A STUDY ON THE REACTION BETWEEN CO₂ AND ALKANOLAMINES IN AQUEOUS SOLUTIONS

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Abstract—Literature data on the rates of reaction between CO₂ and alkanolamines (MEA, DEA, DIPA, TEA and MDEA) in aqueous solution are discussed. These data induced us to carry out absorption experiments of CO₂ into aqueous DEA, DIPA, TEA and MDEA solutions from which the respective rate constants were derived. The experimental technique was similar to that used by Laddha and Danckwerts[30].

The results for DEA and DIPA were analysed by means of a zwitterion-mechanism which was derived from the mechanism originally proposed by Danckwerts[16], and were found to fit the model extremely well.

The reaction rate of CO₂ with aqueous TEA and MDEA solutions shows a significant base catalysis effect which is also reported by Donaldson and Nguyen[17] and Barth et al.[4]. The results for TEA correspond very well with those of Donaldson and Nguyen[17] and Barth et al.[4].

INTRODUCTION
In the last decades, alkanolamines have acquired a well-established position in gas-treating for the removal of the acidic components H₂S and CO₂. Industrially important alkanolamines are: mono-ethanolamine (MEA), di-ethanolamine (DEA), di-isopropanolamine (DIPA) and methyl-di-ethanolamine (MDEA)[27]. In general these amines are used in aqueous solutions, but for particular applications combined solvents are more suited (e.g. water and sulfolane in the Shell "Sulfinol" process)[27].

Due to the high and still sharply rising energy cost involved in the operation of gas-treating plants, the incentive for development of (even slightly) more efficient processes is considerable.

Nowadays large savings in operation and capital cost are obtained by the selective removal of H₂S from CO₂ containing gases, not only in situations where H₂S is the only component to be removed. Even in LNG production, where the CO₂ has to be removed to around 100 ppm so as to avoid plugging of the cryogenic equipment, selective H₂S absorption is economically very attractive, using sophisticated treating schemes as described by McEwan and Marmin[33]. Increasing the selectivity reduces the solvent circulation rate and therefore the steam consumption in the regenerators and, moreover, reduces the dimensions of sulphur recovery and tail-gas units.

The H₂S selectivity in these treating processes depends largely on 3 factors: (1) the kinetics of the reactions between H₂S/CO₂ and alkanolamine solutions; (2) the mass transfer properties of the absorption equipment; (3) the equilibria in H₂S-CO₂-amine systems. In this work we focused our attention on the first factor.

As the reaction between H₂S and aqueous alkanolamines involves only a proton transfer, this reversible reaction can be considered to be infinitely fast[15] and hence the absorption rate is entirely mass transfer controlled under practical conditions[36].

The reactions between CO₂ and alkanolamine solutions, however, proceed at a finite rate, different for the various amines[15, 47]. From a purely kinetic point of view the selectivity for H₂S therefore only depends on the CO₂ reaction rate. Consequently this work deals with the reaction of CO₂ and aqueous alkanolamine solutions.

Until recently the mechanism and kinetics for this reaction were considered to be simple and straightforward[15] for all alkanolamines, although large discrepancies in data have been reported for DEA[1, 17, 18, 19, 22, 26, 43, 47] and TEA[17, 19, 25, 26, 43]. Danckwerts proposed a comprehensive reaction mechanism[16] which is essentially able to cover all kinetic data for both primary and secondary alkanolamines. In addition, Laddha and Danckwerts[30] published experimental results for aqueous MEA and DEA, which seem to support the proposed mechanism.

For tertiary alkanolamines Donaldson and Nguyen[17] suggested a base catalysis of the CO₂ hydration reaction.

In this study we attempt to critically summarize available kinetic data for MEA, DEA, DIPA, TEA and MDEA and present new data on DEA, DIPA, TEA and MDEA.

REVIEW OF LITERATURE KINETIC DATA

General remarks

Comparison of literature data on kinetics of CO₂-alkanolamine reactions is a complicated matter, because the authors use different experimental techniques, physico-chemical data and amine purities. The latter factor is usually overlooked, but even very small amounts of primary amine contaminants in secondary amines and

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primary and secondary amines in tertiary amines can enhance the absorption rate drastically. A mathematical approach for the estimation of the effect of these contaminants is given by, e.g. Jhaeveri[23] and Onda et al.[35], but correction of published kinetics remains ineffective due to lack of data on amine purities.

In order to verify the conditions for pseudo first order absorption experiments and—what is more important—to obtain reaction rate constants from these experiments, the physico-chemical parameter $m_{CO_2} \cdot \sqrt{D_{CO_2}}$ is needed. In literature, the value of this parameter is derived in various ways, as will be demonstrated further, and this may lead to discrepancies in the rate constants calculated from absorption experiments.

As CO$_2$ reacts with aqueous alkanolamines, it is not possible to measure $m_{CO_2}$ or $D_{CO_2}$ in these solutions directly. This problem is usually overcome by measuring these parameters with the non-reacting N$_2$O which is almost identical[1, 12, 29]. Laddha et al.[29] and Laddha and Danckwerts[30] made it plausible that the CO$_2$ parameters can be obtained from N$_2$O measurements by the relation:

$$m_{CO_2} \cdot \sqrt{D_{CO_2}} = 1.43 \cdot m_{N_2O} \cdot \sqrt{D_{N_2O}}. \quad (1)$$

Very often the diffusivity, $D_{N_2O}$ or $D_{CO_2}$, is correlated to the viscosity of the solution at a constant temperature by [40]:

$$D \cdot \eta^n = \text{constant} \quad (2)$$

where usually $0.5 < n < 1$.

