MULTIPHASE LEVEL MEASUREMENT

Improving level control in desalters to aid in use of opportunity crudes
Introduction

Refineries are a complex set of various operating units designed to process crude oil into refined products such as gasoline, diesel, jet fuels and feedstocks for thousands of different products that people use every day. In order to maximize operating margins, refiners rely on pushing the units to maximum throughput. To do this safely and effectively, they must have very good control of the process. Some processes in the refinery industry are a challenge for even the best and most experienced engineers. As Lord Kelvin once said, “If you cannot measure it, you cannot improve it.” and H. James Harrington expanding that thought to say “Measurement is the first step that leads to control and eventually to improvement. If you can’t measure something, you can’t understand it. If you can’t understand it, you can’t control it. If you can’t control it, you can’t improve it.” These statements apply to all refineries, whether they are small or large. The measurement of the density of the fluids is crucial. When we talk about interface level, this is a misnomer, since there isn’t really a defined level between the crude oil and water. There is instead a transition zone where the fluid slowly changes from crude to water. This transition zone is both undefined and under certain circumstances can vary greatly in height (emulsion thickness, see Fig. 2). When interface level is checked by sampling, you will see a variation in amount of oil and water mixed together at different elevations. Many refineries use sampling as their primary method of interface level control in the desalter, due to the lack of confidence in the current technology used. Various technologies have different issues with reliably measuring and thereby controlling the interface level. As refiners try to meet various national, local, and corporate standards, the sampling method itself causes safety risks and environmental impact concerns. What is done with the fluid that must be flushed from the sample lines? With environmental regulations, this oily water has to be collected and treated. How long do you flush the lines? The time it takes to flush line can lead to more fluids that must be collected and treated as well as errors if the line is not flushed long enough. How to limit human error? One person might determine the sample to be all oil whereas another person might determine the sample to be an emulsion. Then, there are safety concerns, these include raising the temperature to 300 °F / 150°C to assist with the separation of the water and salts from the oil. Therefore, sample lines must have coolers to prevent the potential burns to the person taking the sample and with refiners integrating petrochemicals, more benzene can be present causing an inhalation issue.

Fig. 1: Berthold’s EmulsionSENS resulting density profile

Optimize existing desalters

How can we get the most out of our existing desalter? We can ensure that the grids are working to their fullest and that the maximum volume is being utilized in the vessel to increase residence time, ensuring maximum desalting of the oil (Lobo, Kremer, & Cornelius, 2010, slide 23). To ensure that we can achieve this greatest efficiency, control of the interface level becomes crucial. When we talk about interface level, this is a misnomer, since there isn’t really a defined level between the crude oil and water. This transition zone is both undefined and under certain circumstances can vary greatly in height (emulsion thickness, see Fig. 2). When interface level is checked by sampling, you will see a variation in amount of oil and water mixed together at different elevations. Many refineries use sampling as their primary method of interface level control in the desalter, due to the lack of confidence in the current technology used. Various technologies have different issues with reliably measuring and thereby controlling the interface level. As refiners try to meet various national, local, and corporate standards, the sampling method itself causes safety risks and environmental impact concerns. What is done with the fluid that must be flushed from the sample lines? With environmental regulations, this oily water has to be collected and treated. How long do you flush the lines? The time it takes to flush line can lead to more fluids that must be collected and treated as well as errors if the line is not flushed long enough. How to limit human error? One person might determine the sample to be all oil whereas another person might determine the sample to be an emulsion. Then, there are safety concerns, these include raising the temperature to 300 °F / 150°C to assist with the separation of the water and salts from the oil. Therefore, sample lines must have coolers to prevent the potential burns to the person taking the sample and with refiners integrating petrochemicals, more benzene can be present causing an inhalation issue.

