



Proceeding Paper Simulation-Based Techno-Economic Assessment of a Water-Lean Solvent for Natural Gas Sweetening Technology *

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Abstract: Water-lean solvents are thought to deliver promising benefits to include enhanced mass transfer properties, increased absorption capacities and lower solvent regeneration heat duties in natural gas sweetening process. Acid gas (H2S and CO2) removal is an essential piece of the natural gas value chain due to the corrosive effect on pipeline and process equipment, impact on environment and reduction in methane heating value. A number of solvents have been used for this process in the past. However, low acid gas pickup, high cost per unit separation and high regeneration heat duties forms the basis for which we considered a water-lean solvent in this study. This study employs ASPEN HYSIS V12.1 to model natural gas sweetening process of a hypothetical non- associated sour gas well with a novel water-lean solvent (50% wt. MDEA + 30% wt. DIPA + 15% wt. DMSO + 5% wt. H2O). Theoretical solvent screening was carried out to select the most promising waterlean solvent, following a flowsheet design, modeling and result validation. The process economic analysis was carried out using Aspen Process Economic Analyzer to determine the unit separation cost and profitability indicators. Results show that the solubility of CO2 was found to be lower in water-lean solvents. The mass transfer seem not to be better. This was generally difficult to be taken into consideration and assessed properly in Aspen as there is no literature data with DMSO to fit the model parameters. DMSO reacts with H2S, leading to loss of solvent. Although, mass transfer improved with physical co-solvent, acid gas solubility decreased, resulting in more solvent consumption and impacting the capital expenditure. Economic analysis showed that the equipment cost of the proposed solvent is 1.4 M USD/yr higher than that of the aqueous MDEA commercially in use. As such, the it is not considered economically viable.

Keywords: natural gas sweetening; separation process; solvent; absorption; mass transfer

1. Introduction

One of the most significant sources of energy is natural gas, and it can be utilized in a variety of ways as fuel for vehicles, electricity, fertilizer production, and many other chemical industries. [1]. Natural gas has methane as the predominant compound (CH4). [2]. However, it contains several other undesirable impurities such as hydrogen sulfide (H₂S), carbonyl sulfide (COS), carbon disulfide (CS₂) and mercaptans (R-SH) along with the existence of different percentages of nitrogen (N₂), carbon dioxide (CO₂) and other gases [3]. In its processed form, it is one of the most efficient, reliable, and most practical of all energy sources [4]. In power plants, natural gas emits between 50 and 60 percent less carbon dioxide (CO₂) than conventional oil or coal-fired power plants [5]. Natural gas reserves are found in abundance in quite a number of countries in the world, with Russia, Iran, and Qatar topping the list to hold around half of the world's proven reserves [6]. As of April, 2022, Nigeria's proven gas reserves totaled 206.53 trillion cubic feet (Tcf), predominantly in its Niger Delta region, ranked in the top 10 globally and making up around

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Copyright: © 2023 by the authors. Submitted for possible open access publication under the terms and conditions of the Creative Commons Attribution (CC BY) license (https://creativecommons.org/license s/by/4.0/). 3% of the global natural gas reserves [7]. As such, it plays a substantial part in the recent world energy economy and development [8].

Prior to the transmission of natural gas through pipelines, its sweetening, referring to the removal of H₂S and CO₂ (acid gases), is required due to the associated technical difficulties encountered, such as corrosion of pipelines and environmental concerns [10]. The maximum H₂S and CO₂ concentrations allowed in natural gas are 4 ppm and 2 mol%, respectively [11]. There are a number of procedures to eliminate acid gases from unprocessed natural gas, including chemical (absorption) solvents, physical solvents, adsorption, membranes, and cryogenic fractionation. In the chemical (absorption) solvent or amine process which is of interest to us in this study, carbon dioxide and hydrogen sulfide are removed using amine solutions [8]. The H₂S and sulfur compounds are selectively removed by the amine solution during the sweetening process, and some CO₂ is also absorbed because the concentration of CO₂ in the gas must be lower than the permitted concentration for the pipeline specification [1].

