
Question 2: What are the operating constraints in co-processing coker naphtha in a ULSD (ultra-low sulfur diesel) and/or a gas oil hydrotreater unit?

BODOLUS (CVR Energy)

There are some constraints relative to processing coker naphtha. I have outlined a few of them on the slide. Key with coker naphtha are the changes in the process chemistry that occur starting, perhaps, with the dilution of the hydrogen partial pressure due to the vaporization of the naphtha. Vaporization of the naphtha dilutes that vapor space. There is also coker naphtha having a large exotherm. I refer to it a lot with the Operations staff as it is almost like a match for your process. The coker naphtha starts the exotherm off very early at the topmost parts of the bed, and you have to control that exotherm through the rest of the reactor. The exotherm imposes changes in reactor dynamics, in terms of temperature control and any of the other peripheral equipment that might be heat-integrated with the reactor. It also recovers the naphtha material that has been processed in the stripper tower. One issue associated with contamination is bed fouling due to silicon and diolefins. There is a liability with the coker naphtha if the naphtha is routed to the reformer pool because it might not get out all of the nitrogen. You may wind up looking at enhanced nitrogen levels in your reformer area.

Our economic effects: Octane loss in a distillate hydrotreater is high due to saturation of all of the olefins. Coker naphtha processed in an ultra low sulfur gasoline unit will preserve those olefins and result in less octane degradation. Processing coker naphtha comes at a liability of high hydrogen. It takes a lot of hydrogen to process the coker naphtha. In particular, though, in our unit on which we did a revamp in 2008, the rates of the coker naphtha and of other cracked stocks in the unit presented some interesting challenges to the operators. We found that the interaction with other heavy cracked stocks was complicated, so to speak. We look forward to trying to get a predictive model to help the Operations staff run the unit.

Constraints in co-processing Coker Naphtha in a ULSD and/or GOHT unit

➤ **Changes Process Chemistry**

- Dilution of H₂ Partial Pressure Due to Naphtha Vaporization
- Large Exotherm changes unit Heat Balance / Reactor and Stripper
- Contamination and Bed Fouling from Silicon and Di-Olefins
- May result in in-complete nitrogen removal in treated naphtha

➤ **Effects Economics**

- Octane Loss in ULSD unit high vs processing in ULSG unit
- High Hydrogen consumption

➤ **Provides Challenges for Operations Staff**

- Variable Coker Naphtha rate hard to transition
- Interactions with other Heavy Cracked stocks
- Required "Predictive" model to minimize off-spec / giveaway



This slide shows our model covering the two parallel reactors with two beds in each reactor and an inter-bed quench. We had multiple feedstocks coming from multiple sources: some cracked and some straight run, including coker cycle oils and light cycle oils. Of course, the coker naphtha varied in many proportions. I do not want to say on a daily basis; but when we had a lot of different cracked stocks in a unit and the units going up and down were changing the rates, the operators had to jockey all of these particular variables. It is somewhat difficult for them to understand how to change all of the conditions.



Co-Processing Coker Naphtha in ULSD Hydrotreater (Model Development)

- Objective : Develop "Operator Tables" providing guidance on reactor adjustments
- Configuration: 2 Parallel Reactors / 2 Beds / Inter-Bed Quench
- Input Factors : Overall Feed Rate (FdRt) from 14,000 B/d to 32,000 B/d
Coker Naphtha Rate (CNap) from Zero to 6,000 B/d
Percent Heavy Cracked Stocks (HvyCrk) from Zero to 25 %
(Heavy Cracked is mix of FCC LCO and Coker Cycle Oil)
Days On-Line (RunDays) from day 1 to EOR at 4 years
- Data Source: Selected, filtered, stable, on-spec data from first year of operation
- Other Factors: Feed ~ 0.7 Wt % S, 8 ppm Wt Product S, Gas-to-Oil 2200 SCF/B
- Desired Output: Predicted Heater Inlet and Required Overall Reactor W/ABT
Top Bed Delta Temperature (Monitored To Control Quench)



We were looking for an attainable response. In order to cover all of the interactive parameters, we elected to cover those with at least six interactive factors and four primaries squared. When we had about 15 different coefficients in the model, we looked at what the operators monitor. It was a limited slate of furnace inlet, top bed ΔT (temperature differential), and overall WABT (weighted average bed temperature) on the unit to get to spec. The modeling more or less showed quite a few interactive variables and quadratic surface responses. We found that it was not necessarily a linear function of the addition of some of these variables. Over the years, my experience has been that your typical board operator can handle two, three, or four variables at a time. But when you have multiple variables with multiple nonlinear interactions, it is difficult for him to perceive how to change the unit ahead of time preemptively, so a tool is required.