In the market today with all the opportunity crudes that are available, it is important to know how the different crude oils react when mixed together to ensure compatibility. If oils are mixed that are not compatible, they could form emulsions that are very difficult to break (Garrett, Rattanakhambay, Robbins, Wunder, & Yeung, 2016, p. 1). When this happens in the desalter, knowing not only the brine water level is important, but also the top of the emulsion layer to ensure proper desalter operation. There are several technologies that are used to measure the water level in the desalter:

- differential pressure (DP)
- RF absorption probes
- nuclear density profile systems

These can be placed into two basic categories: direct and indirect measurement technologies, with all of them falling into the direct measurement category except the nuclear density profile systems. The direct measurement devices measure a physical characteristic of the fluids (dielectric constant (dK), capacitance, weight, etc.) by directly contacting process fluids. Each one of these technologies have their advantages and disadvantages. These technologies work very well when there is a defined interface between the two fluids, but due to the operational characteristics of the desalter, there might not be a defined interface but a graduation of densities as the fluids separate. When this graduation of densities occurs, the nuclear density profile system is the best option. The nuclear option will provide the user with a density profile of the desalter so that the levels of water and emulsion can be determined, monitored and controlled. In this paper we will briefly explain how DP, RF absorption probes and nuclear density profiler systems measure the interface level in the desalter and some of the issues that each technology may have.
Differential Pressure

DP is an economical way of measuring levels in tanks and vessels and it is the second most common use of pressure measurement behind flow measurements. DP levels use Pascal law to determine level by relating the level, pressure and density of the fluid, simply stated, $P = \rho \times h$, where $P =$ pressure (pascals, Pa), $\rho =$ density of fluid (kg/m$^3$), and $h =$ height of the column of fluid (meters, m). You can solve for $h$ by rearranging Pascal law, $h = \frac{P}{\rho}$ (Meribout, Al Naamany & Al Busaidi, 2011, p. 2). For example, a tank with 2 m of water will exert a pressure of 2000 mmH$_2$O or 0.196 bar. If the fluid was kerosene with a density of 0.82 spg, the pressure exerted will be 1640 mmH$_2$O or 0.161 bar. As you can see the density of the fluid has a great influence on the pressure. If this was a level application calibrated on water to measure a 2 m span and the process fluid changed to kerosene with a density of 0.82 spg, the error of the level output at 50% level would be 9%, or the output would read 41% but actual level would be 50%.

The errors are more compounded for DP levels when trying to measure interface levels in vessels especially if emulsions are present. The difference in pressure being measured is the difference in pressure exerted by the vessel full of the higher density fluid and the vessel full of the lower density fluid. Since two fluids are being measured, the error can be compounded if both fluid densities change. Another point that can cause issue with the DP level measurement is the height of the emulsion. Figure 2 (top) shows level with a relative-ly small transition zone and Figure 2 (below) shows a level with a larger transition zone. Emulsion is not the true word to use in the case of the desalter since emulsion implies a fluid of consistent density, but here we use this term to describe the transition zone from clear oil to clear brine. This zone is actually a gradient change in density from the lighter density fluid to the heavier density fluid. The DP level will only measure the average density across the full span of measurement and cannot tell the operator if the level output is the bottom of this emulsion, the middle or top of the emulsion height.

RF Absorption

Another type of measurement technology that is commonly used in desalters is the RF absorption probe. These RF probes and the nuclear density profiling systems have a similar basis of measurement, which is the absorption of energy. One is the absorption of RF energy and the other is the absorption of gamma energy. The RF energy absorption is based on the dielectric constant (dK) of the fluid and the gamma absorption is based on the material density. Beyond this common principle, they differ greatly. The RF energy detectors are reference devices, meaning that they must be calibrated with oil and water to measure the difference in the amount of energy that is being absorbed. This can lead to some erroneous readings if a refiner changes the crude slate that is being refined since different crudes will absorb different amount of RF energy. For example, if the RF energy probes are calibrated on 26 API oil and water to obtain 0% to 100% water, and the crude is switched to 16 API, the RF energy probes can be reading 15% - 20% water even in 100% crude oil. This is due to the fact that the 16 API oil absorbs more of the RF energy than the 26 API oil, therefore, to ensure proper operation of the desalter the device needs to be recalibrate. Some other issues that can cause errors in the measurement is the probe coating and only measuring the RF energy absorption of the coating and not the material around the probe. In this case, the probes must be removed and cleaned to restore proper operation. As shown in Figure 3 desalter can acquire with large amounts of buildup.

