Effect of Reaction Furnace and Converter Temperatures on Performance of Sulfur Recovery Units (SRUs)

Hamid Reza Mahdipoor, Keyvan Khorsand, Reza Hayati, Hooman Javaherizadeh
Department of Process and Equipment Technology Development, Research Institute of Petroleum Industry,
West Blvd. of Azadi Stadium, Tehran, Iran
mahdipoorhr@ripi.ir

Abstract-The modified Claus process is commonly used in oil refining and gas processing to recover sulfur and destroy contaminants formed in upstream processing. In oil refining, in addition to the typical modified Claus plant feed, i.e. H₂S and CO₂, NH₃, CS₂, and COS are also often present or formed. These contaminants pose a risk of poisoning the catalyst beds, as well decreasing the overall sulfur recovery. In this paper, effect of reaction furnace and converter temperatures on performance of SRUs is described. Then, the common ways for adjusting these important parameters are presented for a typical sulfur recovery unit. The achieved results can be useful for designing the new Claus sulfur recovery units.

Keywords-Claus Process; Furnace and Converter Temperatures

I. INTRODUCTION

The Claus process continues to be the most widely used process for the conversion of H₂S to sulfur [1]. The task of Claus processes is to recover elemental sulfur from hydrogen sulfide and, more generally, from byproduct gases originating from physical and chemical gas and oil treatment units in refineries, natural gas processing, and gasification plants, to quote a few [2]. They consist of a thermal reaction furnace, a waste heat boiler (WHB), and a series of catalytic reactors (converters) and condensers (Figure 1). The reactions occurring in the furnace are numerous, and several authors have attempted to delineate the important ones [3-6]. The overall reaction characterizing the process is as follows [2]:

\[ 2H_2S + O_2 \rightarrow S_2 + 2H_2O \]

Fig. 1 The schematic shape of a typical Claus unit

In the first step or thermal stage, one-third of the H₂S is completely oxidized to SO₂ in the reaction furnace, locating at the front end of plant. A benefit that also occurs is the production of significant quantities of elemental sulfur (S₂) from the thermal decomposition of H₂S. In fact, the sulfur produced in the furnace is 50-60% of the total sulfur condensed in the plant. The main H₂S oxidation reaction is [7],

\[ H_2S + \frac{3}{2}O_2 \rightarrow SO_2 + H_2O \]

The reaction furnace is followed by the waste heat boiler (WHB), where heat is recovered by cooling the furnace product gases [7]. In the second step that the catalytic stage, unreacted H₂S is then combined with SO₂, reacting via eq. 2, over an alumina catalyst to form elemental sulfur in fixed bed reactors by the following reaction [7,8],

\[ 3H_2S + SO_2 \rightarrow S + 2H_2O \]

Since this reaction is exothermic, decreasing the temperature leads the equilibrium reaction toward right hand, i.e. more sulfur yields. On the other hand, low temperatures decrease the reaction rate. Therefore, an appropriate catalyst must be used to increase the reaction rate. However, high sulfur yields still necessitate a multistage process with inter-stage cooling and sulfur condensation [8].

Sulfur formed in each stage of the Claus plant is condensed and recovered to achieve maximum conversion in the catalytic reactors. The unrecovered sulfur, in elemental or combined form (H₂S, COS, CS₂), is combusted to SO₂ in the tail gas incinerator which is then emitted to the atmosphere. Tail gas clean-up units are added sometimes prior to incineration to increase the sulfur recovery and minimize emissions [7].

One of the furnace functions is the destruction of any contaminants what may foul downstream equipments. In oil refinery operations, NH₃ is formed as a byproduct what is then directed to the sulfur recovery facility for destruction [5]. Incomplete pyrolysis or combustion of NH₃ in the furnace results in NH₃ and NO carryover into the catalyst beds. Ammonia can form ammonium salts, which can plug or foul the catalyst beds, other equipments, or piping. Although the formation of SO₃ occurs in the catalyst bed regardless of the presence of NO, the presence of NO in the beds acts as a catalyst for the conversion of SO₂ to SO₃, which in turn causes catalyst sulfation [9]. Of the primary causes of catalyst activity loss, catalyst sulfation is regarded as the most significant [10]. It is therefore critical to convert as much NH₃ to N₂, H₂, and H₂O as possible.

For ammonia destruction, an empirical rule of thumb in industry is that furnace temperature should be greater than 1200-1250 °C [5]. The furnace temperature must be below the temperature limitation of conventional refractories of 1600°C and above the minimum stable furnace temperature of 926°C.
The reaction furnace temperature should not exceed 1380 °C in order not to exceed the maximum temperature limitations of the equipment materials [12].