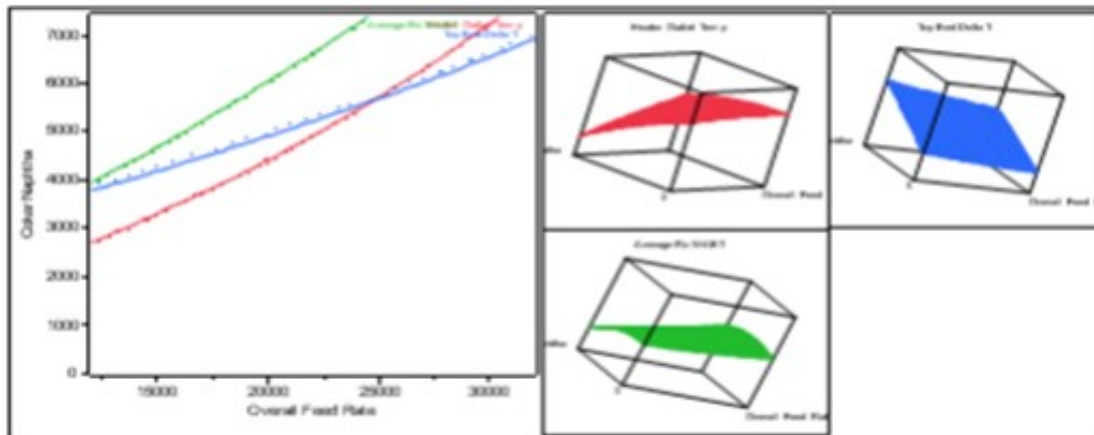
Co-Processing Coker Naphtha in ULSD Hydrotreater (Model Development)

- Model Selection: Interactive Binary Quadratic Model to show non-linearity
- Equation Format:
 - 4 Main Factor Effects
 - 6 Two Factor Interactions
 - 4 Primary Squared Factors
 - 1 Nominal Intercept
 - 15 Total Coefficients
- Response: Tables estimate Furnace Inlet, Top Bed DT and Overall WABT
- Increase in CNap Rate: Decrease Heater Outlet 3 to 5 Deg F per 1 K CNap addition
WABT is parabolic function of CNap addition rate
- Increase Hvy Crk: rates
 - Lower Heater Outlet more at low overall rates – less at high
 - Lower Outlets more pronounced (60 Deg F) at high CNap rates
 - WABT is parabolic function of CNap addition rate



Co-Processing Coker Naphtha in ULSD Hydrotreater (Model Development)

15 Parameter Model Output / 2 variables at a time



These types of graphs are not what you would share with the Operations staff, so we boiled them down to a set of tables. It is an old-school look-up. I mean, if people would say, "Well, just plug the numbers into the computer; it will tell the operators what to do," that would cut the amount of the long-term understanding of the variables and their interactions. So we prepared tables with a date stamp, and then the tables got up periodically. The tables contained most of the variables, so the operators were able to page through and read across on the table. It seemed to help them adjust the unit proactively to be on spec.

Q2



Co-Processing Coker Naphtha in ULSD Hydrotreater (Model Development)

Operator Parameter Table / Key variable look-up

Data Generated For Specific Days On-Line or - 1/2 Day Off For Admin Transfer	Individual Tables Prepared For Heavy Cracked Stocks Ranging From Min (Zero) To Max (25%)						
	Min Coker Naphtha Content			Coker Content 0 NBD to 7 NBD By 1 NBD increments ↔	Max Coker Naphtha Content		
Overall Unit Feed Rate	Header Outlet	1st NBD	Top Blvd UI		Header Outlet	1st NBD	Top Blvd UI
Minimum Rate	Date (Deg F)	Date (Deg F)	Date (Deg F)		Date (Deg F)	Date (Deg F)	Date (Deg F)
Unit Rates # NBD to 32 NBD ↓	Date (Deg F)	Date (Deg F)	Date (Deg F)		Date (Deg F)	Date (Deg F)	Date (Deg F)
Maximum Rate	Date (Deg F)	Date (Deg F)	Date (Deg F)		Date (Deg F)	Date (Deg F)	Date (Deg F)



LEICHTY (Chevron USA, Inc.)

