Title: Estimating an Achievable Target Price to Regenerate Bio-Oils Post Hydrogen Sulfide Removal

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Abstract

Bio-oils offer valuable use as bio-solvents for removing hydrogen sulfide (H\textsubscript{2}S) from natural gas. Preceding bench-scale studies indicate that greater than 90\% of H\textsubscript{2}S can be removed from a gas stream; economic analysis of such a process is necessary to determine solvent regenerative power required and price limits on a to-be-determined solvent regeneration scheme. With a processing goal 1000 kmol/h of sour gas and removing 99.9\% of H\textsubscript{2}S from gas streams at variable feed concentration, design of an absorption unit to process natural gas using bio-oils was carried out through equilibrium stage analysis. Comparison to conventional amine gas treating was used as a cost threshold for gas treatment. The economic viability of using bio-oils as gas sweetening agents depends on capability of regenerating and recycling more than 98\% of the soybean oil bio-solvent to compete with amine gas treating, the most popular industrial method.

Keywords

Biomaterials; bioreactor design; bioseparations; biotechnology; modeling
Introduction

The use of bio-oils – conventional soybean, high oleic soybean, canola, and sunflower – to remove hydrogen sulfide (H\textsubscript{2}S) from a gaseous mixture has been demonstrated\textsuperscript{1} and has potential for use in the natural gas industry to sweeten gas as it is extracted. High concentrations of H\textsubscript{2}S in natural gas render it sour, with sour gas defined as natural gas containing anywhere from 4 ppmv to thousands of parts per million H\textsubscript{2}S.\textsuperscript{2} Bio-oils have demonstrated the ability to remove up to 90% of gaseous H\textsubscript{2}S from a gas stream at the bench-scale\textsuperscript{1} and could be used in a scheme similar to the industrially prevalent amine gas treating methods.\textsuperscript{3,4}

In an effort to evaluate if further investigation into bio-oils as gas sweetening agents could yield a viable process, the economic feasibility must be assessed. The present work aims to determine the number of stages required to remove 99.9% of H\textsubscript{2}S from methane feed gas with varying concentrations of H\textsubscript{2}S, to examine the capital costs, determine the required regenerative power of the bio-oil solvent, and ultimately determine the maximum cost of regenerating and recycling the bio-oils to be competitive with currently implemented gas processing methods.

Methods

In considering the price of using bio-oils as extraction solvents for treating sour gas and removing H\textsubscript{2}S, the cost of such a process would need to be less than or equal to the existing industrial processes for treating sour natural gas. Industrial natural gas production plants using amine gas treating methods typically spend $4/GJ on gas treating.\textsuperscript{5} To draw a comparison between using bio-oils such as conventional soybean or high-oleic soybean oil as the gas sweetening solvent, the factors that impact the cost of the process must be verified. The cost of the absorption unit – an extraction column – is a key part of the capital cost estimation. By constructing equilibrium stage diagrams an extraction column can be designed, which can then be used to estimate capital costs.
Sensitivity analysis, with respect to bio-oil solvent regeneration capability, can also aid in understanding how reliant the process viability is on being able to recycle the bio-oil solvents. Recovered sulfur could be a lucrative byproduct, as sulfur is valued at more than $200/ton as a chemical building block. As a baseline comparison, the overall cost can be set equal to the current cost of amine gas treatment ($4/GJ); from there, the total maximum cost of bio-oil solvent regeneration/sulfur recovery units can be calculated and the feasibility of developing a process within those cost constraints can be evaluated.

To construct the equilibrium stage diagrams, experimentally determined partition coefficients for H₂S in each bio-oil were used. Feed gas concentrations of 40 and 400 ppm H₂S in nitrogen were examined with 99.9% removal of H₂S as the target. A 40 ppm H₂S feed gas concentration would be 2.6 mol% H₂S and 97.4% methane; a 400 ppm H₂S feed gas concentration would be 20 mol% H₂S and 80 mol% methane. These concentrations were chosen to examine the capability of the bio-oil solvents to achieve 99.9% H₂S removal from gas with low and high starting H₂S concentrations, as well as to compare with bench-scale studies carried out using 40 ppm H₂S. A 1000 kmol/h gas feed rate was applied for all simulations, which is on par with flow rates and volumes used industrially in amine gas treating. A graphical solution method was used to determine the optimum flow rate for the bio-oil solvents and the number of equilibrium stages necessary for removing 99.9% of the H₂S from different feed gas concentrations using soybean oil or high oleic soybean oil as the sorption solvent. The Kremser method was then used to evaluate the percent of H₂S absorbed at each theoretical equilibrium stage for different absorbent flow rates and feed gas compositions: the parameters used are shown in Table 1.
Table 1. Flow rates and compositions used for graphical and Kremser methods of equilibrium stage analyses.

| Parameter            | Value                                          |
|----------------------|------------------------------------------------|
| Feed gas flow rate   | 1000 kmol/h                                    |
| Feed gas composition | 40 ppm (2.6 mol%) or 400 ppm (20 mol%) H₂S    |
| Sorbent flow rate    | 96 – 250 kmol/h                                |
| Sorbent composition  | 100% bio-oil: soybean or high oleic soybean    |

To set a maximum total price, comparisons were drawn between similar amine gas treating processes employing an absorption unit and solvent regeneration and recycle units.\(^2\)\(^4\)\(^6\) Industrially popular amine gas treating works similar to the scheme proposed here, with alkylamines (most commonly diethanolamine or monoethanolamine) acting as an H₂S and CO₂ absorbent in an extraction column, while the gases are stripped off in a stripping column to regenerate the amine solvents.\(^3\)\(^5\)

**Results & Discussion**

Economic evaluation of processing sour natural gas using bio-oils is necessary in determining economic viability of such a process before taking efforts to scale up a bench-scale process. Methods to regenerate and recycle the bio-oils and recover sulfur are not yet fully understood, so a total price for the process is set equal to that of existing technologies (such as amine gas treating). After subtracting capital costs and solvent costs from the total price limit, the maximum cost of solvent regeneration and sulfur precipitation was calculated. Discussion regarding the feasibility of developing a process to meet these cost constraints is included.