Since the amine process has been used on a large scale to treat natural gas for more than 80 years, it is thought to be the most widely used procedure in the gas processing industry [12]. Examples of amine processes currently in use include aqueous amine, chemical mixture solvents, and the shell sulfinol process. The solvent price, reboiler and cooling duty can be reduced by using a mixture solvent. The system by a chemical mixture solvent offers better performance than other solvents, especially in comparison to regular aqueous solvents [2]. However, the high energy demand required for regeneration has limited its usage. The most recent and widely used technology for acid gas removal as of now is the Shell Sulfinol process. This employs Diisopropanolamine (DIPA), Sulfolane, and water to offer the main advantages of low solvent circulation rates, low plant cost, and high effectiveness for acid gas removal [13]. Solvent utilization for acid gas removal has witnessed several industrial challenges such as vaporization, high hydrocarbon pickup, low acid gas pickup, high cost per unit separation, and high regeneration heat duties. These technical challenges are generally responsible for low solvent effectiveness, giving room for an increased presence of acid gas in the sweet gas stream. The result of which translates to natural gas pipeline & process equipment corrosion and low gas heating value. As such, the development of high-performance solvents at a lower cost is of critical importance in acid gas removal from natural gas in order to meet the requirements of commodity gas quality standards [14].

With previous studies for CO₂ absorption rate increment in MDEA suggesting the addition of a small amount of primary or secondary amines (such as DEAs or DIPA) to the solution to combine their high reactivity with the little energy requirements for regeneration of tertiary amines (MDEA) [32]. In this study, we will be assessing the techno-economic viability and performance of a water lean solvent containing a mixture of methyl diethanolamine (MDEA), di-isopropanol amine (DIPA), dimethyl sulfoxide (DMSO), and water for the natural gas sweetening process. This expands the analysis carried out by previous researchers while focusing particularly on these proposed amines and organic diluents. Both DIPA and methyl diethanolamine are made commercially and offered for sale. These chemical processes allow the amines to absorb acid gas.

Water-lean solvents, refering to a mixture of organic diluents with amines are thought to deliver benefits in carbon capture and gas sweetening systems, including increased H₂S absorption capacities and lower regeneration heat duties. However, solubility of CO₂ is always lower [15]. The aim of this study is to carry out a Simulation-based Techno-Economic Assessment of a water-lean solvent for the natural gas sweetening process using ASPEN HYSIS V12.1. Theoretical screening of organic diluents and amines was carried out to select the seemingly most promising solvent for the process, following a feed gas composition data collection from a non associated gas field. A design of the flow-sheet and thermodynamic properties modeling of a water-lean solvent based natural gas sweetening plant was done and a replica simulation of the process, replacing the water-lean solvent with an aqueous solvent was carried out to compare results for solvent

performance evaluation. Lastly, the economic viability and performance of the water-lean solvent for the gas sweetening process were evaluated using Aspen Process Economic Analyzer.

2. Process Description

This section discusses the various methods that was employed in carrying out the simulation and economic analysis of this study. The process flowsheet for amine absorption and stripping was designed in ASPEN HYSYS V12.1. Setting up a mixture of the component list and the fluid package forms the initial stage of this simulation. The property package is based on the acid gas-chemical solvent package model and vapor-liquid equilibria (VLE), which was selected to be the H₂S-CO₂-MDEA-DIPA-DMSO-H₂O electrolyte system. The Electrolyte Non-Random Two-Liquid (eNRTL) model for electrolytic thermodynamics and Peng-Robinson (PR) for the estimation of gas phase parameters are the foundations of the acid gas fluid package. For all amine solvents utilized in the industry, including MDEAs, MEAs, and Sulfolane-MDEA, regression has been done using the existing VLE and heat of absorption data [2]. Then, simulation of the Natural Gas Sweetening Plant flowsheet was carried out, and the effect of flow parameters on the process performance was tested by variation. An economic analysis of the process was done using a Microsoft Excel Spreadsheet to determine the solvent separation cost.

In the formulation of the gas sweetening solvents, 30 mol% of diisopropanolamine is added in a blend (DIPA & MDEA), combining their high reactivity with the low energy requirement for regeneration of tertiary amine, 50 mol% of MDEA. 15 mol% DMSO and 5 mol% water solution formed the diluent in this study. Sour feed gas with a composition as shown in Table 1. enters the filter separator V-100, with a 40Kpa of vapour outlet pressure drop to separate solid and liquid impurities from natural gas. At 32.22 °C, this vapour outlet V1 enters from the bottom of the absorption column(s) T-100 containing 0.09 mol% CO₂ and 0.12 mol% H₂S.

