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1 Comparative studies of CO₂ capture solvents for gas-fired power plants:

2 Integrated modelling and pilot plant assessments

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5

6 **Abstract-** Natural gas accounts for 21.3% of the total primary energy supply. In particular, with the shale gas
7 boom, its application has experienced an exponential growth rate. Nevertheless, natural gas has the highest
8 hydrogen to carbon ratio among all hydrocarbons and the dilute (with respect to CO₂) exhausts of its
9 combustion pose considerable thermodynamic challenges for carbon capture. Solvent-based carbon capture
10 can potentially mitigate the carbon emissions. Nonetheless, minimizing the penalties associated with the costs
11 of carbon capture requires detailed understanding of the underlying physical and chemical phenomena. The
12 present research exploits a rigorous methodology based on rate-based distributed modelling and statistical
13 associating fluid theory (SAFT). The important characteristics of this modelling approach include abstract
14 formulation in conjunction with high predictability. The present research aims at establishing the performance
15 of a new solvent (an amine-promoted buffer salt, APBS) in comparison to the baseline solvent, *i.e.*,
16 monoethanolamine (MEA). The features of interest include: (i) developing a high fidelity mathematical model
17 of the carbon capture process, (ii) validating the model with pilot plant data, (iii) quantification of the energetic
18 and technical performances of the solvents using key process indicators (KPIs), and (iv) employing optimization
19 programming for further solvent performance improvement.

20

21 **Keywords:** Solvent-based carbon capture, SAFT, rate-based modelling, model validation, optimization, MEA,
22 APBS, GCCmax.

23

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24 1. Introduction

25 Increasing energy demand and associated pollution have posed an important paradoxical challenge toward the
26 security of energy supply and environmental protection. This has translated into various national and global
27 policies and regulations. The instances of these international regulations include the so-called Europe 2020
28 which targets 20% reductions in greenhouse gases, 20% energy supply from renewables, and 20%
29 improvement in energy efficiency. In the UK, the Climate Change Act 2008 mandates an 80% overall cut in six
30 greenhouse gases by 2050 compared to the 1990 baseline. While integration of renewable energy resources
31 can moderate the problem to some extent, the International Energy Agency (IEA) asserts that in a foreseeable
32 future, fossil fuels remain the dominant source of energy supply, and application of carbon capture
33 technologies will become increasingly urgent (IEA 2014a).

34 Natural gas (NG) accounts for 21.3% of the global primary energy supply (IEA 2014b) and with the shale gas
35 boom, its application has experienced an exponential growth rate. Nevertheless, natural gas has the highest
36 hydrogen to carbon ratio among all hydrocarbons and the dilute (with respect to CO₂) exhausts of its
37 combustion pose considerable thermodynamic challenges for carbon capture. Amongst various carbon capture
38 methods, post-combustion has the advantage of being an end-of-pipe treatment, requiring a minimal of
39 retrofit to the existing energy infrastructure. In particular, solvent-based carbon capture has been already
40 applied in ammonia and natural gas sweetening processes and is deemed to be the most commercially viable
41 technology for end-of-pipe waste treatment of power plants. Furthermore, solvent-based carbon capture has
42 proved to be an efficient method for increasing the yield of biorefineries (Sharifzadeh, et al., 2015) and
43 emission reduction in the iron and steel industry (Ho et al., 2013).

44 In the recent years, developing solvents with superior techno-economic performance has been the focus of
45 academic and industrial researchers. Frimpong et al., (2013) compared the performance of a proprietary
46 amine-carbonate blend for carbon capture from coal-derived flue gas. The comparison with MEA baseline
47 solvent was made based on carbon capture efficiency, energy of solvent regeneration, and overall mass
48 transfer rate. Both solvents were studied at similar experimental conditions. Moser et al., (2011a,b) evaluated
49 the performance of a solvent developed by BASF in a highly instrumented pilot plant in Niederaussem. They
50 used CHEMASIM, an in-house simulation software tool, for data reconciliation and model validation. Their
51 results suggested a high correlation between the reboiler duty, the desorber pressure, solvent circulations

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52 rate, and solvent heat of absorption. Similar observation was made by Bumb et al., (2014) who studied the
53 performance of a novel solvent optimized for coal-fired flue gas, at TNO pilot plant, Netherlands. They studied
54 variation of the desorber top pressure between 0.8-1 barg, while maintaining the reboiler temperature
55 constant. They reported that reboiler heating duty and solvent circulation were both minimum at 0.8 barg.
56 However any further reduction would result in unacceptable condensate rate from the desorber overhead.
57 Mangalapally et al., (2011a,b, 2012) conducted pilot plant studies for four new solvents. They identified the
58 solvent circulation rate as the most influential operating parameter for pilot plant operation. Rabensteiner et
59 al., (2014) studied pilot scale performance of ethylenediamine (EDA) compared to MEA using literature data.
60 Cousins et al., (2012, 2015) studied the performance of concentrated piperazine and MEA in an Australian pilot
61 plant. They observed that at elevated liquid/gas ratio, absorber intercoolers can mitigate the regeneration
62 energy requirements by 10%. They also reported that rich solvent split can result in less reboiler and
63 condenser duties.