The fourth detector may be installed on the water/brine outlet to measure the amount of hydrocarbon that is potentially being discharged from the desalter with the water/brine (Agar Corp, 2018). The RF energy absorption probes are calibrated in the range of 0% to 100% water. The probes measure how much RF energy is absorbed by the material surrounding it. Water absorbs more RF energy than the oil. The RF energy detectors are reference devices, meaning that they must be calibrated with oil and water to measure the difference in the amount of energy that is being absorbed. This can lead to some erroneous readings if a refiner changes the crude slate that is being refined since different crudes will absorb different amount of RF energy. For example, if the RF energy probes are calibrated on 26 API oil and water to obtain 0% to 100% water, and the crude is switched to 16 API, the RF energy probes can be reading 15% - 20% water even in 100% crude oil. This is due to the fact that the 16 API oil absorbs more of the RF energy than the 26 API oil, therefore, to ensure proper operation of the desalter the device needs to be recalibrate. Some other issues that can cause errors in the measurement is the probe coating and only measuring the RF energy absorption of the coating and not the material around the probe. In this case, the probes must be removed and cleaned to restore proper operation. As shown in Figure 3 desalter can acquire with large amounts of buildup.

The RF energy absorption probe (controller) measures in one localized area and only provides information at that specific elevation. This can be used for control of the interface level in a separator if the emulsion stays relatively small. If the emulsion layer were to grow inside the vessel the operator would be blind to this change until other issues occur since the control point RF energy absorption probe might not detect any change. For example, if the control point is set at 50% water and the “emulsion” bands grows from 75 mm to 300 mm, the RF probe can still be reading 50% water but now the top of the emulsion band is 150 mm higher in the vessel than what is being indicated by the RF probe. This is also true of the bottom of the emulsion band, which is now 150 mm lower than indicated. Figure 4 shows different transition zones, all of these levels could measure 50% water even though the top of the emulsion is at different heights.

Fig.2 Different emulsion thicknesses in the vessel

Fig.3 Inside of a desalter with large amounts of buildup

Fig.4 Use of RF absorption probes in vessel with different emulsion heights
Operating companies are starting to rely more on nuclear density profile systems since they can give the operators a view into the desalter itself. A larger global refiner once stated “I understand the nuclear density profiler system is more expensive to install but I know it works a 100% of the time no matter what type of crude we are processing, whereas the other technologies might only operate correctly 85% of the time. This 15% difference cost us more than the price of the nuclear density profile system.” There has been a trend in the large major oil refineries to rely more on nuclear density profile systems due to this reasoning.

Nuclear Density Profiler uses radioactive sources and a radiation-based detector to measure density of fluids inside the vessels at various elevations in order to provide the user with a density profile of the fluids inside the vessel as they separate. The density is measured by the amount of radiation being sensed by the detector. The amount of radiation that is absorbed is constant, predictable and directly proportional to the density of the process fluid. This makes the measurement of the fluid density very reliable. The basic principle of a radiometric based density gauge is that materials attenuate radiation at a given rate and this attenuation of radiation can be calculated by Lambert-Beers law (\( I = I_0 \times e^{-μ} \)). The change in the amount of radiation from a radioactive source can then be used to calculate the density of the fluid that is between the source and detector. This change in the amount of radiation to the detector is measured and then converted to a density value for each elevation. Therefore, operator can basically see the transitions zones in real time as the water/brine separates from the oil.

**Different Arrangements**

There are several types of density profiling systems, all of them have their advantages and disadvantages but all work on the same basic principle stated above. There are four types of arrangements of radiometric density profilers. The main difference between them is the orientation of radioactive sources and detectors. They are listed as follows:
- Internal sources and internal detectors
- Internal sources and external detectors
- One internal source and external detectors
- One external source and external detectors

**Internal sources and internal detectors**

Sources and detectors are located inside the vessel mounted in dip tubes or wells (see Figure 5). Each detector has its own source and the density is measured at each of these elevations to provide the end-user a density profile. This type of profiler typically uses Am-241 as the radioactive source and the source is loaded into the source well by a qualified radiation worker. Am-241 is a lower energy gamma emitter and is therefore very sensitive to small changes in density, but also requires a very short process path (typically not more than 75 mm, this means buildup can be an issue). Buildup can be an issue since the process path is small and a small layer of build is a large percent of the measurement span. For example, if the source and detector wells have 10 mm of buildup, that would equate to 26% of the process being measured would be buildup and not process fluids. Also due to the lower gamma energy, the source and detector wells need to be made out of Titanium, since it has a lower density than steel. Carbon steel attenuated too much of the gamma energy, therefore not enough energy is left in order to make a useful measurement of density. Since the detectors are inside the vessel, cooling is needed for the electronics if the process is heated.