Chevron has two high pressure hydrotreaters that co-process coker naphtha in a diesel service. The other feedstocks are coker light gas oil and FCC (fluid catalytic cracking) light cycle oil. Both units feed 100% cracked feed stocks.

Considerations: Beginning with the feed pump, the addition of coker naphtha must not create a mixture at the feed pump suction conditions such that the vapor pressure results in cavitation. Moving on to the preheat train, we find that diolefins in the feed can result in polymerization fouling. In some cases, an antioxidant injected into the coker naphtha rundown, combined with an antifoulant in the total unit feed, can mitigate the fouling. Also, we found that if the naphtha content is too high in the feed, accelerated fouling can result. We monitor this fouling by calculating the degrees below dew point in the preheat exchangers. This fouling may be due to lower stability or solubility of gums as the mixture partially vaporizes through the exchangers. We have also determined that the presence of LCO (light cycle oil) in the feed can sometimes help keep gums and other polymerization products soluble, thereby mitigating furnace and/or exchanger fouling. In the reactor, silica poisoning of the catalyst can be mitigated through the use of some of the newer generation coker antifoams containing lower silica content. If the silica is from another source, a guard bed may be necessary to achieve the desired run length. With respect to desulfurization of the diesel, there may be a negative space velocity impact if the naphtha feed is added to the existing diesel range feed.

Co-processing Coker Naphtha in Diesel HDS



- Chevron has two high pressure hydrotreaters that co-process Coker Naphtha in a Diesel HDS
- Considerations
 - Feed temperature / pump cavitation
 - Di-olefins foul preheat train and cause reactor ΔP
 - Filtration and Antioxidant/Antifoulant injection
 - Degrees below dew point in heat exchangers
 - LCO in feed can keep gums soluble
 - Silica causes catalyst de-activation - Mitigate with guard bed(s) or reactor catalyst and conditions
 - LHSV effect if naphtha is additive to diesel feed

Additional Considerations: In order to effectively strip H₂S (hydrogen sulfide) from the product and separate the naphtha from distillate, a stripper column followed by a fractionator is an effective configuration. Note that even with this configuration, the naphtha will still contain some H₂S; but, the H₂S content can be made low enough to feed directly to a reformer. Another important aspect to consider is the ability of the relief system to handle the addition of coker naphtha. A relief system study should be performed to ensure adequate capacity. Lastly, but most importantly, the olefins in the coker naphtha cause high heat release in the reactor, temperature control challenges, and high hydrogen consumption. If not properly managed, the effluent train temperature profile could change such that the

location of the metallurgy switch between carbon and stainless steel is no longer adequate to protect against high temperature hydrogen attack and high temperature sulfidation. These two corrosion mechanisms can lead to catastrophic failure. In addition, the higher temperature in the effluent train may reduce the percent liquid water remaining at the continuous water injection point, which can lead to a failure in the reactor effluent air cooler below the 25% minimum guideline.

G22

Coker Naphtha to Diesel HDS



AFPM

American
Fuel & Petrochemical
Manufacturers

The new NPRA.

- **Considerations (continued)**

- Stripper followed by fractionator

- Light naphtha will still have H₂S

- Heavy naphtha can go direct to reformer

- Relief system capacity

- Heat release

- Watch out for max bed temperature limits, high temperature H₂ attack, sulfidation and minimum liquid water remaining at injection point(s)

I think we are letting the panelists off a little too easy, so I will ask a question. Does the coker naphtha that you mentioned include the C5/C6 portion, sometimes called PenHex? If it does, can you comment on a specific experience you have had with that portion of the naphtha?

LEICHTY (Chevron USA, Inc.)