This study aims to investigate the number of stages necessary for 99.9% removal of H₂S from varying feed gas concentrations, the price of the extraction column and initial solvent cost, the potential revenue through conversion of H₂S and sale of sulfur, and impact of solvent lifetime and regeneration. Figure 1 shows a general schematic of the proposed process.
Figure 1. Schematic diagram of a potential process using bio-oil as an extraction solvent to remove H$_2$S from sour natural gas. An extraction column and solvent cost will be the primary initial costs and will be used to set a limit on how much the solvent regeneration/sulfur recovery units could cost in order for the total process to be economically viable.

*Estimation of Stages and Stage Efficiency for Recovering H$_2$S as Concentration Varies*

To determine the number of stages necessary for handling sour natural gas using bio-oil extraction solvents, varying concentrations of H$_2$S in methane were examined (Table 1 in *Methods*). A graphical method was implemented to determine the optimal flow rates of the liquid absorbent feed and the number of stages required to achieve 99.9% removal of H$_2$S from the feed gas. Partition coefficient values ($K$) for soybean oil (SBO, $K = 0.08$) and high oleic soybean oil (HOSBO, $K = 0.1$) were previously experimentally determined for H$_2$S partitioning in bio-oils.'

Two concentrations of H$_2$S were chosen for study: 40 ppm (2.6 mol%), a proximate value for the bench scale equilibrium experiments, and 400 ppm (20 mol%), to evaluate capability to sweeten sour gas streams at the more concentrated end of the spectrum. Figure 2 shows the absorption model framework for graphical evaluation.
Figure 2. A tray column absorption unit operation with a gas feed of either (a) 20 mol% H$_2$S (~400ppmv) or (b) 2.6 mol% H$_2$S (~40ppmv). Based on a feed gas flow rate ($G'$) of 1000 kmol/h, the absorbent flow rate $L'$ and number of stages $N$ necessary to achieve 99.9% removal of the H$_2$S from the gas feed was determined using a graphical method.

For design of an absorption unit akin to the depiction in Figure 2, Equations 1-3 were used to determine the minimum absorbent flow rate, $L'_{min}$, the optimum absorbent flow rate, $1.5L'_{min}$, and the operating lines. The minimum absorbent flow-rate, Equation 1, is a function of the partition coefficient $K_N$ and the flow rate $G'$ of the gas feed. The optimum absorbent flow rate, $L'$, in Equation 2, is recognized to be a multiple of the minimum absorbent flow rate, $L'_{min}$, and is typically estimated to be close to optimal at $1.5 \times L'_{min}$. Depending on which multiple of $L'_{min}$ is chosen for $L'$, different operating lines can be graphed according to Equation 3.

Minimum absorbent flowrate

$$L'_{min} = G'K_N \quad (1)$$

Optimum absorbent flowrate

$$L' = 1.5 \times L'_{min} \quad (2)$$

Absorber operating lines

$$X_0L' + Y_{N+1}G' = X_NL' + Y_1G' \quad (3)$$
The equilibrium line was constructed using an experimentally determined $K_N$ value of 0.1 mg H$_2$S (gas) / mg H$_2$S (oil) for high oleic soybean oil and 0.08 mg H$_2$S (gas) / mg H$_2$S (oil) for soybean oil. The equilibrium line equation used in the graphical solutions is shown in Equation 4. The concentration of H$_2$S in the gas ($X$) and the concentration in the absorbent liquid ($Y$) at any stage ($N$) is a function of the partition coefficient ($K$).

Equilibrium line  

$K_N = \frac{Y_{N+1}/(1 + Y_{N+1})}{X_N/(1 + X_{N+1})}$  

The graphical solution for (a) 20 mol% H$_2$S absorbed by high oleic soybean oil, (b) 2.6 mol% H$_2$S absorbed by high oleic soybean oil, (c) 20 mol% H$_2$S absorbed by soybean oil, and (d) 2.6 mol% H$_2$S absorbed by soybean oil are shown in Figure 3. Table 2 summarizes the conclusions for $1.5L'_{\text{min}}$ and $N$ depending on the absorbent and feed concentration of H$_2$S.
Figure 3. A graphical solution to determine the number of stages necessary for removing 99.9% of H$_2$S from a feed gas of (a) 20 mol% H$_2$S in methane, HOSBO; (b) 2.6 mol% H$_2$S in methane, HOSBO; (c) 20 mol% H$_2$S in methane, SBO; (d) 2.6 mol% H$_2$S in methane, SBO. The operating lines are based on the flow rate of the absorbent feed. The equilibrium line is based on the partition coefficient $K$ of H$_2$S in the absorbent, high oleic soybean oil, previously determined to be 0.1. The graphical solution is completed for the optimum operating line of 1.5$L'_{\text{min}}$, and the number of stages $N$ was determined to be (a) 10.2 (b) 13.3 (c) 9.6 and (d) 14.2 stages.