64 An ideal solvent should have high CO₂ loading, fast reaction kinetics, low vapour pressure, negligible
65 corrosiveness, and high thermal stability, in addition to a lower cost. Often these criteria are interdependent
66 and require establishing a compromise. Oexmann and Kather (2010) provided an analytic analysis of thermal
67 requirements for solvent regeneration heating duty, as a function of (i) solvent sensible heat (energy needed
68 to increase the solvent temperature from absorber temperature to desorber temperature), (2) heat of
69 evaporation (to produced stripping steam in the desorber), and (3) heat of desorption (needed to break the
70 solvent-CO₂ bound). They demonstrated that these solvent characteristics are often interdependent. Solvents
71 with a high heat of absorption often feature a higher kinetic rate. Furthermore, since the desorber can be
72 operated at a higher pressure and the reboiler can be operated as a higher temperature, the water reflux from
73 the desorber overhead is lower; hence the heating duty of the desorber will be also lower. Furthermore, a
74 faster kinetic enables higher CO₂ loading and therefore, resulting in less solvent circulation rate and hence less
75 sensible heat requirement. Nevertheless, the maximum allowable temperature in the desorber depends of the
76 solvent thermal stability against degradation reactions. By comparison solvents with a lower heat of
77 absorption may benefit from operating the desorber in vacuum operating conditions, at the costs of a higher
78 power requirement for CO₂ compression.

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79 More recent studies have focused on solvents with additional functionalities. For example, ammonia is
80 regarded as a cost-effective solvent for CO₂ separation. However, low absorption rates limit the practical
81 application of this solvent. Yang et al., (2014) investigated the effects of adding potassium sarcosinate as the
82 rate promoter to the ammonia solvent. Similarly, Thee et al., (2012) showed that the addition of small
83 quantities of MEA to potassium carbonate solvent results in radical increases in the overall reaction rate.
84 Recently, a new class of solvent is introduced, which exploit phase change in order to limit the energetic
85 penalties of solvent regeneration. For example as shown by Liebenthal et al., (2013), an amine blend can form
86 two liquid phases under certain CO₂ loading conditions. Then, a large energy saving is possible by sending only
87 the CO₂-rich phase to the desorber. Furthermore, Ionic liquids are a promising class of solvents for carbon
88 capture. The tunability of the ionic liquids' structure and properties offers several advantages such as lower
89 energy requirement for solvent regeneration, higher thermal stability, and lower vapour pressure. However,
90 the high production cost of ionic liquids is a barrier that needs to be addressed for their commercialization,
91 (Chen et al., 2014).

92 Finally, process optimization has been the focus of various studies. Lee et al., (2013) investigated the
93 performance of several amine solvents. They demonstrated a strong trade-off between the capital investment
94 and operating costs using Pareto front diagrams. Mores et al., (2012) studied simultaneous optimization of the
95 dimensions of the process equipment and process operating conditions. Three scenarios were considered in
96 which reboiler duty was being optimized with respect to different combinations of reboiler inlet liquid flow-
97 rate, reboiler inlet solvent composition, reboiler pressure, CO₂ loading and CO₂ recovery target. They observed
98 a correlation between the optimal solvent CO₂ loading and the CO₂ recovery targets. In addition, they reported
99 that as CO₂ loading increases, the specific heating and cooling duties (per unit of CO₂ recovered) decrease.
100 Hopkinson et al., (2014) developed a simple method for optimization of a conceptual solvent based on
101 properties of commercial amine solvents. Solvent heat of reaction was applied in order to estimate the
102 sensible and stripping heat of the process. They reported the value of 71 kJ/mol CO₂ for the heat of reaction
103 and the value of 0.1034 kWh/kg CO₂ for the total equivalent work in the case of 90% CO₂ recovery from coal-
104 fired flue gases. Damartzis et al., (2015) proposed a module-based generalized design framework in order to
105 synthesize the optimal process flow diagram. Features of interest included the stream topologies, the heat
106 redistribution and the cascades of desorption columns for several commercially available solvents. Significant

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110 economic improvement (15%-35%) and reductions in the reboiler duty (up to 55%) were observed. Recently,
111 Burger et al., (2015) proposed a hierarchical methodology based on a group contribution method which
112 considers the molecular decisions at the same level of process design decisions. They demonstrated their
113 method for selection of the optimal solvent over a wide range of ethers, for CO₂ separation from a methane-
114 rich mixture.

115 Several common conclusions can be drawn from the aforementioned theoretical and pilot plant studies.

- 116 • In establishing the performance of new solvents, there is always a need for a well-understood and
117 predictable reference solvent such as monoethanolamine (MEA) which facilitates pilot plant testing
118 runs and provides benchmark performance measures.
- 119 • Due to interdependencies of process operating specifications and combinatorial characteristics of the
120 involved parameters, the pilot plant studies were often conducted in conjunction with simulation and
121 optimization software tools.
- 122 • The pilot plant operating conditions are solvent-specific and the performance of each solvent should
123 be studied at its corresponding optimal specifications.
- 124 • There is a need for key performance indicators which are not solvent-specific and are reproducible for
125 different solvents.

126 The present research aims at benchmarking the performance of a new solvent called GCCmax. The GCCmax
127 solvent is a novel solvent recently developed by Carbon Clean Solutions Limited (CCSL) for carbon capture from
128 natural gas combustion exhausts and belongs to the class of amine-promoted buffer salt (APBS) solvents. The
129 conducted research is comprehensive and include, pilot plant studies, model development and validation, and
130 optimization in order to provide guidelines for further solvent optimization. The novel contribution of the
131 present research includes:

- 132 1. Developing high fidelity multi-scale models for post-combustion carbon capture and validating the
133 model for the case of the GCCmax and MEA solvents using pilot plant data.
- 134 2. Quantification of the performance of the solvents using a novel set of key performance indicators
(KPIs), that enables objective and reproducible comparisons.
3. Identification of further opportunities for process optimization in terms of equipment design,
operating conditions, and solvent composition.

