AM-16-44 Alternatives to Flare Gas Recovery for Sour Gas

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Abstract

For many sour gas streams present in a flare header, there may be less expensive options than flare gas recovery. The compressors, liquids management, downstream treating, and other systems required for flare gas recovery can be very expensive to install and/or operate reliably. The presence of sulfur compounds, especially H$_2$S, is often the primary reason why these streams are not sweet-fuel-gas quality and thus can’t continue to be flared. If the stream can be safely and economically treated to sweet-fuel-gas specifications without affecting flare system operation under upset / pressure-relief conditions, then treatment of the flare gas may be economically superior to flare gas recovery processes. This paper reviews the options for flare gas treatment (as an alternative to flare gas recovery) considered for an example oil refiner, including economics, applicable technologies, and techniques that allow gases to be treated under normal operating conditions while not impacting flare function during upset / pressure-relief conditions. Flare gas treatment options including caustic treating, fixed-bed solid H$_2$S scavengers, direct-injection and lead-lag liquid H$_2$S scavengers, and other techniques will be reviewed.

1.0 Introduction

Trimeric Corporation has many years of experience with H$_2$S removal technologies to treat sour gas streams in various industries. Different technologies for H$_2$S removal are often considered depending upon the amount of H$_2$S that needs to be removed, so H$_2$S removal applications are often categorized by the amount of H$_2$S that is removed from the gas stream, usually characterized in terms of long tons of sulfur per day (LTPD). H$_2$S removal technologies that can be used across the full range of possible industrial applications include but are not limited to: non-regenerable solid and liquid H$_2$S scavengers, liquid redox, other regenerable liquid chemical processes, amine/Claus with or without a tail gas treating unit, and numerous others.

An example of a scenario in which an H$_2$S-containing flare gas must be handled in an oil refining setting is provided in this paper. In this example, either H$_2$S must be removed from a flare gas stream to below 60 ppmv to be safely below NSPS Subpart Ja requirements, or a flare gas recovery system must be installed. In this example, the source of H$_2$S is the vent gas of a hydrotreater system that combines with a small natural gas purge stream. Other sweet-gas relief devices are tied into the flare gas system as well. The sulfur load for the flare gas stream is about 51 lb/day or 0.023 LTPD. The flare gas also contains some CO$_2$ and O$_2$ (1% and 0.3%, respectively) with the remaining bulk of the stream being hydrogen and various hydrocarbon species. Table 1 shows the conditions for the example flare gas stream.
At this low sulfur load, it was anticipated that \( \text{H}_2\text{S} \) removal would be less expensive than a flare gas recovery system. The \( \text{H}_2\text{S} \) removal system would need to treat the flare gas stream to sweet-fuel-gas specifications without impacting the flare system function during process upsets when pressure relief systems activate and the flare gas flow rate surges. The \( \text{H}_2\text{S} \) removal system would not be required to remove \( \text{H}_2\text{S} \) from gases resulting from major pressure relief events; instead, a bypass system would be used to route relief events directly to the flare under these conditions. The pressure drop through the \( \text{H}_2\text{S} \) removal system would also need to be low (less than about 5 psi).

This paper first presents an overview of typical flare gas recovery operations for reference. Then, the paper reviews the alternatives to flare gas recovery, namely treatment options for the low-tonnage \( \text{H}_2\text{S} \) removal rates in this example flare gas application. Schematics are used to show the general concepts of how the technologies work. The major advantages and disadvantages for the most promising \( \text{H}_2\text{S} \) removal processes and pertinent economic estimates are summarized. Finally, other techniques that could potentially be used to remove the \( \text{H}_2\text{S} \) from the example flare gas stream, but are not as favorable, are also reviewed.

### 2.0 Flare Gas Recovery (FGR)

A typical flare gas recovery (FGR) system recovers process vent gases that would normally be flared so that they can be used as fuel gas elsewhere in the facility. FGR results in reduced use of purchased fuel gas (recovered gas from the flare displaces purchased fuel, e.g., natural gas), reduced emissions of combustion products at the flare, and reduced flare maintenance costs by extending the life of the associated flare tips due to fewer flaring episodes.

