# CATALYST MANAGEMENT

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### Abstract

Sulphuric acid plants now and then see unexpected increases in  $SO_2$  emissions. Our experience is that it is not always easy for the acid plants to pin point the exact reason for the  $SO_2$  increase.

Sometimes a catalyst bed is sacrificed even though reliable emperature readings and gas analyses could have verified satisfactory activity, hence the leaking heat exchanger is never discovered and after the shutdown the  $SO_2$  emissions are still too high.

This paper will look into some basic catalyst management tools: Good record keeping is a ne-cessity for collection of representative catalyst samples, reliable operating data is an inevitable starting point for all analysis and trending of operating data is a must for plant optimisation and troubleshooting. In addition these catalyst management tools will give a continuous updated knowledge of the catalyst status in each bed providing the basis for a more effective replace-

ment strategy.

The most typical reasons for increasing  $SO_2$  emissions will be presented together with easy guidelines for cross-checking of operating data, temperature optimisation and good record keeping.

### Collection of reliable operating data

Reliable operating data is an inevitable starting point for all analysis. The three most important

control parameters for optimum catalyst performance and early detection of potential problems

are:

Catalyst temperatures inlet and outlet each catalyst bed Gas compositions inlet and outlet of the converter or preferably inlet and outlet each catalyst bed Pressure drops across each catalyst bed

It is important that these parameters are measured on a regular basis. We recommend that pressure drops are measured once per week. The data should be graphed either manually, in a spreadsheet or in a DCS. Any sudden step changes should be investigated to determine the cause.

The reaction of sulphur dioxide with oxygen to form sulphur trioxide is a highly exothermic, reversible reaction, associated with a reduction in gas volume:

$$SO_2 + \frac{1}{2}O_2 \leftrightarrow SO_3 + heat$$

Since the reaction is exothermic, the temperature will rise with approximately 28°C per mole % of SO<sub>2</sub> converted across the catalyst bed, i.e. the total theoretical temperature rise across the converter can easily be calculated from:

 $\Delta$ Ttotal, theoretical = %SO2, inlet x 28°C

The actual temperature rise across the converter can be calculated from the temperature rise across each catalyst bed:

$$\Delta$$
Ttotal, actual <sup>=</sup>  $\Delta$ Tbed 1, actual <sup>+</sup>  $\Delta$ Tbed 2, actual +... $\Delta$ Tbed n, actual

The million dollar question now is: Is the total temperature rise across the catalyst passes in agreement with the SO<sub>2</sub> inlet concentration?

 $\Delta$ Ttotal, actual <sup>=</sup> $\Delta$ Ttheoretical = %SO2, inlet x 28°C

If a disagreement between total actual temperature rise and the  $SO_2$  inlet concentration is ob-served it is important that the reason for this inconsistency is pinpointed and rectified. The following may be considered in the search for possible reasons:

Temperatures should be taken with thermocouples located in the interface between the catalyst and the inert material. Thermocouples located in the gas stream may read 10-20°C too low due to poor heat transfer in the gas phase and heat loss to the surroundings via the thermowell. Thermocouples must be checked and calibrated regularly.

In order to verify temperature homogeneity in the reactor cross section it is recommended that multiple thermocouples are installed (e.g. three thermocouples 120° apart) at the inlet and outlet of each bed. Temperature differences at the inlet to the catalyst layer tend to be amplified through the catalyst bed. A locally lower inlet temperature leads to a lower rate of reaction in that section of the catalyst yielding a smaller conver sion and consequently a smaller temperature increase. Thus, temperature variations at the entrance to a catalyst layer may result in a temperature and concentration spread at the exit meaning an inferior utilisation of the catalyst mass.

Temperature variations may originate from improper mixing of hot and cold process gas before entering the catalyst mass. This may be proved by making holes in the duct and measuring temperature variations by means of travelling thermocouples.

Imperfect gas mixing becomes even more serious if two gases not only are of different temperatures but also of different composition. This will be the case when interbed cooling is performed by injection of cold feed gas or quench air. In these cases cross-sectional temperature as well as concentration variations may occur.

 $SO_2$  measurements should be made on locations that provide gas samples that are representative for the entire gas flow. It is our experience that sample points on the converter wall may produce non-representative gas samples as tiny streams of lower converted gas may channel along the wall, interfering with the samples collected here. More reliable results can be obtained when sampling from points on the inlet ducts and especially the outlet ducts after the gas has been allowed to mix. For this reason, it is preferable that sample points are accessible on the ducts before and after each catalyst bed.