Sada et al. have determined $m_{N_2O}$, $D_{N_2O}$ and $\eta$ for aqueous MEA[41], DEA[45, 46], DIPA[46] and TEA[45, 46] solutions. In Fig. 1 $\ln(D_{N_2O-amine}/D_{N_2O-water})$ is plotted vs $\ln(\eta_{amine}/\eta_{water})$ and according to eqn (2) this should yield a straight line through the origin with slope $n$. For all amines, however, the fit is very poor and, moreover, $n$ depends on the type of amine (e.g. DEA: $n \approx 0.3$, DIPA: $n \approx 0.7$). Therefore, the use of eqn (2) for correlation of diffusivity data in (aqueous) alkanolamine solutions is not recommended, as was already pointed out by Laddha and Danckwerts[30].

The Stokes–Einstein equation, in fact a special case of eqn (2) where $n = 1$, is used by a number of authors for the estimation of diffusivities[18, 20, 47]. This equation grossly overestimates the dependence on the solution viscosity. For this reason Hikita et al.[20] calculate from their absorption experiments a “salting-in” effect for the CO$_2$-solubility in MEA. Most of the data measured, however, unambiguously show a “salting-out” effect[12, 29, 41, 45, 46].

Another approach used by several authors, is to employ the CO$_2$ solubility in water for the interpretation of their experiments[1, 18, 47]. Because obviously $m_{CO_2}$ and $D_{CO_2}$ cannot be correlated separately, in a satisfactory way, we preferred to use the combined parameter $m_{CO_2} \cdot \sqrt{D_{CO_2}}$ as obtained from N$_2$O experiments for the interpretation of literature data and our own data measured at 25°C. The values of $m_{CO_2} \cdot \sqrt{D_{CO_2}}$ are calculated from literature for MEA[12, 30, 41], DEA[30, 45, 46], TEA[45, 46] and DIPA[46], by interpolating the solubilities. These data are complemented by our measurements for DIPA and MDEA (see Table 1). The experimental technique is described elsewhere[8, 9]. All data are summarized in Fig. 2 and polynomial fittings are given in Table 2. These correlations are used throughout this work.

With respect to the reaction rate dependence on the CO$_2$ concentration, invariably a first order rate equation...
A study on the reaction between CO₂ and alkanolamines in aqueous solutions

Table 1. \( m_{\text{N}_2\text{O}} \sqrt{D_{\text{N}_2\text{O}}} \) as a function of [Am] for DIPA and MDEA at 25°C[8]

<table>
<thead>
<tr>
<th>Amine</th>
<th>[Am] (mol/l)</th>
<th>( m_{\text{N}<em>2\text{O}} \sqrt{D</em>{\text{N}_2\text{O}}} \times 10^5 ) m/s¹/₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>DIPA</td>
<td>0.580</td>
<td>2.15</td>
</tr>
<tr>
<td></td>
<td>1.280</td>
<td>1.67</td>
</tr>
<tr>
<td></td>
<td>1.753</td>
<td>1.31</td>
</tr>
<tr>
<td></td>
<td>2.330</td>
<td>1.03</td>
</tr>
<tr>
<td>MDEA</td>
<td>0.511</td>
<td>2.23</td>
</tr>
<tr>
<td></td>
<td>1.060</td>
<td>2.05</td>
</tr>
<tr>
<td></td>
<td>1.440</td>
<td>1.78</td>
</tr>
<tr>
<td></td>
<td>2.056</td>
<td>1.65</td>
</tr>
</tbody>
</table>

is found in literature. In the following we shall, therefore, consider the influence of the amine concentration only.

MEA

Literature sources with kinetic data on aqueous MEA are summarized in Table 3. Wherever possible and necessary, the reaction rate constants at 25°C have been corrected to the standard data for \( \frac{m_{\text{CO}_2} \cdot \sqrt{D_{\text{CO}_2}}}{2.303} \) by using the equations of Table 2. Without any exception a first order reaction rate in MEA has been found regardless of experimental techniques and conditions. The data show fairly good agreement as can be concluded from an Arrhenius plot (Fig. 3). Danckwerts and Sharma's[15] rate constants at 18 and 35°C seem to be fairly high, probably due to the use of too low values of \( m_{\text{CO}_2} \cdot \sqrt{D_{\text{CO}_2}} \) calculated by the Stokes–Einstein equation. The reason of the relatively large scatter in data at 25°C is not quite clear, but could be attributed to different experimental techniques.