We do have coker C5 and C6s (coker PenHex) contained in the coker naphtha that is fed to our distillate hydrotreaters. These low boiling point molecules contain a higher percentage of olefins and are the most reactive; therefore, they are more challenging, from a fouling and reactor heat-release perspective.

BODOLUS (CVR Energy)

Our coke naphtha contains it as well. We process it as we get it.

ALAN WELDON (Hunt Refining Company)

What do you consider a high percentage of cracked coker naphtha versus just diesel?

BODOLUS (CVR Energy)

High would be on the order of 20%, perhaps.

LEICHTY (Chevron USA, Inc.)

Limits could be encountered in several sections of the unit, as discussed in the considerations mentioned previously. If the maximum reactor bed/total temperature rise is the only limiting factor, then 35% coker naphtha can be achievable.

BODOLUS (CVR Energy)

The primary operating constraints in co-processing coker naphtha in a heavy oil hydrotreaters concern dilution of hydrogen partial pressure (due to naphtha vaporization) and the effect that the exotherm has on the unit heat balance. Inclusion of coker naphtha increases the top bed exotherm, often reducing heater duty. Simultaneously, there may be an increase in stripper section duty to lift the product naphtha into the overhead stream.

Primary contamination constraints in coker naphtha include silicon resulting from the decomposition of antifoams used during the coking process. The silicon contamination deposits on the hydrotreating catalyst leading to deactivation and has a severe negative impact on the prospects of regenerating the spent catalyst. Top bed fouling, due to diolefin content of coker naphtha, can also limit run length due to differential pressure build. If the unit is expected to have a steady diet of coker naphtha, catalyst loading options in the top bed can include a silicon trap and/or a layer of controlled diolefin saturation catalyst along with enhanced void bed grading. Note that coker naphtha holding tanks should be protected from oxygen ingress, which produces peroxy free radicals, and long-term storage at elevated temperatures due to olefin/diolefin content.

From an economic standpoint, coker naphtha processing in a distillate hydrotreaters also causes a loss of octane due to saturation of olefins. Consumption of hydrogen will be high and resulting treated naphtha may have residual nitrogen liability. Other processing options, such as co-processing in an ultra-low sulfur gasoline hydrotreater, can preserve octane values and allow treated coker naphtha to enter the gasoline pool (preventing nitrogen liability on the reformer).

Balancing the heat duty and overall rate changes in heavy oil hydrotreaters with variable coker naphtha is not always intuitive to the operations staff. After a more than a year of operations it was decided to develop a transition tool to assist operators in “Predicting” what new heater setting would be required with a rate change in any of the feedstocks (coker naphtha, FCC light cycle oil, coker cycle oil and straight run diesel).

The following slides and tables show how selected process variables were chosen to develop a model for the operator to “Predict” reactor parameters necessary to minimize off-spec product and reduce giveaway.

Co-Processing Coker Naphtha (CNap) in ULSD Hydrotreater (Model Development)