A typical FGR system is shown in Figure 1. The FGR inlet gas is taken downstream of the flare knockout drum and upstream of the liquid seal pot. The flare gas typically exits the main flare header via a branch connection and flows to a liquid-ring compressor (although other compressor types can be used) to increase the pressure of the gas to the desired level for use at the plant. Downstream of the liquid-ring compressor, the compressed gas then flows into a separator that removes condensed hydrocarbon liquids, seal water, and possibly condensed water from the gas. Since the recovered gas is sour in this example, the gas from the separator is sent to a downstream treating system. A water-cooled or air-cooled exchanger is used to remove the heat of compression generated in the system by cooling the water that is recirculated to the liquid-ring compressor. A small blowdown stream is usually generated from the liquid-ring system that typically flows to a sour water system. This blowdown stream is used to control the buildup of dissolved species in the recirculated water, and a small continuous makeup water stream is usually added to the system. Since the feed to the FGR unit comes from the flare header near the flare, very long piping runs (e.g., thousands of feet) may be necessary to

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Units</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flare gas flow rate</td>
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</tr>
<tr>
<td>( \text{H}_2\text{S} ) content</td>
<td>ppmv</td>
<td>6000</td>
</tr>
<tr>
<td>( \text{CO}_2 ) content</td>
<td>mole %</td>
<td>1</td>
</tr>
<tr>
<td>( \text{O}_2 ) content</td>
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</tr>
<tr>
<td>Hydrogen and other hydrocarbon content</td>
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<td>98.1</td>
</tr>
</tbody>
</table>
conduct low-pressure recovered sour gas to the FGR, higher-pressure recovered sour gas from the FGR, recovered hydrocarbon liquids, and/or sour water pipelines to their destinations.

The automated controls of the FGR adjust the flow of low-pressure recovered sour gas to the FGR unit in order to maintain a slight positive pressure on the flare header upstream of the liquid seal pot. This positive pressure ensures that air will not be drawn into the flare header or into the FGR unit. If the volume of flare gas that is relieved into the flare header is greater than the capacity of the FGR unit (for example, as would occur in a major pressure relief event), the pressure in the flare header will increase until it exceeds the backpressure exerted on the header by the liquid seal pot. In this scenario, the excess gas will pass through the seal pot and into the flare where it will be burned. The design of the liquid seal is also very important since it serves as the backpressure control device for the flare header and the low-pressure recovered sour gas to the FGR unit. On the other hand, if the volume of flare gas relieved into the flare gas header is less than the total capacity of the FGR unit, the control system adjusts the flow to the FGR unit. This is accomplished by turning off compressors and/or by recycling the compressor discharge gas back to the suction header via a recycle control valve. The speed of the compressors could also be controlled and adjusted (Baukal, 2013).

Figure 1. Typical Flare Gas Recovery Unit

FGR units are industry proven technology but can be expensive for sour gas streams given the need for compression, liquids management, downstream treating, potentially long recovered-product pipelines, and the control system necessary for consistent, safe operation and reliability. In addition to the high total capital cost and the FGR operating cost, the recovered sour gas, which generally goes into a fuel gas treating system (e.g., an amine unit), may also have negative consequences for the treating system. In this example, the recovered flare gas has 0.3% oxygen, and oxygen is known to cause amine degradation and amine system corrosion (Jones et.al, 2010). Because of these factors, alternative H₂S treating technologies may be more economically advantageous for applications like this example.
There are a large number of processes that have been demonstrated in the industry for the removal of \( \text{H}_2\text{S} \) from gas streams. However, many of these processes are more applicable to cases where the amount of \( \text{H}_2\text{S} \) to be removed is much larger than in this example (0.023 LTPD). \( \text{H}_2\text{S} \) removal and sulfur recovery technologies can generally be categorized in the niches shown in Figure 2. As mentioned previously, the technology niches are based primarily on the amount of sulfur that has to be removed (e.g., the LTPD of elemental S), not the gas flowrate alone.