**Internal sources and external detectors**

An example of this arrangement is Berthold’s EmulsionSENS. Sources are mounted inside the vessel inside a dry well or dip tube (see Figure 6). The sources used can be Cs-137 or Co-60. The shield, used to safely house the radioactive capsules when not inserted into the vessel, is mounted on top of the dry well. Thus a person with basic radiation safety training can safely insert or retract the sources back into the holder and lock them out for vessel entry during maintenance. The energy of Cs-137 is higher than that of Am-241, this allows for the use of standard material such as carbon steel, stainless steel or monel. The process path is also large, typically from 450 mm to 650 mm. Sources and detectors are either aligned at the same elevation or staggered to allow detectors to measure the average density between the elevation of two sources. Detectors are mounted outside of the vessel and can be mounted inside a collimator ensuring that the detector is receiving maximum gamma energy from the source directly across from it to provide the highest accuracy in the density measurement. When the detector is used without a collimator, the detectors can detect small changes in density between the mounted elevations in order to provide a more accurate level reading. Each detector has its own source or is mounted in between the sources and the density is measured at each of these elevations to provide the end-user a density profile. The drywell can be curved to conform to the vessel wall to ensure nearly the same process path at each elevation to provide the proper resolution. With the correct process path, buildup on either the well or vessel wall has a negligible effect on the density output. For example, if the process path is 600 mm, and there is 10 mm buildup on both the vessel and dry well, only about 3% of the process path is affected. If the buildup has density of 950 kg/m³ and the fluid has a density of 920 kg/m³, the error in the density reading would only be 0.9 kg/m³ (0.3% error) if the process span was 700-1000 kg/m³ therefore recalibration would not be required. Since detectors are mounted outside the vessel, no cooling is needed for electronics and maintenance is easy.

**One internal source and external detectors**

A single source is placed inside the vessel and an array of detectors are mounted outside the vessel (see Figure 7). The detectors use an advance algorithm in additional PLC or controller, to determine the density at each elevation depending upon the density of the elevations below and above its physical location. Source used is Cs-137, each single source is limited to approximately 1200 mm span. The source has a source holder mounted on top of the dry well and can be safely retracted by an individual with basic radiation safety training and locked out for vessel entry. If longer span is desired, this can be achieved by using multiple inserting dry wells incl. sources along the length of the vessel.

**One external source and external detectors**

A single source is placed outside the vessel and an array of detectors are mounted above and below the mounted source (see Figure 8). The detectors measure the amount of radiation that is scattered back to the detectors instead of the amount of radiation that is absorbed. The detectors then use an advance algorithm to calculate the density at each elevation using densities of the elevations above and below.
Berthold’s multiphase level measurement system EmulsionSENS consists of radiometric density devices at different elevations through the measurement span (see arrangement “internal sources and external detectors” Figure 1 and Figure 6) and optional evaluation units LB 478. Such an arrangement allows accurate measurement of the density at each of these positions and provides a density profile of the fluids as they separate inside the vessel. This difference in density of the fluids is the main principle of the separation vessel. The density is calculated in each individual detector and then sent to the DCS via 4–20 mA HART, which can then be used to display the density profile on main operators screen on the DCS. By means of an algorithm implemented in the evaluation unit LB 478, interface levels can be calculated from the measured density values. Therefore the density values of the individual detectors are sent to the evaluation unit LB 478 and the calculated interface levels are then transmitted to the DCS via a 4–20 mA signal. This calculated interface level can then be used for automatic control of the water/brine outlet valve. This ensures redundant signal transmission to the DCS. In addition to the calculation, the evaluation unit is also used for numerical and graphical display of both density and level values. This allows the operator to see where the top of the emulsion is located as well as the bottom of the emulsion. Figure 9 shows the break points between clear oil, emulsions and water due to their different densities.

**Improving Level Control**

Refiners have experienced a reduction in upsets in the desalters from 2–6 upsets a year to none for the two years when operating with a nuclear density profiling system. The main reason for this reduction of upsets is the ability to provide automatic control. The brine outlet valve is normal controlled manually by the operator and small changes in valve setting can take up to 6–7 hours to obtain the full level change caused by the valve setting change. This long lag time from setting change to actual level change tends to lead to level being increased too high or too low (large level setting change to actual level change tends to lead to long lag time from setting change to actual level change). This is typical of a density system, the water outlet valve is being continuously adjusted to maintain proper control and therefore minimize the risk of having salt, water and mineral carryover (see Figure 11). This information is not only useful to the operators to control proper level, it is also useful to the chemical treatment companies since it allows them to monitor the effectiveness of chemical treatment. Some refineries are seeing an increase in the use of chemicals to try to manage the emulsions in the desalter. It is not uncommon to see an increase from about 3 ppm of demulsifiers added to about 35 ppm. The cost for chemicals which removes a specific element i.e. Iron is typically about 3 – 5 €/kg.