# Analysis of operating data

When eliable operating data are secured more detailed analyses of the converter operation can begin.

All data must be recorded and plotted to better illustrate possible developments in the performance and to provide the best basis for optimisation and troubleshooting. Any development in temperature rise,  $SO_2$  emission or pressure drop must be followed closely and can in most cases reveal the root cause for changes in the performance.

# Typical reasons for increasing SO<sub>2</sub> emission

# Example 1: Non-optimum inlet temperatures

Temperature optimisation at the inlet of each catalyst bed is crucial for maintaining the lowest possible  $SO_2$  emission. Each set of operating conditions (i.e. inlet flow and  $SO_2$  strength) will require different pass inlet temperatures in order to optimise the operation. Temperatures should therefore be optimised whenever the operating conditions have changed significantly.

Due to the temperature dependence of the equilibrium conversion, the lower the inlet temperature to a catalyst bed is, the higher the equilibrium conversion will be. On the other hand, the lower the catalyst temperature is the lower the rate of reaction will be. For each catalyst bed there will therefore exist an inlet temperature at which the conversion, and correspondingly the temperature rise across the bed:  $T_{out}$  -  $T_{in}$ , is maximum.

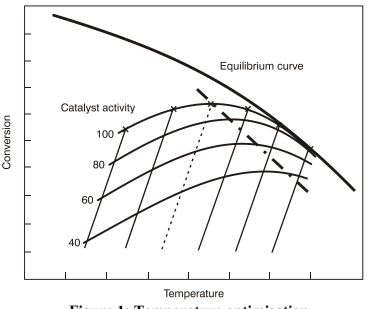


Figure 1: Temperature optimisation

Optimisation of the converter inlet temperature is therefore a trial-and-error search for the inlet temperature at which  $T_{out}$  -  $T_{in}$  and thereby conversion will be maximised.

The necessity of increasing the inlet temperatures as activity declines due to catalyst deactivation is illustrated in Figure 2 which shows the conversion as function of inlet temperature and catalyst activity. It is seen that to each activity level corresponds an optimum inlet temperature. The ----- line shows how the conversion is decreasing if the inlet temperature is kept constant. The optima have been connected by the -- line, which then shows how the optimum inlet temperature increases as activity decreases.

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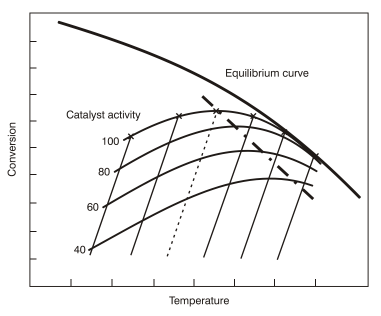


Figure 2: Temperature optimisation due to catalyst deactivation

### What is wrong with my plant?

The SO<sub>2</sub> emissions are increasing. Historical records show that the overall temperature rise is constant. Inlet temperature curves show no significant change for beds 1, 2 and 3. A slow drop of the inlet temperature to bed 4 is experienced, however, the temperature rise across bed 4 is almost constant. Figure 3 shows the SO<sub>2</sub> emission as function of inlet temperature to bed 4 and this figure illustrates the effect of non-optimum temperatures inlet bed 4. The basis is a plant with a 3+1 layout, 11.3% SO<sub>2</sub> and 9.7% O<sub>2</sub> in the feed gas and an overall conversion of 99.8% corresponding to 270 ppm SO<sub>2</sub>. It can be seen from the figure that a non-optimum inlet temperature has a significant effect on the SO<sub>2</sub> emission: A difference of +/- 10°C from the optimum temperature of 420°C results in a 15-20% increase i n SO<sub>2</sub> emission. So, even though the temperature rise across bed 4 is almost constant the non-optimum inlet temperature to bed 4 is the reason for the increasing SO<sub>2</sub> emission.