![Figure 2. Technology Niches for \( \text{H}_2\text{S} \) Removal](image)

For large-scale treating, roughly greater than 20 LTPD, the amine/Claus process combination with a tail gas treating unit is a common selection. For small to medium sulfur loads in a range of 0.1 to 20 LTPD, other continuous regenerable \( \text{H}_2\text{S} \) removal / sulfur recovery technologies are often applied, including liquid redox processes. For processes in which the amount of \( \text{H}_2\text{S} \) to be removed is very small, typically less than about 0.1 LTPD of sulfur, a class of technologies called “scavengers” is most often used. Scavengers are nominally non-regenerable chemicals that typically operate in a semi-batch mode. Although the scavenger processes tend to have a relatively high chemical operating cost when considered on a “per pound of sulfur removed” basis, the capital cost and operating complexity are typically much lower than for the larger-scale processes, resulting in the overall treating cost being lower than the processes targeted for larger sulfur loads.
The most promising technologies that were considered to remove the H$_2$S from the example flare gas stream include: caustic treating, solid scavenging, and liquid scavenging. Each of these processes is discussed separately in greater detail. Other techniques that could be used, but which were not considered further due to unfavorable economic and/or operating conditions, are also presented. Finally, a table comparing the technologies is provided to summarize the options.

### 3.1 Caustic Treating

Sodium hydroxide (NaOH) solutions are often used to scrub H$_2$S from gases for small-sulfur-load applications, although caustic treating is also sometimes applied to medium-scale applications, particularly when the product solution (NaSH) can be marketed. For cases where the caustic is not regenerated, caustic is a “scavenger.” The NaOH reacts with the H$_2$S to form a liquid product with dissolved sodium bisulfide (NaSH) and/or sodium sulfide (Na$_2$S) solutions. If CO$_2$ is in the gas, it will also be scrubbed to form sodium bicarbonate (NaHCO$_3$) and/or sodium carbonate (Na$_2$CO$_3$), which results in additional use of caustic and potential problems with carbonate salt precipitation. At moderate pH, NaSH and NaHCO$_3$ are the predominant species while, at highly alkaline pH, Na$_2$S and Na$_2$CO$_3$ will form.

Caustic scrubbing is done using a variety of equipment including: packed or trayed towers, sparged tanks, in-line contactors, Venturi contactors, or a combination of these devices. Caustic scrubbing can be implemented as a continuous process or in a semi-batch mode.

Caustic scrubbing applied as a continuous process can be designed to minimize the use of caustic. In this mode of operation, the continuous caustic scrubbing tower is designed with two separate loops of recirculating caustic solution. Fresh caustic is fed to a top loop, which operates at a relatively high pH and serves as a polishing section to maximize H$_2$S removal. Caustic overflows from the top section to the bottom section. The bulk of the H$_2$S removal is done in the bottom recirculation loop which operates at lower pH. The continuous dual-loop system requires significant controls and relatively high capital cost. Although the dual-loop system offers low caustic consumption for H$_2$S removal, it is not well suited for gas streams with a relatively large amount of CO$_2$ relative to H$_2$S (Mamrosh et. al., 2013; Mamrosh et. al., 2014).

Caustic scrubbing of H$_2$S in a sparged tank or packed tower also results in the majority of the CO$_2$ in the feed gas being scrubbed. If the gas stream to be treated has large amounts of CO$_2$ relative to the H$_2$S, then the use of a “short-contact-time” contacting system can result in the absorption of the bulk of the H$_2$S, while not absorbing most of the CO$_2$. This concept takes advantage of the fact that H$_2$S is absorbed more quickly than CO$_2$ at high pH, and selective H$_2$S removal can be achieved by limiting the contact between the gas and liquid in a static mixer, Venturi contactor, or other specialized contacting device. However, the complexity and relatively high capital cost of a continuous, short-contact-time system would not be warranted for this small sulfur load of only 0.023 LTPD (Mamrosh et. al., 2008; Mamrosh et. al. 2009).