Objective: Develop "Operator Tables" providing guidance on reactor adjustments

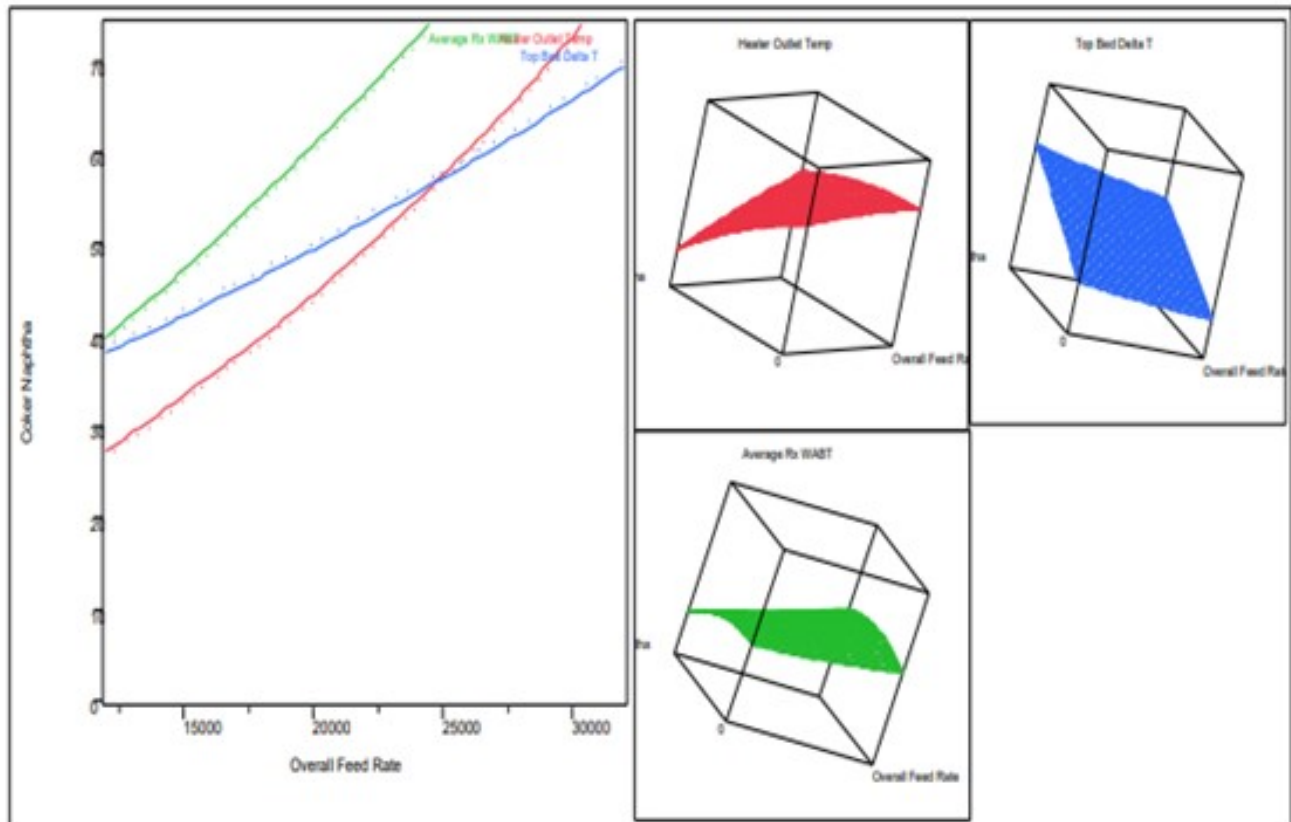
<u>Reactor Configuration:</u>	2 parallel reactors / 2 beds / inter-bed quench
<u>Input Factors:</u>	Overall feed rate (FdRt) from 14 kbpd to 32 kbpd Coker naphtha rate (CNap) from 0 to 6 kbpd Percent heavy cracked stocks (HvyCrk) from zero to 25% (Heavy cracked is mix of FCC LCO and coker cycle oil.) Days online (run days) from Day 1 to EOR at four years
<u>Data Source:</u>	Selected, filtered, stable, on-spec data from first year of operation
<u>Nominal Factors:</u>	Feed ~0.7 wt% S, 8 wppm Product S, gas-to-oil 2,200 scfb (standard cubic foot per barrel)
<u>Desired Response:</u>	Predicted heater inlet and required overall reactor WABT Top bed delta temperature (monitored to control quench)
<u>Model Selection:</u>	Interactive Binary Quadratic Model to show nonlinearity
<u>Equation Format:</u>	4 main factor effects 6 two factor interactions 4 primary squared factors 1 nominal intercept 15 total coefficients
<u>Response Matrix:</u>	Tables estimate furnace inlet, top bed DT (ΔT ; temperature differential) and overall WABT
<u>Increase in CNap:</u>	Decrease heater outlet 3°F to 5°F per 1 K CNap addition WABT is parabolic function of CNap addition rate.
<u>Increase SR Only:</u>	Increase heater outlet depending on existing CNap charge Small increase at low CNap rates (less than 3 K B/d) Large increase at high CNap rate (more than 3 K B/d)
<u>Increase Heavy Crack:</u>	Lower heater outlet more at low overall rates – less at high rates Lower outlets more pronounced (60°F) at high CNap rates WABT is parabolic function of CNap addition rate.

