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Effect of absorption enthalpy on temperature-swing CO₂ separation process performance

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Abstract

We model a CO₂ absorption process to elucidate the rationale for the search for a solvent with an enthalpy of absorption (ΔH) of low magnitude. We explore the relationship between ΔH and the system’s performance. While in general a lower magnitude appears to provide better system performance because it permits the stripper temperature to be decreased, as the magnitude drops below its value for monoethanolamine amine (MEA), 80 kJ/mol, the required solvent mass flow rate must increase precipitously and/or the flue gas must be cooled significantly. We argue that the associated parasitic pumping and cooling loads, as well as the increased capital cost, may set a practical lower limit on the magnitude of the enthalpy of absorption that is not very different from that of MEA.

Keywords: TSA systems; CO₂ capture process performance; enthalpy of absorption; MEA.

1. Introduction

As power plant scale CO₂ post-combustion capture systems around the world move from pilot project to full scale commercial deployment, many of the classical solvent (or sorbent) absorber/stripper systems currently on the market are likely to be deployed. In such temperature or pressure swing absorber (TSA or PSA) systems, the final process design presumably optimizes levelized costs based on solvent/sorbent properties and a host of more site-specific variables. Here, we restrict our analysis to TSA systems with solvents, but the spirit of the discussion also applies to PSA systems with sorbents.

The role of the solvent is key, and a good solvent is generally characterized by the following properties [1, 2]: (i) high selectivity for CO₂, (ii) low propensity to degrade over time, (iii) high maximum solvent loading, (iv) low lifetime cost, (v) wide envelope of possible operating conditions (pressure, temperature), and finally, (v) low enthalpy of absorption (ΔH). The importance of this final property is often directly linked to the energy penalty that the CO₂ scrubber system imposes on the power plant. On the one hand, a ‘large’ ΔH (i.e. large magnitude) is indicative of a solvent with a high affinity for CO₂, but on the other hand, for any amount of heat that is released in
the absorber, an even larger quantity of heat must be supplied to the stripper. As a consequence, much research is devoted to the search for a solvent with a ‘small’ $\Delta H$ (i.e. small magnitude), see for example [2–4].

The heat provided to the stripper is often broken into three separate pieces: (1) sensible heat to bring the stream of rich amine up to the operating temperature of the stripper; (2) enthalpy of reaction to reverse the CO$_2$ absorption; and (3) heat to vaporize water in the stripper to drive the desorption reaction forward by removing CO$_2$ as it is released out of the stripper. How much sensible heat and ‘stripping’ heat is required depends rather indirectly on $\Delta H$; furthermore the heat supplied to the stripper does not make up the entire energy penalty of the CO$_2$ scrubber system. Direct cooling loads, and the electric energy required to compress the CO$_2$ after separation are also main contributors. Clearly, system performance depends on more factors than just $\Delta H$, and in some cases a large $\Delta H$ proves to be the more favorable choice [5, 6]. In this work we aim to gain a better understanding of the relevant importance of $\Delta H$, by modeling a generic TSA system and keeping all solvent properties constant except $\Delta H$.

2. Thermodynamic Background

As discussed in [7], in the most thermodynamically abstract sense, a temperature-swing CO$_2$ absorption & separation process (TSA) can be depicted as an engine that takes heat from a hot reservoir ($T_H$), and uses this heat ($Q$) to internally do work ($W$) on a stream of flue gas (see figure 1). In separating the flue gas into its main components, nitrogen and carbon dioxide, the heat engine rejects heat to a cold reservoir (the environment, at temperature $T_e = T_L$).

