Solvent selection and design for CO$_2$ capture - how we might have been missing the point

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Carbon Capture and Sequestration (CCS) is a vital technology for the cost-effective mitigation of anthropogenic CO$_2$ emissions. However, a key obstacle to its deployment at large scale remains its cost - both capital and operating costs. In this context, the development of improved sorbents is a key research priority. Consequently, there is a vast global effort to develop new materials for this purpose, with literally thousands of new materials having been proposed since the beginning of the millennium. One common element of these contributions is that they focus on the equilibrium capacity of the material to absorb CO$_2$ and rarely, if ever, other key factors such as transport properties. To date, the majority of this effort has cost significant amounts of time and resources and has almost exclusively focused on developing sorbent with either increased CO$_2$ capacity and/or reduced heat of regeneration. Given that sorbent regeneration largely dictates operational cost, this would, on the surface, appear rational. However, it is vital to recall that the cost structure of $/MWh of electricity generated is composed of contributions to both capital and operational cost. Consequently, this single-minded focus on equilibrium CO$_2$ capacity and heat of regeneration excludes the contribution of transport and kinetic properties which determine equipment size and thus capital cost. Therefore, in order to develop sorbents which will result in a non-negligible cost reduction, it is essential to move beyond equilibrium-based metrics of sorbent performance. In this paper, we present a new methodological approach for sorbent screening which explicitly includes rate-based phenomena. Our approach uses both monetised and non-monetised performance indicators. Our results suggest that whilst equilibrium CO$_2$ capacity is a key determinant of process performance, transport properties (e.g., viscosity) and other thermophysical properties (e.g., heat capacity) have a significant effect on the capital cost, and thus on the price of the carbon captured. The key contribution of this work is the identification of the minimum set of thermophysical and kinetic parameters which must be reported in order to justify the claim of adequacy for a new sorbent for CO$_2$ capture in particular and gas separations in general.

Broader context

Separation processes represent a large portion of industrial activity, accounting for 40% to 70% of total CAPEX and OPEX, and, in particular, for 22% of all industrial energy use. Reducing the cost and improving the efficiency of separation units is therefore vital to improving the performance of a wide range of industries. In terms of mitigating climate change, carbon capture and storage (CCS) is currently the most economically viable option to decarbonise the energy sector in a carbon-constrained world. However, the large scale deployment of this technology is generally considered to be hampered by the significant cost associated with this technology. With sorption-based processes being the most mature, the capital and operating cost of these processes is dictated by the thermophysical properties of the sorbent selected. Therefore, understanding the relation between the full set of solvent characteristics and the cost of the process is essential for a more cost effective implementation of CCS in particular and the performance improvement of separation processes throughout the industrial sector in general.

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1 Introduction

In 2014, the Intergovernmental Panel on Climate Change (IPCC) found that the cost of limiting the concentration of CO₂ equivalents in the atmosphere to 430-480 ppm would be 138% higher without a large scale deployment of carbon capture and storage (CCS) in the 2-degree scenario (2DS). CCS technology is therefore key for the decarbonisation of the global economy. At the time of writing, 17 large demonstration plants are in operation, 5 projects are in executing phase and another 5 are projected to be commissioned by 2020. Most of the CCS projects in operation encompass a wide range of CO₂ sources, including natural gas processing (9 projects), hydrogen production (2 projects) and power generation (6 projects). Nonetheless, these plants will only capture a fraction of the total CO₂ that is required to meet the 2DS targets. The current focus on the so-called 1.5 °C target associated with the 2015 Paris COP only serves to increase this ambition.

However, a common theme of all scenarios is that the commercial scale deployment of CCS is significantly behind schedule, relative to what was anticipated in the first decade of this millennium. It is commonly held that the capital and operating cost associated with CCS is a key impediment to its deployment. Accounting for approximately 60 - 70% of the cost per tonne of CO₂ avoided, the capture step is the most costly and energy intensive step of the CCS chain. Consequently, there is an intense focus on reducing the cost of the capture process. Whilst the factors affecting the actual cost of a commercial-scale CCS project are many and varied, one key area where the research community can contribute is in the development of new materials and processes for the CO₂ capture. Chemical and physical absorption, adsorption, cryogenic separation, chemical and calcium looping, and membranes are the technologies which are under active investigation for carbon capture. Nonetheless, chemical absorption using aqueous solutions of alkanolamines is the most mature technology for CO₂ capture, and the only one to have been deployed commercially in power plants thus far.

The cost per tonne of CO₂ captured ($/tCO₂) is an aggregate of the equipment and plant construction, financing, fuel and other utilities cost. The size and operating cost of the process equipment is function of the thermophysical and chemical properties of the solvent. However, within the research community, there is an overwhelming focus on equilibrium properties which dictate the operating cost, to the exclusion of the properties which determine the size and thus capital cost of process equipment. A substantial volume of work is dedicated to increasing the gas carrying capacity (i.e., the CO₂ loading) or decreasing the enthalpy of absorption/desorption of the sorbent. This approach neglects the combined effects that the larger set of physicochemical, thermodynamic and transport properties of solvents have on the performance of the capture process, and therefore, on the cost. For example, the focus on reducing the enthalpy of absorption neglects the contribution of other properties such as the heat capacity and the enthalpy of vaporisation of the solvent to the thermal requirements in the regeneration step. For example, Oexmann and Kather argue that processes using solvents with higher heat of absorption can benefit from a higher pressure in the reboiler, resulting in lower power requirement in the compression stage prior to the CO₂ transport. Even in the event that a solvent reduces the energy per ton of CO₂ captured (GJ/tCO₂) of the capture process, it does not mean that it reduces the overall cost of capture ($/tCO₂), or more importantly, the cost of producing low carbon product, ($/MWh). A solvent with a low energy of regeneration could have inferior mass transfer characteristics, and thus could actually increase the $/MWh cost, despite having an improved GJ/tCO₂ figure. The size of the process units is directly linked not only to the thermodynamic but also to the transport properties of the solvent such as viscosity, thermal conductivity and diffusivity. The key question, therefore, is how these sorbent properties conspire to define a feasible parameter space which corresponds to a meaningful reduction in the total cost of carbon capture.