Component	Content (mol %)
Methane (CH4)	0.69000
Ethane (C ₂ H ₆)	0.06000
Propane (C3H8)	0.00960
i-Butane (i-C4H10)	0.00260
n-Butane (n-C ₄ H ₁₀)	0.00290
i-Pentane (i-C5H12)	0.00140
n-Pentane (n-C5H12)	0.00120
n-Hexane (C ₆ H ₁₄)	0.00180
n-Heptane (C7H16)	0.00650
n-Octane (C ₈ H ₁₈)	0.00140
Nitrogen–N2	0.00160
CO ₂	0.09210
H ₂ S	0.12000
H ₂ O	0.01200

Table 1. Composition of the feed gas.

Figure 1 shows the process simulation of the natural gas sweetening plant.



Figure 1. Natural Gas Sweetening Process Model in ASPEN HYSIS.

As shown in Figure 1, the amine solution of 574,000 kmol/hr is fed from the top of the absorption column at 5900 kPa while maintaining a temperature of 50.0 °C. With the solution in close counter-current contact, the input gas V1 ascends through the column from the bottom of the absorber (T-100). The absorption column consists of 18 stages with a top and bottom pressure of 5600 kPa and 5900 kPa, respectively. The wet sweet gas from the sulfur removal unit at the top of the absorption column (T-100) with a temperature of 50.16 °C is sent to the gas scrubber V-101 (gas-liquid (flash) separator) to separate the liquid droplets and heavy key hydrocarbons that purified gas brings out from the tower. The wet sweet gas from the vapour outlet V3 is then fed to a pressure relief valve (VLV-103), reducing the pressure to 4800 kPa. Rich amine from the bottom of the absorber column (T-100) is fed to a valve (VLV-100) with a pressure drop of 200 kPa and mixed with the liquid outlet L3 from the flash vessel to be fed to a horizontal flash separator (V-102) at 5600 kPa and 82.33 °C. Following the exit of V4 from the flash separator, a pressure relief valve (VLV-102) reduces the pressure to 4900 kPa at 79.63 °C. This is combined with the vapour fraction V3 from (V-101) using (MIX-101) and fed to the bottom stage inlet of a second absorber column (T-101) at 46.49 °C, 4800 kPa, and 5413000 kg/hr. The column operates with a top stage inlet amine recycle of 50 mol % MDEA, 30 mol % DIPA, 15 mol % DMSO, and 5 mol % water at 4700 kPa and 51 °C, with a bottom and top stage pressure of Pn, 4650 kPa, and P1 of 4550 kPa respectively. The sweet gas then enters to the user with a final methane mole fraction of 83.11%, 0.062 mol % CO2 and 0.0051 mol % H2S respectively. Rich amine pumped by P-100 out of the bottom of the rich amine flash vessel exchanges heat with the hot lean amine coming from the bottom of the amine regenerator (stripper column) through the lean/rich heat exchanger (E-100), with a cold and hot fluid pressure drop of 34 kPa and 2 kPa respectively. The outlet rich amine stream is fed to the amine regenerator at a temperature of 88 °C. The stripper column full reflux condenser type with a 20-stage operation operates at a condenser & reboiler pressure of 241.3 kPa and 262.0 kPa, respectively, enabling the hydrocarbons, H₂S, and CO₂ to be stripped out of the rich solution. The amine feed to the regenerator enters at an inlet stage of 6, allowing for the column to operate with a parameter solver of 0.1 fixed damping factor and a stage numbering top-down. The hot lean amine with a hot fluid outlet temperature of 96 °C and a mass flowrate of 1,150,400 kg/hr. from the heat exchanger is pumped by a hot lean amine pump (P-102) until the pressure outlet reaches 350 kPa and sent to a hot lean amine air cooler (E-101) with an outlet temperature of 48.59 °C. The outlet stream is then pumped

with (P-101) to a further pressure of 5900 kPa in preparation for recycling following a stream split into two branches employing the SET blocks, SET1 and SET2 for temperature and pressure respectively, with a source from the initial feed amine. The resulting amine stream is then fed to the make-up vessel (MAKEUP-100) in combination with the water make-up, accounting for the loss as a result of vaporization due to temperature increase in the regeneration unit. The outlet stream is recycled (RCY-1) as a top feed stream for the second absorber column (T-101).

