QUANTITATIVE SPECIATION OF THE LIQUID PHASE BY FTIR SPECTROSCOPY IN THE SYSTEM AMP-PZ-CO$_2$-H$_2$O

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This work reports a multivariate calibration partial least square regression (PLS) model to quantify the liquid-phase concentrations of 2-aminoo-2-methyl-1-propanol (AMP), piperazine (PZ), bicarbonate, PZ monocarbamate, and PZ dicarbamate during the absorption/desorption process by Fourier Transform Mid-Infrared spectroscopy (mid-FTIR). The model could predict 33 different concentrations ranging from 0 to 40 wt.% AMP, 0 to 15 wt.% PZ, and 0 to 12 wt.% total CO$_2$ with relative errors lower than 10%, and 87% of the variance of all samples have been represented (R$^2$), except for PZ dicarbamate. A comparison of a single PLS2 model was made with multiple PLS1 models, one for each chemical species present in the liquid phase. The latter had better predictions and made it possible to differentiate CO$_2$ from its chemical forms, allowing a better understanding of the CO$_2$ capture processes.

Keywords: amine blend; carbamate; carbon capture; infrared spectroscopy.

INTRODUCTION

The CO$_2$ capture process is one important and challenging issue of the moment, motivating intense research and investments.\textsuperscript{1-5} The principal technology used to reduce CO$_2$ emissions is the chemical absorption process using aqueous amine solutions\textsuperscript{6} due to its higher removal rate and efficiency in dilute gas streams.\textsuperscript{1-3} The main drawback of this process is the high energy required to regenerate the solvent.\textsuperscript{7,8}

The use of the 2-aminoo-2-methyl-1-propanol (AMP) and piperazine (PZ) mixture as a solvent has the advantage of AMP requiring a low energy demand for its regeneration, despite the moderate rate of absorption with CO$_2$. And PZ provides higher reaction velocity at the expense of average energy demand for desorption.\textsuperscript{2,4} This mixture of solvents was extensively studied, highlighting its advantages.\textsuperscript{4,5,9-12}

The determination of an accurate mass transport coefficient is necessary to design, optimize and evaluate different capture processes.\textsuperscript{15} Thus, it is required to know the composition of the chemical components present in both phases to characterize the process. Fourier transform infrared (FTIR) spectroscopy integrated with the multivariate regression method allows the simultaneous quantification of the chemical species in a fast, non-invasive manner, and ensures the integrity of the samples, ideal characteristics for online monitoring of the CO$_2$ capture process.\textsuperscript{14,15}

FTIR has been used to study CO$_2$ capture processes. Mergler et al.\textsuperscript{16} identified the spectrum bands associated with reaction products of the MEA-CO$_2$-H$_2$O system. Geers et al.\textsuperscript{17} built a PLS model to estimate the CO$_2$, SO$_x$, and β-alanine concentration from FTIR spectra in a post-combustion capture pilot plant. Diab et al.\textsuperscript{18} used FTIR spectroscopy to speciate the liquid phase in an aqueous solution of diethanolamine (DEA) and CO$_2$ systems for different CO$_2$ loadings, mass fractions, and temperatures. van Eckeveld et al.\textsuperscript{19} developed inverse least-squares models for the online liquid analysis of the MEA-CO$_2$-H$_2$O system using FTIR spectroscopy, density, conductivity, refractive index, and sonic speed measurements as input data. van der Ham\textsuperscript{20} included solvent degradation products of an MEA-CO$_2$-H$_2$O system in the calibration model and obtained worse predictions. Kachko et al.\textsuperscript{15} built a model to quantify MEDEA, PZ, and CO$_2$ concentration based on the NIR spectra, density, pH, conductivity, sound velocity, and refractive index. Kachko et al.\textsuperscript{20} compared the prediction of captured CO$_2$ concentration in aqueous MEA solutions using Raman, NIR, and ATR-FTIR spectroscopy. du Preez et al.\textsuperscript{21} developed an FTIR spectroscopy method to study the CO$_2$ reaction with monoethanolamine (MEA) in an n-propanol solvent. They collected real-time reaction kinetic data using an attenuated total reflectance (ATR) probe confirming that the zwitterion reaction mechanism accurately describes the MEA reaction in a non-aqueous system.