Given the small sulfur load of this example flare gas application, it was assumed that the cost of using complex equipment to operate continuously would be greater than the savings realized by more efficiently utilizing the caustic. Therefore, the use of a simpler semi-batch scheme for caustic treating of the flare gas was chosen as the system that would likely have the lowest capital cost and operating complexity. A schematic of such a system, a sparged tank system, is shown in Figure 3. (Note: the
sparged tank might potentially act as a liquid seal device.) If the pressure drop through a sparged tank is higher than desired, then another way to implement a semi-batch system is to use a packed or trayed contactor with a large sump at the bottom to allow a relatively large volume of caustic solution to be charged to the system, and a small pump to recirculate the caustic from the sump over the packing. This style of scrubber is sometimes used in modified Claus SRU backup service in some refineries (Mamrosh et. al., 2013; Mamrosh et. al., 2014).

Figure 3. Schematic of Semi-Batch Caustic Treating

In a semi-batch process, two sparged tanks and a knockout tank are included in the design. The two sparged tanks would be identical, and consist of a vertical vessel with a gas sparger near the bottom. One tank would be on-line (treating gas), while the other tank would be off-line. The treated gas from the on-line tank would be routed to a generously sized knockout vessel equipped with a demisting pad with a thorough water wash system. The on-line tank would be used until the solution pH dropped to a predetermined level (~pH 12 for this example case) that is determined based on when the system would either i) potentially lose the ability to produce sweet gas, or ii) possibly begin to precipitate solids. An on-line \( \text{H}_2\text{S} \) monitor could also be used to help judge the point at which the caustic batch must be changed. The flare gas flow would then be switched to the second tank while the spent caustic is drained and replaced in the first tank.

The caustic treating unit should be located near the flare and flare knockout. In the event of a major upset causing the flare gas flow rate to increase significantly, the bulk of the flow would bypass the scrubber system via a properly designed seal pot. Alternately, if a device to impart flow through the caustic system (e.g., a scrubbing-liquid driven eductor) is used instead of a seal pot, then the bulk of the flow would pass through the open line to the flare header. This is similar to the flare gas recovery
compressor taking the normal flow, but allowing emergency flow from a major relief event to bypass to the flare. It may also be possible to use some other special device in place of the emergency-flow-bypass seal pot, for example a buckling-pin valve, rupture disk, or other highly reliable device.

The main advantage of caustic treating for this example flare gas application is that the chemical costs are low, and, depending on the number of added features, the capital cost can be low. Even though a large portion of the caustic is wasted due to the co-absorption of CO₂, the chemical cost is still reasonably low ($60,000/year) because caustic is an inexpensive chemical (cost: $500-1500/dry ton). A semi-batch-mode, sparged tank system is simple to operate with relatively modest capital expenses. Even though the caustic will need to be changed out approximately weekly, the change-out process is relatively easy. If the unit is operated properly and the caustic replaced when required, the spent solution should be a single-phase liquid that can be pumped to a waste collection system. Finally, fresh caustic (~20°Be would be used because it has the lowest freezing point of caustic solutions) can be easily added from a tank truck, a local tank, or from a refinery caustic supply line if available.

The main disadvantage of using a semi-batch caustic treating for H₂S removal is the waste of caustic due to nearly all of the CO₂ scrubbing into solution. The caustic also cannot be fully utilized (e.g., it was not allowed to drop below ~pH 12 for this case) to avoid the precipitation of sodium bicarbonate salts, and avoid the breakthrough of H₂S in the treated gas. The greater the ratio of CO₂ to H₂S in the feed gas, the greater the portion of caustic that will be wasted in scrubbing CO₂ instead of H₂S. Although the precipitation of salts should not occur under normal operating conditions, unusual conditions (such as very high CO₂ to H₂S ratios) or if the solution is not changed out when required, could result in salts formation. The formation of salts in the process can result in the plugging of the vessel and lines, which could lead to the need to shut down the system for a cleanout. Finally, the process produces a relatively large quantity (1.3 to 1.4 million pounds/year) of poor-quality spent caustic solution (also referred to as NaSH solution, or sulfidic caustic) that needs to be disposed of either through reuse in an existing refinery caustic unit, sale to a third party, or as a waste stream. NaSH solution that has a low NaSH concentration and/or high Na₂S and Na₂CO₃ concentrations is typically considered to be of low value. NaSH solutions containing a high concentration of NaSH (e.g., 25 – 40 wt%) and low levels of Na₂S and Na₂CO₃ (e.g., less than a few percent each) can be valuable, marketable products, but the spent caustic that would be generated in this system would not meet the specifications usually required for outside sale.

