure 16 Effect of reboiler temperature on amount of CO<sub>2</sub> desorbed

There is a decrease and increase in regeneration energy as the reboiler temperature 570 increases from 105 °C to 125 °C. From Figure 16a,b (with and without motor 571 energy) the regeneration energy decreases as the reboiler temperature increases 572 from 105 °C to 115 °C for Case 1 and 2, but this behaviour changes when the 573 reboiler temperature exceed 115 °C. This is because at higher temperature we 574 expect increase in water vapour flow rate which results in increase in regeneration 575 energy because of heat of vaporisation of water. Also Figure 19b shows how the 576 MEA composition in the vapour stream increases with increase in reboiler 577 temperature this means increase in heat of vapourisation. Also from Figure 15b the 578 loading different between the rich-MEA and lean-MEA stream is wide for Case 1 579 than Case 2 that is why it has higher reboiler duty requirement. **Figure 18** shows 580 how the reboiler duty increases with increase in reboiler temperature. The decrease 581

and increase in the regeneration energy can be further explained by **Figure 20 a,b** and **Figure 21** where the reboiler heat duty requirement is split into the heat of vapourisation, sensible heat and heat of reversible reaction. Increase in reboiler temperature leads to increase in heat of vapourisation and sensible heat while heat of reversible reaction decreases. The decrease in heat of reversible reaction is because of increase in rate of reaction as temperature increases.





591



592



Figure 18 Effect of reboiler temperature on reboiler duty







а

b

596 Figure 20 Effect of reboiler temperature on (a) heat of vapourisation (b) sensible heat





Figure 21 Effect of reboiler temperature on heat of reversible reaction

599

# 5 Comparison between RPB based intensified and PB based regenerator

#### 601 5.1.1 Justification for the case study

This study was carried out to provide a comparison under some fixed conditions such as Rich-MEA flowrate, pressure, temperature, rich-MEA loading and lean-MEA loading between intensified regenerator and conventional regenerator.

#### 605 5.1.2 Setup of the case study

For this study, **Table 10** presents the input conditions for the conventional and intensified regenerator. The rotor speed for the intensified regenerator is kept constant at 1000 rpm. Regeneration efficiency was kept constant at 37.16 % for both the conventional and the intensified regenerators.

610	Table 10 Process	conditions for	Conventional	and RPB	regenerator
				••••••	

Description	Conventional	RPB regenerator
	regenerator	
	R-MEA	Lean-MEA
Rich-MEA temperature (°C)	104	104
Rich-MEA pressure (kPa)	202.650	202.650
Rich-MEA flowrate (kg/s)	0.300	0.300
Rich-MEA loading	0.482	0.482
(mol CO <sub>2</sub> /mol MEA)		
Mass-Fraction (wt%)		
H <sub>2</sub> O	56.072	56.072
CO <sub>2</sub>	11.328	11.328
MEA	32.600	32.600

#### 611 5.1.3 Results and discussion

The results in Table 11 show a 44 times packing volume reduction in RPB 612 regenerator compared to conventional PB regenerator without sumps. Using the 613 assumption given by Agarwal et al. [23] that the casing volume of RPB is 4.5 times 614 the rotating packing volume, the volume reduction compared to conventional PB 615 regenerator is found to be 9.691 times smaller. The height of transfer unit (HTU) for 616 conventional PB regenerator is calculated as 20.8 cm while for the RPB based 617 intensified regenerator is 1.7 cm. The smaller HTU in RPB regenerator leads to its 618 smaller size compared to conventional packed column. Wang et al. [20] performed 619 preliminary technical and economic analysis for intensified PCC process compared 620 with conventional PCC process. Initial prediction on the capital cost of the whole 621 intensified PCC process can reduce by 1/6 (i.e. 16.7%) compared with the same 622 capacity conventional PCC process. The 9.691 times reduction in the volume of 623

intensified regenerator reported here confirmed to the possibility of having 16.7%cost reduction for intensified PCC process.

Description	Conventional	RPB regenerator
	PB regenerator	
Height of packing (m)	3.700	0.371 (r <sub>o</sub> )
		0.152 (r <sub>i</sub> )
diameter (m)	0.476	0.167 axial depth
Packing Volume (m <sup>3</sup> )	0.659	0.015
Packing volume reduction		43.933 times
Volume of unit (m <sup>3</sup> )	0.659 <sup> a</sup>	0.068 <sup>b</sup>
Volume reduction factor		9.691 times
Specific area (m <sup>2</sup> /m <sup>3</sup> )	151	2132
Void fraction	0.980	0.760
Lean-MEA loading (mol CO <sub>2</sub> /mol MEA)	0.303	0.303

Table 11 Comparison between conventional and RPB stripper

627 <sup>a</sup> Excluding sump

<sup>628</sup> <sup>b</sup> Using the assumption given by Agarwal et al [23]

629

#### 630 6 Conclusions

Intensified regenerator using RPB technology was modelled in this study. The steady
state model was implemented by linking Aspen Plus<sup>®</sup> and visual FORTRAN. The
model developed was validated with experimental data reported in Jassim *et al.* [24]
and Cheng et al. [25]. The model validations show good agreement with the
experimental data.

Process analysis on the effect of rich-MEA flow rate, rotational speed and 636 reboiler temperature on CO<sub>2</sub> regeneration efficiency and regeneration energy 637 638 were performed. For the given stripper (fixed in physical size), the study shows that an increase in the rich-MEA flow rate leads to an increase regeneration 639 energy. There is an increase in the regeneration efficiency as the rotor speed 640 increases but the regeneration energy decreases as the rotor speed increases 641 642 since mass and heat transfer is enhanced at higher rotor speed. Reboiler temperature was varied from 105 °C to 125 °C, the results show a decrease in 643 regeneration energy at reboiler temperature between 105 °C to 120 °C, but when the 644 reboiler temperature exceeds 120 °C the regeneration energy begins to increase. 645 Under the same process conditions, RPB based intensified stripper/regenerator has 646 647 volume reduction of 9.691 times compared to conventional PB based stripper/regenerator. RPB stripper/regenerator shows great potential for application 648 as a stripper and has much smaller size compared to conventional stripper which 649 means reduction in capital cost. 650

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