FTIR was used to identify the amines, carbamates, and bicarbonate during the CO$_2$ absorption in aqueous solutions of AMP\textsuperscript{22,23} And for cyclic amines (e.g., PZ).\textsuperscript{24} Ermatchkov et al.\textsuperscript{25} used 1HNMR spectroscopy to speciate an aqueous PZ-CO$_2$ system. Kachko et al.\textsuperscript{14} monitored an absorption process using an AMP-PZ solvent. They used NIR spectroscopy and physical data to build a chemometric model. However, they did not work with CO$_2$-saturated amines. Li et al.\textsuperscript{26} developed a rigorous thermodynamic model in Aspen Plus using the electrolyte non-random two liquid activity coefficient (e-NRTL) model for the system PZ-AMP-CO$_2$-H$_2$O and compared to NMR speciation. Zanone et al.\textsuperscript{3} simultaneously quantified AMP, PZ, and total CO$_2$ in all its chemical forms in the liquid phase. However, they did not distinguish carbamates, carbonates, or soluble CO$_2$.

The main drawback of the absorption using aqueous solutions of amines to absorb CO$_2$ is the high energy needed to regenerate the solvent.\textsuperscript{4,27} This energy is directly related to the stability of the chemical products formed during the reaction with the amines. Carbamates formed due to reaction with primary amines (e.g., MEA) are more stable than carbamates formed from secondary amines (e.g., DEA), for example. Consequently, less energy is required to regenerate the latter.\textsuperscript{28} Furthermore, this means that knowing the CO$_2$ concentration in these solvents is not enough, as the different chemical forms of CO$_2$ will have an impact on the energy needed and the desorption rates.

This study aimed to develop an FTIR spectroscopy method for online speciation of the liquid phase of the chemical absorption/desorption process of the AMP-PZ blend solvent over the entire
We used potentiometric titration (TitroLine easy, Schott Instruments) using 0.5 M HCl to obtain the concentration of AMP and PZ from samples 1 to 13 (non-carbonated pure amines standard samples) in triplicate. Since the equivalence points of AMP and PZ overlap, we calculated the concentration reference value based on the gravimetric data corrected by the purity of the individual amines according to the previous analysis procedure.\textsuperscript{9,20,30} After the carbonation, the titration resulted in the total CO\textsubscript{2} absorbed in all its chemical forms.\textsuperscript{9,20,30}

The titration method could not distinguish among the different CO\textsubscript{2} species (i.e., bicarbonate, PZ mono- and dicarbamate). And there was no access to other analytical equipment to overcome this problem (e.g., NMR spectroscopy). AMP reacts with CO\textsubscript{2} to form AMP carbamate and rapidly reacts with another base resulting in bicarbonate and protonated AMP.\textsuperscript{9,23,31} Additionally, the carbamate formation is ten times lower than the bicarbonate.\textsuperscript{31} Therefore, we considered that all CO\textsubscript{2} quantified in the saturated solutions of AMP was bicarbonate.

Furthermore, PZ reacts with CO\textsubscript{2} to form PZ monocarbamate (PZCOO\textsuperscript{-}), which could react again with CO\textsubscript{2} to form PZ dicarbamate (OOCPZCOO\textsuperscript{-}).\textsuperscript{9,23,34} Consequently, all CO\textsubscript{2} quantified in the saturated solutions of PZ was considered as only PZ dicarbamate. And, we considered that the first spectrum acquired (first 30 seconds of CO\textsubscript{2} absorption) referred to PZ monocarbamate only.

Models

We generated multiple PLS models, one model for each chemical species: non-reacted AMP, non-reacted PZ, bicarbonate, PZ monocarbamate, and PZ dicarbamate. The region used was obtained as reported in past work\textsuperscript{9} using a changeable size moving window (CSMW)\textsuperscript{32} using a script written in MATLAB. The same region was implemented in the models built in the equipment software iC IR\textsuperscript{TM} v4.2.26 to validate the MATLAB script and allow online quantification of the samples.

Altogether, 568 spectra samples were used divided into 270 spectra training-set and 298 spectra validation-set using a leave-one-out cross-validation. It was used 20 spectra for all 26 solutions described in Table 1 randomly separated into the training and validation sets, and 10 spectra for the 7 carbonated solutions prepared from the reaction of samples 20 to 26 (Table 1) with CO\textsubscript{2}. From the solutions with CO\textsubscript{2}, only the saturated sample spectra were randomly chosen to be in the training set, while the partially carbonated solution was all present in the validation set.

All data were mean-centered, and other pretreatments were made specifically for each species, as detailed in Table 4. Also, the wavenumber range used in the PLS model was different for each species. The minimum root-mean-square error of prediction (RMSEV) determined the number of factors.