The data most recently published, i.e. those of Laddha and Danckwerts[30], Donaldson and Nguyen[17] and Alvarez-Fuster et al.[1], fit the equation by Hikita et al.[19]:

\[
\log_{10} k_2 = 10.99 - 2152/T \quad (1./\text{mole.sec})
\] (3)

extremely well. We concluded therefore that these literature sources provide the reaction rate constant of the CO₂-MEA reaction with good accuracy and that no additional kinetic data on MEA seem to be required.
DEA

The kinetic data of the CO$_2$-DEA reaction are given in Table 4. The reaction rate expressions found in literature vary widely in dependence on the DEA and OH$^-$ concentrations. As pointed out by Danckwerts[16], the principal reason for this disagreement is the fact that the reaction mechanism is far more complicated than most authors assumed. With respect to this point we deemed it useful to provide additional data on the CO$_2$-DEA reaction over a wide range of conditions.

DIPA

Only two investigations on reaction rates of CO$_2$ with DIPA have been published in open literature (see Table
A study on the reaction between CO\textsubscript{2} and alkanolamines in aqueous solutions

\[ \text{TEA} \]

The rate constants for the CO\textsubscript{2}-aqueous TEA solutions show in general a first order dependence on the TEA-concentrations (see Table 6). The results of Jørgensen and Faurholt\cite{25} and Jørgensen\cite{26} at pH ~ 13, however, include a hydroxyl ion term caused by monoalkyl carbonate formation, which is not found at pH < ~ 11\cite{17, 19}. The rate constants at 25°C vary from 2.8 l./mole.sec\cite{17, 4} up to 50 l./mole.sec (Hikita et al.\cite{19}). Because of these discrepancies we provide some data on the CO\textsubscript{2}-TEA reaction.

\[ \text{MDEA} \]

In the open literature only one single investigation is published which contains only little information on the reaction of CO\textsubscript{2} in aqueous MDEA solutions\cite{4}. Barth et al.\cite{4} also present some information on reaction mechanism and rate but they used a relatively high pK\textsubscript{a} value in the evaluation of their experiments (pK\textsubscript{a} = 8.65 instead of pK\textsubscript{a} = 8.52 at 25°C as reported by Perrin\cite{37}). This results in an overestimation of the contribution of the hydroxyl ions to the measured reaction rate and in a too low contribution by MDEA. Therefore we also included MDEA in our test programme.

**THE REACTION MECHANISM**

In aqueous solutions of primary and secondary alkanolamines, the following reactions with CO\textsubscript{2} occur (see e.g. \cite{15, 16}):

- carbamate formation:
  \[ \text{CO}_2 + 2R_1R_2\text{NH} = R_1R_2\text{NCOO}^- + R_1R_2\text{NH}_3^+ \]  
  \[ (4) \]

- bicarbonate formation:
  \[ \text{CO}_2 + \text{OH}^- = \text{HCO}_3^- \]  
  \[ (5) \]

- carbonic acid formation:
  \[ \text{CO}_2 + \text{H}_2\text{O} = \text{HCO}_3^- + \text{H}^+ \]  
  \[ (6) \]

- alkylcarbonate formation:
  \[ \text{CO}_2 + R_1R_3\text{C} = \text{C} = \text{NH} + \text{OH}^- \]  
  \[ \text{OH} \]  
  \[ R_1R_3\text{C} = \text{C} = \text{NH} + \text{H}_2\text{O} \]  
  \[ (7) \]
Tertiary alkanolamines do not react directly with CO$_2$ according to reaction (4), because they lack the free proton. Yet they combine with CO$_2$ in aqueous solutions by the reactions (5)-(7).

In principle, each of the reactions (4)-(7) for primary and secondary, and (5)-(7) for tertiary alkanolamines, contributes to the measured overall reaction rate constant $k_{ov}$. Some neglections are justified, however. The formation of carbonic acid from CO$_2$ and H$_2$O by reaction (6) is very slow ($k = 0.026$ sec$^{-1}$ at 25°C[38]) compared to the reactions (4) and (5) at pH > 9.5 as will be shown later. Hence reaction (6) is not incorporated in $k_{ov}$. By using the data of Jensen et al.[22] and Jørgensen[26], it can be calculated that the alkylcarbonate reaction (7) contributes negligibly to the CO$_2$ absorption rate for MEA[48] and DEA at pH < 12. As this condition is amply fulfilled in our experiments, we neglected this reaction for DEA as well as for DIPA.

On the other hand, the contribution of the bicarbonate formation reaction (5) to the absorption rate can be substantial at low concentrations of secondary amines and is essential for tertiary amines, as will be demonstrated further. This reaction is well documented, e.g. by Pinsent et al.[8], and its forward rate constant is expressed by:

$$\log_{10} k^* = 13.635 - \frac{2895}{T} \text{ (l./mole.sec).} \quad (8)$$
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Table 5. Literature data on the reaction between CO₂ and aqueous DIPA

<table>
<thead>
<tr>
<th>reference</th>
<th>temperature [°C]</th>
<th>[DIPA] mole/l</th>
<th>k₂ l/mole.s</th>
<th>Eₚₑᵢₓ [kJ/mole]</th>
<th>Experimental technique</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sharma [47]</td>
<td>15</td>
<td>1.0</td>
<td>230</td>
<td>~ 41.8</td>
<td>Laminar jet</td>
</tr>
<tr>
<td>Danckwerts and Sharma [15]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Groothuis [18]</td>
<td>25</td>
<td>2.0</td>
<td>450 - 550</td>
<td>~ 41.8</td>
<td>Stirred cell</td>
</tr>
<tr>
<td>Sharma [47]</td>
<td>25</td>
<td>1.0</td>
<td>400 - 440</td>
<td>~ 41.8</td>
<td>Laminar jet</td>
</tr>
<tr>
<td>Danckwerts and Sharma [15]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sharma [47]</td>
<td>35</td>
<td>1.0</td>
<td>680</td>
<td>~ 41.8</td>
<td>Laminar jet</td>
</tr>
<tr>
<td>Danckwerts and Sharma [15]</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*: corrected for m_CO₂ \sqrt{D}_{CO₂} \text{ in table 2.}