Table 2. Optimum flow rate ($1.5L'_{\text{min}}$) and number of stages ($N$) calculated for different feed conditions of H$_2$S (2.6 mol% or 20 mol%) and different absorbing liquids (HOSBO or SBO) using the graphical solution method shown in Figure 3.

| $1.5L'_{\text{min}}$ (kmol/h) | HOSBO | SBO |
|-------------------------------|-------|-----|
| 20 mol% H$_2$S                | 120   | 96  |
| 2.6 mol% H$_2$S               | 146   | 117 |

| $N$                           | HOSBO | SBO |
|-------------------------------|-------|-----|
| 20 mol% H$_2$S                | 10.2  | 9.6 |
| 2.6 mol% H$_2$S               | 13.3  | 14.2|
Additionally, the Kremser method\textsuperscript{7,8} was used to determine the amount of H\textsubscript{2}S sorbed at each theoretical equilibrium stage \( N \) using Equations 5 and 6. In Equation 5, the separation factor \((A)\) is a function of the absorbent flow rate \((L'\text{, varied from } 101 – 250 \text{ kmol/h based on the graphical solutions})\), the feed gas flow rate \((V = 1000 \text{ kmol/h})\), and the partition coefficient \((K)\) of the solute \((\text{H}_2\text{S})\) in the system. For modeling an absorption unit, the fraction of solute absorbed can be calculated as a function of the separation factor \(A\) and number of stages in the unit \(N\) as shown in Equation 6.

\[
\text{Separation Factor} \quad A = \frac{L'}{K_iV} \quad (5)
\]

\[
\text{Fraction solute absorbed} \quad Fraction \text{ solute absorbed} = \frac{A^{N+1} - A}{A^{N+1} - 1} \quad (6)
\]

Tables 3 and 4 display results for percent of solute absorbed at each stage, \(N\), using 1-15 stages and varied absorbent flow rates, \(L'\). Table 3 is based on using high oleic soybean oil \((K = 0.1)\) as the absorbent while Table 4 shows results for conventional soybean oil \((K = 0.08)\).
Table 3. Percent absorption of H₂S using high oleic soybean oil as the absorbent. Percent absorption of H₂S depends on the absorbent flow rate \( L' \) (varied from 101 – 250 kmol/h), the partition coefficient \( (K = 0.1) \), feed gas flow rate \( (V = 1000 \text{ kmol/h}) \), and is shown for each stage \( N \), up to 15 stages. The values represent the fraction of solute (H₂S) absorbed, and ≥99.9% is shaded green, ≥99.0% is shaded yellow, and <99.0% is shaded orange. The target is greater than 99.9% absorption of the H₂S.

| N   | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8  | 9  | 10 | 11 | 12 | 13 | 14 | 15 |
|-----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| L  (kmol/h) | 101 | 50.2% | 67.0% | 75.4% | 80.4% | 83.7% | 86.1% | 87.9% | 89.3% | 90.4% | 91.4% | 92.1% | 92.8% | 93.3% | 93.8% | 94.2% |
|     | 125 | 55.6% | 73.8% | 82.7% | 87.8% | 91.1% | 93.4% | 95.0% | 96.1% | 97.0% | 97.7% | 98.2% | 98.5% | 98.8% | 99.1% | 99.3% |
|     | 146 | 59.3% | 78.2% | 87.0% | 91.8% | 94.7% | 96.5% | 97.7% | 98.4% | 98.9% | 99.3% | 99.5% | 99.7% | 99.8% | 99.8% | 99.9% |
|     | 175 | 63.6% | 82.8% | 91.0% | 95.1% | 97.3% | 98.5% | 99.1% | 99.5% | 99.7% | 99.8% | 99.9% | 100.0% | 100.0% | 100.0% | 100.0% |
|     | 200 | 66.7% | 85.7% | 93.3% | 96.8% | 98.4% | 99.2% | 99.6% | 99.8% | 99.9% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% |
|     | 225 | 69.2% | 88.0% | 94.9% | 97.8% | 99.0% | 99.6% | 99.8% | 99.9% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% |
|     | 250 | 71.4% | 89.7% | 96.1% | 98.4% | 99.4% | 99.8% | 99.9% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% |

Table 4. Percent absorption of H₂S using conventional soybean oil as the absorbent. Percent absorption of H₂S depends on the absorbent flow rate \( L' \) (varied from 101 – 250 kmol/h), the partition coefficient \( (K = 0.1) \), feed gas flow rate \( (V = 1000 \text{ kmol/h}) \), and is shown for each stage \( N \), up to 15 stages. The values represent the fraction of solute (H₂S) absorbed, and ≥99.9% is shaded green, ≥99.0% is shaded yellow, and <99.0% is shaded orange. The target is greater than 99.9% absorption of the H₂S.

| N   | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8  | 9  | 10 | 11 | 12 | 13 | 14 | 15 |
|-----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| L  (kmol/h) | 96  | 54.5% | 72.5% | 81.4% | 86.6% | 89.9% | 92.3% | 93.9% | 95.2% | 96.1% | 96.9% | 97.5% | 97.9% | 98.3% | 98.6% | 98.9% |
|     | 101 | 55.8% | 74.1% | 83.0% | 88.1% | 91.4% | 93.6% | 95.2% | 96.3% | 97.2% | 97.8% | 98.3% | 98.7% | 99.0% | 99.2% | 99.4% |
|     | 117 | 59.4% | 78.3% | 87.1% | 91.9% | 94.7% | 96.5% | 97.7% | 98.4% | 98.9% | 99.3% | 99.5% | 99.7% | 99.8% | 99.8% | 99.9% |
|     | 125 | 61.0% | 80.0% | 88.7% | 93.2% | 95.8% | 97.4% | 98.4% | 99.0% | 99.3% | 99.6% | 99.7% | 99.8% | 99.9% | 99.9% | 100.0% |
|     | 156 | 66.1% | 85.2% | 92.9% | 96.5% | 98.2% | 99.1% | 99.5% | 99.8% | 99.9% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% |
|     | 200 | 71.4% | 89.7% | 96.1% | 98.4% | 99.4% | 99.8% | 99.9% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% |
|     | 250 | 75.8% | 92.8% | 97.7% | 99.3% | 99.8% | 99.9% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% | 100.0% |

The graphical solution indicated 146 kmol/h would be an optimal absorbent feed flow rate when using high oleic soybean oil. Using the Kremser method, it is evident that as absorbent flow rate increases, the number of stages required to attain 99.9% absorption of H₂S decreases. However,
there may be other negative impacts of increasing solvent flow rate, such as increased solvent use and solvent cost.