Co-Processing Coker Naphtha (CNap) in ULSD Hydrotreater

Operator Parameter Table / Key Variable Look-up

<i>Data Generated For Specific Days On-Line or - 1.2 Deg F Per Month thereafter</i>	Individual Tables Prepared For Heavy Cracked Stocks Ranging From Min (Zero) To Max (25%)						
Overall Unit Feed Rate	Min Coker Naphtha Content			<i>Coker Content 0 KBD to 7 KBD By 1 KBD Increments</i> ↔	Max Coker Naphtha Content		
	Heater Outlet	Rx WABT	Top Bed DT		Heater Outlet	Rx WABT	Top Bed DT
Minimum Rate	Data (Deg F)	Data (Deg F)	Data (Deg F)		Data (Deg F)	Data (Deg F)	Data (Deg F)
<i>Unit Rates 14 KBD to 32 KBD</i> ↓	Data (Deg F)	Data (Deg F)	Data (Deg F)		Data (Deg F)	Data (Deg F)	Data (Deg F)
Maximum Rate	Data (Deg F)	Data (Deg F)	Data (Deg F)	Data (Deg F)	Data (Deg F)	Data (Deg F)	

15 Parameter Model Output / 2 variables at a time



KASPAR VOGT and STEVE MAYO (Albemarle Corporation)

Coker naphtha is typically high in sulfur, nitrogen, diolefins and silicon. It often also contains significant amounts of arsenic and in some cases particulates from the coker and iron scale. Treating coker naphtha is much more challenging than straight run naphtha, even when it is blended with straight run naphtha. For these reasons, refiners are sometimes tempted to treat coker naphtha in higher pressure units with a heavier feedstock such as diesel or VGO (vacuum gas oil). This can be done successfully, but there are caveats to consider before embarking down such a path:

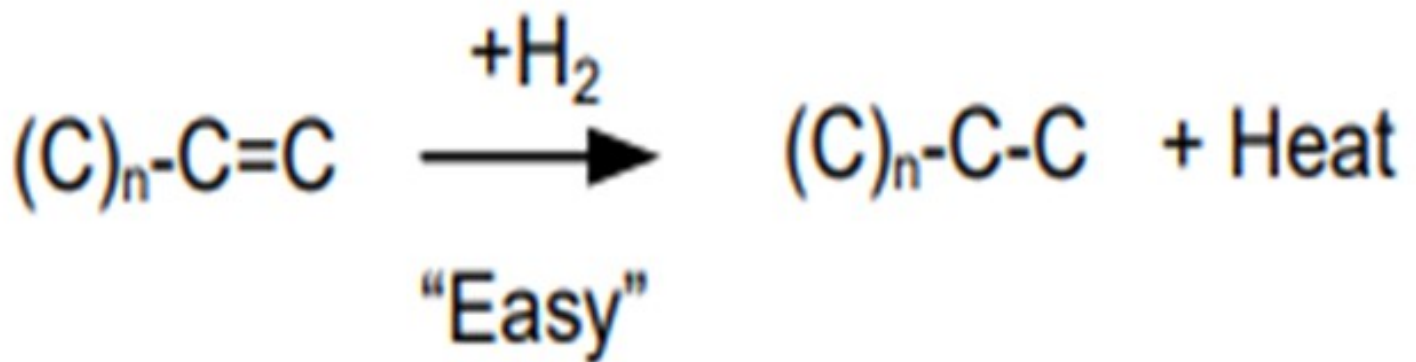
- The addition of a lighter feed component that fully vaporizes will reduce hydrogen partial pressure, potentially reducing the unit's performance with regard to its primary feedstock.
- The introduction of olefins and diolefins may result in higher exotherms, as well as increased hydrogen consumption. As a consequence, hydrogen partial pressure will be reduced and potentially also the operating window for the unit. When possible, coker naphtha used as a liquid quench can alleviate its exotherm by heat of vaporization. A downside is introduction of catalyst poisons in the lower beds of the hydrotreater.
- Poisons such as As (arsenic) and Si (silicon) will be introduced to the unit and may require a more extensive guard bed system to handle them. A larger portion of the reactor devoted to guard bed catalyst will reduce the volume available for main bed catalyst and could impact cycle length.
- In particular for ULSD, the most difficult sulfur compounds require hydrogenation before they can be desulfurized. Adding coker naphtha, with its typically high N (nitrogen) content, increases organic nitrogen into the reactor and may increase the level of nitrogen inhibition of catalyst active sites. This disproportionately affects hydrogenation sites and effectively reduces the rate of removal of the most difficult sulfur compounds.
- VGO units, with their typically lower LHSV (liquid hourly space velocity) and higher hydrogen partial pressure, are better equipped to handle the introduction of coker naphtha. However, the naphtha will operate in the vapor phase and may encounter diffusion issues with the VGO operating in liquid phase, reducing the rate of sulfur and nitrogen removal.
- The fractionated naphtha stream will be too high in mercaptans (due to recombination) to send directly to the reformer and will need to be post-treated