3. Results

In this section, the result of the study is presented, analyzed, and discussed in accordance with the aim, which is to carry out a Simulation-based Techno-Economic Assessment of a water-lean solvent for natural gas sweetening process. The amine absorber utilized a novel solvent, designed to achieve higher absorption capacity and reduce heat regeneration tasks in a heat integrated flowsheet of a gas sweetening plant, as described in Sections 2 and 3 of this study.

In an attempt to verify the authenticity and validity of the simulation results, key parameters from the simulation model are authenticated against guidelines from the Gas Processing Supplier Association (GPSA).

Figure 2 shows that increase in absorber solvent flowrate decreases the composition of carbon dioxide and hydrogen sulfide in the sweet gas. This implies that; the more the solvent flow, the more efficient the acid gas loading and the less the carbon dioxide and hydrogen sulfide content in the sweet gas. Hence, solvent flowrate is inversely proportional to carbon dioxide and hydrogen sulfide content in the sweet gas.



Figure 2. H₂S and CO₂ against solvent flowrate (kmol/hr).

Figure 3 presents the effect of feed gas flow rate on the amount of H₂S and CO₂ in sweet gas stream. As seen, increase in feed gas flowrate leads to increased impurity H₂S and CO₂ content in the sweet gas stream. This is because as feed gas flow rate is increasing, the chances of acid gas and solvent molecule reaction becomes limited due to the flow velocity. Also, as we increase the feed gas flowrate, it means that the initial sour acid gas content is also increasing by mass, hence, an increment of impurities should be expected in the sweet gas stream.



Figure 3. Effect of feed gas flow rate on the amount of H₂S and CO₂ in sweet gas stream.

4. Conclusions

In line with the aim of this study being to carry out a Simulation-Based Techno-Economic Assessment of a Water-Lean Solvent for Natural Gas Sweetening, a comparison of aqueous solvent and the novel solvents in this study was carried out. Solubility of CO₂ was found to be lower in the water-lean solvent under study. The mass transfer seem not to be better. This was generally difficult to be taken into consideration and assessed properly in Aspen as there is no literature data with DMSO to fit the model parameters. Because water-lean solvents are not well represented in Aspen Plus, experimental data was employed to validate the models, evaluating the performance of the water-lean solvent against aqueous MDEA. In the water-lean solvent, the cycli capacity was found to be smaller, moving towards lower loadings. The vapour liquid equilibrium (VLE) curves moved to the left in loading CO₂ partial pressure as with the aqeous MDEA solvents. However, aqueous MDEA outperforms the proposed waterlean solvent under study. Although mass transfer improved with physical co-solvent in the waterlean solvent, the acid gas solubility decreases. Hence, more solvent will be needed and larger towers required, impacting the capital expenditure (CAPEX). Dimethyl Sulfoxide (DMSO) was also found to react rapidly with H₂S. As such, acting as a co-solvent for this process led to rapid solvent degradation. The water-lean solvent treated the sour gas to a removal of 0.7 mol of acid gas per mol of solvent and showing a final sweet gas CO₂ and H₂S concentration of 4.1 mol% and 0.142 mol% respectively. This does not meet the gas pipeline specification or permissible limit of 4% mol CO₂. Especially given that the solubility of CO₂ is always lower in waterlean solvents. Total utility energy saving of approximately 1.9×10^5 KJ/h was recorded in the aqueous MDEA plant against the waterlean solvent. Economic analysis showed that the total CAPEX savings from aqueous MDEA acid gas removal unit over the waterlean solvent under study is \$6900/MMSCFD and the proposed waterlean solvent is \$1.4 M/yr higher in terms of equipment cost than aqueous MDEA. As such, the waterlean solvent under study is not considered economically viable in comparison to current industrial practices.

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