Table 6. Literature data on the reaction between CO₂ and aqueous TEA and MDEA

<table>
<thead>
<tr>
<th>amine</th>
<th>reference</th>
<th>temperature (range) [°C]</th>
<th>[A] mole/l</th>
<th>k₁° [CO₂] l/mole.n⁻¹.s⁻¹</th>
<th>Eₚₑᵢₓ [kJ/mole]</th>
<th>Experimental technique</th>
</tr>
</thead>
<tbody>
<tr>
<td>TEA</td>
<td>Nikita et al. [19]</td>
<td>20-40</td>
<td>0.005-0.0076</td>
<td>10 - 10.72 - 25841/\text{TEA}</td>
<td>51.5</td>
<td>Rapid mixing method</td>
</tr>
<tr>
<td>TEA</td>
<td>Jørgensen and Faurhol [25]</td>
<td>0</td>
<td>0.1, 0.2, 0.5</td>
<td>5930 [TEA][OH⁻] [16]</td>
<td>-</td>
<td>Competitive reaction with 0.1 M NaOH</td>
</tr>
<tr>
<td>TEA</td>
<td>Jørgensen [26]</td>
<td>10</td>
<td>0.2, 0.4</td>
<td>12730 [TEA][OH⁻] [16]</td>
<td>-</td>
<td>Competitive reaction with 0.2 M NaOH</td>
</tr>
<tr>
<td>TEA</td>
<td>Barth et al. [41]</td>
<td>25</td>
<td>0.02 - 0.2</td>
<td>2.85*[TEA]</td>
<td>-</td>
<td>Stopped flow method</td>
</tr>
<tr>
<td>TEA</td>
<td>Donaldson and Nguyen [17]</td>
<td>25</td>
<td>0.1 - 1.0</td>
<td>2.7 [TEA]</td>
<td>-</td>
<td>Facilitated transport in aqueous TEA membranes</td>
</tr>
<tr>
<td>TEA</td>
<td>Sada et al. [43]</td>
<td>25</td>
<td>0.50 - 1.60</td>
<td>16.8 [TEA]</td>
<td>-</td>
<td>Wetted wall column</td>
</tr>
<tr>
<td>MDEA</td>
<td>Barth et al. [4]</td>
<td>25</td>
<td>0.02 - 0.2</td>
<td>not conclusive</td>
<td>-</td>
<td>Stopped flow method</td>
</tr>
</tbody>
</table>

In our mechanistic model we assume that the carbamate formation from primary and secondary alkanolamines by reaction (4) takes place in accordance with the approach given by Danckwerts [16], derived from Caplow's work [10]. The reaction steps successively involve the formation of a "zwitterion":

\[ \text{CO}_2 + R_1R_2NH \xrightarrow{k_2} R_1R_2NH^+\text{COO}^- \]  \quad (9)

and the subsequent removal of the proton by a base B (base catalysis):

\[ R_1R_2NH^+\text{COO}^- + B \xrightarrow{k_0} R_1R_2NCOO^- + BH^+. \]  \quad (10)

For this mechanism Danckwerts [16] derived the forward reaction rate equation at quasi-steady state:

\[ r_{CO_2} = k_2[R_1R_2NH] \frac{R_1R_2NH^+\text{COO}^-}{1 + \Sigma k_0[B]} \]  \quad (11)

where \( \Sigma k_0[B] \) indicates the contribution to the proton removal step (10) by all bases present in the solution.

In a recent second publication on this subject, Laddha and Danckwerts [30] considered only the amine as a base in eqn (11), neglecting contributions by H₂O and OH⁻. In several articles [1, 2, 10, 22, 25, 26] these species are shown to have a pronounced effect on the rate of reaction (4). In our model we considered, therefore, incorporation of H₂O and OH⁻ as bases in (11) to be...
essential. Rate equation (11) then becomes:

\[
2 < Ha \ll E, \quad \text{(16)}
\]

and:

\[
E_t = \sqrt{\frac{D_{CO_2}^T}{D_{Am}}} + \sqrt{\frac{P_{CO_2}^T}{P_{CO_2}^T}} \cdot m_{CO_2} \cdot A. \quad \text{(18)}
\]

The CO₂-absorption rate is then described by:

\[
J_{CO_2} \cdot A = \sqrt{k_{ov} \cdot D_{CO_2} \cdot m_{CO_2} \cdot P_{CO_2}^T \cdot A}. \quad \text{(19)}
\]

In our closed reactor with a gas volume \( V_g \), the pressure-time relation can easily be obtained from an in-stationary mass balance and becomes:

\[
\ln P_{CO_2}\_t = -m_{CO_2} \cdot A \cdot \sqrt{k_{ov} \cdot D_{CO_2} \cdot t} + \ln P_{CO_2}\_i - a. \quad \text{(20)}
\]

The overall-reaction rate constant, \( k_{ov} \), was determined from the slope in a ln \( P_{CO_2} \)-time plot in the region where condition (16) was met.