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135 The rest of the manuscript is organized as follows. Firstly, the research methodology is discussed. The features
136 of interest include the applied modelling approach and underlying assumptions. Then, the results of model
137 validation are presented and compared for the baseline MEA solvent and the new GCCmax solvent. The
138 comparison is made based on several novel key process indicators which quantify the difficulty of separating
139 CO₂ from the flue gas. These “technical” KPIs are used as substitutes for economic indicators because using the
140 latter to compare alternative solvents would suffer from assumptions around the cost of capital, the capital
141 estimation method and the cost of utilities, limiting the reproducibility of the results. The last part of the paper
142 explores the opportunities for further improvement of the new solvent performance through optimization
143 with respect to process equipment, operating conditions, and the solvent composition.

144 **2. Methodology**

145 The research methodology is based on a multi-scale process modelling and optimization. Firstly, a rigorous
146 process model was developed based on accurate understanding of the involved physical and chemical
147 phenomena and rate-based modelling of gas-liquid contactors. Then, the developed model is validated using
148 pilot plant data. Finally, the validated model is optimized for various design and retrofit scenarios.

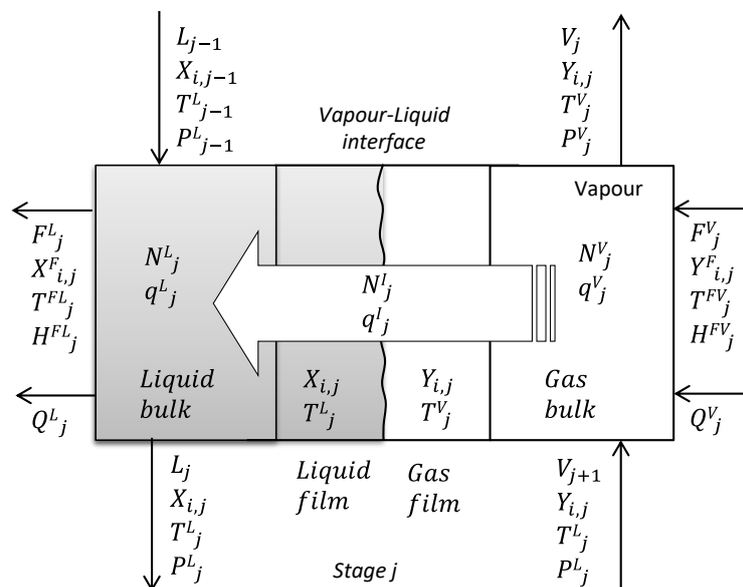
149 **2.1. Process model development**

150 The accurate modelling of the solvent-based CO₂ capture processes for the purposes of solvent benchmarking
151 and comparison requires thorough understanding of the underlying physical and chemical phenomena
152 involved in absorption and desorption of CO₂ into and from the working solvent. The methods for modelling
153 gas-liquid contactors can be broadly classified into equilibrium and rate-based models. Early-stage attempts
154 for modelling gas-liquid separation processes were based on the assumption that at each separation stage, the
155 exiting phases are in equilibrium conditions. Such assumptions were also incorporated into graphical methods
156 (McCabe et al., 2004). However, it was widely observed that for most industrial applications assuming
157 equilibrium composition of the exiting phases results in unrealistic predictions. The first attempt to overcome
158 this limitation was to consider an efficiency measure for each separation stage, (McCabe et al., 2004), which
159 can be applied to calculate the actual number of stages from the theoretical number of stages. Application of
160 empirical efficiency correlations has been promising for binary and close-boiling separation systems. However,
161 it was widely observed that for multicomponent systems, the efficiency of separation stages can vary from 5%
162 for absorption systems with high viscosity, high molecular weight solvents to 120% in single-liquid-pass large-

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163 diameter distillation trays (or packed segments) because of the crossflow effect (Seader et al., 2013). Due to
 164 difficulties associated with uncertainties in the stage efficiency method, intensive research was devoted to
 165 developing non-equilibrium models based on transport rates (Waggoner and Loud, 1977; Krishna and Standart
 166 1979; Taylor and Krishna, 1993; Krishnamurthy and Taylor, 1985; Taylor et al., 1994). The rate-based models
 167 are founded on the two-film theory, as shown in Fig. 1. Here, thermodynamic equilibrium is assumed only at
 168 the interface of vapour and liquid phases. Unlike equilibrium-based models, the exiting vapour phase is
 169 superheated and the exiting liquid phase is subcooled and they have different temperatures. The exchanged
 170 mass and energy between phases depend on the driving forces, transport coefficients, and the interfacial area.
 171 Often, both convective and diffusive transport phenomena are involved and component-coupling effects
 172 should be considered (Seader et al., 2013). Various empirical correlations for calculating the mass transfer
 173 coefficient are proposed by researchers for random packing (Onda et al., 1968; Bravo and Fair, 1982) and
 174 structured packing (Bravo et al., 1985, 1992). Finally, the bulk liquid and gas phases may have different flow
 175 configurations such as plug or mixed flows.



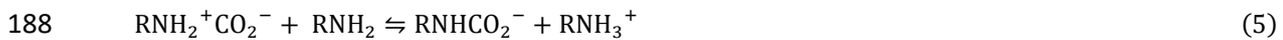
176
 177 Fig. 1. Two-film model used for modelling rate-based absorption and desorption of CO₂ into and from solvent,
 178 (Zhang et al., 2009).