If the disposal of this spent caustic as a waste material is required, it might be considered hazardous because of the high pH. Neutralization of the high pH by the addition of a strong acid may not be practical, because if this is done some of the H₂S would be liberated from the spent caustic solution. More sophisticated caustic scrubber designs (e.g., the short-contact-time system mentioned previously) may be able to produce higher-quality product (high NaSH content with less Na₂S and/or Na₂CO₃), and that high-quality product can often be sold.

3.2 Solid Scavenger Treating

A number of commercially available solid chemicals can be used as H₂S scavenging agents (Fisher et. al. 1999). Several types of solid scavengers are based on the use of iron oxide (Fe₂O₃) to react with H₂S in the presence of water to form iron-sulfur compounds and water. The iron oxide is a granular solid or a solid mixture on an inert substrate such as clay. Sufficient residence time of the gas with the scavenger solid is required to ensure that the H₂S can be removed to ppm levels.
Other solid materials can also be used for H$_2$S scavenging. These include solid scavengers containing potassium permanganate and copper or zinc oxides. Generally, solid scavengers that are not based on iron oxide are more expensive than the iron-oxide-based materials. Most iron oxide based scavengers require that some water be present in the feed gas stream in order to function at their full capacity. When dry gas streams are being treated (and water cannot be added), capacity can be greatly reduced, or more expensive mixed metal oxide scavenger products may be required.

One of the older solid scavengers is iron sponge, which is made by depositing iron oxide on wood chips. It is an inexpensive product, but the spent material can be difficult to handle because it can be pyrophoric and must be kept wet after removal. Exposure of the spent material to oxygen results in exothermic side reactions that can cause the substrate to catch fire. Iron-oxide-based scavengers that are not pyrophoric generally use a non-flammable substrate material. These materials are not classified as pyrophoric but can still get hot when removed from service, so it is generally recommended to remove them wet to reduce thermal hazards.

A system using non-pyrophoric iron oxides was considered for this example refinery flare gas application. A schematic diagram is shown in Figure 4. Two vessels are used in this process. One vessel would be on-line treating gas. The off-line vessel would be available for changing out the spent adsorbent, and then serve as a spare for the vessel in service. Water or steam is required to saturate the gas stream to increase the lifetime of the adsorbent. This can be done with a direct steam injection into the line or by adding a recirculating water spray. Because the flare gas flow rate could vary, control of the steam rate might be difficult and require significant instrumentation; a water spray could be simpler and less expensive. A pump could be used to transfer the water / condensate from these vessels. A seal pot, Venturi/ejector, or other special device (buckling pin, rupture disk, and/or relief valve) could be used to bypass the process during major flow events. It may also be necessary to monitor the treated gas H$_2$S or another surrogate parameter to demonstrate removal.

The main advantages of using solid scavenger technology are that the process has been used in industry for a number of years, is considered to be reliable, and the process is also simple since it requires just a single bed on line and one off line with minimal control requirements. As a result, the capital cost for the process is relatively low. The pressure drop through the system can be designed to be less than 1 psi. It is also possible to design the solid scavenging system to have a long time between changeouts (e.g., up to 8 months at one known refinery fuel gas application). Based on the sulfur load for this example flare gas stream and the assumed vessel sizes, the solid scavenger material would only need to be changed out every 70 to 180 days. This would result in a replacement cost ranging from $50,000 to $150,000/year depending on the specific product chosen. The spent product can usually be disposed of as a non-hazardous solid waste. Lastly, the effectiveness of solid scavengers is mostly independent of gas flow rate (below the design flow rate); therefore, there are typically few problems with turndown.
A major disadvantage of this process is that the spent material has no value and must be disposed of, likely in a landfill. Changing out the solid scavenger material must be done manually (unlike the caustic treating product, which can be pumped or vacuumed out) and requires more effort, time, and equipment. The change-out can be messy and the material is often not free-flowing. During cleanouts, water is generally used to keep the material from becoming too hot and to help flush the material out of the vessel. About 150,000 to 300,000 pounds/year of solid scavenger material would be used in this application, and the number of change-outs required would be 2–6 per year. Another drawback of the process is that the feed gas needs to be saturated with water. If the system is operated with a gas that is dry, or nearly dry, then the solid scavenger will work, but the capacity will be lower resulting in more change-outs of material.