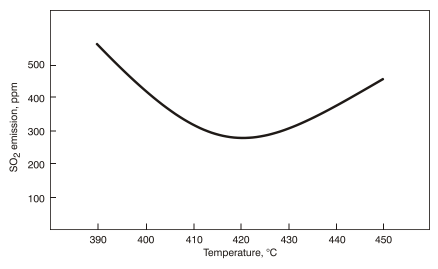


Figure 3: Effect of non-optimum inlet temperature to bed 4. Basis: 3+1 layout, 11.3% SO<sub>2</sub> and 9.7% O<sub>2</sub> in the feed gas. SOR conversion = 99.8% corresponding to 270 ppm SO<sub>2</sub>

### Example 2: Increased SO<sub>3</sub> level inlet bed 4

The presence of SO<sub>3</sub> at the inlet of the catalyst pass located after the intermediate absorption tower (IAT) can originate from either poor performance of the IAT or from leaking reheat-exchangers. In many double absorption plants the cold gas from the IAT is reheated with the exit gas from the third pass before entering the final catalyst pass(es). This means that SO<sub>2</sub> gas containing no SO<sub>3</sub> is reheated with the partly converted SO<sub>2</sub> gas containing high amounts of SO<sub>3</sub>. The amount of SO<sub>2</sub> on an SO<sub>3</sub>-free basis is the same in the two gas streams and the leak thus cannot be detected by a Reich's test. The gas coming from the IAT may contain acid carry over and the result is corrosion and leaks in the hot and/or cold reheat-exchangers. Consequently an SO<sub>3</sub> increase will be experienced at the inlet of the final catalyst pass(es). This has a negative effect on the equilibrium and the equilibrium curve is pushed downwards as illustrated in Figure 4. The final result is a lower conversion in the catalyst passes after the IAT and consequently increasing SO<sub>2</sub> emissions.

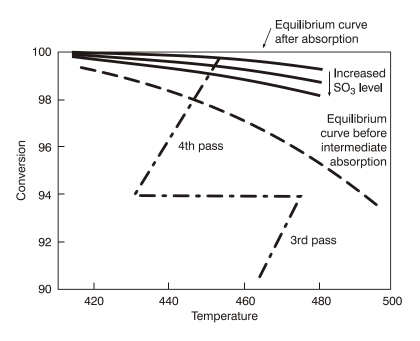


Figure 4: The SO3 concentration effect on the equilibrium curve after IAT

When a gas containing SO<sub>3</sub> contacts humid air, white acid fume is formed immediately. This phenomenon can be used in detecting the presence and origin of SO<sub>3</sub> downstream the IAT. The gas leaving the IAT should be clear. If this gas is visible, too much SO<sub>3</sub> is slipping through the IAT. If the gas leaving the IAT is clear, but the gas entering the last pass(es) is visible there is a heat exchanger leak. If there is more than one heat exchanger between the IAT and the converter, the leaking exchanger can be determined by checking the gas inlet and outlet of each heat exchanger. The heat exchanger that has clear gas at the inlet and visible gas at the outlet is the leaking heat exchanger (Figure 5).

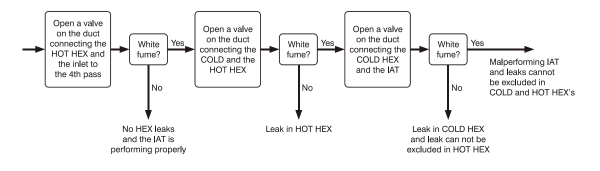


Figure 5: Visual detection of S03 leaks

What is wrong with my plant?

The  $SO_2$  emissions are increasing! Historical records show that the overall temperature rise is almost constant. Inlet temperature curves show no changes for beds 1, 2, 3 and 4.

The operating data show no signs of changes in the  $SO_2$  concentration in the feed gas or pressure drop across the individual beds.

The next step is to check the inlet gas to the catalyst pass after the IAT for visible gas. Figure 6 shows the SO<sub>2</sub> emission as function of IAT efficiency and this figure illustrates the effect of increased SO<sub>3</sub> levels inlet bed 4. The basis is a plant with a 3+1 layout, 11.3% SO<sub>2</sub> and 9.7% O<sub>2</sub> in the feed gas, an IAT efficiency of 99.9% and an overall conversion of 99.8% corresponding to 270 ppm SO<sub>2</sub>. It can be seen from the figure that a poor working IAT or leaking reheat-exchangers illustrated as low IAT efficiency has a significant effect on the SO<sub>2</sub> emission.