The answer to this question can only be obtain by a monetised process performance assessment. With this in mind, it is vital to identify the minimum set of properties which affect the cost of the carbon capture. Defining and optimising the operational envelop of those properties provides a space for evaluating the opportunities to design new solvents. It is equally important to have a reliable method for discriminating among candidate molecules using monetised and non-monetised key performance indicators (KPI’s). Whilst the use of non-economic KPI’s is useful for a preliminary sorbent screening effort, economic KPI’s will likely determine the final viability of a candidate molecule.

In this paper we present a new methodology to assess the process performance of sorbents for carbon capture. We have developed a software tool that evaluates a range of selected KPI’s based on the description of the reaction kinetics, thermodynamics, and transport properties of the solvent. Our tool includes a description of the chemical equilibria, mass and energy balances, solvent and energy requirements, and heat- and mass-transfer rates, as a function of the key operating parameters (KOPs). The novelty of this work is that our approach collapses the what is otherwise a very large variable space to a reduced number of selected KPI’s. Following this strategy, solvents can be evaluated given that their key properties have been fully determined or predicted. Our results clearly indicate that, in addition to CO₂ capacity, the other solvent properties which have a primary effect on the price of the carbon capture are viscosity and heat capacity.

The remainder of this paper is structured as follows: Section reviews the current state of the art of chemisorption for carbon capture. Section presents the methodological and modelling approach developed in this study and the process model. Section analyses the results on the effect on the selected indicators of

† The CO₂ capture plants from power generation in operation at the time of writing in 2017 are Boundary Dam in Canada, Petra Nova in United States. The capture method in both cases is chemical absorption using amine.

‡ 40 Mtpa will be captured with new the capacity installed in 2017. The capacity required by 2040 is 4000 Mtpa.
the characteristics of the solvent and the fluegas. Finally, Section 5 presents the conclusions of our work.

2 Chemical Solvents for carbon capture

Monoethanolamine (MEA) is a fast reacting alkanolamine, and was first proposed for gas separations by R. R. Bottoms in 1930,23 and is generally used as the benchmark against which all other CO₂ capture processes are compared, having been proven at both pilot and commercial scale. The reaction between the CO₂ and the MEA involves the formation of a carbamate salt.25 The stoichiometry of this reaction limits the loading capacity to 0.5 moles of CO₂ per mole of the amine.26 The viscosity of its aqueous solution is similar to that of water.27 However, it is a relatively volatile compound28 leading to partial vaporisation of the amine and subsequent release into the environment. Further, large amounts of energy are required to break the stable carbamate bond (∼85 kJ mol⁻¹),22,29,33 and desorb the CO₂ from the solvent. The total thermal energy requirements are estimated in the range of 3.5 to 4 GJ per tonne of CO₂, as a function of process configuration, and is composed of the contributions to the heat duty of regeneration; i) the energy required to break the chemical bond between the CO₂ and the amine, ii) the energy required to generate stripping steam in the reboiler and iii) the sensible heat required to raise the temperature to that of the regeneration.22,23

Secondary amines, e.g., diethanolamine (DEA), and tertiary amines, e.g., triethanolamine (TEA), and N-methyl-diethanolamine (MDEA), have also been investigated for carbon capture. Secondary and tertiary amines tend to have both a reduced rate of reaction and energy of regeneration. Sterically hindered compounds were first proposed by Sartori and Savage in 1983.33 At the time, they were proposed as a means to increase the carrying capacity of the solvent above that of standard primary alkanolamines. The primary driver for this was the relatively low stability of the carbamate salt, leading to commensurately greater concentrations of free amine in solution, thus moderating the otherwise limiting effects of steric hindrance on reaction rates. Versteeg et al. performed extensive experimental screening campaigns to determine the relationship between the structure and the CO₂ absorption,34,35 and desorption.36 They showed that the absorption capacity increases with the chain length between the amine and the hydroxyl functional group, in contrast to the rate of absorption, which it was found to decrease with the chain length. Puxty et al. screened seventy-six different amine based on their CO₂ absorption capacity.37 Their work included primary, secondary, and tertiary amines; alkanolamines; polyamines of a mixed or single type; cyclic and aromatic amines; amino acids; and sterically free and sterically hindered amines.37

Some of those amine which have increased CO₂ carrying capacity relative to that of MEA exhibit steric hindrance local to the amine functional group. Further, evaluation of other properties, e.g., absorption rate as a function of temperature, energy requirement of the process and their recyclability and stability, is required to determine the potential of the amines, as noted by the authors.37 Polymamines and diamines were also found to possess high CO₂ loading and recuperation capacity,38 but their process performance were not investigated further by the authors. Heterocyclic amines such as piperidine and piperazine have also been extensively assessed.37,39 Alternative classes of solvents continue to be proposed in the literature, including blended solutions,40,43 biphasic solvents,44,46 poly(ethylene glycol) ethers,47 ionic liquids,48,50 and deep eutectic solvents.49 However, despite this plethora of activity, very few of these solvents have made it "out of the lab". In fact, there are relatively few solvents which have been proposed for CO₂ capture and have been deployed at large scale. These include commercial solvents such as Shell's Cansolv technology claim that the process requires only 2.3 GJ/tCO₂.52 Moreover, Fluor’s Econamine,53 and MHI’s proprietary hindered amine KS-155,55 exhibit similar performance.

It is clear whilst there have been decades of extensive effort in solvent screening and property measurements with some attempts to develop links between solvent properties, or chemical structure, and process performance, this linking has always been relatively tenuous and indirect. The technologies that have thus far been commercially deployed represent only a very limited subset of the technologies under investigation. Given that it typically takes several decades for a technology to move "out of the lab", there is therefore an imperative to develop a technology screening approach that will allow technologies to "fail quickly", thus avoiding years of costly experimentation and prioritising effort on the most promising technologies. Addressing this challenge is the purpose of our current study.

3 Model Development

We have developed a software tool for the rapid screening of potential chemical solvents on the basis of their impact on process performance and cost, as opposed to solvent-specific metrics such as CO₂ loading or energy of regeneration. Using an archetypal chemisorption process topology, illustrated in Fig. 1, it estimates the size of the units and the energy requirements as a function of the thermochemical characteristics of the solvents. This includes the chemical equilibria description, the mass and energy balances, the solvent and energy requirements, etc. The model involves the computation of non-monetised and monetised process performance indices. The non-monetised indices include the degree of capture, the rate of recovery of the solvent, the height of the absorption column, the area of the heat exchangers and the heat and work requirements of the process. The monetised indices selected include the annualised capital expenditure (CAPEX), operating expenditure (OPEX), and total annual cost (TAC).