In this most abstract formulation, it is clear that the maximum efficiency of the internal heat-to-work engine is limited by the Carnot efficiency,

$$\frac{W}{Q_{net}} = \eta_C = 1 - \frac{T_L}{T_H},$$

and the hot reservoir’s temperature corresponds to the temperature of the stripper. The minimum work required to separate a fraction $f$ of CO$_2$ from a gaseous stream with CO$_2$ mol fraction $x$ is ([7])

$$W_{\text{min}} = RT \frac{1}{x} \left[ (1-f)x \ln \left( \frac{1-f}{1-fx} \right) + (1-x) \ln \left( \frac{1-x}{1-fx} \right) - x \ln(x) - (1-x) \ln(1-x) \right],$$

where $R$ is the universal gas constant, and $T$ is temperature. For typical conditions of $x = 0.12$ and $f = 0.9$, $W_{\text{min}} \approx 8$ kJ/mol. Given that the heat that is provided to run this CO$_2$ -separation machine is typically taken from the steam cycle of a power plant where it would have been used to generate power, it is appropriate to consider the loss of exergy or availability associated with the CO$_2$ scrubber [8]. A natural figure of merit for a CO$_2$ separation system is thus the exergy loss per mol of CO$_2$ captured and compressed. By looking at exergy (i.e. $Q_{\text{heat}} \eta_C$) rather than just $Q_{\text{heat}}$, we place an appropriate premium on high temperature heat.

For a given CO$_2$ exit partial pressure, the performance of the CO$_2$ scrubber varies with $\Delta H$. This is demonstrated in figure 2 where we have plotted the equilibrium vapor pressure of CO$_2$ over a solution of CO$_2$ solvent for different temperatures (see (3)). The curve corresponding to $\Delta H = -80$ kJ/mol corresponds to equilibrium data for mo-

![FIG. 1. Abstracted representation of a thermal CO$_2$ separation engine.](image-url)
noethanolamine (MEA) from [9, 10], while the other curves use some data for MEA (i.e. the entropy of the overall reaction, and the concentration & loading dependence) but alter \( \Delta H \) (see Table 1 in [9]). Note that the y-axis cutoff for the logarithm of the partial pressure \( (p_{CO_2}) \) is directly related to \( \Delta S^0/R \), where \( \Delta S^0 \) is the entropy of the overall dissolution / absorption reaction at standard conditions. This is important to note since most solvents have very similar \( \Delta S^0 \) (for the family of the alkanolamines, for example, the spread is less than 9% [9]). This entropy change is dominated by the entropy loss of the CO2 molecule upon going from an ideal gas state to a liquid state relatively low entropy, and cannot be expected to vary greatly from one liquid to another. Thus we set the point on the y-axis from which all other curves, regardless of \( \Delta H \), emanate. If one chooses the operating partial pressures of CO2 in a TSA absorber and stripper respectively; a particular \( \Delta H \) will set the required operating temperatures as shown in figure 2. The associated Carnot efficiency increases with increasing \(|\Delta H|\), since the 'hot' temperature \( T_H \) increases accordingly, but the cooler environment temperature (\( T_e \)) to which heat will eventually be rejected stays the same. From observation of figure 2 we can understand why the chilled ammonia process (with \( \Delta H \approx -60 \) kJ/mol, [11]) necessarily runs at an absorber temperature close to 273 K, but we also see that this means that it will have a relatively low internal Carnot efficiency. Of course, this ‘first-order’ view of the process neglects the fact that solvent loading can impact the \( p_{CO_2} \) vs. \( \Delta H \) curve, as well as other process considerations. However, the role of \( \Delta H \) in setting overall process performance is not obvious.

\[ 10^{\log_{10}(c) \Delta S^{0}/R+c} \]

\[ p_{CO_2} \text{ in Pa} \]

\[ 0 \quad 0.001 \quad 0.002 \quad 0.003 \quad 0.004 \]

\[ 0 \quad 10^4 \quad 10^5 \quad 10^6 \]

\[ T = 486 \text{ K} \quad T = 474 \text{ K} \]

\[ T = 399 \text{ K} \quad T = 387 \text{ K} \]

\[ T = 291 \text{ K} \quad T = 289 \text{ K} \]

\[ p_{CO_2} = 1 \times 10^3 \text{ Pa} \]

\[ p_{CO_2} = 1.22 \times 10^4 \text{ Pa} \]

\[ p_{CO_2} = 1.22 \times 10^4 \text{ Pa} \]

**FIG. 2.** Partial pressure of CO2 over an amine solution with different \( \Delta H \). The required temperatures in the absorber and stripper are set by \( \Delta H \) and the required operating pressures (which are assumed to be equal to the equilibrium \( p_{CO_2} \)).