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179 2.2. Reaction kinetics and thermodynamics

180 A full space rate-based representation of CO₂-solvent gas-liquid contactors requires detailed modelling of
181 underlying chemical reactions, column hydraulics, as well as mass and heat transfer phenomena. For the case
182 of CO₂ capture using alkanolamines, this would require modelling a series of speciation reactions as follows
183 (Mac Dowell et al., 2010):



190 where for the case of MEA, R=HO-CH₂-CH₂- . In the scheme above, reaction (1) represents water hydrolysis,
191 reactions (2) and (3) are concerned with carbonic acid and bicarbonate formation, respectively. Reaction (4)
192 represents the Zwitterion formation and reaction (5) is base catalysis. The two latter reactions are highly
193 coupled as carbamate and protonated amines are tightly bonded. Therefore, reaction (4) and (5) can be shown
194 abstractly as:



196 Reaction (7) is a reversible acid-base neutralization reaction. In this reaction, absorption of CO₂ in
197 alkanolamine solvent is exothermic; hence, it is possible to liberate CO₂ by heating the CO₂-rich solvent
198 mixture and driving the reaction to left. Therefore, in theory, it is possible to regenerate the alkanolamine and
199 recycle it for reuse in the absorber. In practice, a small amount solvent make-up is supplied to compensate the
200 losses associated with the solvent escape and degradation reactions.

201 Various thermodynamic models were proposed by researchers in order to describe the thermophysical
202 properties of the mixture of CO₂ absorbed into alkanolamines. These methods can be broadly categorized to
203 non-rigorous and rigorous methods. In non-rigorous methods, equilibrium constants are considered for the
204 aforementioned reactions and are reconciled against experimental data. The major drawback of non-rigorous
205 methods is the lack of precision for closing energy balances and predicting speciation of components¹⁷. In
206 addition, the rigorous methods can be classified to (1) activity-based models (derived from Gibbs free energy),

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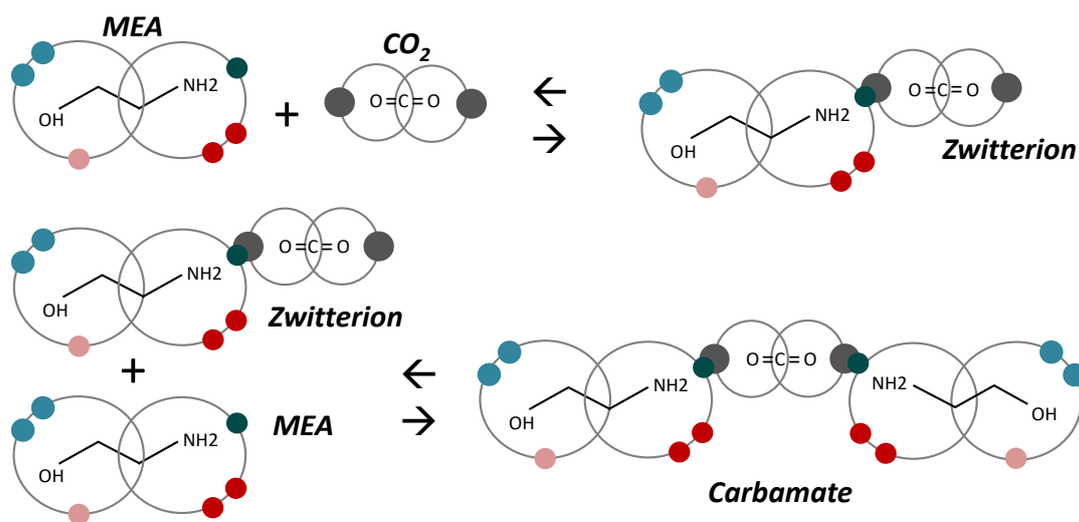
207 and (2) equation of state models (derived from Helmholtz free energy). A frequently used activity-based model
208 is Electrolyte Non-random two-liquid model (E-NRTL) in which the activity coefficient expressions of the
209 original NRTL model are modified, and certain constraints regarding local like-ion repulsion and
210 electroneutrality are imposed (Chen and Song, 2004). An alternative activity based model is the Extended
211 UNIQUAC model where the original UNIQUAC model was modified to account for ionic interactions (Faramarzi
212 et al., 2009). A limitation of the aforementioned activity-based models is that they only describe the liquid
213 phase and a separate equation of state (e.g., Soave–Redlich–Kwong equation of state by Faramarzi et al.,
214 2009) should be applied for modelling the vapour phase. Alternatively, researchers attempted to use
215 equations of states which can consider the presence of ionic components (Huttenhuis et al., 2008). More
216 recently, application of equations of state which are able to accommodate chemical equilibria such as
217 association between molecules, has been the focus of various research groups (Fürst and Renon, 1993; Button
218 and Gubbins, 1999). Here, the treatment is based on statistical associating fluid theory (SAFT), in which the
219 free Helmholtz energy is correlated to the intermolecular association between the molecule segments
220 (Chapman et al., 1989; Mac Dowell et al., 2010):

$$221 \quad \frac{A^{mix}}{NkT} = \frac{A^{IDEAL}}{NkT} + \frac{A^{MONO}}{NkT} + \frac{A^{CHAIN}}{NkT} + \frac{A^{ASSOC}}{NkT} \quad (8)$$

222 In above, N refers to the number of molecules, k is the Boltzmann constant and T is the mixture temperature.
223 On the right-hand-side, the first term represents the ideal Helmholtz free energy and the three other terms
224 refer to the residual contribution due to monomers interactions, formation of chains, and intermolecular
225 interaction between associating sites. Then, given the Helmholtz free energy of the mixture, A^{mix} , it is possible
226 to calculate other mixture properties (e.g., $\mu_i = (\delta A / \delta N_i)_{T,V,N_{i \neq j}}$). The association contribution is based on
227 thermodynamic perturbation theory (TPT) (Wertheim, 1984), where by using intermolecular potential models
228 (e.g., square well with variable range) and by adjusting their parameters (well depth and range), it is possible
229 to fully characterize the strength of the associating sites.