3.1 Absorption and desorption

The absorber and desorber columns are characterised by the rate of capture n₁, and lean loading of the solvent. The columns are represented as rate-based adiabatic packed columns. The model comprises the mass and energy balances, equilibrium relations, the rate equations required to describe mass transfer. Non-equilibrium rate-based models show a good agreement with pilot plant data,56,57 in contrast to the results obtained using equilibrium-based models.58 Since the rate-based approach was originally proposed,59 and refined,60,61 the trend in the literature as been to develop increasingly detailed models, with some
studies going as far as to include rigorous simulations of the reactive absorption in the liquid film, discretizing both the column and the diffusion film. However, owing to mathematical stiffness, such complex models would be unsuitable for use as a screening tool as is our current focus. Thus, in this study, we have developed a process model which includes a physically based description of heat and mass transfer, but is sufficiently computationally tractable for our purposes.

3.1.1 Mass rate transfer equations.

The ultimate goal of a carbon capture plant is to capture a given amount of CO₂ which is defined by the mass balance in the column:

\[ \nu_{i}^{in} \cdot y_{i}^{in} - n_{i}^{abs} = \nu_{i}^{out} \cdot y_{i}^{out} \]  

(1)

where \( \nu_{i}^{in} \) and \( \nu_{i}^{out} \) are the mole flowrate of inlet and outlet gas, \( y_{i}^{in} \) and \( y_{i}^{out} \) are the molar fraction of the component in the inlet and outlet gas respectively, and \( n_{i}^{abs} \) is rate of capture expressed as mole flowrate.

The mass transfer has been modelled following the two-film theory as reviewed for carbon capture by Wilcox et al. Non-reactive thermodynamic equilibrium is assumed to be attained at the gas-liquid interface. If absorption occurs at low pressure, it can be assumed that it obeys Henry’s law, so the molar fraction of the solute in the liquid interface with the physical solvent (\( x_{i}^{phys} \)), assuming ideal gas and liquid, would be calculated as:

\[ y_{i}^{L} = H_{i} x_{i}^{phys} \]  

(2)

where \( y_{i}^{L} \) is the molar fraction of the solute in the gas interface and \( H_{i} \) is the Henry’s coefficient. The mass transfer resistance is considered to occur in the vicinity of the gas-liquid interface or fluid film. The liquid mass transfer coefficient (\( k_{L} \)) have been estimated using the Onda’s correlations for randomly packed columns:

\[ k_{L} = \left( \frac{D_{L}}{HL_{S}} \right)^{1/3} = 0.0051 \left( \frac{R_{e}^{2/3} \left( a_{p} d_{p} \right)^{0.4}}{Sc_{L}^{0.5}} \right) \]  

(3)

where \( Sc_{L} \) is the Schumidt number for the liquid phase, and \( Re_{L} \) is the Reynolds number using the wet interfacial area (\( a' \)) calculated as a correction of the packing specific area (\( a_{p} \)) using Onda’s correlation:

\[ a' = a_{p} \left( 1 - \exp \left( -1.45 \left( \frac{\sigma}{\sigma_{c}} \right)^{0.75} R_{e}^{0.1} F_{r}^{0.05} W_{0}^{0.2} \right) \right) \]  

(4)

where \( \sigma \) is the surface tension of the liquid and \( \sigma_{c} \) is the critical surface tension of the packing. \( F_{r} \) and \( W_{0} \) are the Froude and Weber dimensionless numbers for the liquid phase.

The calculation of the rate of diffusion is ordinarily one of the most computationally intensive elements of an absorption model, typically employing a Maxwell-Stefan formulation. However, in the context of this study, such an approach would run counter to our aim of developing a high throughput screening tool. Thus we include a description of the limiting diffusivity, i.e., the diffusion of the unreacted base to the reaction plane. Therefore, the diffusivity coefficient has been estimated using:

\[ D_{L} = x_{B} D_{B,H,O}^{0} + x_{H,O} D_{H,O,B}^{0} \]  

(5)

where the diffusivity coefficients at infinite dilution were estimated with the Wilke-Chang correlation:

\[ D_{j, i}^{0} = 7.4 \times 10^{-8} \left( \frac{\phi M_{j}}{V_{i}} \right)^{1/2} \left( \frac{\mu_{i}}{V_{i}^{0.6}} \right) \]  

(6)

where \( M_{j} \) is the molecular weight of solvent \( j \), \( V_{i} \) is the molecular volume of the solute \( i \), and \( \phi \) is the association factor of the solvent. \( \phi \) is chosen as 2.6 for water and 2.26 for the base.

The enhancement factor \( E \) is defined as the ratio of the average rate of absorption into an agitated liquid in the presence of reaction to the average rate of absorption without reaction. Kenig et al. discretised the film region and demonstrated that the chemical reaction occurring at the film considerably enhances the mass transfer of CO₂ within the film region. In the cases where the reactions is instantaneous, the enhancement factor is only a function of the diffusion, and consequently, dividing the film in segments does not improve the result. Therefore, for those cases when the reaction is instantaneous, the enhancement factor can be estimated by:

\[ E = 1 + \frac{D_{B} c_{B}}{D_{L} c_{CO_{2}}} \]  

(7)

where \( D_{B} \) is the diffusivity coefficient of the binder component in the aqueous phase, \( c_{CO_{2}} \) is the molar concentration of CO₂ in the liquid interface, \( c_{B} \) is the molar concentration of the binder component, and \( D_{L} \) the diffusion coefficient of component \( i \) in the liquid phase. For a fast pseudo-first order reaction, the enhancement factor can be estimated as:

\[ E = \frac{\sqrt{D_{k} k_{2} c_{B}}}{k_{L}} \]  

(8)

where \( k_{2} \) is the pseudo-first order reaction constant.

Dimensions of the column. The total number of moles \( N_{CO_{2}} \) absorbed or desorbed in the process columns is calculated from the total contact area which is a function of the size of the tower and the type of packing:

\[ N_{CO_{2}} = J_{CO_{2}} \varepsilon a' A_{T} Z_{T} \]  

(9)
where $\varepsilon$ is the void fraction of the packing, $A_F$ is the cross-section area of the column and $Z_T$ the height of the absorber or desorber tower.