### 3. Model Development

#### 3.1. Process chemistry

Our model is a straightforward mass- and heat balance based on equilibrium thermodynamics, and implemented in MATLAB. It operates with a solvent that is ‘MEA-like’, meaning that the CO2 absorption reaction chemistry (and related mass balance) is akin to that of CO2 and MEA, i.e.,

\[ 2\text{MEA}_{(aq)} + \text{CO}_2\text{ (aq)} \rightarrow \text{MEAC}_{(aq)} + \text{MEA}_{H(aq)} + (\Delta H) \tag{2} \]

where MEA is a place holder for any other similar solvent. In the case of MEA, MEAC is more accurately the
carbamate associated with MEA (i.e. HO(CH2)2NHCOO−) while MEAH is the protonated solvent molecule (i.e. NH2(CH2)2OH+). This process chemistry was chosen since there is a lot of data available for MEA; but the model is easily altered for any other reaction stoichiometry.

3.2. Process description

Actual TSA processes are decidedly more complex than the abstract depiction presented in figure 1, but they are certainly subject to the same fundamental thermodynamic limits. In order to extend the abstract analysis of the system in figure 1 to include ΔH, we zoom in on the dotted box to separate out the typical subsystems separately; as in figure 3.

Warm moist flue gas exiting a wet flue gas desulfurization (FGD) unit at T0 = 328K [12] is first cooled down to T1 in the pre-conditioning unit which removes heat (QPC) by heat exchange with the cooler environment (i.e. cooling water at Te = TL). In the case where T1 < Te, further refrigeration work is required (Wref), and such a refrigerator is assumed to have a COP = T1/(Te − T1), powered by a heat engine that runs between the hot (TH) and cold (TL) sinks. The pre-conditioned flue gas then enters the adiabatic absorber.

In the absorber, CO2-rich flue gas is contacted with lean amine (θ = 0.05) and the exiting streams of CO2-lean flue and CO2-rich amine are assumed to be at equal temperatures (T3 = T4), as set by the release of ΔH in the absorber. We also specify the pressure in the absorber (p3) to be just over 1 bar, and perform all calculations for a flow of 1 mol of CO2 per second entering in stream 1, while assuming a 90% capture fraction. We assume that the partial pressure of CO2 in the flue gas entering the absorber is equal to the equilibrium saturation partial pressure of CO2 associated with the exiting rich amine stream (i.e. streams 1 and 4). In particular, we use the empirical equilibrium correlation between CO2 partial pressure and solvent temperature, concentration, and loading as documented by Gabrielsen et al. [9]

\[
p_{CO2} = \exp \left( A + \left( \frac{-ΔH}{RT} + C_{a0}θ \right) \frac{θ^2}{(1−θ)^2} \right) \text{ (in kPa)}, \tag{3}
\]

where \(A \propto ΔS^0 / R\), a measure of the overall change of entropy associated with the process of dissolution and absorption; \(C = -7 187\), \(a_0\) is the initial concentration of amine (mol MEA/mol solvent, roughly 0.1123 for an MEA solution
of 30% by weight as used in this work) and \( \theta \) is the solvent loading (mol CO\(_2\) /mol MEA, for MEA \( \theta_{\text{max}} = 0.5 \)).

We further take the water vapor partial pressure to be \( p_{3,\text{H}_2\text{O}} = x_{4,\text{H}_2\text{O}} p_{\text{H}_2\text{O},\text{sat}}(T_3) \), where \( x_{i,j} \) denotes the mol fraction of component \( j \) in stream \( i \), and \( p_{\text{H}_2\text{O},\text{sat}}(T) \) is the water saturation pressure at temperature \( T \) given by the Antoine equation. Consequently, since \( T_3 > T_1 \), some water is ‘lost’ in the absorber, and this is replenished by make-up water entering at temperature \( T_e \) in stream 2b. Using (3) together with a heat and mass balance that accounts for water loss, the model calculates the required solvent flowrate, loading, and the temperature in the absorber.