The apparent reaction rate constant for the carbamate formation reaction (4), \( k_{app} \), is calculated from the overall rate constant \( k_{ov} \) by correction for the contribution of the bicarbonate formation by reaction (5):

\[
k_{app} = k_{ov} - k_{ov} \cdot [OH^-] = \frac{k_2[R,\, R_2NH]}{k_{R_2}\, [R,\, R_2NH]} \quad \text{(sec⁻¹)}. \quad \text{(21)}
\]

The experimental set-up is shown in Fig. 4. The absorption experiments were carried out in a \( \sim 10 \) cm i.d. all glass, thermostatted stirred-cell reactor at 25.0°C (see Fig. 5). The reactor consists of upper and lower parts which seal on ground flanges. The glass stirrer is equipped with molten in magnets and is driven externally at 60 rpm. The absolute pressure in the reactor is recorded by a mercury pressure indicator and is read by a kathetometer to an accuracy of \( \sim \)0.02 mm Hg.

Before the experiments the freshly prepared solutions were degassed under vacuum in a separate glass vessel to strip off all inert gas contaminants. After degassing, the solution was fed under vacuum into the stirred-cell reactor and the vapour-liquid equilibrium was allowed to establish. The pressure was read and this value, \( P_{H_2O} \), was used to calculate the actual \( P_{CO_2} \) used in eqn (20) from the measured total pressure \( P_{tot} \) by:

\[
P_{CO_2} = P_{tot} - P_{H_2O}. \quad \text{(22)}
\]

The \( P_{H_2O} \) measured proved to be proportional to the
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Fig. 4.

Fig. 5.

mole fraction water at all amine concentrations, according to Raoult's law.

After equilibration, pure CO$_2$ was introduced into the reactor. By means of the amount of CO$_2$ both initial pressure $P_{CO_2,0}$ and the average CO$_2$-liquid load for the experiments could be adjusted. By this substantial CO$_2$ liquid-load primary (and secondary) amine contaminants, if present, were substantially converted to carbamates and their effects on the measured absorption rate could be strongly reduced. On the other hand, the average [OH$^-$] during the experiments could be adjusted by variation of the liquid-load.

The decrease of $P_{CO_2}$ during the experiment caused $E_i$ to increase steadily (see eqn 18) and, therefore, condition (16) could always be met in part of the pressure-time curve, even if a high initial $P_{CO_2}$ was used to obtain a
substantial average CO₂ liquid-load (see Fig. 6 for a typical example). At least 10 pressure-time readings in the region where condition (16) was met were used to obtain the overall reaction rate constant \( k_{ov} \). Duplicate runs varied only 3% in \( k_{ov} \), at the maximum.

After the experiment, the solution was analysed for total amine by standard potentiometric titration with 0.500 M hydrochloric acid and for total CO₂ by the method described by Verbrugge [49] and Jones et al. [24]. From this final CO₂ content and the pressure-time readings an average CO₂ liquid-load during each experiment was calculated. The equilibrium model described by Blauwhoff and van Swaaij [6] was used for calculation of hydroxyl ion and actual free amine concentrations from average CO₂ liquid load and total amine concentration. This model includes, among others, equilibrium reactions (4)–(6), and the carbamate hydrolysis equilibrium. At the relatively low CO₂ loadings in our investigations the effect of this latter equilibrium on the amount of the actual free amine concentration used for the calculations is, however, generally negligible.

The influence of the CO₂ liquid-loading in the experiments on the rate of the reverse of reaction (4) and with that its effect on the measured absorption flux \( J_{CO₂} \) was checked for DIPA using the numerical solution technique described by Cornelisse et al. [14]. For this check, calculated molefluxes with and without CO₂ liquid-loads in the experimental range, having the same real free amine bulk concentrations (obtained with the equilibrium model) were compared. The difference in molefluxes amounted to 1% at the maximum for the combination of the lowest DIPA concentration and the highest CO₂ liquid-load. The effect of the reversibility on measured reaction rate constants could, therefore, be neglected.

**RESULTS**

**DEA and DIPA**

For both DEA and DIPA, three series of CO₂ absorption experiments (A, B and C) were carried out at 25°C. A range of amine concentrations was applied within each series (see Tables 7 (DEA) and 8 (DIPA)). In the A- and B-series in almost constant average CO₂...
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Table 7. Experimental results for DEA at 25°C (present investigation)

<table>
<thead>
<tr>
<th>[DEA] mole/1</th>
<th>[CO₂]¹⁺⁺</th>
<th>[OH⁻]⁻⁻</th>
<th>[H₂O]⁻⁻</th>
<th>k_{app} s⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0.509</td>
<td>1.19</td>
<td>3.15</td>
<td>52.6</td>
<td>199.5</td>
</tr>
<tr>
<td>0.516</td>
<td>1.15</td>
<td>3.33</td>
<td>52.6</td>
<td>213.1</td>
</tr>
<tr>
<td>0.900</td>
<td>1.15</td>
<td>5.67</td>
<td>50.6</td>
<td>466.0</td>
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<td>11.50</td>
<td>44.0</td>
<td>1545</td>
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<tr>
<td>2.308</td>
<td>2.59</td>
<td>7.44</td>
<td>43.1</td>
<td>1760</td>
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</table>

Liquid concentration was realized in the way previously described. The measured apparent rate constants have been plotted logarithmically versus \(\ln [R_1R_2NH]\) in Fig. 7. Two important features can be observed: (1) the apparent reaction order in amine, \(n_{app}\), is \(-1.3\) increasing with amine concentration to \(-1.5\) for DEA and \(-1.6\) to \(-1.8\) for DIPA; (2) the apparent rate constant decreases with rising CO₂ liquid-load.