The molar flux of CO$_2$ in the interface $J_{CO_2}$ can be therefore calculated using:

$$J_{CO_2} = k^0_{ij} E \left(c^0_{CO_2} - c^B_{CO_2}\right)$$  \hspace{1cm} (10)

Considering that CO$_2$ reacts across the film, it is reasonable to assume that the concentration of molecular CO$_2$ in the bulk liquid is negligible.$^{[22]}$ The $0$ indicates that no chemical reaction is considered at the interface, and only physical absorption is occurring. For fast pseudo-first order, the size of the column can be estimated combining equations (8), (9) and (10) giving:

$$N_{CO_2} = \sqrt{D_L k_F c^0_{CO_2}} e_{d} A_T Z_T$$  \hspace{1cm} (11)

This approach reduces the numerical complexity of the absorption model in order to permit the rapid evaluation of the performance of a sorbent, but still capturing the influence of the reaction kinetics, the mass transfer across the gas-liquid interface, the thermodynamic equilibrium and the hydrodynamics of the packing.

**Energy balance equations.** The absorber has been modelled as an adiabatic column$^{[21,22]}$ where the gas and the liquid transfer heat through the absorption of CO$_2$ and the vaporisation of water:

$$\Delta H^I = \Delta H^L - \sum_j n_j \Delta h_j^{I,G}$$  \hspace{1cm} (12)

where $h_j^{I,G}$ is the specific enthalpy of phase change, e.g., the absorption or desorption of CO$_2$ and the heat of vaporisation of water, and $\Delta H^I$ and $\Delta H^L$ are the change of enthalpy in the liquid and vapour phase respectively, which are estimated using:

$$\Delta H^I = 0.5 \left( \sum_i \left( C_{p,G}^{in} + C_{p,G}^{out} \right) (T^I - T^{in}) \right)$$  \hspace{1cm} (13)

$$\Delta H^L = 0.5 \left( \sum_i \left( C_{p,L}^{in} + C_{p,L}^{out} \right) (T^L - T^{in}) \right)$$  \hspace{1cm} (14)

where $C_p$ is the heat capacity at the corresponding temperature $T$ of the inlet and outlet liquid and vapour streams.

Pilot plant experiments show that the temperature across the desorber height can be estimated as the average value between the temperatures of the lean solvent entering the absorber and the rich solvent leaving the absorber$^{[24]}$ and therefore, the stripper has been modelled as an isothermal system.$^{[25]}

### 3.2 Thermodynamic modelling.

The solubility of CO$_2$ in alkanolamines is calculated using the equilibrium model proposed by Gabrielsen et al.$^{[26]}$ where the gas is considered ideal, and the Henry’s coefficient is combined with the chemical equilibrium constant.

$$P_{NCO_2} = K_{CO_2} c_{CO_2} \left( \frac{\alpha_0 \theta}{(\alpha_0 (1 - \theta))} \right)$$  \hspace{1cm} (15)

where $P$ is the total pressure, $\alpha$ is the mole fraction of amine in the solution, $\theta$ is the CO$_2$ loading, $z$ is the ratio of amine per mole of CO$_2$ (e.g., $z = 2$ for MEA) and $K_{CO_2}$ is the combined Henry's law and chemical equilibrium. $K_{CO_2}$ is estimated as a function of the temperature, the amine concentration and the loading:

$$\ln K_{CO_2} = A + B \frac{1}{T} + C \alpha_0 \theta + D \sqrt{\alpha_0 \theta}$$  \hspace{1cm} (16)

where $A$, $B$, $C$, and $D$ are amine-specific parameters.$^{[25]}$ Following the approach described by Kale et al.$^{[27]}$ Henry’s law is used to describe the physical vapour-liquid equilibrium of CO$_2$ and N$_2$ in the interface. The Henry’s law constant of a solute in a blend can be estimated from the Henry's law constants of the solute in each of the pure solvents.$^{[27]}$ The Henry’s coefficient of reactive solvents is estimated using the so-called N$_2$O analogy, where dinitrogen monoxide (N$_2$O) is used as a non-reactive gas to estimate the properties of the gases in liquids such as diffusivity and solubility.$^{[28,29]}$ The extended Raoult’s law for MEA and water is applied to determine the vaporisation of both components into the gas stream using the activity coefficients reported by Park et al.$^{[30]}$

### 3.3 Heat Exchangers

The process model includes two heat exchangers without phase change, i.e., the lean-rich heat exchanger and the lean cooler, and two heat exchangers with phase change, i.e., the condenser and the reboiler. In all cases they have been modelled as shell-and-tube units, with the condenser being considered to be vertical condenser and the reboiler being modelled as a thermosyphon. The general equation for the heat transfer area is:

$$Q = U A_{HE} \Delta T_{LM}$$  \hspace{1cm} (17)

where $Q$ is the heat exchange rate, $A_{HE}$ is the area of heat exchange, and $\Delta T_{LM}$ is logarithmic mean temperature difference of the two streams exchanging heat. The overall heat transfer coefficient $U$ is the reciprocal of the sum of the different heat transfer resistances. For a tube heat exchanger is typically estimated as:

$$\frac{1}{U} = \frac{1}{h_i} + \frac{1}{h_o} + \frac{2}{h_d}$$  \hspace{1cm} (18)

where $h_i$, $h_o$, and $h_d$ are the heat transfer coefficient of the fluid inside the tube, the heat transfer coefficient of the fluid outside the tubes (in the shell) and the heat transfer coefficient of the dirt, respectively. $t$ is the thickness of the tube, and $k_w$ is the thermal conductivity of the heat exchanger wall.

**Single phase heat exchangers.** The heat rate in a heat exchanger where no phase change occurs is given by the sensible heat:

$$Q = m C_p \Delta T$$  \hspace{1cm} (19)

where $m$ is the mass flowrate of the stream, $\Delta T$ is the temperature difference between the inlet and the outlet of the stream, and $C_p$ is the arithmetic average heat capacity in the range $\Delta T$.