3.3 Liquid Scavenger Treating

Liquid scavengers are also available to remove H₂S from small-scale applications. One commercially available liquid scavenger is a class of chemicals called triazines. Triazines have three nitrogen atoms that take the place of carbons in a six member ring. The triazines used for H₂S scavenging typically have hydroxyethyl side groups, are water soluble, and are usually not considered to be hazardous. The triazine chemical is usually mixed with water and may also contain some alcohol and alkanolamines. H₂S reacts with the triazine to make water soluble products, such as dithiazine and bisdithiazine. The spent solution is normally all single-phase liquid, although solids can form if the solution is over utilized.
Triazine is frequently used in the natural gas production and treatment industries to remove small amounts of H$_2$S to pipeline specification (< 4ppm). It is also practiced in the refinery setting, although it is not as common. Some refiners have been known to use direct injection scavenging (see Figure 5) on flare gas to temporarily remove H$_2$S so as to lower the SOx emissions from a flare; the applications that the authors know of do not achieve deep H$_2$S removal, although deep removal is possible in a properly designed system.

In direct injection, the process is fairly simple since the equipment consists of only a chemical injection pump, a means of introducing the scavenger into the sour gas pipeline, a length of pipe to allow for gas/liquid contact, and a downstream device for separating spent or excess scavenging agent from the treated gas. The amount of gas/liquid contact time is dependent on the type of contacting device, the gas velocity, and residence time. Because of this, the degree of mixing and efficiency is sensitive to changes in gas flow (Fisher et. al., 2003; Fisher et. al. 2005; Jamal et. al. 2009; Leppin et. al. 2009). Direct injection scavenging typically has a low capital cost; however, initial calculations showed that long pipe lengths would be required to reduce the H$_2$S in this example refinery flare gas to the required 60 ppmv level.

If not enough pipe length is available, then a sparged tower could be used instead to meet the treatment specification. With a reasonable liquid level above the sparger and an efficient sparger design, the pressure drop requirement of 5 psi could also be met. Because of this, a semi-batch process similar to that for semi-batch caustic treating could be implemented (see Figure 6). Storage for the fresh and spent triazine would be required. An H$_2$S monitoring device or other surrogate method may be necessary to demonstrate removal.

Although the capital costs for using triazine scavenger in a semi-batch mode would be low, the chemical costs for replacing the spent triazine would be significantly higher than the other H$_2$S treating options considered in this study. Based on a typical cost of triazine, the annual chemical cost for

**Figure 5. Schematic of Direct Injection H$_2$S Scavenging**

![Diagram of Direct Injection H$_2$S Scavenging](image-url)
treating this example flare gas application would be $300,000 to $325,000/year, which is 3 to 5 times higher on average than the other options (caustic and iron oxide scavengers). About 750,000 to 800,000 pounds of spent triazine solution would need to be disposed of per year.

Other liquid scavenging materials that may be commercially available include sodium nitrite, bleach, and peroxide. Sodium nitrite solution has significant historical use in H₂S scavenging; its use results in the formation of solid elemental sulfur, so the spent solution is a slurry.

Bleach scavenging is usually done in a continuous mode of operation in a packed tower with the addition of fresh makeup chemicals. Bleach consists of an aqueous solution of sodium hypochlorite; sometimes excess caustic is used to maintain the solution alkalinity and enhance the absorption of H₂S. Bleach scrubbing oxidizes the H₂S to sulfates (Na₂SO₄). It is not known if bleach scrubbing can be done in a simple sparged vessel, plus the high cost of bleach would result in treatment costs that are approximately 2-5 times that of caustic or iron oxide-based scavenging.