The first point confirms that the reaction mechanism is not simple and straightforward indeed, and indicates that more complicated schemes, like the one proposed in this paper, are needed to explain the results.

The second point can be explained only by a decrease of the hydroxyl ion concentration with increasing CO₂ liquid load because its influence on real free amine concentration, \([R_1R_2NH]\), is incorporated using the equilibrium model mentioned earlier[6]. This effect is in line with the mechanism proposed and the reaction rate equation (12) derived. In the experiments by Laddha and Danckwerts[30], however, the average CO₂ liquid-load was not varied systematically, as can be concluded from their experimental description. Hence this influence could probably not be discerned and was consequently not incorporated in their reaction rate equation.

We analysed the proposed reaction mechanism by fitting \(k_{app}\) (eqn 21) to all our experimental results, minimizing the sum of least squares by a linear regression technique. In this way we determined the values of \(k_2, k_2k_{\text{H}_2\text{O}}, k_2k_{\text{OH}^-}\) and \(k_2k_{R_1R_2NH}\) (see Table 9). The average deviation between the experiments and the fitted model is extremely small.

Some very important conclusions can be drawn from the results in Table 9, which hold both for DEA and DIPA: firstly, \(1/k_2\) is less than 10% of \(k_1\) at all amine concentrations. For this reason, the exact
values of $k_2$ could not be determined, but the minimum values seem to be $-5 \times 10^{-11}$ mole/sec, which is in the same order of magnitude as the rate constant for the CO$_2$–MEA reaction. This implies that the rates of formation of the zwitterions for DEA and DIPA are at least in the same range as for MEA.

Secondly, it follows:

$$k_{-1} \geq \frac{1}{k_{-1} \Sigma k_i[\beta]}$$  \hspace{1cm} (23)$$
indicating that the larger part of the zwitterions is reverted to CO$_2$ and amine and that only a small part is converted to carbamate. This limited stability is the essential difference with the CO$_2$–MEA reaction. Using (23), rate eqn (12) can be simplified to:

$$r_{10} = \frac{k_2}{k_{-1}} [\text{CO}_2][R,R_2\text{NH}]$$

$$\times \{k_{\text{H}_2\text{O}}[\text{H}_2\text{O}] + k_{\text{OH}^-}[\text{OH}^-] + k_{R_1R_2\text{NH}}[R_1R_2\text{NH}]\}$$

(mole/l/sec).  \hspace{1cm} (24)$$

This rate equation is basically different from the one used by Laddha and Danckwerts[30]:

$$r_{10} = \frac{[\text{CO}_2][R,R_2\text{NH}]}{1410 + \frac{1}{1200[R,R_2\text{NH}]}}$$

(mole/l/sec).  \hspace{1cm} (25)$$

At very low amine concentrations, condition (23) is fulfilled and our rate equation (24) tends to a first order dependency of $r_{10}$ in amine, $k_{\text{H}_2\text{O}}[\text{H}_2\text{O}]$ becoming the
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Equation (25) by Laddha and Danckwerts[30], however, tends to a second order dependency in amine. Our eqn (24) agrees, therefore, with the trend found by the British Gas Corp. (mentioned by Laddha and Danckwerts[30]). On the other hand, a second order in amine is predicted by eqn (24) at high amine concentrations, provided conditions (23) is still met, which is in agreement with Nunge and Gill[34]. Rate expression (25) of Laddha and Danckwerts predicts in this case a first order in amine.

Thirdly, comparing the relative magnitudes of the "partial rate constants",

\[
k_{2k_{H2O}} \frac{k_{2k_{OH}}}{k_{-1}}, \frac{k_{2k_{OH}}}{k_{-1}} \text{ and } \frac{k_{2k_{R_{1}R_{2}NH}}}{k_{-1}}
\]

(Table 9) an increase is found with increasing basicity of the base in the proton removal step (10). This Brønsted effect is clearly shown in Fig. 8 (DEA: \(pK_a = 8.88 \) at 25°C[37]; DIPA: \(pK_a = 8.88 \) at 25°C[7]). It is also significant that the "partial rate constants" for DEA are larger than for DIPA, most likely caused by the larger

Table 9. Fitted values of kinetic constants

<table>
<thead>
<tr>
<th></th>
<th>(k_2)</th>
<th>(k_{2k_{H2O}})</th>
<th>(k_{2k_{OH}})</th>
<th>(k_{2k_{R_{1}R_{2}NH}})</th>
<th>average deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEA</td>
<td>5800</td>
<td>5.34</td>
<td>7.05 \times 10^{-4}</td>
<td>228</td>
<td>3.3</td>
</tr>
<tr>
<td>DIPA</td>
<td>5300</td>
<td>0.813</td>
<td>4.17 \times 10^{-4}</td>
<td>144.6</td>
<td>3.4</td>
</tr>
</tbody>
</table>
sthetic hindrance for DIPA in the base-catalysed proton removal step (10).