### Table 1. Concentration of the non-carbonated samples

| Sample | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8  | 9  | 10 | 11 | 12 | 13 |
|--------|----|----|----|----|----|----|----|----|----|----|----|----|----|
| AMP (wt.%) | 5  | 10 | 15 | 20 | 25 | 30 | 35 | 40 | 0  | 0  | 0  | 0  | 0  |
| PZ (wt.%)  | 0  | 0  | 0  | 0  | 0  | 0  | 3  | 6  | 9  | 12 | 15 |
| Sample    | 14 | 15 | 16 | 17 | 18 | 19 | 20 | 21 | 22 | 23 | 24 | 25 | 26 |
| AMP (wt.%) | 28 | 26 | 24 | 22 | 20 | 18 | 15 | 30 | 0  | 0  | 21 | 12 | 5  |
| PZ (wt.%)  | 2  | 4  | 6  | 8  | 10 | 12 | 0  | 4.5| 9  | 9  | 3  | 15 |
RESULTS AND DISCUSSION

Figure 2 shows the spectra of all 33 different concentrations solutions (26 concentrations shown in Table 1 plus 7 carbonated solutions) used to build the PLS models.

Spectra information

Table 2 summarizes the wavenumber, the band attribution, and the species concerned. We used these as the starting window for the CMSW method to build the PLS models. Table 4 specifies the final region chosen.

Samples

The titration in triplicate of all samples is on the supplementary material. Comparing the weighed mass and the titration concentration allowed us to predict the AMP and PZ purities as 91±1% and 98±1%, respectively. The titration standard deviation error was 0.08 wt.%.

PLS2 model

We already developed a single PLS2 model to predict AMP, PZ, and CO2 absorbed in all its chemical forms (i.e., carbamates and carbonates). This model was obtained with a single-window at 1690-846 cm⁻¹ without any pretreatment using 8 factors. Altogether, 568 spectra samples had their concentration predicted with relative errors lower than 10% and 93% of the variance of the samples represented. Although the data variance could be well reproduced (R² > 0.999 and Q² > 0.9), as shown in Table 3, it had a drawback in predicting the PZ concentration in some of the CO2 saturated solutions. This probably happens due to the PZ peak at 1326 cm⁻¹ that gets overlapped by the PZ carbamates formation. Table 3 shows the training coefficient of determination (R²), the validation coefficient of determination (Q²), the root-mean-square error of calibration (RMSEC), and the root-mean-square error of prediction (RMSEP).

Table 3. Coefficient of determination and root-mean-square errors for the PLS2 model

| Component | Factors | R²      | Q²      | RMSEC | RMSEP |
|-----------|---------|---------|---------|-------|-------|
| AMP       | 9       | 0.99999 | 0.93252 | 0.1530| 0.825 |
| PZ        | 9       | 0.99988 | 0.98172 | 0.1400| 0.904 |
| CO2       | 9       | 0.99999 | 0.99398 | 0.0186| 0.223 |

PLS models

We developed one PLS model for each chemical component identified in the liquid phase: AMP, PZ, HCO3⁻, PZCOO⁻, and OOCPPZCOO⁻. The models’ specifications are shown in Table 4.
The region in 1303-836 cm\(^{-1}\) contained all the information used in all models, meaning we could gather more information with less data than the PLS2 model. Figure 3 shows the regions used in the model.

The AMP model used the C-N stretching band (\(\nu\)) region, which decreased and shifted, distinguishing from the \(\nu\)C-O as AMP got protonated to AMPH\(^+\). The PZ was predicted based on the disappearance of the medium stretching (\(\nu_{\text{m}}\)) of the CN band. The PZ carbamates models (PZCOO\(^-\) and \(\text{OOCPZCOO}^-\)) used the symmetrical and asymmetrical \(\nu\)N-COO\(^-\) bands. As both overlaps, we applied a second derivative pretreatment to distinguish them. Finally, the bicarbonate model used the out-of-plane deformation \(\gamma\)CO\(_3\) mode.

Altogether, 568 spectra samples had their concentration predicted with relative errors lower than 10% and 87% of the variance of all samples represented. Table 5 shows the \(R^2\), \(Q^2\), RMSEC, and RMSEP. The model drawback is the PZ dicarbamate with half \(R^2\) compared to the other species. Probably, the assumption that the CO\(_2\) saturated PZ solutions contained only PZ dicarbamate is weak. Probably dicarbamate converted to bicarbonate at rich loadings.\(^{26}\)

Figure 4 (a) compares the concentration predicted by the PLS model and the titration reference value. The residues are shown in Figure 4 (b) and seem random. Since the titration does not distinguish the CO\(_2\) absorbed into its different chemical forms, we only present in Figure 4 the amines and total CO\(_2\) concentration.