3.4 Other Techniques

Other technologies exist that could be used to remove H₂S from flare gas streams. Contacting the flare gas stream with amine solution is one example. At the given sulfur load, installing a dedicated
continuous operating amine unit would likely have a very high relative capital cost. However, if the refinery had an existing amine system it might be possible to tie into that system. A small portion of the amine could be drawn from the main lean-amine line and used in a conventional tray or packed column to remove the H\textsubscript{2}S from the flare gas. The rich amine would then be routed back to the main rich amine line in the amine plant. Some key items to consider with using this method are whether the lean amine is lean enough to remove the H\textsubscript{2}S at the flare gas stream operating temperature and pressure, with the low pressure being particularly significant. The amine contacting tower would need to be designed to handle fluctuations in the flare gas flow rate and/or have an effective bypass system, like some of the designs shown previously. Also, the composition of the flare gas stream would need to be considered as oxygen and other contaminants can significantly degrade the amine solution and cause amine-system corrosion (Jones et.al, 2010), and potentially also limit the amount of H\textsubscript{2}S that can be removed from the system.

The removal of H\textsubscript{2}S from gases using redox chemistry is well known and widely practiced in industry. Typically, a water soluble metal or a water soluble metal-chelate is reacted with the H\textsubscript{2}S to form elemental sulfur. The elemental sulfur is filtered from the solution and the metal is regenerated to its reactive state using oxygen (air). These processes are typically used for sulfur loads above those associated with H\textsubscript{2}S scavenging, and below that associated with the use of amine/Claus (0.1 to 20 LTPD). Redox processes are typically designed as continuous processes, with H\textsubscript{2}S removed from the gas in one unit operation, and the solution regenerated in another. However, if the gas stream to be treated contains a substantial amount of oxygen, the H\textsubscript{2}S removal and solution regeneration can be done in one vessel; this may improve the economic viability of the redox processes into the lower sulfur load region. A batch liquid redox process was examined for this example case, since the flare gas stream has enough oxygen to meet the stoichiometric oxygen requirement of the reaction (1/2 mol O\textsubscript{2} per mole H\textsubscript{2}S). However, there is not a significant amount of excess O\textsubscript{2} in the gas. The low O\textsubscript{2} to H\textsubscript{2}S ratio is lower than typical and may have kinetic or equilibrium limitations to the system that would constrain regeneration of the solution. For this reason, a batch redox process was not considered to be a candidate technology for H\textsubscript{2}S removal from the example refinery flare gas stream.

### 3.5 Overall Comparison of H\textsubscript{2}S Treating Technologies and Flare Gas Recovery

A comparison of the key alternative H\textsubscript{2}S treating technologies and the FGR option are presented in Table 2 below. The comparison provides an indication of estimated capital and/or operating expenses as well as other intangibles, such as material handling and treatment limitations, which should be taken into consideration when evaluating and selecting an appropriate technology. All of the H\textsubscript{2}S treating technologies are operating in simple batch mode for this low sulfur tonnage application.

As shown in the table, the solid scavenger technology had the lowest capital cost estimate of the H\textsubscript{2}S treating technologies and the flare gas recovery system. The chemical costs estimated for this process are on average 50% higher than the least expensive option (caustic treating). The solid scavengers produced the least amount of waste compared to the other H\textsubscript{2}S treatment processes. The main disadvantage to the solid scavengers is that the changeouts, although less frequent than the other
Table 2. Comparison of H₂S Treating Technologies and Flare Gas Recovery for an Example 0.023 LTPD Flare Gas Stream