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Fig. 2. Schematic representation of Carbamate formation

232 The idea is shown in Fig. 2 for the case of reaction (4) and (5), (Mac Dowell et al., 2010). In the new approach,
233 the rate of reactions, the concentration of intermediate zwitterion, and its thermophysical properties are not
234 formulated anymore. Instead, CO₂ and MEA are represented as associating molecule chains with two and six
235 associating sites, respectively. Then, the concentration of CO₂ in association with two MEA molecules
236 represents the actual CO₂ loading of the solvent at different temperatures and pressures. The combination of
237 rate-based modelling and representation of chemical reactions using statistical associating fluid theory (SAFT)
238 provides a consistent modelling approach. The justification is that for solvents such as MEA and GCCmax, the
239 rate of reaction is significantly faster than the heat and mass transfer phenomena. Therefore, the knowledge
240 of the rate of reactions at the gas-liquid interface is unnecessary and the chemical equilibrium sufficiently
241 describes the actual physical system behaviour. It is notable that in the case of solvents for which the reaction
242 kinetics are not fast enough, this modelling approach may introduce approximations.

243 This modelling approach offers several advantages; firstly unlike activity-based models, the same equation of
244 state is used to describe both liquid and vapour phases. Secondly, the chemical equilibria are treated at the
245 same level as phase equilibria. Furthermore, this approach results in significant model reduction because the
246 speciation of intermediate ions is not included in the mathematical formulation and the uncertainties
247 associated with their thermophysical parameters are disentangled from problem formulation. Most of all, the
248 aforementioned approach establishes a connection between the chemical and physical behaviour of the
249 mixture and the molecular structure of the involved materials. This is of particular importance to new solvents

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250 as the required information can be acquired from the available data for the molecular segments of associating
251 sites from literature experimental data.

252 In the present research, rate-based gas-liquid contactors were developed based on a combination of two-film
253 theory and statistical associating fluid theory for potentials of variable range (SAFT-VR). The applied software
254 tools were advanced modelling library gas-liquid contactors (AML-GLC) and gSAFT toolboxes developed by
255 Process System Enterprise Ltd (PSE). The parametric values of thermodynamic models for the GCCmax solvent
256 are obscured in order to respect the confidentiality agreements with Carbon Clean Solutions Limited (CCSL)
257 and Process System Enterprise (PSE). The underlying mathematical modelling of the gas-liquid contactors were
258 similar to (MacDowell et al., 2013). The SAFT thermodynamic model and its application for modelling CO₂
259 capture solvent are described by MacDowell et al., (2010).

260 3. Pilot plant operation and model validation

261 In order to ensure effective benchmarking and model validation, two sets of pilot plant runs were conducted
262 using the monoethanolamine (MEA) and GCCmax solvents. MEA served as the baseline reference solvent. The
263 pilot plant studies were conducted in the US National Carbon Capture Center (NCCC) located in Alabama, USA.
264 The capture process is shown in Fig. 3, which is based on a more detailed process flow diagram in (Gayheart et
265 al., 2013). The column specifications for the capture process in the National Carbon Capture Center (NCCC) are
266 reported in Table 1. The process description is as follows. Firstly, the pressure of the pre-scrubbed and cooled
267 flue gas is increased via a blower and then is sent to the absorber column. In the absorber column, flue gas is
268 brought in contact with the lean solvent, and the CO₂ is chemisorbed into the solvent. The absorption process
269 is exothermic and the released heat increases the solvent temperature. The solvent enriched with the
270 absorbed CO₂ is sent to the desorber column where the carbon dioxide is released through heating. Before,
271 recycling the hot lean solvent to the absorber column for reuse, there is an opportunity for heat integration to
272 the rich solvent. Additional cooling will be provided using cooling water. The solvent and water losses are
273 compensated with adequate make-ups.

274 **Table 1. Column Specifications (Gayheart et al., 2013).**

Column	Packing	Height of each bed (m)	Diameter (m)	Number of beds
Absorber	Mellapakplus M252Y- Steel	6.1	0.66	3
Desorber	Mellapakplus M252Y- Steel	6.1	0.61	2