MEREDITH LANSDOWN, BRIAN WATKINS, and BRIAN SLEMP (Advanced Refining Technologies)

Co-processing coker naphtha in ULSD service can have several undesirable effects on the performance of the hydrotreater and the catalyst if the system was not properly designed to handle it. In general, coker stocks have a higher level of olefins present from the coking process. Once in the hydrotreater these olefins will quickly get saturated (Figure 1) consuming additional hydrogen and generating extra heat. As a general rule of thumb, one mole of hydrogen is required per mole of carbon/carbon double

bond, or between five and 10 times the bromine number reduction in standard cubic feet of hydrogen per barrel (scfb). This additional heat (130 BTU/scf (British thermal unit per standard cubic foot) to 160 BTU/scf hydrogen consumed), if not spread out through a decent portion of the catalyst bed, will initiate the subsequent reactions creating a much higher temperature rise than expected. This excess temperature can also speed up the coking or polymerization mechanism which will lead to an increase in pressure drop. This can set an upper limit as to how much coker naphtha can be processed either by a need to limit the heat rise, or from too much hydrogen consumption that could starve the downstream catalysts.

Figure 1. Olefin Saturation



A system that is properly size and activity graded will be extremely important when co-processing coker naphtha in a diesel unit. ART utilizes a grading system to help mitigate pressure drop build-up. ART's GSK-19 is a 19-mm inert ring with a very high void fraction used for trapping large particulates and is placed at the top of the reactor. GSK-9 is loaded next and is a 9-mm macro-porous ring that traps iron, as well as other finer particulates that can increase pressure drop. ART also utilizes two other types of active grading, GSK-6A and GSK-3A, which are smaller rings with a small number of active metals present in order to begin any olefin saturation reaction, as well as provide additional void space at the top of the reactor. Underneath the grading options, it is recommended to use a layer of ART's AT724G or AT734G which can provide both additional olefin saturation, additional void fraction for pressure drop mitigation, as well as a trapping mechanism for silicon (and arsenic), which is another concern with co-processing coker naphtha in a ULSD unit.

Another major concern is that coker naphtha can also bring silicon into the unit which is a permanent poison for hydrotreating catalyst. A silicon guard, such as ART's AT724G or AT734G, should be loaded in the reactor to mitigate silicon poisoning. Silicon pickup is temperature dependent, and at the higher temperatures ULSD units are operating at, silicon pickup in the order of 16 wt% to 25 wt% could be

expected with AT724G or AT734G. If arsenic is present in the coker stocks, the use of AT734G is preferred as it will have the same silicon pickup as AT724G and will also protect the active catalyst against arsenic poisoning.

A third concern is the high degree of vaporization of the coker naphtha. ULSD hydrotreaters are typically designed such that their feed distribution system will contain liquid, and the additional gas present from the coker naphtha may cause some systems to perform poorly giving rise to maldistribution. In order to minimize feed vaporization and poor distribution tray utilization, the coker naphtha should be mixed with the other feed streams at a temperature where it is still liquid before feeding to the charge heater. The recovery system should also be evaluated for the increase in naphtha that will be present so that the downstream equipment is not overloaded.

A final consideration would be that additional coker naphtha in a diesel can generate incremental dry gas products such as methane and ethane. These products will increase in concentration in the recycle gas loop, causing a decrease in the hydrogen partial pressure for the hydrotreater. It will also increase the molecular weight of the recycle gas, which can lead to compressor capacity limitations. These additional products can also lead to incremental stripper off-gas and related problems.