Another way of examining the system is to set a flow rate and compare the amount of H$_2$S absorbed by the different absorbing bio-oils. In this method, the absorbent flow rate ($L$), feed gas flow rate ($V$) are fixed and the absorption factor $A$ is dependent on the $K$ value of H$_2$S in each bio-oil. Figure 4 shows the comparison between the bio-oil absorption capacity at different equilibrium stages for two absorbent flow rates, 125 kmol/h and 200 kmol/h.

![Figure 4](image)

**Figure 4.** The percent of H$_2$S recovered at each equilibrium stage is dependent by each oil at absorbent flow rates (L) of (a) 125 kmol/h and (b) 200 kmol/h. The percent of H$_2$S recovered is a function of the absorbent flow rate, the feed gas flow rate ($V = 1000$ kmol/h), and the $K$ value for H$_2$S in each oil (0.08 for SBO, 0.1 for HOSBO), as shown in Equations 5 and 6.

As seen in Figure 4 and in Tables 3 and 4, each oil is capable of attaining 99.9% H$_2$S absorption after some number of stages, $N$. In the case of the lower flow rate, as shown in Figure 4a, the difference between the oils is larger (root mean square deviation is 0.037) while requiring 14 stages for both oils to achieve > 99.0% H$_2$S sorption. At a higher flow rate (Figure 4b), the oils’ H$_2$S absorption more quickly converges (root mean square deviation is 0.018) and only five stages are required to attain > 99.0% sorption of H$_2$S. A balance must be struck between number of stages, optimal flow rate, and the resulting bio-oil solvent use. For the remaining analysis, an absorbent
flow rate of 120 kmol/h soybean oil was chosen to limit solvent use while attaining 99.9% removal of H$_2$S in the least number of stages.

*Column Design and Capital Costs*

A significant cost of developing a natural gas processing facility that used bio-oils as the absorbing solvent would be the cost of the absorption unit. Based on the flow rates determined in the previous section (1000 kmol/h feed gas flow rate; 120 kmol/h bio-oil solvent flow rate) and the size and daily processing needs of existing gas treatment methods like amine gas treating, a sieve-plate tray column is recommended as the absorption unit. Other cost considerations would include piping, installation, and any other units for solvent storage, etc.

To design an extraction column to meet the needs for 1000 kmol/h of feed gas, 120 kmol/h of liquid absorbent, and 14 trays (determined by the equilibrium stage calculations: Equations 1-6; Figures 3-4; Tables 3-4), flooding capacity of the column must be calculated.

Flooding capacity is a function of the density of the liquid and vapor phases and a flooding capacity factor, $C$. Equations 7 – 12 were used to characterize column flooding capacity. Equations 7-12 were solved sequentially. Equation 7 – for the entrainment flooding capacity ($F_{LV}$) was first. In Equation 7, $L$ and $V$ are the liquid and vapor feed flow rates respectively, $M_L$ and $M_V$ are the molar masses of the liquid and vapor, and $\rho_L$ is the liquid density and $\rho_V$ is the vapor density. The entrainment flooding capacity ($F_{LV}$) is then used to solve Equation 8, to find the ratio of the active tray area ($A_d$) to the total tray area ($A$). This ratio from Equation 8 is then used to solve Equation 9 and find the factor for ratio of vapor hole area to tray active area. Next, Equation 10 was used to find the surface tension factor, $F_{ST}$, which is a function of the surface tension of the liquid, $\sigma$. Having solved equations 7-10, Equation 11 for the flooding capacity parameter, $C$, can be solved. In Equation 11, $C$ is the flooding capacity parameter, $F_{ST}$ is the surface tension factor solved for in
Equation 10, $F_F$ is a foaming factor assumed to be 1.0, $F_{HA}$ is the factor for ratio of vapor hole area to tray active area from Equation 9, and $C_F$ can be graphically estimated as a function of $F_{LV}$ and plate height, which in this case was assumed to be 24 inches and gave a $C_F$ of 0.35. Finally, Equation 12 for flooding velocity is solved. Flooding velocity, $U_f$, is a function of the flooding capacity parameter, $C$ (Equation 11), and the liquid density, $\rho_L$, and the vapor density, $\rho_V$. Table 5 shows the parameters used in Equations 7-12.

### Entrainment flooding capacity

$$F_{LV} = \left(\frac{LM_L}{VM_V}\right)\left(\frac{\rho_V}{\rho_L}\right)^{0.5}$$  \hspace{1cm} (7)

### Ratio of active tray area ($A_d$) to total tray area ($A$)

$$\frac{A_d}{A} = 0.1 + \left(\frac{F_{LV} - 0.1}{9}\right), \quad 0.1 \leq F_{LV} \leq 1.0$$  \hspace{1cm} (8)

### Factor for ratio of vapor hole area to tray active area

- $F_{HA} = 1.0$ for $A_d/A_a \geq 0.10$
- $F_{HA} = 5(A_d/A_a) + 0.5$ for $0.06 \geq A_d/A_a \leq 0.1$  \hspace{1cm} (9)

### Surface tension factor

$$F_{ST} = \left(\frac{\sigma}{20}\right)^{0.2}$$  \hspace{1cm} (10)