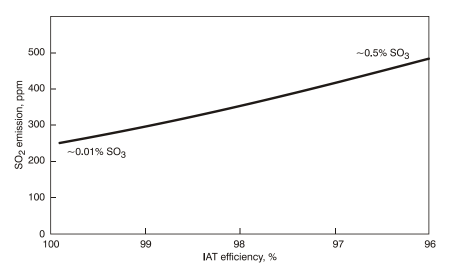


Figure 6: SO2 emission as function of IAT efficiency. Basis: 3+1 layout, 11.3% SO2 and 9.7% O2 in the feed gas. 99.9% IAT efficiency. Start-of-run conversion = 99.8% corresponding to 270 ppm SO2

# **Example 3: Channelling**

Uneven gas flow distribution through the catalyst bed or channelling may be caused by: A collapsed grid causing a hole in the catalyst bed; high gas velocities inlet the converter may cause catalyst movements and possible "holes" in the catalyst bed; and dust build-up in the top of the catalyst bed will reduce the "available" catalyst area.

Channelling will cause non-uniform temperature and concentration profiles at the exit of the bed. A drop in the outlet temperature will only be detected if the thermocouple(s) are located right below the "channelling" area(s). Sometimes the temperature rise is apparently as expected and sometimes an average temperature rise is seen corresponding to a mixture of converted and partly unconverted gas. The average conversion taking place in the bed will be lower than expected i.e. a higher concentration of  $SO_2$  will enter the following bed and give a temperature rise in the following beds which is higher than expected.

Figure 7 illustrates the effect on temperature rise in a four-bed converter with and without channelling. The dark grey colour illustrates the temperature rise across each bed during a three year period in case of no channelling. A slightly decrease in the temperature rise across each bed is caused by the normal catalyst deactivation. The light grey colour illustrates the temperature rise across each catalyst bed during a two-year period in case of channelling caused by dust build-up in the top of the first bed. The figure clearly shows that as the dust deposition develops in bed 1 and the degree of channelling increases the missing conversion in the first bed (shown as a drop in the temperature rise) will show up as increasing temperature rises in the lower beds.

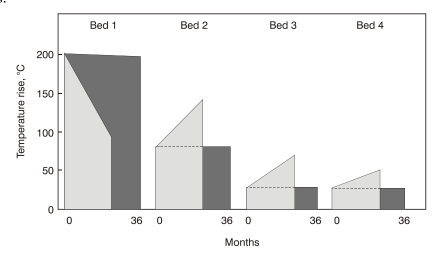


Figure 7: The effect of channelling in bed 1 on temperature rises. Dark grey: No channelling. Light grey: Channelling due to dust deposition. Basis: 3+1 layout, 11.3% SO<sub>2</sub> and 9.7% O<sub>2</sub> in the feed gas. SOR conversion = 99.8% corresponding to 270 ppm SO<sub>2</sub>

To distinguish between the different channelling reasons it is important to study the speed of the change in temperature rise: Is it developing over time or is it a sudden change taking place overnight? In addition, pressure drop measurements may support such investigations: Is the pressure drop increasing due to dust deposition or does it decrease due to a "hole" in the catalyst bed or a dropped grid?

# What is wrong with my plant?

The SO<sub>2</sub> emissions are increasing! Historical records shows that the overall temperature rise has increased and that  $\Delta$ Ttotal, actual  $^{>}\Delta$ Ttheoretical. The records also show that the temperature increase across bed 2 has increased and historical pressure drop curves across each bed show a sudden decrease in pressure drop across bed 1. These signs indicate that a grid has dropped in bed 1 causing low conversion in bed 1, which is partly compensated for in bed 2.

## Example 4: Heat exchanger leak after final catalyst bed

In many metallurgical plants the cold feed gas from the drying tower is as a first step heated with the exit gas from the final catalyst pass. This means that strong unconverted  $SO_2$  feed gas is heat exchanging with highly converted gas from the final catalyst pass only containing a few hundred ppm  $SO_2$  or less. The feed gas is coming directly from the drying tower and often acid carry over problems are experienced. The result is corrosion and tube leaks in this very critical cold-gas-heat-exchanger. The higher pressure  $SO_2$  rich feed gas will leak into the lower pressure  $SO_2$  lean exit gas causing an increase in  $SO_2$  emissions which very often increase over time.