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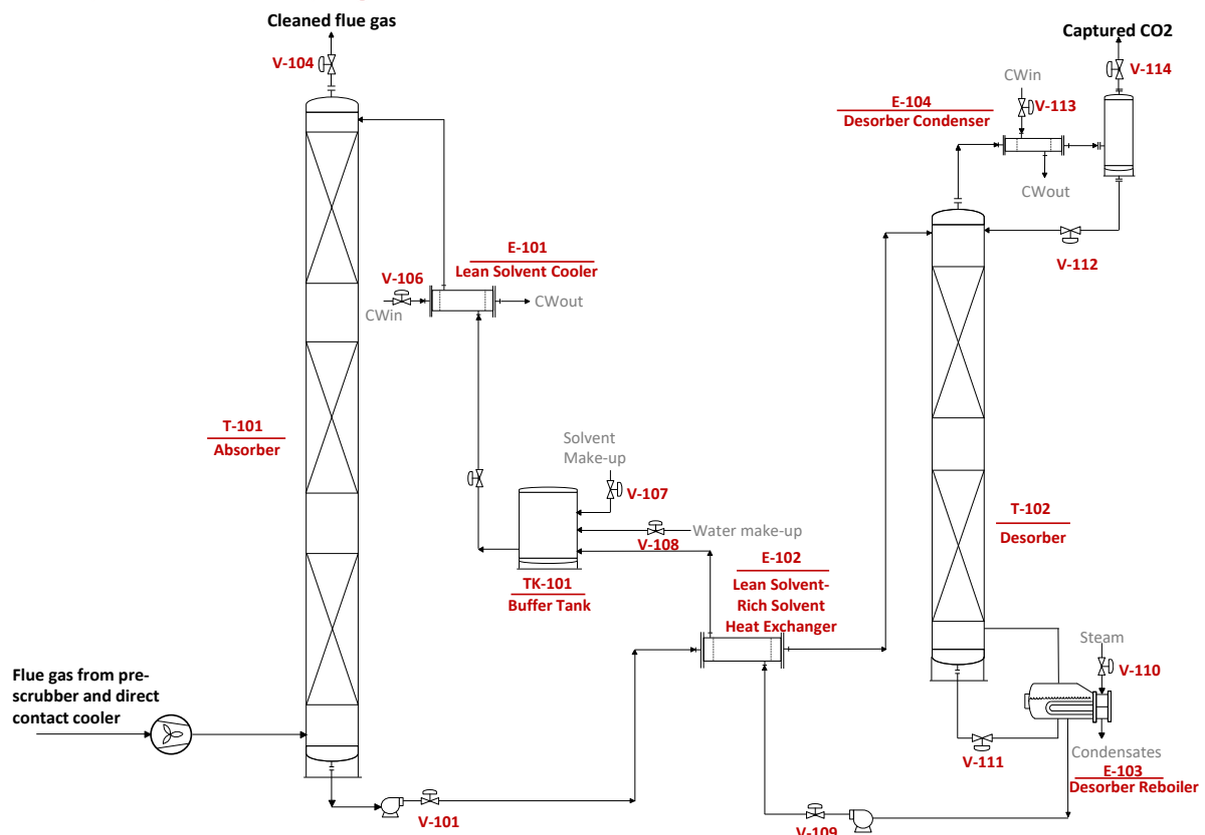


Fig. 3. A representative flow diagram of the US Natural Carbon Capture Center (NCCC) CO₂ capture pilot plant, Alabama, US (Gayheart et al., 2013).

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279 Table 2 shows the results of model validation. Since the NCCC pilot plant was not previously operated under
280 natural gas exhaust conditions, the model validation was conducted based on historical data for a scenario of
281 coal-fired exhausts. The last column in Table 2 reports the prediction of the model, when the process is
282 operated under natural gas exhaust conditions. Table 2 shows a very good agreement between pilot plant data
283 and simulation results, in terms of captured CO₂, and the solvent composition. Minor discrepancies in the
284 consumed steam are deemed to be associated with heat losses and temperature indicator errors. Table 3
285 reports the results of the GCCmax solvent model validation under natural-gas-fired conditions. Two sets of
286 pilot plant data were used, which essentially are different with respect to the lean solvent temperature
287 entering the absorber top. The justification was due to the fact that in different parts of the world, cooling
288 water may be supplied at different temperatures. Again the model predictions are in very good agreement
289 with the pilot plant data with respect to the captured CO₂ and the solvent concentrations, giving confidence in
290 the model's predictive capabilities. Minor discrepancies in the required steam and temperatures were
291 attributed to lack of insulation or temperature measurement errors.

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Table 2. Model validation for MEA baseline solvent.

		Alabama-coal	Alabama-coal	Alabama NG
		Pilot Plant	Simulation	Simulation
Flue gas to the absorber				
Nitrogen + Oxygen	Mass Fraction	0.786	0.786	0.909
Carbon Dioxide	Mass Fraction	0.165	0.165	0.067
Water	Mass Fraction	0.049	0.049	0.024
Total flowrate	kg/s	0.628	0.628	0.628
Flue Gas Temperature - Absorber Inlet	K	316.56	316.56	316.56
Lean Solvent - Absorber Inlet				
Amine	Mass Fraction	0.297	0.298	0.296
CO ₂	Mass Fraction	0.063	0.059	0.059
Water	Mass Fraction	0.640	0.643	0.645
Total	kg/s	2.52	2.51	1.3
Lean solvent temperature	K	316.15	316.15	316.15
Intercoolers outlet temperature	K	316.15	316.15	-
Reboiler Steam				
Steam pressure	bar	2.92	2.92	2.92
Steam temperature	K	405.6	405.6	405.6
Steam flowrate	kg/s	0.18	0.15	0.07
Lean-Rich Heat Exchanger				
Lean in	K	388.87	388.9	389
Lean out	K	331.4	338.9	330.8
Rich in	K	327.91	330.4	321.7
Reboiler temperature	K	385.6	388.0	389.0
Absorber bottom pressure	bar	1.1	1.17	1.16
Absorber top pressure	bar	1.04	1.04	1.04
Desorber bottom (reboiler) pressure	bar	1.71	1.71	1.16
Desorber top pressure	bar	1.69	1.70	1.70
General specifications				
CO ₂ capture efficiency	%	91.84	91.85	90
Inter-stage Cooling		Yes	Yes	No

295

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296

Table 3. Model validation for GCCmax solvent.