### Flooding Capacity Parameter

$$C = F_{ST} F_F F_{HA} C_F$$  \hspace{1cm} (11)

### Flooding Velocity

$$U_f = C \left(\frac{\rho_L - \rho_V}{\rho_V}\right)^{0.5}$$  \hspace{1cm} (12)
Table 5. Parameters used in tray diameter and flooding capacity calculations for design of an absorption column.

| Parameter                                      | Parameter Value | Source (if applicable)               |
|------------------------------------------------|-----------------|--------------------------------------|
| Temperature                                    | 298 K           | -                                    |
| Pressure                                       | 110 kPa         | -                                    |
| V, vapor flow rate                             | 1000 kmol/h     | -                                    |
| L, liquid flow rate                            | 120 kmol/h      | -                                    |
| $M_V$, molecular weight of vapor phase with 20% H$_2$S and 80% CH$_4$ | 32.4 kg/kmol   | Calculated based on feed gas composition |
| $M_L$, molecular weight of soybean oil         | 920 kg/kmol     | Patzek 2009                          |
| $\rho_V$, density of vapor phase with 20% H$_2$S and 80% CH$_4$ | 1.438 kg/m$^3$ | Calculated using ideal gas law       |
| $\rho_L$, density of soybean oil               | 916 kg/m$^3$    | Sahasrabudhe et al. 2017             |
| $F_f$, foaming factor                          | 1.0             | Seader & Henley, 1998                |
| $\sigma$, surface tension                      | 29.4 dynes/cm   | Sahasrabudhe et al. 2017             |
| Tray height                                    | 24 inches       | -                                    |
| $f$, flooding capacity                         | 0.8             | -                                    |

Equations 7-12 must be solved prior to calculating the required column diameter since column diameter is a function of flooding velocity, flooding capacity, surface tension and active tray area. Ultimately, column diameter ($D_T$) can be calculated using Equation 13 as a function of the gas flow rate ($V$) and molar mass ($M_V$), the flooding velocity ($U_f$) multiplied by a flooding factor ($f$), ratio of tray active area to area ($A_d/A$) and the density of the vapor phase ($\rho_V$). For Equation 13, $V$, $M_V$, and $\rho_V$ are all given in Table 5. $A_d/A$ was calculated using Equation 11, and $U_f$ was calculated using Equation 7. The flooding capacity $f$ must be chosen based on desired column performance. A good rule of thumb is that flooding capacity is often best around 60%, but anywhere from 40-90% is reasonable. In this case, $f = 0.8$ is used, as that is reasonable for a gas absorption column and will help keep the column size smaller, which keeps costs lower. 

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7 In this case, $f = 0.8$ is used, as that is reasonable for a gas absorption column.
Column diameter

\[ D_T = \left( \frac{4VM_Y}{fU_f\pi(1 - \frac{A_d}{A})\rho_Y} \right)^{0.5} \] (13)

Finally, column diameter was calculated as 1.96 m or 6.39 ft; for 14 trays and 24 inches between each tray, column height would be 28 ft. Using a column diameter of 2 m and column height of 8.5 m (6.4 ft and 28 ft), the total column volume would be 84.2 m³.

Equation 14 relates the cost of a carbon steel trayed column \((C_p^0)\) to a column size attribute \(A\), volume in m³) and three constants, \(K_1, K_2,\) and \(K_3\).\(^{12}\) The constants are obtained from Turton et al. 2012 Table A-1.\(^{12}\) \(K_1 = 3.4974, K_2 = 0.4485, K_3 = 0.1074,\) and \(A = 84.2 \text{ m}^3\).

Equipment Cost Equation

\[ \log C_p^0 = K_1 + K_2 \log(A) + K_3[\log(A)]^2 \] (14)

Cost Index Adjustment

\[ C_2 = C_1 \left( \frac{I_2}{I_1} \right) \] (15)

However, Equation 14 is normalized for material pricing in 2001. To obtain an estimate for current pricing, a cost adjustment using the Chemical Engineering Plant Cost Index (CEPCI) was used. In Equation 15, the 2019 cost \((C_2)\) is equal to the 2001 cost \((C_1 = C_p^0\) from Equation 14) multiplied by the ratio of the 2019 CEPCI \((I_2 = 607.5)\) to the 2001 CEPCI \((I_1 = 394)\).\(^{12,13}\) The cost of the extraction column, with the desired specifications, is estimated at $131,600.

**Solvent Cost**

Regeneration of the solvent (soybean or high oleic soybean oil) will be critical to lowering operating costs and developing the present method into a viable process that can compete economically with amine gas treating. The upper acceptable limit for the cost of regeneration will be explored in this discussion. Examining annual solvent cost as a function of the percent of solvent that is regenerated and recycled gives insight into the viability of using bio-oils as extraction
solvents. Soybean oil is examined as a case study representative of similar costs and trends with other bio-oils.

Soybean oil availability is dependent on soybeans harvested, and about 10% (by mass) of soybeans harvested is converted into soybean oil, annually, as shown in Figure 5. Soybean oil accounts for 55% of vegetable oil consumption in the United States, with canola oil taking 14% of the market share and other vegetable oils taking less than 10% each.\(^\text{14}\) 68% of soybean oil is used in food products, 25% in biodiesel and for bioheat, and 7% goes to industrial uses including solvents, paints, plastics and cleaners.\(^\text{14}\) The U.S. Soybean Check-off advertises a “Fuel vs. Food: You don’t have to choose” message, indicating a national effort to find new industrial and non-food uses of soybean oils.\(^\text{15}\) Soybean oil prices are shown in Figure 6.