### What is wrong with my plant?

The SO<sub>2</sub> emissions are increasing! Historical records show no changes in temperatures, SO<sub>2</sub> concentration in the feed gas or pressure drop that can explain this increase. Also the test for visible gas at the inlet of the catalyst pass after the intermediate absorption tower is negative. The only development in the operating data is the increasing SO<sub>2</sub> emission indicating that the root of the problem must be found downstream the converter. In order to locate the problem, SO<sub>2</sub> measurements are carried out simultaneously on the lean SO<sub>2</sub> gas at the inlet and outlet of the cold-gas-heat exchanger. An SO<sub>2</sub> concentration at the inlet of the heat-exchanger corresponding to the outlet of bed 4 pinpoints a leaking heat exchanger.

## Good catalyst record keeping

When loading the catalyst, it is important to calculate the depth of the catalyst layer(s) and supporting material for each bed and to mark the converter wall plainly. Good record keeping is invaluable in connection with future screening and catalyst sampling activities especially when a catalyst bed consists of several catalyst layers of different type or age. Precise knowledge about the interface between two catalyst layers e.g. standard catalyst and a more expensive caesium catalyst or new and old catalyst will ease the unloading and prevent mixing of the two catalyst types which would reduce the effect of the caesium catalyst. In addition, this knowledge is essential when collecting catalyst samples as otherwise wrong and expensive conclusions regarding future replacement strategies may be drawn. We also recommend that all inspections in the converter and all phenomena observed is documented with photos and/or videos which can aid in troubleshooting and operator training programmes.

# Spent catalyst samples

Activity testing of spent catalyst samples in combination with evaluations of the catalyst performance based on operating data is a very efficient tool to follow the catalyst history. However, as described above activity information will only have value if exact knowledge about catalyst type, age and location are known for each sample.

In order to obtain representative catalyst samples from the converter beds we recommend collecting samples according to the below guidelines:

As the first pass has been exposed to the highest  $\Delta T$  as well as possible contaminants in the feed, it is recommended to sample from three depths down through the catalyst bed for characterising the performance of the bed.

Four samples should be collected from each of the layers: top, middle and bottom. The top samples should be collected from a depth of 5-10 cm (not including the inert material). Likewise, the bottom samples should be collected 5-10 cm from the bottom. The four samples from each layer should be collected 90°C apart, e.g. in the d irections: N, E, S and W and then combined to one sample for testing, i.e. three composite samples representing the top, middle and bottom layers of bed 1 are sent for testing.

For the second pass it is adequate to test composite samples representing the top and bottom layers. For the lower passes a single composite sample from each pass is sufficient. This sample should be collected 5-10 cm from the top of the catalyst bed.

It is important that the sample containers are marked to identify the sample location (bed No and layer) and that each sample is followed by information on catalyst type, age and any possible observations or pictures regarding dust depositions etc.

### Conclusion

Reliable operating data is required for all performance analyses. The operating data should be measured and monitored on a regular basis. If any discrepancy exists between  $\Delta$ Ttotal, actual and %SO2, inlet it is important that the reason for this inconsistency is pinpointed and rectified.

When reliable operating data is obtained a more detailed analysis of the converter operation can begin: All data should be recorded and plotted to better illustrate possible trends in the performance and to provide the best basis for troubleshooting and optimisation. Any changes in temperature rise, SO<sub>2</sub> emissions and pressure drop should be investigated thoroughly and in most cases it will reveal the root cause of changes in the plant performance.

Good catalyst record keeping is invaluable for planning future catalyst screening and sampling activities, especially when a catalyst bed consists of various catalyst layers of different type or age.

Activity testing on spent catalyst samples in combination with evaluations of the catalyst performance is a very efficient tool for catalyst management. However, activity information will only have value if exact knowledge about catalyst type, age and location is available for each sample.

A deeper understanding of the actual status of the catalyst in each pass of a sulphuric acid converter is important knowledge when troubleshooting, optimising plant performance and developing the most cost-effective replacement strategy.

# The Author



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- Ms. Lene Hansen holds a B.Sc. in chemical engineering and joined the Catalyst Division of Haldor Topsoe in 1994.
- In 1996 she was appointed Product Manager for Topsoe's sulphuric acid catalysts and travelled all over the world with Topsøes portable gas analysis unit the TOPGUN and gained a substantial hands-on experience in trouble-shooting on a wide range of sulphuric acid plants
- In 2004 she took up the position as General Manager for Topsoe's sulphuric acid catalysts