From the absorber, the CO\(_2\)-rich amine goes through an ideal heat exchanger (i.e. no heat lost to the environment) where its temperature is raised to \( T_4 \) by the hot lean amine exiting the stripper (using heat capacity data from [13]). The warm rich amine then enters the stripper where the CO\(_2\) is stripped out of solution. We specify the temperature in the stripper (i.e. \( T_s = T_{5a} = T_6 \)), and provide enough \( Q_{\text{heat}} \) to reach the required temperature, reverse the absorption reaction, and provide stripping steam. The model calculates the stripper pressure, and the mass ratio of CO\(_2\) to H\(_2\)O in stream 6, as well as \( Q_{\text{heat}} \). In order to do this, we again assume that the partial pressure of CO\(_2\) in stream 6 is equal to the saturation partial pressure of CO\(_2\) in the rich incoming amine (stream 4) at temperature \( T_{3b} \).

The CO\(_2\) + H\(_2\)O stream exiting the stripper is cooled to \( T_e \) (cooling load \( Q_{\text{cond}} \)); and the condensed water is returned at temperature \( T_e \) to the stripper. The CO\(_2\) then exits the scrubber system, and is compressed isothermally to \( p_F = 100 \) bar (i.e. \( W_{\text{comp}} = \eta_{\text{CO}_2} R T \ln(p_F/p_I) \), which assumes CO\(_2\) to act as an ideal gas [4]). For the final figure of merit of the overall system, we include this compression work. Simultaneously, the lean (but warm) amine is cooled (\( Q_{C2} \)) from \( T_{5b} \) to reach \( T_2 \), closing the solvent cycle.

4. Results and Discussion

4.1. Model performance

Figures 4 and 5 indicate typical model outputs showing (i) solvent flowrate, (ii) solvent loading, (iii) stripper pressure, and (iv) CO\(_2\)/H\(_2\)O mass ratio in stream 6 respectively; as a function of the enthalpy of reaction (\( \Delta H \)) and the prescribed stripper temperature. As the magnitude of the absorption enthalpy drops from 90 to 60 kJ/mol, CO\(_2\) loading in the rich solvent drops precipitously. In order to take up enough CO\(_2\) to achieve 90% capture, the solvent flow rate must rise precipitously. This result can be understood using figure 2. To attain the maximum possible CO\(_2\) loading of the rich amine exiting the absorber, its saturation CO\(_2\) pressure must be no greater than 12.2 kPa (the lower horizontal dashed line). This requirement sets a maximum permissible exit temperature that depends on the value of \( \Delta H \) and is indicated as the intersection of the various curves with this dashed line. When the magnitude of \( \Delta H \) drops enough that this temperature drops below the inlet flue stream temperature \( T_1 \), the rich amine exit stream cannot leave maximally loaded. So as the magnitude of the absorption enthalpy drops significantly below the value of 80 kJ/mol characteristic of MEA, the requisite pumping hardware and its parasitic load must be increased substantially. Although we do not include these costs in this model, we point out that this criterion may determine a practical lower limit on the

FIG. 4. Flowrate of solvent and resultant solvent loading (rich phase) \((T_1 = 313 \text{ K})\).
magnitude of the absorption enthalpy near that of MEA.

Figure 5A shows the pressure in the stripper \( (p_s) \) that would result from a given choice of \( \Delta H \) and \( T_s = T_6 \). The figure shows that stripper pressure depends mainly on temperature, and the pressure becomes excessive as \( T_s \geq 400 \) K. Actual systems are expected to operate in this temperature range. Figure 5B shows the \( \text{CO}_2/\text{H}_2\text{O} \) mass ratio that is to be expected at a given \( (T_s, \Delta H) \) combination. Again, for typical operating conditions (i.e. \( \Delta H_{\text{MEA}} = -82 \) kJ/mol \( \text{CO}_2 \) [9]) our model tends to overestimate this \( \text{CO}_2/\text{H}_2\text{O} \) ratio. From figures 5A and 5B, we learn that our model’s dependence on equilibrium data may not yield quantitatively accurate results for these elevated temperatures and dynamic conditions. However, we expect qualitative trends to persist.