The fluid heat transfer coefficient value depend on the flow regimes, which is a function of the heat exchanger internals and arrangements and on the fluid properties. For turbulent flow, the heat transfer coefficient for the inside fluid has been estimated
via:

\[ Nu = CR e^{0.5} Pr^{0.14} \left( \frac{\mu}{\mu_w} \right) \]  

(20)

where \( C = 0.023 \) for non-viscous liquids and 0.027 for viscous solvents, \( Nu, Re \) and \( Pr \) are the dimensionless numbers of Nusselt, Reynolds and Prandtl respectively, which are calculated as:

\[ Nu = \frac{h d_t}{k} \]  

(21)

\[ Re = \frac{u d_t \rho}{\mu} \]  

(22)

\[ Pr = \frac{C_p \mu}{k} \]  

(23)

where \( d_t \) is the internal diameter of the tube, \( k \) is the thermal conductivity of the fluid, \( u \) is the linear velocity of the fluid, \( \rho \) is the density and \( \mu \) and \( \mu_w \) are the viscosity at the fluid temperature and at the wall temperature respectively.

The heat transfer coefficient of the shell side is calculated using the Kern’s method:

\[ Nu_b = j_R e^{0.5} Pr_b^{0.14} \left( \frac{\mu}{\mu_w} \right) \]  

(24)

where \( j_R \) is the shell-side factor. The value of \( j_R \) can be found elsewhere.

Two-phase heat exchangers. The heat rate in a phase change heat exchanger where is given by both the sensible heat and the latent heat:

\[ Q = m \Delta h \Delta T + \sum m_j \Delta h_{lj}^{j-8} \]  

(25)

where \( m_j \) is the mass of component \( j \) that changes phase, and \( \Delta h_{lj}^{j-8} \) is the specific enthalpy of phase change, e.g., the heat of vaporisation. The heat associated with the absorption or desorption of \( CO_2 \) might be included in this term if applicable. The condenser has been modelled as a horizontal exchanger with condensation in the shell. The heat transfer coefficient of the condensate (\( h_c \)) for a tube bundle is given by:

\[ h_c = 0.95 k_f \left[ \frac{p_L (p_l - \rho_v) k_f}{\mu_L T_v} \right]^{1/3} \]  

(26)

where \( g \) is the gravitational acceleration (9.81 m s\(^{-1}\)) and \( T_v \) is the condensate flow per unit length of tube. The condensate density (\( \rho_L \)), viscosity (\( \mu_L \)) and thermal conductivity (\( k_f \)) are evaluated at the evaluated the mean of the temperature of condensation and the tube-wall temperature.

The reboiler has been modelled as a thermosyphon using the model proposed by Chen for forced-convection boiling. This method considers that the heat transfer of the boiling stream is the sum of the contribution of convective and nucleate boiling. The convective boiling coefficient can be estimated using equation (20) corrected by a two-phase flow factor based on the Lockhart-Martinelli two-phase flow parameter. The nucleate boiling coefficient is estimated using the Forster and Zuber correlation.

### 3.4 Economic indices

A key objective of this work is to relate the cost of the carbon capture plant to the properties of the solvents. Our model uses the Total Annualised Cost (TAC) as the index against which to assess the economic performance of the plant:

\[ TAC = CRF \sum \text{CAPEX} + \sum \text{OPEX} \]  

(27)

where the \( CRF \) is the capital recovery factor and is given by:

\[ CRF = \frac{i(1 + i)^n}{(1 + i)^n - 1} \]  

(28)

which is a function of the discount rate \( i \) and the annuity period \( n \). A discount rate of 10% and an annuity period of 25 years have been assumed. The capital cost, CAPEX, for each unit has been estimated using correlations that link the cost to key properties of the unit, e.g., the area of a heat exchanger, or the dimensions of the column. For example, the installed cost of an absorption column and heat exchanger is given by equations (29) and (30) respectively.

\[ C_{abs} = 1.281[f_1 C_b + V_p C_p + C_{ps}] C_t \]  

(29)

where \( f_1 \) refers to the material of construction and is assigned a value of 2.1 for SS316 and \( C_b \) is a function of the weight of the shell, given by:

\[ C_b = 1.218 \exp(6.629 + 0.1826(\ln W) + 0.02297(\ln W)^2) \]  

(30)

where \( W \) is the weight of the shell. In this work, a thickness of 2 mm was assumed. The product \( V_p C_p \) corresponds to the cost of column internals where \( V_p \) is the volume of packing required and \( C_p \) is the specific cost of the packing used. \( V_p \) is directly calculated via the column sizing equations and \( C_p \) is assigned a value of $76.6/ft\(^3\), representative of the packing used in this study. Finally, \( C_{ps} \) is given by:

\[ C_{ps} = 3000 \left( \frac{0.7396}{d_f} \right)^{0.7068} \]  

(31)

where \( D \) and \( Z_T \) are the column diameter and tan-to-tan height, respectively.

In the case of heat exchangers,

\[ C_{HX} = 1.218(f_d f_m f_p C_b) C_t \]  

(32)

where \( f_d, f_m, \) and \( f_p \) are functions of the heat exchanger type, pressure range and material of construction, respectively. \( C_b \) is an explicit function of the heat transfer area, \( A \), and is given by:

\[ C_b = \exp[8.821 - 0.30863(\ln A) + 0.0681(\ln A)^2] \]  

(33)

In both equations (29) and (32) \( C_t \) is a cost multiplier to account for the cost of installation and is assigned the value 2.1 and 1.9 for stainless steel absorption columns and heat exchangers, respectively.
respectively. More details on these equations and their derivation can be found in Couper.  

Thus, it is possible to make a direct connection with solvent thermophysical or kinetic properties and system cost. For example, the viscosity of the solvent influences the diffusivity (equations 5 and 6) which, in turn influences the flux (equation 10) and the height of the column (equation 11). Finally, this dictates the installed cost of the absorber (equation 29) and thus the TAC of the entire system (equation 27). A thorough evaluation of the sensitivity of key process design variables and TAC to solvent properties is presented in Section 4.

The cost derived from the electricity consumption and from the heat requirements have been estimated separately. The short-run marginal cost (SRMC) of the electricity has been used assuming a price of coal of $50/t CO₂ and the price per tonne of CO₂ emitted is $70/tCO₂. The cost of steam has been estimated using the SRMC assuming that the efficient of the boiler is 90%. The annualised costs, i.e., annual CAPEX, OPEX and TAC, are then divided by the tonnes of CO₂ captured in one year. The costs of compression of the CO₂ for transport and storage have not been included in this work. A previous study shows that the cost of CO₂ compression and dehydration is on the order of $20 - 25/tCO₂.  