![Figure 5. Soybean and soybean oil production in the United States, 1991 – 2018. Soybean oil production is about 10% (by mass) of soybean production overall and has continued to rise from 2010 – present.](image)
On average from 2014 – 2018, the price of one metric ton of soybean oil was $679, with yearly prices shown in Table 6. Natural gas plants typically operate year-round with 5-10% of forced downtime for maintenance and outages, both scheduled and unscheduled. Figure 7 shows the solvent cost in millions of U.S. dollars per year, depending on if the absorption unit is operating continuously or if the plant anticipates 5-10% of the year that the absorption unit would be offline and solvent would not be required. The price is based on the 2014-2018 average of $679/metric ton of soybean oil but could increase in future years.

**Table 6.** U.S. soybean oil production and average $/ton of soybean oil from 2014 – 2018. The five-year average is $679/metric ton of soybean oil.

| Year | Soybean Oil (Million Metric Tons) | Average $/ton of soybean oil |
|------|-----------------------------------|------------------------------|
| 2014 | 9.7                               | $697                         |
| 2015 | 10.0                              | $659                         |
| 2016 | 10.0                              | $716                         |
| 2017 | 10.8                              | $661                         |
| 2018 | 11.1                              | $661                         |
Figure 7. Solvent cost of using soybean oil ($679/ton) depending on the amount of solvent that can be recovered and re-used, and the anticipated downtime of the absorption unit of the plant. For a plant operating 95% of the year, solvent cost with no recovery would amount to $624 million/year, but even 50% of solvent recovery would result in a solvent cost of $312 million/year.

Adding Value through Sulfur Recovery

In addition to post-absorption recovery of H$_2$S, which would require the bio-oil solvents to be regenerated and recycled, conversion of H$_2$S to elemental sulfur would allow for sulfur to be a secondary product of the process and generate revenue. Sulfur is a valuable chemical building block used to make sulfuric acid and other commodity chemicals and products. The annual demand for sulfur in the United States exceeds 12.7 megatons, and 36% of that is imported annually.$^6$ The current selling price of sulfur is around $200/ton, and has remained stable above $150/ton since 2014.$^6$

A relationship can be developed to predict the revenue generated in recovering and selling sulfur. This relationship is based on the selling price of sulfur ($200/ton), the amount of sulfur recovered out of the potential amount of sulfur recovered (assumed 90%), portion of the year the plant is
online (90 – 100% of the year, assuming the absorption is running at full capacity for all the time online), and the feed gas concentration of H₂S. This relationship is displayed in Figure 8.

![Graph showing relationship between feed gas concentration of H₂S and revenue.]

**Figure 8.** Assuming that 90% of all H₂S processed is successfully captured and converted to sulfur, revenues from $1.2 - $10 million annually can be anticipated from sulfur sales. This is dependent on the concentration of H₂S in the feed gas being processed throughout the year, as well as the portion of the year the plant is online and operating (assuming the absorption unit is at full capacity and online for the same percent of the year as the plant). Processing more sour gas (400 ppm) would lead to higher revenues than less sour gas (40 ppm). If the plant and absorption unit are online and full capacity 95% of the year, and the average feed gas concentration for the year was 200 ppm H₂S, an annual revenue of approximately $5.6 million could be expected. This would correspond to selling 28,000 tons of sulfur per year.

The H₂S recovered from the natural gas must be captured and harnessed in some way, as regulations prevent the emission of this hazardous gas to the atmosphere. Conversion to sulfur for sale as a chemical building block is a potential lucrative option, and the Claus process for converting gaseous H₂S to elemental sulfur is well established.
**Determination of the upper limit for the cost of solvent regeneration**

The amount of CH\(_4\) available will be higher when the concentration of H\(_2\)S is lower, as shown in Figure 9. The most recent price for industrial natural gas is $3.54/thousand ft\(^3\) as of February 2020\(^1\) and is used for the revenue calculations shown in Figure 9, assuming that the absorption unit is able to process 1000 kmol/h of natural gas when online, and that 100\% of the CH\(_4\) in the feed gas is recovered.

![Figure 9. Annual revenue from processing natural gas.](image)

However, a more sour gas gives less CH\(_4\) but more H\(_2\)S, and overall it is more profitable to process a more sour gas (Figure 10). Even as the amount of near-pure methane recovered decreases, the increased recovery of sulfur makes up for the loss and in fact is more valuable.
Figure 10. Revenue generated by the designed absorption unit operating with 5% downtime annually, 100% recovery of CH₄, and 90% recovery and conversion of H₂S to elemental sulfur. As feed gas becomes increasingly sour, although amount of sweet gas for sale will decrease, sales of sulfur will increase and increase the total revenue.

Based on 1000 kmol/h gas entering the extraction column, 5% downtime annually (but otherwise at full capacity), an average feed gas concentration of 200 ppm H₂S (with 90% sulfur recovery and 100% CH₄ recovery), and 100% solvent regenerated, costs and revenues are shown in Table 7.

Table 7. Costs and projected revenues for the designed absorption unit processing 1000 kmol/h gas with 5% downtime annually, an average feed gas concentration of 200ppm, with 100% recovery of CH4 (primary sweet gas component) and 90% recovery and conversion of H₂S to elemental sulfur. The known costs can be used to set a limit for regeneration of the bio-solvent and precipitation of sulfur to stay competitive with other methods.

| Sources                        | Amount             |
|--------------------------------|--------------------|
| Costs                          |                    |
| Fixed                          | Extraction Column  | $131,600            |
| Variable                       | Extraction Solvent | $8,000,000 / year   |
| Fixed                          | Regeneration/Precipitation Unit(s) | TBD |
| Revenues                       | Sweetened natural gas | $20,280,000 / year |
|                                | Sulfur             | $5,610,000 / year   |
Total profits for the plant will be revenue from selling sulfur and sweet gas minus the fixed costs of capital investments, maintenance and operating costs. While the cost of the extraction column was estimated here, the maximum cost of a regeneration/precipitation unit is to be determined, there may be other costs for heat exchangers, reboilers and condensers, and other installation parts. Bryan Research & Engineering found that the absorption and stripping columns of amine gas treatment facilities typically comprise less than 45% of the fixed capital costs. Another study of natural gas plants in Canada using amine gas sweetening methods found that absorbers are typically 10% of the fixed equipment cost, a regeneration/stripping column 22%, and other equipment (heat exchanger, reboiler, condenser, misc.) comprise the remainder of the costs. The study also notes that for most natural gas plants, the sweetening process only accounts for 3% of the capital expenses; liquefaction and other operations within the process account for the remainder. Calculated costs for capital expenses are shown in Table 8.