Figure 5 shows the prediction of the mono- and dicarbamate of PZ and bicarbonate in the 5 wt.% AMP 15 wt.% PZ blend during the carbonation process. At the beginning of the reaction, there was a rapid formation of PZCOO\(^-\) which then reaches a maximum concentration. At the same time, monocarbamate is transforming into \(\text{OOCPZCOO}^-\) at a lower rate and rapidly after the monocarbamate reaches the maximum concentration. This faster dicarbamate formation is associated with the total consumption of free PZ. Although the \(\text{OOCPZCOO}^-\) prediction is not reliable, this profile agreed with published works.\(^{9,23,24}\)

In the blend, AMP acts as a base that catalyzes the PZ reaction with CO\(_2\), then HCO\(_3^-\) does not have a significant concentration. Moreover, a higher spectral noise present at wavenumbers below 900 cm\(^{-1}\) reflect a higher deviation on the bicarbonate concentration prediction.

### Models comparisons

In this section, we compare our newly developed PLS models

| Component | Factors | \(R^2\) | \(Q^2\) | RMSEC | RMSEP |
|-----------|---------|---------|---------|--------|--------|
| AMP       | 10      | 0.99999 | 0.99733 | 0.1268 | 1.5400 |
| PZ        | 7       | 0.99999 | 0.99398 | 0.1804 | 0.6517 |
| PZCOO\(^-\) | 1      | 0.90358 | 0.81104 | 0.82579 | 1.5529 |
| \(\text{OOCPZCOO}^-\) | 1      | 0.45810 | 0.31482 | 0.43613 | 1.3462 |
| HCO\(_3^-\) | 1      | 0.99398 | 0.98015 | 0.25516 | 1.3963 |

### Table 4. PLS specifications for each species model

| Species concerned | Region (\(\text{cm}^{-1}\)) | Pre-Treatment | Factors |
|-------------------|-----------------------------|---------------|---------|
| AMP               | 1082-1026                  | 9 points Savitz-Golay | 10      |
| PZ                | 1140-1080                  | 9 points Savitz-Golay | 7       |
| PZCOO\(^-\)       | 1303-1250                  | second derivative | 1       |
| \(\text{OOCPZCOO}^-\) | 1303-1250                | none           | 1       |
| HCO\(_3^-\)       | 845-836                    | second derivative | 1       |
with the previous PLS2 model. Although a univariate model would be much simpler and would work as multivariate models to quantify the pure amines. Thanks to the peaks shift and strong overlapping during the carbonation process, the multivariate model had better predictions for the carbamates.

Although both models, PLS and PLS2, had a similar prediction of the concentrations of AMP and PZ, the calibration model uses only the non-carbonated (first point) and the saturated solution (last point). If we compare the middle points, the concentration during the partial carbonation process as shown in Figure 6, the prediction of these amines’ concentration was indeed different. We chose these samples as they represent the worst case.

Figure 7 compares the AMP concentration prediction of two PZ solutions. The PLS model performed much better, as no AMP existed on these solutions. The same happened in all PZ solutions, more intensely in the concentrated ones, as shown in Figure 7 (a) and (b). The PZ monocarbamate may be associated with this deviation on the PLS2 model as the AMP prediction happens only at the beginning of the carbonation process. As the partially carbonated solution were all present in the validation set, it can be noted the large increase in the mean error of prediction shown in Tables 3 and 5.

When comparing the PZ prediction in the AMP solutions (Figure 8), the same behavior happens. The points used in the calibration model correctly predicted the zero PZ concentration on both models. When comparing the partial carbonation, the PLS2 failed to predict no PZ.
We could infer that the PLS2 model attributed the CO₂ reaction products formed to the quantification of one of the amines due to a broader region used in the model. The PLS model used smaller windows for each species than the PLS2 model. Even gathering all regions together, it was still smaller. So the model had less information and noise, which lead to a better prediction during the carbonation process. Also, we could have overfitted the PLS2 model.

CONCLUSIONS

Using only FTIR spectroscopy allowed the quantification of AMP, PZ, total CO₂, PZCOO⁻, and HCO₃⁻ enabling online monitoring of the absorption/desorption process of a CO₂ capture process.

The PLS models showed a significant improvement in predicting the intermediate points of the absorption process compared to the previous PLS2 model. Furthermore, this work used a narrow spectrum region and could extend the quantification of the carbonated reaction products of the amine blend CO₂ reaction.

The differentiation of the CO₂ chemical forms allows a better understanding of the absorption/desorption process in real-time and process optimization since the different CO₂ species need distinctive regeneration temperatures and have varying reaction rates.

SUPPLEMENTARY MATERIAL

The spectra absorbance, concentration, and titration of each sample are available at http://quimicanova.sbq.org.br as an Excel file, with open access.

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