		Data Set 1	Data Set 1	Data Set 2	Data Set 2
		Pilot Plant	Simulation	Pilot Plant	Simulation
Absorber inlet gas stream					
Nitrogen + Oxygen	Mass Fraction	0.895	0.895	0.896	0.896
Carbon dioxide	Mass Fraction	0.065	0.065	0.065	0.065
Water	Mass Fraction	0.040	0.040	0.039	0.039
Temperature	K	313.1	313.1	312.8	312.8
Total flowrate	kg/s	0.995	0.995	0.9951	0.9951
Absorber					
Absorber top pressure	bara	1.160	1.160	1.160	1.160
Absorber bottom pressure	bara	unavailable	1.224	unavailable	1.221
Absorber outlet CO ₂ concentration	Mass Fraction	0.0053	0.0056	0.004	0.005
Lean solvent - absorber inlet temperature	K	304.1	304.1	325.4	325.4
Rich solvent - absorber outlet temperature	K	318.1	316.5	318.7	316.1
Lean solvent flowrate	kg/s	0.857	0.857	0.756	0.756
Desorber (regenerator)					
Desorber Bottom Temperature	K	388.6	388.6	395.4	395
Desorber Top Pressure	bara	1.701	1.708	2.031	2.03
CO ₂ stream	kg/s	0.0578	0.0611	0.060	0.0605
Reboiler Steam					
Steam pressure	bar	3.606	3.605	4.075	4.075
Steam temperature	K	402.8	402.8	408.5	408.5
Steam condensate Temperature	K	401.9	401.9	407.8	407.5
Steam flowrate	kg/s	0.091	0.078	0.086	0.0823
Lean-Rich Heat Exchanger Temperatures					
Lean solvent in	K	387.4	388.6	394.2	395.0
Lean solvent out	K	325.0	322.6	324.7	327.4
Rich solvent in	K	318.9	316.5	319.9	316.1
Rich solvent out	K	380.4	379.9	384.0	379.9
Lean solvent concentration					
GCCmax Solvent	Mass Fraction	0.410	0.410	0.439	0.439
Water	Mass Fraction	0.536	0.541	0.504	0.515
CO ₂	Mass Fraction	0.054	0.049	0.057	0.046
General specifications					
CO ₂ Capture Efficiency	%	89.1	91.7	92.50	92.13
Inter-stage Cooling		No	No	No	No

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299 The validated pilot plant model was used to extract several technical key process indicators (KPIs) which are
300 important measures that quantify the difficulties associated with CO₂ separation from the flue gas in terms of
301 the required heating and cooling duties, required packing, and solvent circulation. These measures are scaled
302 with respect to the amount of pure CO₂ captured, to become independent of the pilot plant throughput and
303 enable comparisons. Table 4 provides the comparisons between the solvents. This table reveals that compared
304 to the MEA reference solvent, the GCCmax solvent requires 25.4-29.4% less steam for solvent regeneration in
305 the reboiler, per ton of CO₂. In addition, compared to the MEA reference solvent, the GCCmax solvent requires
306 73-84.4% less cooling water in the condenser and intercooler. The volume of packing in the absorber and
307 desorber columns also shows improvements in the case of the GCCmax solvent. Finally, the GCCmax solvent
308 circulation is 58.8-64.7% less than MEA solvent circulation which implies significant reductions in the electricity
309 power needed for solvent pumping.

310 **Table 4. Key process indicators (KPIs) for the GCCmax solvent and baseline MEA solvent**

Key Process Indicators (KPIs)	Unit	MEA	GCCmax (Data1)	GCCmax (Data2)
Heating duty	(MJ/ton CO ₂)	3986	2813	2975
Cooling duty	(MJ/ton CO ₂)	5644	1524	884
Volume of packing	(m ³ /ton CO ₂ hr ⁻¹)	46.619	45.64	45.91
Solvent circulation flow rate	(ton solvent/ton CO ₂)	34	14	12

311

312 **4. Solvent performance optimization**

313 The last part of the present research was concerned with model-based optimization of the carbon capture
314 process. Having validated the process model using pilot plant data, we wished to explore the improvement
315 opportunities and the significance of various investment and operational decisions. Three scenarios were
316 studied:

317 In Scenario (1), the dimensions of process equipment and operating conditions were optimized
318 simultaneously. This scenario represents the case where the plant is built for the first time.

319 In Scenario (2), the dimensions of process equipment are fixed and only the operating conditions are
320 optimized. This study is concerned with the scenario where an existing process is adapted to a new
321 solvent.

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322 In Scenario (3), similar to the second scenario, the dimensions of process equipment are fixed. However,
323 operating conditions and the GCCmax solvent blend composition are optimized simultaneously. This
324 scenario highlights the implication of the solvent optimization for energy saving.