In Figs. 9 and 10 our fitted rate equation lines for the A- and B-series of DEA and DIPA are shown together with literature data from Tables 4 and 5 at 25°C and our rate constant equation (21), extrapolated to a zero CO$_2$ liquid-load.

To DEA the following remarks apply. Neither the data of Sada et al. [3], nor those of Hikita et al. [19] and Sharma [7] agree well with our apparent rate constants and all data exceed our measurements, as well as those of Laddha and Danckwerts, significantly. The reason for these discrepancies cannot be derived from the original articles but might be caused by primary amine contaminants. On the other hand, Groothuis' [18] single data point coincides with our value at zero CO$_2$ liquid load. Laddha and Danckwerts' measurements [30] fall between our line calculated for zero CO$_2$ liquid-load and the experimental one for an average ~0.010 moles/L CO$_2$ load. This might indicate a small and varying CO$_2$ load in their experiments, but this was not specified in their work. The rate expression by Blanc and Demarais [5] agrees fairly well with the results in our experimental concentration range (0.39 < [DEA] < 2.3 mole/L), although their reaction order in DEA is lower (= 1) and constant. Part of this effect may be attributed to the varying CO$_2$ liquid load along their wetted-wall column. This affects the hydroxyl ion concentration at the different DEA concentrations in a systematic way and hence the bicarbonate reaction (5) contributes differently for each amine concentration to the overall absorption rate. This reaction (5) has a large influence on the overall reaction rate at low amine concentrations. In Fig. 11 the results of Blanc [5], corrected for reaction (5), are compared with the calculated values of our model.

A very strong support for our mechanism was obtained by extrapolating the work on DEA of Jørgensen [26] at 0°C and Jensen et al. [22] at 18 to 25°C. In their experiments a high pH was realized by the addition 0.1-0.3 M NaOH. Our calculations by rate equation (21) and their extrapolated measurements are shown in Table 10 and agree well if the extrapolation of (21) over ~3 pH units is taken into account. This agreement cannot be achieved by any other rate expression given in literature (e.g. [19, 30]) as these do not incorporate the effect of [OH$^-$] on the reaction rate.

For DIPA both Sharma's [47] and Groothuis' [18] results are relatively high, possibly due to contamination by primary amines as mentioned earlier.

**TEA and MDEA**

Also for TEA and MDEA, absorption experiments were carried out at 25°C with different amine concentrations and CO$_2$ liquid loads (see Tables 11 and 12 and Fig. 12).

Due to the absorption of CO$_2$ the hydroxyl ion concentration decreased continuously during an experiment and with that the contribution of the bicarbonate reaction (5) to the absorption rate decreased significantly. The resulting in $P_{CO_2}$-time plot, therefore, did not show a linear relation as predicted by eqn (20) because of the declining $k_{ov}$. This problem was overcome by calculating $k_{ov}$, for each interval between two pressure-time readings and individual correction of this $k_{ov}$ for the bicarbonate reaction (5) by eqn (21) to obtain an instantaneous $k_{app}$. These apparent rate constants did not vary over a large pressure range and hence were independent of CO$_2$ liquid load and [OH$^-$].

Had monoalkylcarbonate formation (7) been one of the prevailing reactions, an influence of [OH$^-$] would have been expected during the experiments. Using the third order rate constant for this reaction with TEA as extrapolated by Donaldson and Nguyen [17] ($k_3 = 1.53 \times 10^{10}$ 1$^3$/mole'sec) from the experiments of Jørgensen and Faurholt [25] and Jørgensen [26], the contribution of this reaction during absorption would have decreased from ~1 to ~0.2 sec$^{-1}$ during some of our experiments. This pseudo first order rate constant is in the same range as the measured ones and the decrease would have been observed if this occurred.

Based on our measured results, therefore, we conclude that no or negligible monoalkylcarbonate formation had taken place. As an independent check we carried out experiments similar to those of Chan and Danckwerts [11] but no substantial reaction products of CO$_2$ and TEA, e.g. monoalkylcarbonate were detected.
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The only way to explain the phenomena described above is by a base catalysis of the CO₂ hydration reaction (6) as proposed by Donaldson and Nguyen[17]. The essence of this catalysis is assumed to be a hydrogen bonding between the real free amine and water which increases the reactivity of water towards CO₂:

\[
R_3NH^+ + HCO_3^- \rightarrow H + R_3NH + CO_3^{2-}.
\]  

(26)

The reaction rate constants measured are \(k_2 = 2.9\) l./mole.sec for TEA and \(k_2 = 4.8\) l./mole.sec for MDEA respectively.