Table 8. Capital expenditures and operating expenditures based on the design of the absorption column, solvent cost, and volumes of gas processed annually. Relationships based on empirical data collected from Canadian natural gas plants and described in the literature are used to calculate costs of equipment and labor.

| Capital Expenditures         | $ USD     | Calculation method                                      |
|------------------------------|-----------|---------------------------------------------------------|
| Equipment Costs              | $43,866,508 |                                                        |
| Sweetening Equipment Costs   | $1,316,000 | Absorption is 10% of total sweetening equipment costs   |
| Absorption Equipment         | $131,600   | Calculated based on column size, type, number of trays  |
| Regeneration Equipment       | $289,520   | 22% of sweetening costs                                 |
| Heat Exchangers & Reboilers  | $736,960   | 56% of sweetening costs                                 |
| Misc.                        | $210,560   | 16% of sweetening costs                                 |
| Other Equipment              | $42,550,667| Sweetening is 3% of total equipment costs; other costs come from liquefaction |
| Labor Costs                  | $263,200,000| 6x equipment costs                                      |

| Operating expenditures       | $ USD     | Calculation method                                      |
|------------------------------|-----------|---------------------------------------------------------|
| Solvent (99% recovery)       | $8,000,000/year | Calculated based on soybean oil prices, as shown in Figures 6-7. |
Based on a study of Canadian natural gas plants, natural gas is typically processed at a cost of $8/GJ – where $4/GJ is allocated to pipeline/source costs, leaving $4/GJ for actual treatment and liquefaction of the gas to be sold as liquefied natural gas. Based on the simple model in Table 8, which roughly estimates the capital costs and neglects all parts of operating expenditures other than solvent recovery, and assuming the plant has only 5% downtime but otherwise runs at maximum capacity, a continuous cash flow diagram is shown in Figure 11.

Figure 11. Continuous cash flow diagram assuming a 20-year plant and equipment lifetime. Building a new plant would require a large capital investment and the plant would not begin to see profits until year 12. This model also assumes solvent regeneration and re-use is at 99% and that the cost of processing the gas is even with that of amine gas treating.

After 12 years, the plant would begin to profit. Based on the analysis, over a 20-year plant lifetime, the gas processing cost would be $3.7/GJ, just under the $4/GJ needed to be competitive with amine gas treating and other methods. However, the present model makes several assumptions, including: solvent regeneration and re-use is at 99%, solvent prices do not increase, cost estimates in Table 8 and relationship between cost of different items (i.e. labor = 6x capital costs) are accurate, that natural gas pipelines prices do not increase, that process gas prices do not decrease,
that the absorption operation of the plant is running at full capacity all but 5% of each year, and neglects operating costs. A sensitivity analysis showing the relationship between solvent recovery and the cost of processing the gas is shown in Figure 12. Based on the calculations presented, solvent regeneration and re-use would have to exceed 98% to ensure the costs remain low enough for the overall expense of treating the gas to be comparable with amine gas treating. Increased solvent recovery significantly decreases the cost of treating the gas.

![Figure 12](image)

**Figure 12.** Sensitivity analysis of the relationship between solvent regeneration and recovery and the cost of processing gas. The cost of amine gas sweetening methods is approximately $4/GJ natural gas. To be competitive, the process designed and presented here would need to achieve greater than 98% solvent regeneration and re-use.

As of yet, a definitive method for stripping the H₂S from the soybean oil and bio-solvents and regenerating the bio-oils to be used again is unknown. However, there are a number of methods for recovering H₂S gas in aqueous solutions and oils, well-described in the literature.²¹–²⁴ Most methods fall into one of three categories: chemical precipitation, chemical oxidation, or biological oxidation.²¹–²⁴ The key to process viability would be to develop a method that maximizes the percent of the bio-solvent that can be regenerated and re-used. An alternative to the lofty goal of
98% solvent regeneration and recovery is to find other avenues in which costs can be reduced. While the price of sweetening is well established in Table 8, the cost of other equipment is more loosely correlated and estimated, and a full process simulation including other equipment needs may give better insight into the overall process cost and the solvent regeneration percent that must be obtained.