325 In the first scenario, the objective function was the Total Annualized Costs because:

$$326 \quad Total\ Annualized\ Costs = \frac{Fixed\ Capital\ investment}{Plant\ effective\ Life} + Total\ Annual\ Energy\ costs \quad (9)$$

327 However, since in the second and third scenarios the equipment dimensions are fixed, the first right-hand-side
328 term of the above equation is constant and only Total Annual Energy Costs are optimized. In equation (9), the
329 value of 5 years were considered for the plant life cycle, in order to incorporate interest rate and the time
330 value of money. The costs of process equipment were calculated according to the costing correlations
331 provided in (Couper et al., 2012). Lang factor of 6 was considered for estimating the total capital investment
332 (Peters et al., 2004). The considered utility costs were 37.02 \$/MWh for electricity (Electricity Wholesale
333 Market 2015), 0.048 \$/tonne for cooling water (Ulrich and Vasudevan, 2006), and 14.5 \$/tonne for steam. Two
334 additional constraints were considered:

$$335 \quad Carbon\ capture = 90\% \quad (10)$$

$$336 \quad Reboiler\ temperature < 120\ ^\circ C \quad (11)$$

337 The first constraint ensures that 90% CO₂ is captured from the flue gas. The second constraint limits the
338 maximum reboiler temperature in order to avoid solvent degradation. Table 5 presents the results of
339 optimization. A comparison between Scenario (1) in which process equipment was being optimized and the
340 other two scenarios, suggests that the size of the pilot-plant absorber and desorber columns was larger than
341 the required values. This is an intuitive observation, because pilot plants are often to an extent oversized
342 in order to allow a variety of solvent testing. In the first scenario, the heating and cooling duties are slightly
343 higher than the other two scenarios, as they were optimized in conjunction with capital investment. In the
344 other two scenarios, the process equipment items were fixed and the optimizer took the opportunity to
345 exploit process equipment for energy saving. It is notable that there is a significant energy saving when the
346 solvent composition and operating conditions were optimized simultaneously, which illustrates the
347 importance of solvent control and optimization during process operation.

348

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Table 5. Optimization results

		Scenario (1)	Scenario (2)	Scenario (3)
Optimization objective function		Total annualized costs	Annualized energy costs	Annualized energy costs
Optimization variables		<i>Equipment size and operating conditions</i>	<i>Operating conditions only</i>	<i>Operating conditions and solvent composition</i>
Flue gas to absorber				
H ₂ O	mass fraction	0.0388	0.0388	0.0388
CO ₂	mass fraction	0.0651	0.0651	0.0651
N ₂	mass fraction	0.7320	0.7320	0.7320
O ₂	mass fraction	0.1640	0.1640	0.1640
Pressure	Pa	1.22E+05	1.22E+05	1.22E+05
Temperature	K	312.8	312.8	312.8
Flowrate	kg/s	0.995	0.995	0.995
Absorber diameter	m	0.5	0.66	0.66
Absorber height	m	9	18.3	18.3
Lean Solvent to absorber				
Flowrate	kg/s	0.787	0.874	0.739
Temperature	K	324.2	319.9	320.2
Composition				
H ₂ O	mass fraction	0.5390	0.5390	0.5470
CO ₂	mass fraction	0.0517	0.0517	0.0444
GCCmax	mass fraction	0.4093	0.4093	0.4086*
Reboiler temperature	K	398	398	398
Reboiler steam pressure	Pa	4.07E+05	4.07E+05	4.07E+05
Reboiler steam temperature	K	408.5	408.5	408.5
Reboiler Steam flowrate	kg/s	8.325E-2	7.22E-02	6.282E-2
Lean-Rich heat exchanger				
Lean in	K	398	398.0	398.0
Lean out	K	325.3	321.0	321.2
Rich in	K	317.6	316.3	317.2
Rich out	K	386.7	390.0	390.0
Condenser temperature	K	301.3	301.3	301.5
Desorber condenser duty	W	58909.5	49195.7	43096.1
Lean solvent cooler duty	W	2152.89	2374.1	1936.87
The total cooling duty	W	61062.39	51569.8	45032.97
The heating (Reboiler) duty	W	176845	152554	132781
CO₂ Capture		90%	90%	90%

350

351

* The mass fractions of the ingredients of the GCCmax solvent were optimized, with a variation of over 50% in some mass fractions.

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352 **5. Conclusions**

353 This paper presented integrated modelling and pilot plant studies in order to benchmark alternative carbon
354 capture solvents, along with key performance indicators (KPIs) to support comparison. The proposed
355 methodology was used to study the technical and energetic performances of GCCmax, a new amine-promoted
356 buffer salt (APBS) solvent, developed by Carbon Clean Solutions Limited (CCSL). In order to obtain meaningful
357 results, a high fidelity model was developed based on rate-based modelling of absorption and desorption
358 columns, in addition to modelling thermodynamic and reaction phenomena based on statistical associating
359 fluid theory (SAFT). The developed model was validated using pilot plant data, with very good agreements
360 between predicted and reconciled data. Detailed comparisons were made based on several key process
361 indicators which quantify the difficulties associated with CO₂ capture and include: the required heating duty,
362 cooling duties, volume of column packing and the solvent circulation rate. The comparisons between key
363 process indicators (KPIs) established the superior performance of the new GCCmax solvent in comparison with
364 the MEA baseline solvent. The last part of this paper explored the opportunities for further improvement of
365 the solvent performance via optimization. The studied scenarios were comprehensive and included (1)
366 designing a new process, (2) retrofitting an existing process using a new solvent and (3) optimizing the solvent
367 composition and the operating conditions of an existing process. While the presented methods are applied for
368 the new GCCmax solvent and the MEA reference solvent, the research methodology is general in nature and
369 can provide an effective standard for solvent comparisons.

370 The present research illustrated the application modelling and pilot plant studies for optimization of the
371 solvent performance. In the future studies, we will apply the developed model for scale up and power plant
372 integration based on the methodology of integrated process design and control (Sharifzadeh 2013a) using
373 inversely controlled process models (Sharifzadeh 2013b; Sharifzadeh and Thornhill 2012, 2013).

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