4 Results and Discussion

This section has been structured as follows: we first investigate the effect of a range of thermo-physical and chemical properties of the solvents on selected monetised and non-monetised indices. The minimum set of properties required for solvent evaluation using our methodology are: thermodynamic equilibria of CO₂, reaction constant(s), viscosity, density, heat capacity, thermal conductivity and surface tension. For that, we evaluate a carbon capture plant treating 900 kg s⁻¹ of flue gas containing 12% (v/v) of CO₂ from a 800 MW supercritical pulverised coal power plant (ScPC). We assume that the plant runs for 335 days per year at 100% capacity. The capture rate is set to 90% of the carbon emitted by the power plant. We aim to identify and rank those thermophysical properties that have the most prominent impact on the cost of the capture plant. Then, we explore the trade-offs that those properties have on the monetised indices via a comprehensive sensitivity analysis.

Next, we examine in more detail the CO₂ transport from the gas phase into the liquid in the absorber by analysing the effect of thermodynamics, reaction kinetics and transport properties on the molar flux across the interface. Further, we study the effect of the concentration of CO₂ in the gas phase by analysing how the flux is affected when the flue gas treated comes from a ScPC plant and from a combined cycle gas turbine (CCGT). The concentration of CO₂ in the CCGT is assumed to be 4% (v/v). Finally, we compare the process performance of the carbon plant dedicated to treat the flue gas from that of a ScPC with that of CCGT considering the same net power output for both power plants.

We have selected 30 wt% MEA as the benchmark solvent. Table 1 shows its properties at 303 K. Our model assumes that the coefficient between the new value and that of the benchmark is constant across the temperature range studied. This is, if the property doubles at 303 K with respect the benchmark, it also doubles at 393 K. We have fixed the rich loading to 0.47 and the lean loading to 0.3. The results show that the TAC of the capture process is $51/tCO₂, which gives a total cost of the CO₂ capture of $73/tCO₂ once the costs of CO₂ compression and dehydration are accounted. This is in line with current projections of the cost of carbon capture at industrial scale for Petra Nova Carbon Capture Project, whose cost has been estimated to be $70/tCO₂.

| Property                        | Value  |
|--------------------------------|--------|
| Equilibrium constant (kPa)     | 0.015  |
| Henry’s coefficient (kPa m⁻³ kmol⁻¹) | 4560   |
| Reaction constant (m³ kmol⁻¹ s⁻¹) | 8008   |
| Density (kg m⁻³)               | 1168   |
| Viscosity (mPa s)              | 2.51   |
| Heat capacity (kJ kmol⁻¹ s⁻¹)   | 89.7   |
| Thermal conductivity (kJ m⁻³ s⁻¹ K⁻¹) | 4.75·10⁻⁴ |

4.1 Evaluation of the thermophysical properties of solvents

Using our modelling tool, we have identified the following properties that can affect the process performance of the carbon capture: density, viscosity, surface tension, heat capacity, thermal conductivity, heat of absorption which might inherently include the heat of reaction in case of reactant solvents, and CO₂ solubility, which might be expressed in terms of loading, Henry’s coefficient or a generic equilibrium constant. A sensitivity analysis has been carried out to determine the aftermaths of varying each of the aforementioned properties on the annualised CAPEX, OPEX and TAC of the carbon capture plant. Every property has been screened individually by varying its value within sensible ranges that were selected based on typical values of solvents.

Equilibrium constant. The equilibrium constant of the CO₂-amine-H₂O system is the main thermodynamic property of the system. It is a non-linear function of the temperature, the partial pressure of CO₂ the concentration species in the mixture and the Henry’s coefficient of CO₂ in the liquid phase. The latter is related with the theoretical physical solubility in the solvent if no reaction would take place between the CO₂ and the solvent components. Note that the equilibrium constant and the Henry’s coefficient are inverse functions of the gas solubility. For decades, the search of a more effective solvent for carbon capture has been largely driven by improving the value of the equilibrium constant. A more CO₂-philic solvent reduces the solvent flowrate, the size of the units and the heat requirements which are related to the sensible heat. Fig. 2 illustrates the increase in size of the absorption column when the equilibrium constant (see eq. 16) increases, i.e., the solubility of CO₂ decreases. The cost associated to the change of the equilibrium constant is shown in Fig. 3.

At higher equilibrium constant, the CO₂ has a lower tendency to stay in the liquid phase. This has an opposite effect in the CAPEX and in the OPEX. The size of the process units increases as shown in Fig. 2 increasing the capital cost. However, a lower solubility, i.e., a weaker solvent, facilitates the recovery of the solvent, and therefore reducing the heat requirements in the reboiler. This is in accordance with the van’t Hoff equation, which can be derived.
from the Gibbs-Helmholtz equation, where the enthalpy of solution of a gas in a solute is related to the equilibrium constant. An increase in the CO\textsubscript{2} capacity, i.e., lower equilibrium constant, results in larger enthalpies of solutions for strong chemical solvents, as discussed in the work of Mathias and O’Connell.93

**Viscosity.** The impact of the viscosity of the solvent on the absorber height is illustrated in Fig. 4 and on the annualised CAPEX, the OPEX and the TAC is shown in Fig. 5. For reference, the viscosity of a 30 wt% MEA solvent at 313 K is 2.5 mPa·s.27 If a solvent presents a viscosity of 16 mPa·s, the total cost of the equipment is estimated to be more than double that of the benchmark plant. At this viscosity, the height of the absorber required is 133 m, as seen in Fig. 4 compared to the 50 m absorber required for the benchmark viscosity. The viscosity has a significant impact on the hydrodynamics of the absorber, particularly on the hold-up.93 The mass transfer of the gas in the liquid phase is hindered at higher viscosities. Therefore, longer contact times are required to achieve the same degree of separation, resulting in taller towers. Furthermore, heat exchangers are greatly affected by the viscosity, with a larger impact on those units which work at lower temperatures, e.g., the lean cooler, wherein even moderate viscosities result in low Reynolds numbers. At higher viscosities, the flow in the tubes develops from turbulent to laminar regime, changing radically the hydrodynamics and the heat transfer performance. In the case of turbulent regime, the fluid flow obviously advances in the axial direction across the tube, but additionally, it is constantly undergoing multidirectional swirling and mixing. As a consequence, the liquid surface in contact with the tube walls is continuously renewed. The convective contribution to the heat transfer prevails over the conductive contribution, enhancing the heat transfer in these systems. In contrast, in the case of laminar flow, the fluid only flows in the axial direction in parallel layers which do not mix. Thus, conduction is the leading mechanism for heat transfer. The effect of the regime transition from turbulent to laminar on the cost of the process is illustrated in Fig. 5. The discontinuities in the costs correspond to the viscosities at which the flow regime experience this transition in the cooler (18 mPa·s) and the heat exchanger (36 mPa·s). The annualised CAPEX increases from $48/t\textsubscript{CO}_2$ to $54/t\textsubscript{CO}_2$ as a result of the transition from turbulent to laminar flow in the cooler, i.e., the CAPEX increases by a 12.5%.