**Conclusion**

Soybean oil and high oleic soybean oil were used as case studies to examine viability of using bio-oils as solvents for extracting \( \text{H}_2\text{S} \) from sour natural gas, with an aim of designing an absorption operating that could process 1000 kmol/h of natural gas and remove 99.9% of \( \text{H}_2\text{S} \) from feed gas ranging from 40 – 400 ppm \( \text{H}_2\text{S} \). Graphical methods and the Kremser method examined the absorption unit and found a trayed tower with 14 stages, a 2 m diameter and 8.5 m height, could successfully meet these goals with a soybean oil absorbent flow rate of 120 kmol/h. The cost of such an extraction column was estimated along with other capital costs, and the dependence on the economic viability of such process hinges on capability of regenerating and recycling more than 98% of the soybean oil bio-solvent to stay competitive with amine gas treating, the most popular industrial method. The potential of the proposed process being economically favorable looks unlikely. However, we show here where cost-cutting measures could be developed and implemented to reduce processing cost so that bio-based oils are able to compete with current technologies.
Notation

\( A \) Separation factor
\( A_d/A \) Ratio of active tray area to total tray area
\( C \) Flooding capacity parameter
\( C_F \) Flooding capacity velocity
\( CH_4 \) Methane
\( CO_2 \) Carbon dioxide
\( D_T \) Column diameter
\( f \) Flooding capacity
\( F_F \) Foaming factor
\( F_{HA} \) Factor for ratio of vapor hole area to tray active area
\( F_{LV} \) Entrainment flooding capacity
\( F_{ST} \) Surface tension factor
\( G' \) Feed gas flow rate
\( HOSBO \) High-oleic soybean oil
\( H_2S \) Hydrogen sulfide
\( K \) Partition coefficient
\( L \) Liquid flow rate
\( L' \) Absorbent flow rate
\( L'_{min} \) Minimum absorbent flow rate
\( L'_{opt} \) Optimal absorbent flow rate
\( M_L \) Molar mass of liquid
\( M_V \) Molar mass of vapor
\( N \) Number of stages
\( SBO \) Conventional soybean oil
\( U_f \) Flooding velocity
\( V \) Vapor flow rate
\( X_N \) Concentration of H2S in the gas phase at stage \( N \)
\( Y_N \) Concentration of H2S in the liquid phase at stage \( N \)
\( \sigma \) Surface tension
\( \rho_L \) Liquid density
\( \rho_V \) Vapor density
Declarations

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Authors’ contributions
ECB and ASE designed the study. ECB performed all of experiments and carried out the initial analysis. ECB and ASE wrote the manuscript. All authors read and approved the final manuscript.

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References

1. Brace EC, Engelberth AS. Assessing Viability of Soybean Oils to Remove Hydrogen Sulfide from Natural Gas. ACS Sustain Chem Eng. 2020;8(25):9377-9384. doi:10.1021/acssuschemeng.0c01991

2. Acid Gas Removal. Natl Energy Technol Lab.

3. Eow JS. Recovery of sulfur from sour acid gas: A review of the technology. Environ Prog. 2002;21(3):143-162. doi:10.1002/ep.670210312

4. Polasek JC, Bullin JA, Iglesias-Silva GA. Using Mixed Amine Solutions for Gas Sweetening.; 2006.

5. Raj R, Suman R, Ghandehariun S, Kumar A, Tiwari MK. A techno-economic assessment of the liquefied natural gas (LNG) production facilities in Western Canada. Sustain Energy
6. Sulfur Recovery and Tail Gas Treating. *Natl Energy Technol Lab.*

7. Seader JD, Henley EJ. *Separation Process Principles.* 1st ed. John Wiley & Sons; 1998.

8. Kremser A. Theoretical Analysis of Absorption Process. *Natl Pet News.* 1930;22(21):43-49.

9. Oliver ED. *Diffusional Separation Processes: Theory, Design, and Evaluation.* John Wiley and Sons; 1966.

10. Patzek TW. *A First Law Thermodynamic Analysis of Biodiesel Production From Soybean.*

11. Sahasrabudhe SN, Rodriguez-Martinez V, Farkas BE. International Journal of Food Properties Density, viscosity, and surface tension of five vegetable oils at elevated temperatures: Measurement and modeling. 2017. doi:10.1080/10942912.2017.1360905

12. Turton R, Bailie RC, Whiting WB, Shaeiwitz JA, Bhattacharyya D. *Analysis, Synthesis, and Design of Chemical Processes.* 4th ed. Upper Saddle River, NJ: Prentice Hall; 2012.

13. Jenkins S. 2019 Chemical Engineering Plant Cost Index Annual Average. *Chem Eng.* March 2020.

14. U.S. Soybean Oil: U.S. Production History |.

15. Food + Fuel | U.S. Soy.

16. Fakhry R. The Myth of the 24/7/365 Power Plant. *Nat Resour Def Counc.* February 2019.

17. Dalrymple D, Trofe T, States) JE-CEP ;(Unite., 1989 undefined. An overview of liquid redox sulfur recovery. *osti.gov.*

18. Pandey RA, Malhotra S. Desulfurization of Gaseous Fuels with Recovery of Elemental Sulfur: An Overview. *Crit Rev Environ Sci Technol.* 1999;29(3):229-268.
19. U.S. Natural Gas Prices.

20. Ochieng R, Berrouk AS, Peters CJ, Slagle J. *Amine-Based Gas-Sweetening Processes Prove Economically More Viable than the Benfield HiPure Process.*

21. Zhang L, De Schryver P, De Gusseme B, De Muynck W, Boon N, Verstraete W. Chemical and biological technologies for hydrogen sulfide emission control in sewer systems: A review. *Water Res.* 2008;42(1-2):1-12. doi:10.1016/J.WATRES.2007.07.013

22. Shiraishi Y, Tachibana K, Hirai T, Komasawa I. Desulfurization and Denitrogenation Process for Light Oils Based on Chemical Oxidation followed by Liquid-Liquid Extraction. *Ind Eng Chem Res.* 2002;41(17):4362-4375.

23. Yu G, Lu S, Chen H, Zhu Z. Oxidative Desulfurization of Diesel Fuels with Hydrogen Peroxide in the Presence of Activated Carbon and Formic Acid. *Energy & Fuels.* 2005;19(2):447-452.

24. Syed M, Soreanu G, Falletta P, Béland M. Removal of hydrogen sulfide from gas streams using biological processes -A review. *Can Biosyst Eng.* 2006;48(21).