In the Claus process, other sulfur compounds will be formed, such as carbon disulfide (CS₂) and carbon oxysulfide (COS), and these compounds can often contribute from 20 to 50% of the pollutants in the tail-gas [13,14]. Furthermore, presence of O₂ traces in the CS₂-H₂O mixture caused a decrease in the activity of alumina and titania catalysts due to sulfate formation [15]. Therefore, COS and CS₂ should be hydrolyzed in the catalytic converter [16,17], as shown below:

\[
\text{COS} + \text{H}_2\text{O} \leftrightarrow \text{H}_2\text{S} + \text{CO}_2
\]

\[
\text{CS}_2 + 2\text{H}_2\text{O} \rightarrow 2\text{H}_2\text{S} + \text{CO}_2
\]

The temperature of the first catalytic reactor is maintained at about 350°C to hydrolyze COS and CS₂, while that of the subsequent reactors is just above the sulfur vapor dew point [18]. Transition metal oxides can be used to modify gamma-alumina to form a catalyst that is effective at temperatures higher than the dew point of sulfur [19-21]. However, thermodynamics provide a strong incentive to operate the catalytic converters at low temperature [22, 23] as a lower temperature should increase the exothermic reaction efficiency. Therefore, the temperature of the process gas at the inlet of the catalytic converters should be such that the effluent gas temperature is about 14-17°C higher than the expected outlet sulfur dew point and high enough for hydrolysis of COS and CS₂ for the first catalytic converter only [24] (about 350°C).

In this paper, the temperature of reaction furnace of a typical Claus Sulfur Recovery Unit is adjusted to ensure suitable NH₃ destruction. Moreover, the inlet temperatures of SRU converters are determined such that the proper conversion can be achievable without any processing problems. The process temperatures are important in designing the Claus sulfur recovery units.

II. SRU REACTION FURNACE

As mentioned before, since only one-third of H₂S convert to SO₂ in the reaction furnace, the unreacted H₂S and inert gases consume the released energy from this exothermic reaction and the furnace reach to a thermal equilibrium. It decreases the temperature of reaction furnace. For this reason, a split-flow reaction furnace design (Figure 2) is applied to process the gas streams containing considerable amounts of ammonia. In this method, all the combustion air and all the ammonia containing sour gas are mixed with a portion of the acid gas stream and inserted to the first zone of the furnace and the remaining acid gas is then mixed with the products of the combustion from the first zone in zone 2 of the reaction furnace [25,26].

For illustrating the split-flow method, the Claus unit of a typical refinery is considered. The SRU feedstock consists of 74 Kmol/h acid gas from amine sweetening unit at 46°C and 1.3 bar, together with 38 Kmol/h sour gas from sour water treating unit at 90°C and 1.5 bar. The molar compositions of these streams are presented in Table 1. The sour gas containing 275 Kg/h ammonia and therefore, the furnace temperature should increase, up to the ammonia destruction temperature. For this reason, a case study is performed in which the acid gas split into two zone. Figure 3 illustrates the achieved results. The horizontal axis indicates the ratio between the flow rate of acid gas entered to zone 1 and the flow rate of acid gas entered to zone 2. The vertical axis represents the changes of the first zone temperature.

\[
\text{COS} + \text{H}_2\text{O} \leftrightarrow \text{H}_2\text{S} + \text{CO}_2
\]

If combustible gases enter to the furnace first zone as their stoichiometric coefficients, the reaction furnace temperature will be maximized, for no excess gas or excess oxygen (excess air) is there. The maximum point of this curve is related to this condition. Before this point, the amounts of combustible gases (including H₂S) are less than needed one-third of total H₂S and therefore, temperature is decreased. Regarding to Figure 3, if the split-flow ratio was equal to 0.8, a temperature around 1350°C would be achievable for zone 1 of the reaction furnace. This temperature guarantees the complete destruction of NH₃.

If more expensive refractory with better resistance was applied, the furnace would be operate at higher temperatures by decreasing this ratio (see Figure 3).

III. SRU REACTORS (CONVERTERS)

The temperature of the process gas at the inlet of the catalytic converters should be such that the effluent gas
temperature is about 14-17°C higher than the expected outlet sulfur dew point. For this purpose, the outlet streams of two converters in the above plant are investigated. Table II presents pressure, temperature, and compositions of outlet streams of these converters. These data are achieved by the assumption of 265°C for temperature of the first converter inlet stream and 195°C for temperature of the second converter inlet stream.

**TABLE II THE OUTPUT CONDITIONS OF CLAUS CONVERTERS**

<table>
<thead>
<tr>
<th>Component (mass%)</th>
<th>Converter 1</th>
<th>Converter 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>S</td>
<td>28.0</td>
<td>22.9</td>
</tr>
<tr>
<td>H₂O</td>
<td>22.9</td>
<td>28.0</td>
</tr>
<tr>
<td>CO₂</td>
<td>0.6</td>
<td>0.7</td>
</tr>
<tr>
<td>SO₂</td>
<td>2.7</td>
<td>0.8</td>
</tr>
<tr>
<td>H₂S</td>
<td>0.2</td>
<td>0.2</td>
</tr>
<tr>
<td>N₂</td>
<td>56.4</td>
<td>65.7</td>
</tr>
<tr>
<td>Sulfur</td>
<td>14.4</td>
<td>3.8</td>
</tr>
</tbody>
</table>

Figure 4 represents the variations of vapor phase fraction vs. outlet temperatures of two Claus reactors. According to this figure, dew point of the first converter outlet stream is equal to 253°C which is 57°C less than its assumed temperature from Table 2. With respect to criterion of 14-17°C, the temperature of 270°C seems to be appropriate for the outlet temperature of first converter. However, this temperature should be increased to ensure hydrolysis of COS and CS₂ in the first reactor. Therefore, the assumption of 265°C is a good value for the temperature of the first converter inlet stream. Furthermore, dew point of second converter outlet stream is equal to 210°C which is 15°C less than its assumed temperature value from Table II and takes the above criterion in the consideration.

**REFERENCES**