Higher values of viscosity increases pump power as a result of
changes in the hydraulic behaviour of the solvent associated with higher friction factor and pressure drop, thus increasing the operational challenges. Yang et al. reported that a mixture of aqueous MEA with an ionic liquid blocked the pipelines and their pumps stopped operating when they used a mixture with a viscosity of 16 mPa·s. However, the cost related to the electric requirements represents 5% of the total OPEX of the process. Thus, an increase in plant electricity requirements does not correspond to a significant increase in process costs. The results show that the cost of the electric requirements is $1/tCO₂ at the viscosity of the benchmark, whereas at 40 mPa·s the cost is $2.7/tCO₂ relative to an operating cost of $56/tCO₂ at this viscosity; a negligible amount.

**Heat Capacity.** The heat capacity is the only property that has a pronounced effect on both the CAPEX and the OPEX as illustrated in Fig. 6. As a reference, the heat capacity of 30% MEA in water at 303 K is 89 kJ mol⁻¹ K⁻¹ or 3.6 kJ kg⁻¹ K⁻¹. The increase on the TAC with the heat capacity is obvious. Higher values of heat capacity of the solvent result in higher heating and cooling requirements, i.e., in the OPEX. As a result, larger contact areas are required in the heat exchangers, increasing the CAPEX. Conversely, the overall heat transfer coefficient increases with the heat capacity. However, this increase is minor, and its contribution towards the reduction of the CAPEX in negligible compared to the increase driven by a larger area. Less evident is, though, the increase of CAPEX at low values of heat capacity. At lower heat capacity, the temperature of the solvent further increases as it moves downward the absorber given the same enthalpy of reaction. The capacity of the solvent to absorb CO₂ declines with temperature. Therefore, the rich loading of the solvent decreases, requiring more solvent to capture the same amount of CO₂. This leads ultimately to larger units, and consequently to higher CAPEX.

**Density.** Density has a limited impact on the CAPEX and minor effect on the OPEX, as shown in Fig. 7. The fact that it is the only property that decreases the cost as it increases is noteworthy. For example, if there was a solvent with a density double that of water, it would reduce the units cost by 20%.

**Thermal conductivity and surface tension.** We found that varying the thermal conductivity and the surface tension of the solvents have a minor effect on the overall cost of the carbon capture plant compared to the effect of the properties investigated above. Having said that, the surface tension is expected to play a more important role in the case of non-reactive solvents. In conclusion, viscosity and heat capacity are the primary thermo-physical properties that together with the equilibrium constant impact the mass transfer and the energy transfer. The feasible parameter space which corresponds to a reduction in the cost of carbon capture with respect to the benchmark is identified in Fig. 8. It is clear that any potential solvent for carbon capture should not only improve the CO₂ carrying capacity, but should also present a reduced value of viscosity and heat capacity and an increased density with respect to those of the benchmark. Nevertheless, these properties cannot be tuned independently from each other as they are a function of the chemical structure and its functional properties and the cost of the capture plant using the benchmark solvent, i.e., 30% MEA.

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**Fig. 6** Effect of heat capacity of the solvent on the annualised CAPEX, the OPEX and the TAC of the carbon capture plant.

**Fig. 7** Effect of the density of the solvent on the annualised CAPEX, the OPEX and the TAC of the carbon capture plant.

**Fig. 8** Relative effect on the TAC as a function of the relative density, viscosity, heat capacity and equilibrium constant with respect the same properties and the cost of the capture plant using the benchmark solvent, i.e., 30% MEA.
groups. Therefore, any molecular design of prospective candidate molecules should take this into account. Functionalising the molecular structure intending to improve the CO\textsubscript{2} carrying capacity might counteract the aim of reducing the cost of carbon capture. For example, increased functionalization (increasing carrying capacity) might also increase viscosity.

In the next sections we explore the trade-offs between the viscosity, the equilibrium constant and the heat capacity. Frequently, the effort to design solvents with a higher CO\textsubscript{2} leads to chemical structures that contains atoms or functional groups that not only interact more efficiently with the CO\textsubscript{2} but also present intra- or intermolecular interactions, increasing the viscosity of the system. Fig. 9 shows the effect of both properties on the monetised indices of the process. The effect of the viscosity in the flow regime inside the heat exchanger discussed above (see Fig. 5) is captured again in Fig. 9.

The discontinuity of the evolution of the CAPEX with the viscosity correspond to the points where the flux transitions from turbulent to laminar in the heat exchangers. Interestingly, the heat capacity can displace the laminar regime front towards lower viscosities, as shown in Fig. 10. At the heat capacity of the benchmark, the first transit to laminar flow in the cooler occurs at viscosities 7 times higher than the viscosity of the benchmark. When the heat capacity doubles, the laminar front is found at 4.5 times the viscosity of the benchmark. This is an important observation as the heat capacity is usually overlooked in the screening, selection and design of solvents for carbon capture.