Comparing these rate constants together with the pKₐ's of TEA (pKₐ = 7.76 at 25°C[37]) and MDEA (pKₐ = 8.52 at 25°C[37]) a positive effect of amine basicity on catalytic activity is observed. As the pKₐ values of TEA and MDEA are lower than the actual solution pH (>9.5), the larger part of the amine is not protonated and therefore catalytically active. Donaldson and Nguyen[17] showed experimental results with triethylamine which they observed to have a negligible catalytic activity on reaction (26). The reason for this is the high pKₐ of triethylamine (pKₐ = 10.75 at 25°C[37]) which in their solution of pH ~ 9.5 results in a very low concentration of catalytically active unprotonated triethylamine.

Our rate constant measured for TEA corresponds very well with the result obtained by Donaldson and Nguyen[17] (\(k_2 = 2.0 - 2.8\) l./mole.sec) and Barth et al.[4] (\(k_2 = 2.85\) l./mole.sec). Compared to Hikita et al.[19] (\(k_2 = 50.1\) l./mole.sec at 25°C) and to Sada et al.[43] (\(k_2 = 16.8\) l./mole.sec at 25°C) our results and those of Donaldson and Nguyen are substantially lower. As for DEA and DIPA the reason for this very large difference can not be made clear from their articles but possibly originates from primary and secondary amine contaminants which accelerate the CO₂ absorption rate. In Donaldson and Nguyen, Barth et al.'s[4] and in our experiments this problem is avoided or at least reduced.

Fig. 9.
Fig. 10.

\( \text{ln } k \approx \text{Constant} \cdot \text{ln } [\text{DIPA}] \)

- Groothuis [8]
- Sharma [47]
- Our model

\( [\text{CO}_2] = 0.027 \text{ mole/l} \)

\( [\text{CO}_2] = 0.037 \text{ mole/l} \)

Fig. 11.

\( \text{ln } k \approx \text{Constant} \cdot \text{ln } [\text{DEA}] \)

- Blank and Demara [5]
- Our model
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Table 10. Comparison of our kinetic model with reaction rate constants extrapolated from Jørgensen [26] and Jensen et al. [22]

<table>
<thead>
<tr>
<th>Initial concentrations</th>
<th>Jørgensen [26] 0°C</th>
<th>Jensen et al. [22] 18°C</th>
<th>Extrapolation</th>
<th>Our model 25°C</th>
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<tr>
<td>0.1</td>
<td>0.1</td>
<td>0.0859</td>
<td>0.0949</td>
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<tr>
<td>0.2</td>
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<tr>
<td>0.1</td>
<td>0.2</td>
<td>0.0874</td>
<td>0.1931</td>
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</table>

Table 11. Experimental results for TEA at 25°C (present investigation)

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<tr>
<td>0.524</td>
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<td>0.667</td>
<td>0.0026</td>
<td>0.0114</td>
<td>11.4</td>
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<td>0.0097</td>
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<td>0.963</td>
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Table 12. Experimental results for MDEA at 25°C (present investigation)

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<td>0.0073</td>
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<td>34</td>
<td>14</td>
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<tr>
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<td>0.0224</td>
<td>36</td>
<td>16</td>
<td>5.4</td>
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<tr>
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<td>0.0162</td>
<td>51</td>
<td>27</td>
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by the substantial CO₂ liquid load which "neutralizes" the contaminants.

CONCLUSIONS

The available literature data provide a good basis for the calculation of the rate constant of the CO₂-MEA reaction, which is first order in the amine.

Our data measured on DEA and DIPA are very well described by the rate expression derived from the proposed mechanism. The results of Laddha and Danckwerts[30] agree well with our measurements. In their model, however, they do not incorporate H₂O and OH⁻ as bases and so arrive at a reaction rate expression which depends only on the amine concentration. Our data showed a significant dependence on [OH⁻] and, therefore, we incorporated [OH⁻] in our reaction mechanism. The good extrapolation of our model to the high pH measurements of Jørgensen[26] and Jensen et al.[22] is a strong support for the proposed mechanism.

With TEA and MDEA, no alkylcarbonate formation was observed. A base catalysis mechanism of the CO₂ hydration reaction fits the phenomena observed and is in agreement with Donaldson and Nguyen[17].

NOTATION

A interfacial area, m²
Am amine
B base
D diffusion coefficient, m²/sec
Eₐ infinite enhancement factor defined by eqn (18)
Ha Hatta number defined by eqn (17)
J mole flux, mole/m²sec

k₁ first order reaction rate constant, sec⁻¹
k₂ reaction rate constant sec eqn (9), sec⁻¹
kₚ app apparent rate constant defined by eqn (21), sec⁻¹
kₜ, kₜ₂, kₜ₃₉, kₜ₂OH⁻ rate constants defined by eqns (11) and (12), l./mole.sec
kₜₚ₄ liquid phase mass transfer coefficient, m/sec
kₜₚ₉₄ reaction rate constant for bicarbonate formation defined by (8), l./mole.sec
kₜₜ₉ overall rate constant defined by (13), sec⁻¹
m dimensionless solubility
nₚ app apparent reaction rate order in amine
P pressure, Pa
rₑ₀ rate of reaction (4), mole/l.sec
R gas constant, l.Pa/mole °K
t time, sec
V volume, m³
η viscosity, Pa.sec
v stoichiometric coefficient

REFERENCES

A study on the reaction between CO₂ and alkanolamines in aqueous solutions