PAUL CECCATO (Criterion Catalysts & Technologies)

Several factors should be evaluated when considering the co-processing of coker naphtha in either ultra-low sulfur diesel or FCC pretreat units. Changes to the hydrogen partial pressure, hydrogen consumption, reactor fouling, reactor temperature profiles, fractionation capabilities, and catalyst deactivation will impact both types of operations. Ultimately, the selection of where to co-process coker naphtha is based on the relative impacts of these factors on each unit.

Co-processing coker naphtha reduces hydrogen partial pressure due to the vaporization of the naphtha range material and increased hydrogen consumption through saturation of its high diolefin/olefin content. A reduction in hydrogen partial pressure directionally reduces heteroatom conversion, aromatic saturation, and catalyst stability. In a ULSD operation, treating the more sterically hindered sulfur species with less hydrogenation activity becomes more difficult, limiting the feed endpoint and the percentage of cracked stock in the feed blend. In an FCC pretreat unit, reductions in nitrogen and aromatic conversions directionally lower FCC conversion.

As the hydrogen partial pressure decreases, aromatic equilibrium and condensation occur at lower temperatures. For a ULSD unit, cycle life may shorten due to increased coking and insufficient hydrogenation activity at elevated temperatures. In FCC pretreat operations, cycle life also declines due to accelerated catalyst deactivation as larger aromatics condense to coke at lower temperatures.

Saturation of the diolefins and olefins introduced with coker naphtha consumes significant hydrogen, depleting excess hydrogen at the reactor outlet. ULSD or FCC pretreat units operating with marginal hydrogen treat gas-to-makeup ratios may experience hydrogen starvation and accelerated catalyst deactivation. Insufficient excess hydrogen may also result in lower bed pressure drop issues when processing cracked stocks in either the ULSD or FCC pretreat units or high asphaltene feeds in the FCC pretreat unit due to rapid coking.

Polymerization of the diolefins/olefins in the coker naphtha may lead to increased reactor fouling and upper bed pressure drop. Typically, neither ULSD nor FCC pretreat units are designed with low temperature lead reactors for mild saturation of the highly reactive diolefins. Competition for active catalyst sites at the top of a reactor may leave diolefins and olefins unsaturated and exposed to elevated temperature which drives polymerization.

Heat release from diolefin and olefin saturation is significant and may result in a more ascending temperature profile in a single bed reactor and increasing quench requirement in a multi-bed reactor. Maintaining preferred equal bed outlet temperature profiles can become difficult as quench is diverted upward in the reactor. Radial temperature spreads in reactors with older reactor internals may increase with the increased axial delta T (temperature differential) and push established operating limits, constraining the reactor inlet temperature.

Delayed coker naphtha typically contains varying quantities of silicon from antifoam addition. Si is a poison to hydrotreating catalyst with a larger impact on the denitrification function. Reactor loadings can be adjusted to include high capacity Si trap catalyst at the expense of high activity catalyst volume. In a ULSD unit, Si induced catalyst deactivation impacts cycle life. In the FCC pretreat unit which typically contains more catalyst volume, the Si impact is less; but when combined with other metals induced deactivation, cycle life may be impacted or a reduced.

Lastly, co-processing coker naphtha requires sufficient fractionation capabilities to recover the naphtha product. Poor fractionation in a ULSD unit may result in heavy naphtha contamination of the diesel product for lower cetane. Poor fractionation in the FCC pretreat unit may result in insufficient naphtha recovery and the reprocessing of an increased diesel draw through the ULSD unit.

The impacts of co-processing coker naphtha in a ULSD or FCC pretreat units should be evaluated individually and compared to unit constraints and objectives. Ultimately, the selected unit will be capable of minimizing the impacts.

Print as PDF:

Tags

[Aromatics](#)

[Blending](#)

[Catalysts](#)

[Coker](#)

[Compressors](#)

[Delayed Coking](#)

[Fouling](#)

[Hydrogen](#)

[Octane](#)

[Operations](#)

[Poisoning](#)

[Process](#)

[Reactor Vessel](#)

[Reforming](#)

[Regenerator](#)

[Tankage](#)

[ULSD](#)