Note that the white areas in figures 5 and 6 correspond to unphysical combinations of \( T_s \) and \( \Delta H \). In each of these figures, the white patch in the lower left hand corner represents a zone where \( T_s \) is too low to accommodate the relatively large magnitude of \( \Delta H \). As \( T_s \) is decreased, so is the required minimum stripper temperature; this is related to the lower flowrate (and consequently higher solvent loading) that comes with a decrease in \( T_s \). The white patch on the right of each figure (independent of \( T_s \)) stems directly from the maximum accessible \( \text{CO}_2 \) partial pressure in the absorber at a given \( \Delta H \) and \( T_s \), as depicted in figure 2. Further shifts in the regions of ‘unphysical’ solutions can be accomplished by altering the \( \Delta S^0 \) of the overall reaction, and/or the concentration dependence in (3).

4.2. System performance as a function of \( \Delta H \)

As discussed, we define a figure of merit (FOM) for the \( \text{CO}_2 \) capture system by considering the total exergy (or availability) that is ‘consumed’ in the separation and compression of \( \text{CO}_2 \), per mol of \( \text{CO}_2 \). Specifically, we calculate

\[
\text{FOM} = \frac{Q_{\text{heat}}\eta_c(T_s) + W_{\text{comp}} + W_{\text{ref}}}{n_7},
\]

where \( W_{\text{ref}} \) is the work required to refrigerate the flue gas when \( T_s < T_6 \), and \( \eta_c(T_s) = 1 - T_c/T_s \) while \( n_7 \) is the number of moles of \( \text{CO}_2 \) that are captured and compressed by the quantities of work and heat presented here. Clearly, a complete exergy analysis would include more contributions such as the kinetic energy in the various flows under consideration, along with additional losses that occur in the system such as frictional losses and pump- and heat exchanger efficiencies, etc. However, the figure of merit in (4) provides, in a thermodynamic sense, the most ‘fair’ measure of system performance, since the loss of availability represents the useful work that could have been obtained from the heat if it had not been applied to \( \text{CO}_2 \) separation.

Armed with the figure of merit as defined in (4), we interrogate our model to learn about system performance as a function of \( T_s \), \( \Delta H \), and \( T_1 \). Figure 6 shows that system performance is a rather complex function of \( \Delta H \), even in the absence of further complicating factors such as solvent degradation issues etc. In general, system performance
improves with decreasing $\Delta H$, but for any given $T_s$ local minima exist, see for instance figure 6B at $T_s = 380K$ for $\Delta H$ between -75 and -90 kJ/mol CO$_2$. The complex nature of the ‘$\Delta H$, FOM’ -landscape is the result of different non-linear contributions to the figure of merit (4). Notably, as stripper pressure increases, the amount of work required for subsequent compression decreases. When designing actual systems, however, solvent degradation at high pressure, and the capital cost associated with high operating pressures, may prove to be excessive.

Comparing figures 6A and 6B, we see the significant effect that $T_1$ has on system performance. Across the board, a lower $T_1$ seems to result in better system performance, but note that chilled systems have a greater cooling load, as shown in figure 7. Some of the heat carried away by the cooling streams indicated in figure 3 may have further beneficial use, but most of the heat is going to be very low grade and will thus directly impact cooling water pumping requirements at the power plant.

In summary, we have modeled a typical TSA CO$_2$ scrubbing system using an equilibrium thermodynamic approach. Subsequent interrogation of the model with respect to the role of the enthalpy of absorption reveals the relationship between $\Delta H$ and the system’s performance. While in general a lower magnitude of the absorption en-
thalpy appears to provide better system performance because it permits the stripper temperature to be decreased, as the magnitude of ΔH drops below its value for MEA, 80 kJ/mol, the required solvent mass flow rate must increase precipitously and/or the flue gas entering the system must be chilled significantly. The associated parasitic pumping and cooling loads, as well as the increased capital cost, while not included in the present model, may set a practical lower limit on the magnitude of the enthalpy of absorption that is not very different from that of MEA.

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