4.2 The competition between equilibrium, transport and kinetic properties.

The absorption column might be considered the core of the capture process. Thus, solvent selection is particularly important here. A solvent with high CO\textsubscript{2} capacity per unit volume will serve to reduce the amount of solvent required, reducing both the CAPEX and OPEX. Moreover, the foregoing results emphasise the importance of maximising rates of mass transfer in order to reduce the size and capital cost associated with absorption column. Gas-liquid interphase mass transfer is a complex function of simultaneous transport and thermodynamic driven phenomena occurring across and at both sides of the interface. This includes the transport of the CO\textsubscript{2} through the gas-liquid interface, and then its diffusion entangled with the chemical reaction towards the bulk liquid. Understanding the contribution of the different parameters that determine the rate at which CO\textsubscript{2} is transferred from the gas to the liquid phase is imperative for the design of new solvents. Fig. 11 shows the contrasting impact of diffusivity ($D$) and reaction constant ($k_2$) for five different archetypal solvents with Henry's coefficients of 1000, 3000, 5000, 8000 and 10000 kPa-m\textsuperscript{3} kmol\textsuperscript{-1}, 4 mol\% (left) and 4 mol\% (right) respectively. Note than by definition, Henry’s coefficient and gas solubility are inversely related. Solvents with particularly high CO\textsubscript{2} solubility are represented by $H = 10000$ kPa-m\textsuperscript{3} kmol\textsuperscript{-1}, whereas low CO\textsubscript{2} solubility solvents usually present Henry’s coefficient values higher than 10\textsuperscript{4} kPa-m\textsuperscript{3} kmol\textsuperscript{-1}. In the case of the high gas-phase CO\textsubscript{2} concentration, it is evident that liquid phase diffusivity is the rate limiting step in the absorption of CO\textsubscript{2}, with reaction kinetics playing a secondary role.
This is in line with previous results and reinforces the typical approximation of treating chemical reactions in these systems as equilibrium, as opposed to using a full reaction kinetic description. However, if a solvent has a lower CO$_2$ solubility, then the influence of diffusivity and reaction rate are of the same order. To illustrate this, let us perform the following thought experiment with two solvents, solvent A with a Henry’s coefficient of 3000 kPa·m$^3$·kmol$^{-1}$, and a solvent B with a significant higher CO$_2$ solubility exemplified in Fig. 10 by the Henry’s coefficient of 1000 kPa·m$^3$·kmol$^{-1}$. Let us assume that solvent A presents a high diffusivity, e.g., $10^{-8}$ m$^2$·s$^{-1}$, and solvent B, which presents a markedly higher CO$_2$ solubility, has a hindered diffusivity coefficient below $10^{-9}$ m$^2$·s$^{-1}$. Solvent A outperforms the CO$_2$ flux of solvent B by a 154% despite the lower CO$_2$ capacity because of its favoured transport properties.

**Effect of the CO$_2$ concentration in the flue gas.** Thus far, we have considered the performance of solvents in the context of an exhaust gas with a relatively high CO$_2$ concentration, e.g., a coal fired power plant. However, it is increasingly recognised that it will also be necessary to capture CO$_2$ from dilute sources, such as the exhausts arising from gas-fired power stations. Therefore, in this section, we investigate how a low concentration exhaust gas, representative of that of a combined cycle gas turbine (CCGT), will affect the design of the carbon capture plant. Natural gas has a significantly higher hydrogen/carbon ratio, and thus the energy density of natural gas is higher than coal. Consequently, the carbon intensity of CCGT is significantly lower. The emission from coal range between 0.825 and 1.035 tons of CO$_2$ per MWh, whereas emissions from natural gas are in the range of 0.35-0.4 tons of CO$_2$ per MWh.

The typical flue gas from a coal plant contains 10-15% (v/v) of CO$_2$, whereas the content of CO$_2$ of the flue gas from a gas plant is 3-5% (v/v). The impact that this much more dilute exhaust gas has on solvent performance is illustrated in Fig 10b. In this context, the dominance of Henry’s constant and liquid phase diffusivity on the rate of CO$_2$ absorption is effectively eliminated. Here, each of CO$_2$ solubility, diffusivity and chemical kinetics play approximately equivalent roles. This implies that retaining the kinetics in sorbent screening is important and that treating these reactions as equilibrium is likely to be a poor approximation.

### 5 Conclusions

We have presented a process performance indexed methodology to assess potential of chemical solvents for carbon capture. Monetised (e.g., CAPEX, OPEX and TAC) and non-monetised (e.g., equipment size) KPI’s are evaluated as a function of selected KOP’s of the process and the characteristics of the solvent. The KOP’s include the solvent flowrate as a function of the exhaust gas composition, the rate of absorption and the SRMC of the steam available. The characteristics of the solvent are the kinetics of the reaction, the thermodynamic, and physico-chemical properties and transport properties of the solvent, i.e., reaction constant, equilibrium constant, heat of reaction, molecular weight, density, heat capacity, viscosity, thermal conductivity and surface tension of the solvent. Historically, most effort on solvent development has focused on enhancing the CO$_2$ absorption and reducing the heat of reaction, neglecting the effect of the other properties on the process cost, particularly on the capital cost contribution to the overall cost of CO$_2$ capture ($$/t_{CO_2}$). The properties that have a primary importance on the TAC of CO$_2$ capture are, in order of prominence:

1. Viscosity
2. Equilibrium loading of CO$_2$
3. Reaction kinetics
4. Heat capacity
5. Heat of absorption
6. Density
7. Surface tension

It must be clearly recognised that of these, the first three properties are significantly more important than the other four. The fact that solvent viscosity is found to be of primary importance confirms our hypothesis that, in order to rationally screen, select and design sorbents for chemical processes in general, and CO$_2$ capture in particular, a whole systems approach is necessary. Otherwise, it is impossible to fully appreciate the impact of solvent thermophysical properties on process performance and therefore
costs. The absence of such an approach to date has led to an over emphasis of CO₂ solubility in solvent selection and design, potentially limiting progress in this area. One of the main drawbacks to a "designer solvent" approach is the impact of solvent functionalisation on physical properties, or the unintended consequences that may accompany a sole emphasis on CO₂ alisation on physical properties, or the unintended consequences to a "designer solvent" approach is the impact of solvent functionally limiting progress in this area. One of the main drawbacks is normally higher, our model identifies this as a minor impact, and is likely offset by the inevitable increase in heat of absorption and therefore regeneration energy. Our model clearly demonstrates that this creates an ever greater pressure on increased solubility to offset the reduced mass transport, thereby leading to a "death spiral" of offsetting improvements in one property at the expense of another. Only a holistic approach to system design, with all solvent properties properly weighted, can lead to an optimal (or possibly even improved) solvent design.

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