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<th>Process</th>
<th>Advantages</th>
<th>Disadvantages</th>
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<th>Changeout Rate (days)</th>
<th>Waste Produced (K lb/yr)</th>
<th>Chemical Costs ($K/yr)</th>
<th>Relative Capital ($)</th>
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</thead>
<tbody>
<tr>
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<td>-- Changeouts considered by some operators to be time consuming and difficult</td>
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<td>70-180</td>
<td>150-300</td>
<td>50-150</td>
<td>1</td>
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<tr>
<td></td>
<td>-- Selective to H₂S adsorption</td>
<td>-- Steam or water injection needed if flare gas is dry</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>-- Low pressure drop</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Least frequent changeouts</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Waste solid usually not hazardous</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Liquid Scavenger</td>
<td>-- Simple process</td>
<td>-- High chemical costs</td>
<td>5</td>
<td>14</td>
<td>750-800</td>
<td>300-325</td>
<td>--</td>
</tr>
<tr>
<td></td>
<td>-- Selective to H₂S adsorption</td>
<td>-- Precipitation possible if over-spend scavenger</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Spent liquid typically a single aqueous phase</td>
<td>-- Pressure drop limit pushed</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Cleanout / turnaround easy</td>
<td>-- Odor with spent solution</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Caustic</td>
<td>-- Established in industry for other low-pressure gases</td>
<td>-- May not be possible to put waste to sewer</td>
<td>5</td>
<td>7</td>
<td>1300-1400</td>
<td>30-100</td>
<td>1.5</td>
</tr>
<tr>
<td></td>
<td>-- Simple process</td>
<td>-- CO₂ also scrubbed, wasting caustic</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Caustic is inexpensive</td>
<td>-- Chance for salts to form if mis-operated</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Changeouts / turnaround easy</td>
<td>-- Spent caustic waste can be considered hazardous</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Spent caustic may be reusable at other refinery caustic scrubbers</td>
<td>-- High waste volume</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>-- Solid elemental sulfur formation possible due to presence of O₂</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Flare Gas Recovery</td>
<td>-- Established in industry</td>
<td>-- High capital cost</td>
<td>na</td>
<td>na</td>
<td>na</td>
<td>na</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>-- Lower flare emissions</td>
<td>-- Long lines may be necessary to send recovered gas and liquid to destination</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Lower fuel gas costs</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Lower flare maintenance</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- Familiarity to refinery staff</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- No chemical usage required</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>-- No spent chemicals to dispose of</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Notes:
1) The table does not include the operating costs (which can be high given the power for compression) and the value of the recovered fuel for the FGR option.
2) K = 1000
options, are considered by some operators to be rather messy and time intensive compared to the other
treating options; saturation of the feed gas with water is also required, if low chemical costs are desired.

The liquid scavenger process had significantly higher chemical costs (3-5 times greater on
average) than the other two H₂S treatment options. It also did not appear to provide any clear
advantages in capital cost, complexity or reliability. It would probably be more competitive for cases
where direct injection into an existing pipeline would be sufficient for reaching the desired H₂S removal
efficiency, which would result in a relatively low capital cost.

Caustic treating had the lowest chemical costs of the technologies considered, but the capital
costs for the process were about 50% higher than with solid scavenging. The higher capital costs may
be a result of the need for a higher complexity of controls (e.g., pH probes, differential pressures, and
level) and special material of construction due to potential corrosion issues with the caustic. Caustic
treating produces a significant amount of spent caustic that must be disposed of, or potentially reused at
the refinery if other caustic units are operated on site. Disposal costs are not included in the table.

The alternative H₂S treating options appear economically competitive to the flare gas recovery
system for this low sulfur tonnage, example flare gas case. The capital expense for the FGR process is
about 3 times greater than the lowest alternative H₂S treating option (solid scavenging). The FGR
process requires costly compression, controls, and long pipe lengths to communicate the recovered gas
and liquid streams to their final destination. Additional treatment of the sour liquid and recovered gas
streams is still also required. The potential benefits are reduced emissions and fuel gas costs; some flare
maintenance cost savings may be achieved as well.

4.0 Summary

Alternative H₂S treatment options are economically and technically viable compared to flare gas
recovery for the example considered in this paper. It is important to evaluate the technologies on a case-
by-case basis because the feed gas conditions (e.g., sulfur loading, hydrocarbon content, impurities, flow
rate, temperature, and pressure), treatment specification, and other parameters (e.g., pressure drop, upset
conditions) can impact the technology selection. For this example, refinery flare gas stream, the sulfur
tonnage is low (0.023 LTPD), making the capital costs of the alternative H₂S treating technologies less
because simple batch operation H₂S scavengers can be used. The chemical costs for the H₂S scavengers
are reasonable given the associated savings in capital cost. The compressor, controls, long liquid and
recovered gas pipelines, and potential effects on fuel gas treating systems make the FGR unit less
economically competitive for this gas stream because of capital cost.

5.0 References

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2) Fisher, K.S., J.E. Lundeen, D. Leppin, “Fundamentals of H₂S Scavenging for Treatment of
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