If a refinery is 270.000 barrels a day (B/D), at 3 ppm the chemical cost is 436 € per day or 159,200 € annually. At 35 ppm the chemical cost would be 5,100 € a day or 1,857,300 € a year, an increase of almost 1.7 million Euros a year. When using the nuclear density profiling system, reducing the chemical usage by 25% (from 35 ppm to 26 ppm) can have a savings of approximately 500,000 € a year. In addition to the reduction of upsets and the chemical savings, a refiner can greatly increase the reliability of downstream equipment of the desalters by severely limiting the amount of corrosion caused by salt carryover and extending the life of catalyst by increasing the efficiency of the desalter. All in all, the profitability of a refiner can be significantly increased.

**References**


**Figures**

- **Fig. 9** Example of a density profile
- **Fig. 10** Typical density curve with manual adjustment of water outlet
- **Fig. 11** Typical density curve with automatic adjustment of water outlet
Outstanding long-term stability

A reliable measurement is vital for the operation of a process and is therefore, our highest priority. Berthold’s detectors operate consistently irrespective of changes in ambient temperature. Even drastic temperature shifts, e.g. from winter to summer don’t influence the measurement drift. Due to various patented technologies for detector stabilization and the use of cosmic radiation as an external reference source, the detector’s output has an accuracy of under 0.001% per °C temperature change. Apart from employing these cutting-edge technologies in our detectors Berthold is also the only supplier that compensates degradation caused by natural aging. The result: many years of operation without the need for recalibration or maintenance and a measurement that you can absolutely rely on!

Protected against X-Ray interference (XIP, RID)

Non-destructive testing e.g. for weld inspections can become really distressing if nothing is done to protect the radiometric measurement against interfering radiation. Every Berthold detector employs the X-ray Interference Protection (XIP), whereby the system is able to detect interference. As a result the measurement value is locked before a false level signal can be communicated. By no means are Berthold detectors harmed by the excessive radiation and automatically return to normal operation after the disturbance is over. By employing Berthold’s unique Co-60 rod sources in combination with our patented Radiation Interference Discrimination (RID) feature, it is even possible to continue the measurement despite non-destructive testing is being carried out. This secures a safe process and makes you independent from actions that might even occur in neighboring plants.

Minimal source activity

Berthold detectors are highly sensitive to gamma radiation. With a scintillation crystal of 150 x 150 mm the SuperSENS is the most sensitive detector on the market. Due to their excellent efficiency the detectors can be operated with very low source activities, which is important for our customer’s HSE programme and also a major cost saving factor. In fact Berthold detectors can be retrofitted on existing measurements where the source has become too weak to work with the current detector.

SIL2 / SIL 3 certified

The SENseries LB 480 detectors are certified for use in SIL2 applications. Even SIL3 is achieved with homogenous redundancy. The certificate covers all measurement applications, from high level or low level alarms to continuous level and density measurement. Thus safe operation of critical processes is guaranteed.
THE EXPERTS
IN MEASUREMENT TECHNOLOGY

Berthold Technologies stands for excellent know-how, high quality and reliability. The customer is always the focus of our solution. We know our business!
Using our varied product portfolio, our enormous specialized knowledge and extensive experience, we develop suitable solutions together with our customers for new, individual measurement tasks in a wide variety of industries and applications. Berthold Technologies is specialised in radiometric process measurements for 70 years. This is our core competence with state-of-the-art and cutting edge products and solutions covering a vast range of industries and applications.

We are here for you – worldwide!
The engineers and service technicians from Berthold Technologies are wherever you need them. Our global network assures you fast and above all competent and skilled assistance in case of need. No matter where you are, our highly qualified experts and specialists are ready and waiting and will be with you in no time at all with the ideal solution for even the most difficult measurement task.

Berthold Technologies GmbH & Co. KG
Calmbacher Straße 22 · 75323 Bad Wildbad · Germany
+49 7081 1770 · industry@berthold.com · www.berthold.com